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## (54) TREATING AND PREVENTING MICROBIAL INFECTIONS

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#### (57) ABSTRACT

The invention provides methods for treating or preventing microbial (eg, bacterial) infections and means for performing these methods. In particular, treatment of infections requiring rapid and durable therapy is made possible, such as for treating acute conditions such as septicemia, sepsis, SIRS or septic shock. The invention is particularly useful, for example, for treatment of microbes such as for environmental, food and beverage use. The invention relates inter alia to methods of controlling microbiologically influenced corrosion (MIC) or biofouling of a substrate or fluid in an industrial or domestic system. The invention also useful for the treatment of pathogenic bacterial infections in subjects receiving a treatment for a disease or condition, such as a transplant or a treatment for cancer, a viral infection or an autoimmune disease.

#### 19 Claims, 9 Drawing Sheets

Specification includes a Sequence Listing.

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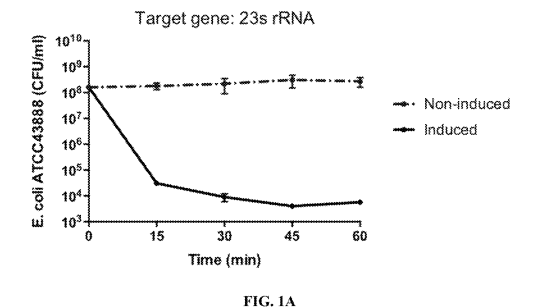
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\* cited by examiner



 Dilution
 0
 1
 2
 3
 4
 5
 6

 Non-induced
 Induced
 Induced

FIG. 1B

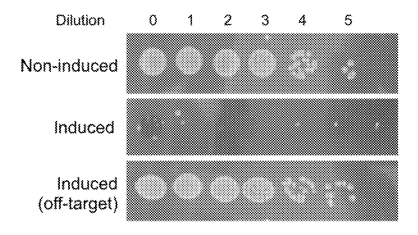


FIG. 2

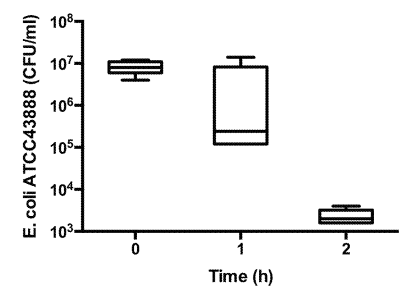


FIG. 3

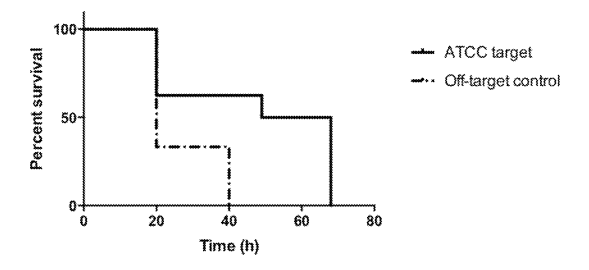


FIG. 4

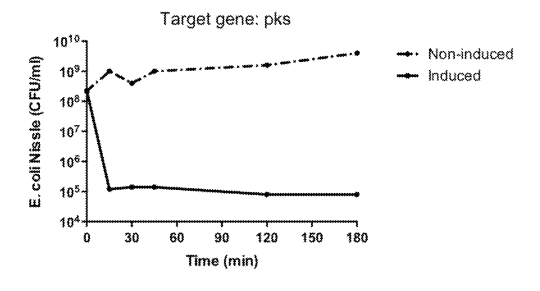


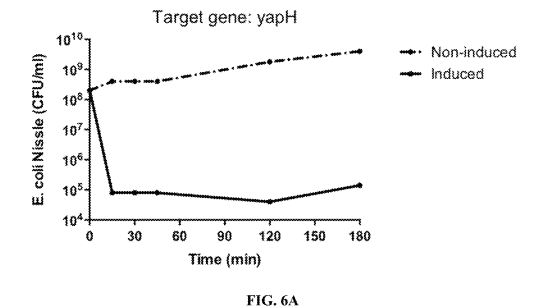
FIG. 5A

Dilution 0 1 2 3 4 5 6

Non-induced

Induced

FIG. 5B



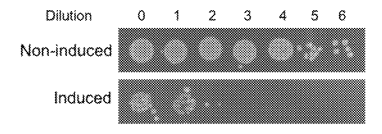
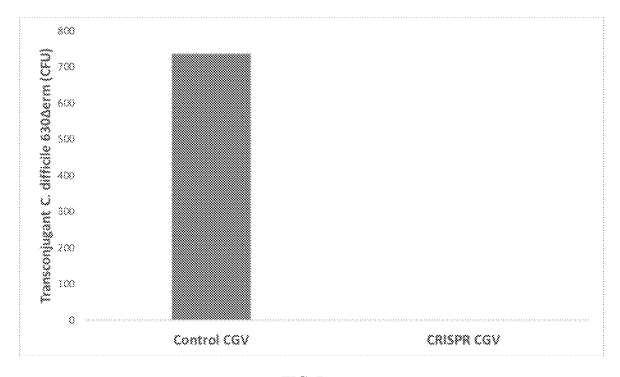
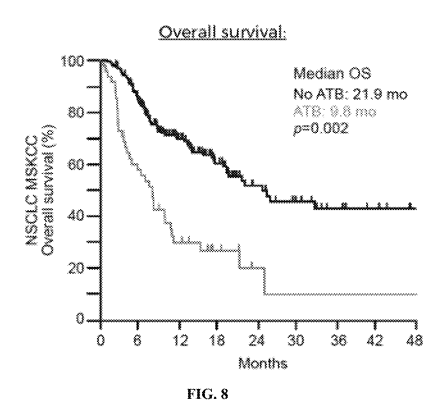


FIG. 6B



**FIG.** 7



# Progression-free survival by fecal diversity:

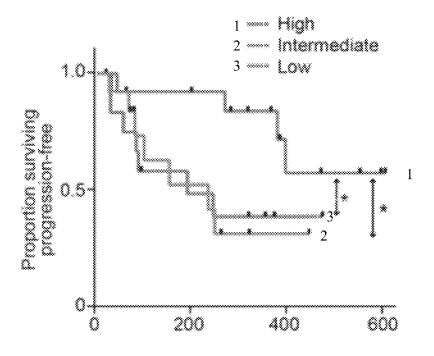


FIG. 9A

### Progression-free survival by low/high abundance of certain bacteria:

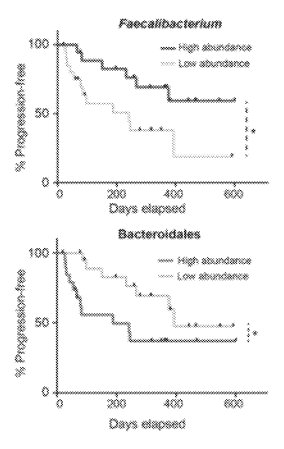


FIG. 9B

# TREATING AND PREVENTING MICROBIAL INFECTIONS

# SUBMISSION OF SEQUENCE LISTING ON ASCII TEXT FILE

The content of the following submission on ASCII text file is incorporated herein by reference in its entirety: a computer readable form (CRF) of the Sequence Listing (file name: 786212000300SEQLIST.txt, date recorded: Apr. 30, 2018, size: 21 KB).

#### TECHNICAL FIELD

The invention provides methods for treating or preventing microbial (eg, bacterial) infections and means for performing these methods. In particular, treatment of infections requiring rapid or durable therapy is made possible, such as for treating acute conditions such as septicemia, sepsis, SIRS or septic shock. The invention is also particularly useful, for example, for treatment of microbes for environmental, food and beverage use. The invention relates inter alia to methods of controlling microbiologically influenced corrosion (MIC) or biofouling of a substrate or fluid in an 25 industrial or domestic system.

The invention also useful for the treatment of pathogenic bacterial infections in subjects receiving a treatment for a disease or condition, such as a transplant or a treatment for cancer, a viral infection or an autoimmune disease.

#### BACKGROUND

Septicaemia is an acute and serious bloodstream infection. It is also known as bacteraemia, or blood poisoning. 35 Septicaemia occurs when a bacterial infection elsewhere in the body, such as in the lungs or skin, enters the bloodstream. This is dangerous because the bacteria and their toxins can be carried through the bloodstream to a subject's entire body. Septicaemia can quickly become life-threatening. It 40 must be rapidly treated, such as in a hospital. If it is left untreated, septicaemia can progress to sepsis.

Septicaemia and sepsis aren't the same. Sepsis is a serious complication of septicaemia. Sepsis is when inflammation throughout the body occurs. This inflammation can cause 45 blood clots and block oxygen from reaching vital organs, resulting in organ failure. The US National Institutes of Health (NIH) estimates that over 1 million Americans get severe sepsis each year. Between 28 and 50 percent of these patients may die from the condition. When the inflammation 50 occurs with extremely low blood pressure, it's called septic shock. Septic shock is fatal in many cases.

The increase in average age of the population, more people with chronic diseases, on immunosuppressive drugs, and increase in the number of invasive procedures being 55 performed has led to an increased rate of sepsis. People over 65 years old, particularly those who have health issues, are even more susceptible to sepsis than any other group. According to a study published in 2006, while people aged 65 years and older make up about 12% of the American 60 population, they make up 65% of sepsis cases in the hospitals.

Septicaemia is caused by an infection in a part of the body. This infection is typically acute. Many types of bacteria can lead to septicaemia. The exact source of the infection often 65 can't be determined. The most common infections that lead to septicaemia are:

2

urinary tract infections

lung infections, such as pneumonia

kidney infections

infections in the abdominal area

Bacteria from these infections enter the bloodstream and multiply rapidly, causing acute infection and immediate symptoms.

People who are already in the hospital for something else, such as a surgery, are at a higher risk of developing septicaemia. Secondary infections can occur while in the hospital. These infections are often more dangerous because the bacteria may already be resistant to antibiotics. There is a higher risk of developing septicaemia if the subject:

has severe wounds or burns

is very young or very old

has a compromised immune system, which can occur from diseases such as HIV or leukaemia

has a urinary or intravenous catheter

is on mechanical ventilation

is receiving medical treatments that weakens the immune system, such as chemotherapy or steroid injections

The symptoms of septicaemia usually start very quickly. Even in the first stages of the illness, a person can look very sick. They may follow an injury, surgery, or another localized (eg, confined to one location) infection, like pneumonia. The most common initial symptoms are:

chills

elevated body temperature (fever)

very fast respiration

rapid heart rate

More severe symptoms will begin to emerge as the septicaemia progresses without proper treatment. These include the following:

confusion or inability to think clearly

nausea and vomiting

red dots that appear on the skin

reduced urine volume

inadequate blood flow (shock)

be carried through the bloodstream to a subject's entire body. Septicaemia can quickly become life-threatening. It was be rapidly treated, such as in a hospital. If it is left untreated, septicaemia can progress to sepsis.

Septicaemia that has started to affect the organs or tissue function is an acute medical emergency. It must be rapidly treated at a hospital. Many people with septicaemia are admitted to a hospital's ICU for treatment and recovery. It is recommended to never take a "wait and see" approach or try to treat the problem at home. It is crucial to get to the hospital right away if the subject is showing signs of septicaemia.

Septicaemia has a number of serious complications. These complications may be fatal if left untreated or if treatment is delayed for too long.

Septic Shock

One complication of septicaemia is a serious drop in blood pressure. This is called septic shock. Toxins released by the bacteria in the bloodstream can cause extremely low blood flow, which may result in organ or tissue damage. Septic shock is an acute medical emergency. People with septic shock are usually cared for in a hospital's intensive care unit (ICU). The patient may need to be put on a ventilator, or breathing machine, if in septic shock.

Acute Respiratory Distress Syndrome (ARDS)

Another complication of septicaemia is acute respiratory distress syndrome (ARDS). This is a life-threatening condition that prevents enough oxygen from reaching your lungs and blood. According to the National Heart, Lung, and Blood Institute (NHLBI), ARDS is fatal in about one-third of cases. It often results in some level of permanent lung damage. It can also damage the brain, which can lead to memory problems.

Sepsis

Sepsis occurs when the body has a strong immune response to the infection. This leads to widespread inflammation throughout the body. It is called severe sepsis if it leads to organ failure. People with chronic diseases, such as 5 HIV or cancer, are at a higher risk of sepsis. This is because they have a weakened immune system and cannot fight off the infection on their own. Sepsis causes millions of deaths globally each year and is the most common cause of death in people who have been hospitalized. The worldwide incidence of sepsis is estimated to be 18 million cases per year. In the United States sepsis affects approximately 3 in 1,000 people, and severe sepsis contributes to more than 200,000 deaths per year. Sepsis occurs in 1-2% of all  $_{15}$ hospitalizations and accounts for as much as 25% of ICU bed utilization.

Early diagnosis is necessary to properly manage sepsis, as initiation of rapid therapy is key to reducing deaths from severe sepsis. Within the first three hours of suspected 20 sepsis, diagnostic studies should include white blood cell counts, measuring serum lactate, and obtaining appropriate cultures before starting antibiotics, so long as this does not delay their use by more than 45 minutes

The most common primary sources of infection resulting 25 in sepsis are the lungs, the abdomen, and the urinary tract. Typically, 50% of all sepsis cases start as an infection in the lungs.

Speed of treatment is essential. Two sets of blood cultures (aerobic and anaerobic) should be taken without delaying 30 the initiation of antibiotics. Cultures from other sites such as respiratory secretions, urine, wounds, cerebrospinal fluid, and catheter insertion sites (in-situ more than 48 hours) can be taken if infections from these sites are suspected. In severe sepsis and septic shock, broad-spectrum antibiotics 35 (usually two, a β-lactam antibiotic with broad coverage, or broad-spectrum carbapenem combined with fluoroquinolones, macrolides, or aminoglycosides) are conventional. However, combination of antibiotics is not recommended for the treatment of sepsis without shock and in immuno- 40 compromised persons unless the combination is used to broaden the anti-bacterial activity. The administration of antibiotics is important in determining the survival of the person. Some recommend they be given within one hour of making the diagnosis, stating that for every hour of delay in 45 the administration of antibiotics, there is an associated 6% rise in mortality.

Early goal directed therapy (EGDT) is an approach to the management of severe sepsis during the initial 6 hours after diagnosis. It is a step-wise approach, with the physiologic 50 goal of optimizing cardiac preload, afterload, and contractility. It includes giving early antibiotics.

Neonatal sepsis can be difficult to diagnose as newborns may be asymptomatic. If a newborn shows signs and symptoms suggestive of sepsis, antibiotics are immediately 55 started and are either changed to target a specific organism identified by diagnostic testing or discontinued after an infectious cause for the symptoms has been ruled out.

Approximately 20-35% of people with severe sepsis and 30-70% of people with septic shock die. The Surviving 60 Sepsis Campaign (SSC) is a global initiative to bring together professional organizations in reducing mortality from sepsis. Antibiotics are administered within two hours of admission/diagnosis. For every hour a patient is denied antibiotic therapy after the onset of septic shock, the 65 patient's chance of survival is reduced by 7.9% (Survivesepsis.org 2005)

There is, therefore, a need for a rapid treatment of acute microbial infections, such as bacterial infections associated with septicaemia, sepsis or septic shock. It would also be advantageous if the treatment is durable for many hours. Rapid and durable treatment of microbes is also desirable for is for controlling microbiologically influenced corrosion (MIC) or biofouling of a substrate in industrial and domestic

Acute bacterial infections can, in certain circumstances, be health-threatening or even life-threatening. This may be the case, for example, in cancer patients, transplant patients or other subjects. The need for the treatment of the bacterial infection can become urgent and indeed an immediate focus of attention in the medical care. It would be useful to provide methods of treating such pathogenic bacterial infections in a way that does not adversely undermine the efficacy of the cancer or other separate therapy to which the patient also needs to respond.

#### SUMMARY OF THE INVENTION

The invention provides a solution by using the action of programmable nuclease cutting of microbe genomes; this is different from the metabolic inhibitor and other mechanisms of action used by beta-lactams and other conventional antibiotics for treating infections. The targeted cutting provides selective microbe killing or reduction of growth or proliferation to treat or prevent infection. Moreover, the inventors have surprisingly found a substantial killing (by several logs) can be achieved very rapidly (eg, within 15 minutes) and sustainable effects can be achieved (eg, for more than 1 hour, and even around 3 hours after treatment commenced) in some embodiments. Thus, the invention provides the following configurations.

In a First Configuration

A programmable nuclease for use in a method of treating a microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated.

In a Second Configuration

A plurality of viruses (eg, phage or phagemids for producing phage) for use with a programmable nuclease in a method of treating a microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and viruses wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated; wherein each virus comprises a copy of a nucleic acid that

encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject;

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wherein the viruses are capable of infecting microbes comprised by the subject to deliver thereto the nucleic acid.

In a Third Configuration

A composition comprising a plurality of nucleic acids for programming a programmable nuclease in a method of 5 treating a microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain 10 are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and the nucleic acids wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut 15 and microbial infection of the subject is treated;

wherein each nucleic acid encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject.

In a Fourth Configuration

A CRISPR/Cas system comprising a nuclease according to the invention for use in the method of treatment, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9) and the system comprises one or more guide RNAs or DNA encoding one or more guide RNAs, wherein each guide RNA is capable of programming the Cas nuclease to cut a target site comprised by the genomes of the microbes.

In a Fifth Configuration

A method of treating a microbial infection of a subject, 30 wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the 35 microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated.

In a Sixth Configuration

A method of treating a microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that 45 have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and a plurality of viruses wherein the nuclease is programmed to cut the target 50 site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated; wherein each virus comprises a copy of a nucleic acid that encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to 55 program the nuclease to cut the target site in microbes comprised by the subject; wherein the viruses are capable of infecting microbes comprised by the subject to deliver thereto the nucleic acid.

In a Seventh Configuration

A method of treating a microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first 65 species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising

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exposing the subject to the nuclease and a plurality of nucleic acids wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated; wherein each virus comprises a copy of a nucleic acid that encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject; wherein each nucleic acid encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject.

In a Eighth Configuration

Use of a nuclease, plurality of viruses, system, guide RNA, DNA or vector of the invention, in the manufacture of a composition for carrying out a method of treatment as defined herein, wherein the subject is an organism other than a human or animal.

In a Ninth Configuration

Use of a nuclease, plurality of viruses, system, guide RNA, DNA or vector of the invention, in the manufacture of a composition for carrying out an ex vivo or in vitro a method of treatment of a microbial infection of a substrate, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the substrate, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the substrate is treated.

In a Tenth Configuration

Use of a programmable nuclease in the manufacture of a composition for carrying out an ex vivo method of treatment of a microbial infection of a substrate, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the substrate, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the substrate is treated.

In any Configuration:

For example, the infection is an acute infection. For example, the infection is an acute infection that is rapidly treated. For example, the infection is treated rapidly—for example, the method comprises reducing the infection at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment. For example, the treatment is durable—for example, the reduction in infection persists for at least 30 minutes immediately after the first 30 minutes of the treatment. Also, optionally a reduction of the infection by at least 100-fold or 1000-fold is maintained for at least 60 minutes (eg, at least 120 minutes) after commencement of the treatment. Exemplification is provided below which surprisingly demonstrates these, such as a rapid killing that was durable around 3 hours after treatment commenced. For example, the method improves survival of the subject, or improves survival rates in humans or human patients suffering from infection by the microbes of the first species or

The invention also provides a solution to the need for effective treatment of pathogenic bacterial infections in subjects undergoing a cancer or other, separate therapy which must also be efficacious. Thus, the invention further provides:—

In an Eleventh Configuration

A method for treating a pathogenic bacterial infection in a human or animal subject caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a programmable nuclease that is programmed to cut the target site, wherein the subject is suffering from a further disease or condition other than the pathogenic bacterial infection and the method comprises administering a therapy to the subject for treating or preventing the further disease or condition, wherein the nuclease treats the infection and the therapy is efficacious in the presence of the programmed nuclease to treat or prevent the disease or condition.

In a Fo

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) that are *E coli*, *Pseudomonas aeruginosa* or *Klebsiella* bacteria, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer;

Wherein

- (a) The immunotherapy comprises administering to the 35 patient an anti-PD-1/PD-L1 axis antibody optionally selected from pembrolizumab, nivolumab, atezolimumab, avelumab and durvalumab; and
- (b) The cancer is selected from melanoma; renal cell carcinoma; bladder cancer; a solid tumour; non-small 40 cell lung cancer (NSCLC); forehead and neck squamous cell carcinoma (HNSCC); Hodgkin's lymphoma; a cancer that overexpresses PD-L land no mutations in EGFR or in ALK; colorectal cancer and hepatocellular carcinoma.

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the 50 cutting is carried out using a programmable nuclease that is programmed to cut the target site, wherein the subject is suffering from a cancer and the method comprises administering a cancer therapy to the subject for treating the cancer, wherein the nuclease treats the infection and the 55 cancer therapy is efficacious in the presence of the programmed nuclease to treat the cancer.

In a Twelfth Configuration

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first 60 species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the 65 method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nucle-

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ase treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer.

In a Thirteenth Configuration

A programmable nuclease for use in the method of the invention.

In a Fourteenth Configuration

A CRISPR/Cas system comprising a nuclease according to the 13<sup>th</sup> Configuration for use in the method of the 11<sup>th</sup> or 12<sup>th</sup> Configuration, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9) and the system comprises one or more guide RNAs (gRNAs) or DNA encoding one or more guide RNAs, wherein each guide RNA is capable of programming the Cas nuclease to cut a target site comprised by the genomes of first bacteria.

In a Fifteenth Configuration

A guide RNA or a DNA encoding a guide RNA for use in the system or method of treating a pathogenic bacterial infection.

In a Sixteenth Configuration

A nucleic acid vector comprising the guide RNA or DNA. In a Seventeenth Configuration

A pharmaceutical composition comprising a first nucleic acid vector (or a plurality thereof) encoding the nuclease and a second nucleic acid vector (or a plurality thereof) encoding the guide RNA.

#### BRIEF DESCRIPTION OF THE DRAWINGS

FIGS. 1A-1B show time-kill curves for *Escherichia coli* (EHEC) ATCC43888 strain harboring the CGV system. FIG. 1A shows that CRISPR induction killed 99.98% of the population in 30 minutes (black line). Growth in absence of induction is shown in dashed lines. CRISPR was induced at time-point 0 and monitored until 60 minutes. FIG. 2B shows dilution series  $(10^1 - 10^6)$  of drop spots  $(5~\mu l)$  on LB agar plates of *E. coli* ATCC43888 harboring the CGV system after 30 minutes of induction.

FIG. 2 shows CRISPR killing of target strain *Escherichia coli* (EHEC) ATCC43888 in *Galleria mellonella* larvae. *G. mellonella* larvae were delivered injections of bacteria behind the final left proleg. Approximately 1 h after the injection, CRISPR inducers were administered behind the final right proleg. Larvae were incubated at 37° C. for 2 h and sacrificed. Control bacteria carrying an off-target single guide RNA plasmid were also injected in the control group.

FIG. 2 shows CRISPR-kill curves of *Escherichia coli* (EHEC) ATCC43888 in *Galleria mellonella*. *G. mellonella* larvae were delivered injections of bacteria behind the final left proleg. Approximately 1 h after the injection, CRISPR inducers were administered behind the final right proleg. Larvae were incubated at 37° C. and sacrificed at 0, 1, and 2 h after induction.

FIG. 3 shows E. coli ATCC43888 count over time.

FIG. 4 shows Kaplan-Meier survival curves of *Galleria mellonella* larvae infected with *Escherichia coli* (EHEC) ATCC43888. CRISPR induction significantly improves survival of the larvae (black line) compared to the off-target control carrying an off-target single guide RNA plasmid (dashed line).

FIGS. **5**A-**5**B show time-kill curves for *Escherichia coli* Nissle 1917 harboring the CGV system targeting pks. FIG. **5**A shows that CRISPR induction killed 99.98% of the population in 15 minutes (black line). Growth in absence of induction is shown in dashed lines. CRISPR was induced at time-point 0 and monitored over 3 h. FIG. **5**B shows dilution

series  $(10^1-10^6)$  of drop spots  $(5 \mu l)$  on LB agar plates of E. coli Nissle 1917 harboring the CGV system after 15 minutes of induction.

FIGS. 6A-6B show time-kill curves for Escherichia coli Nissle 1917 harboring the CGV system targeting yapH. FIG. 5 6A shows that CRISPR induction killed 99.98% of the population in 15 minutes (black line). Growth in absence of induction is shown in dashed lines. CRISPR was induced at time-point 0 and monitored over 3 h. FIG. 6B shows dilution series ( $10^1$ - $10^6$ ) of drop spots (5  $\mu$ l) on LB agar plates of E. 10 coli Nissle 1917 harboring the CGV system after 15 minutes of induction.

FIG. 7 shows complete killing of transconjugant C. difficile. The complete precision killing of Clostridium difficile using a gRNA-encoding CRISPR array that was delivered 15 from a probiotic carrier bacterial species by conjugative plasmids as vectors is shown. A carrier bacterium (E. coli donor strain containing the vectors was mated with Clostridium difficile which was killed upon delivery of the designed array. This harnessed the endogenous Cas3 20 a microbial infection (eg, an ancute bacterial infection) of a machinery of Clostridium difficile. A 100% killing of Clostridium difficile cells was achieved and is shown in this

FIG. 8 shows the antibiotic treatment during ICI therapy has fatal outcomes. Kaplan Meier curve for overall survival 25 of a validation cohort from the Memorial Sloan Ketterin Cancer Center including n=239 advanced NSCLC patients treated with anti-PD-L1/anti-PD-1 mAb who received (ATB, n=68) or not (no ATB, n=171) antibiotics (ATB) two months before the injection of immune checkpoint blockade. 30 There was a medial overall survival of 21.9 months in the absence of antibiotic treatment, compared to an overall survival of 9.8 months with antibiotic treatment. So, the median overall survival in patients treated with classical antibiotics is <50% (or >12 months shorter) that of patients 35 not receiving antibiotic treatment.

FIGS. 9A and 9B show that the gut microbiome modulates the efficacy of anti-PD-1 inhibition in melanoma patients (from Gopalakrishnan et al, Science 2018, 359, 97-103).

#### DETAILED DESCRIPTION

The approach of the present invention is different from conventional antibiotic approaches. The present invention 45 utilizes targeted cutting of microbial genomes using programmed nucleases, whereas conventional antibiotics rely upon metabolic processes and cell replication cycles—and the inhibition of these—for their activity. By focusing instead on nuclease cutting, the invention surprisingly 50 achieves very quick and efficient microbial killing that also is remarkably durable. This is demonstrated in experiments below with different microbes, different nucleases and different delivery approaches. Typically, 99-100% killing was surprisingly observed many times and killing of 3-4 logs 55 was very quickly achieved and with lasting duration.

The invention provides methods for treating or preventing microbial (eg, bacterial) infections and means for performing these methods. In particular, treatment of infections requiring rapid therapy is made possible, such as for treating 60 acute conditions such as septicemia, sepsis, SIRS or septic shock. As explained herein, a rapid response is vital to address microbial infection in many settings. Speed is of the essence for many infection scenarios, such as acute infections requiring hospital admission. Benefits of the invention 65 can be one or more of: the reduction in the spread, severity or progression of the infection in the subject; reduction in the

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development, severity or progression of symptoms of the infection (eg, sepsis or septic shock); and an increase in the likelihood of survival in human or animal patients.

The invention uses programmable nuclease cutting of microbe genomes. The targeted cutting provides selective microbe killing or reduction of growth or proliferation to treat or prevent infection, as opposed to more broad-spectrum microbial killing of several different species as seen with conventional antibiotics. Selective killing is advantageous to leave beneficial microbes untargeted by the treatment, which may be beneficial to the patient. Moreover, the inventors have surprisingly found a substantial (by several logs) killing can be achieved very rapidly (eg, within 15 minutes) and sustainable effects can be achieved (eg, for more than 1 hour) in some embodiments. As exemplified below, the inventors surprisingly could remarkably achieve a fast and durable killing for around 2-3 hours.

Thus, the invention provides the following aspects:-

A programmable nuclease for use in a method of treating subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated.

Another aspect provides: A programmable nuclease for use in a method of rapidly treating an acute microbial (eg, bacterial) infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the subject is rapidly treated.

Another aspect provides: A programmable nuclease for use in a method of treating a microbial (eg, bacterial) infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and a nucleic acid that programs the nuclease to recognise and cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated.

Another aspect provides: A programmable nuclease for use in a method of rapidly treating an acute microbial (eg, bacterial) infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and a nucleic acid that programs the nuclease to recognise and cut the target site, whereby

genomes of the microbes comprised by the subject are cut and acute microbial infection of the subject is rapidly

Another aspect provides: A programmable nuclease for use in a method of durably treating a microbial (eg, bacte-5 rial) infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are durably 10 killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is 15

Another aspect provides: A programmable nuclease for use in a method of durably treating a microbial (eg, bacterial) infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the 20 nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are durably killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the 25 nuclease and a nucleic acid that programs the nuclease to recognise and cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated.

Another aspect provides: A programmable nuclease for 30 use in a method of durably treating an acute microbial (eg, bacterial) infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected 35 the subject, whereby microbes of the first species or strain are durably killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and a nucleic acid that programs the genomes of the microbes comprised by the subject are cut and acute microbial infection of the subject is treated.

Surprisingly, as exemplified below, a durable effect of several logs (eg, 3 or 4 logs) using a nuclease (as opposed to conventional means for conventional antibiotic killing) 45 was observed around 3 hours after the first exposure of bacteria with a programmed nuclease. This aspect of the invention, therefore, makes possible dosing regimens for less frequent exposure to a programmed nuclease (ie, less frequent administration of a programmed nuclease, pro- 50 grammable nuclease and/or nucleic acid for programming the nuclease). For example, a Cas and gRNA (or DNA encoding a gRNA) for programming the nuclease are administered with a programmable nuclease (eg, a Cas 9 or Cas3) to the subject at a first time (T1) and at a second time (T2); 55 or gRNA (or DNA encoding a gRNA) is administered on T1 and T2 for programming an endogenous Cas nuclease (eg, a Cas9 or Cas3) of bacteria of said first species or strain, wherein the programmed endogenous Cas cuts the genomes of the bacteria to kill the bacteria or to reduce growth or 60 proliferation, thus treating the infection. Such less frequent dosing is convenient for the healthcare practitioner and patient, as well as provides for economical therapy. Thus, optionally, the nuclease and/or nucleic acid is administered to the subject on T1 and T2, wherein T2 is at least 2, 3, 4, 65 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 24 hours after T1. For example, T2 is 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 24

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hours after T1. For example, T2 is 2-7 hours after T1. For example, T2 is 1 hour after T1. For example, T2 is 2 hours after T1. For example, T2 is 3 hours after T1. For example, T2 is 4 hours after T1. For example, T2 is 5 hours after T1.

Optionally, the nuclease (eg, programmed nuclease) and/ or a nucleic acid that programs the nuclease to recognise and cut the target site is administered to the subject on T1 and T2, wherein T2 is at least 1 hour (eg, 1, 1.5, 2, 2.5 or 3 hours) after T1.

Another aspect provides: A Cas nuclease for use in a method of treating a microbial (eg, bacterial) infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable with a guide RNA (gRNA) to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising administering to the subject said a nucleic acid, wherein the nucleic acid is the gRNA or a DNA encoding the gRNA, thereby programming the nuclease to recognise and cut the target site of the microbes comprised by the subject, whereby genomes of the microbes are cut and microbial infection of the subject is treated, wherein the method comprises administering the nucleic acid to the subject on at a first time (T1) and at a second time (T2), whereby the subject is exposed to programmed nuclease on T1 and T2, and wherein T2 is no less than 1 hour after T1.

Optionally, T2 is no less than 2 hours after T1; optionally, T2 is no less than 3 hours after T1; optionally, T2 is no less than 4 hours after T1; optionally, T2 is no less than 5 hours after T1; optionally, T2 is no less than 6 hours after T1; optionally, T2 is no less than 7 hours after T1; optionally, T2 is no less than 8 hours after T1; optionally, T2 is no less than 9 hours after T1; optionally, T2 is no less than 10 hours after T1; optionally, T2 is no less than 11 hours after T1; optionally, T2 is no less than 12 hours after T1; optionally, T2 is no less than 13 hours after T1; optionally, T2 is no less than 14 hours after T1; or optionally, T2 is no less than 24 hours nuclease to recognise and cut the target site, whereby 40 after T1. Additionally or alternatively: Optionally, T2 is no more than 7 hours after T1; optionally, T2 is no more than 12 hours after T1; optionally, T2 is no more than 24 hours after T1; optionally, T2 is 2-7 hours after T1; optionally, T2 is 24 hours after T1; optionally, T2 is 7 hours after T1; optionally, T2 is 6 hours after T1; optionally, T2 is 5 hours after T1; optionally, T2 is 4 hours after T1; optionally, T2 is 3 hours after T1; optionally, T2 is 2 hours after T1; optionally, T2 is 1 hour after T1. For example, T2 is 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 or 24 hours after T1. For example, T2 is 1-7 hours after T1; or T2 is 2-7 hours after T1; or T2 is 3-7 hours after T1; or T2 is 4-7 hours after T1; or T2 is 5-7 hours after T1; or T2 is 6-7 hours after T1.

Optionally, the method comprises reducing the infection at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment. Optionally, the method comprises reducing the infection at least 1000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment. Optionally, the method comprises reducing the infection at least 10000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.

Optionally, the method comprises reducing the infection such that the reduction in infection persists for 30 minutes immediately after the first 30 minutes of the treatment. Optionally, the method comprises reducing the infection such that a reduction in infection by at least 100-fold persists for 30 minutes immediately after the first 30 minutes of the treatment. Optionally, the method comprises reducing the

infection such that a reduction in infection by at least 1000-fold persists for 30 minutes immediately after the first 30 minutes of the treatment. Optionally, the method comprises reducing the infection such that a reduction in infection by at least 10000-fold persists for 30 minutes immediately after the first 30 minutes of the treatment.

Optionally, the method comprises reducing the infection at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein a reduction in infection by at least 100-fold persists for 30 minutes immediately after the first 30 minutes of the treatment. Optionally, the method comprises reducing the infection at least 1000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein a reduction in infection by at least 1000-fold persists for 30 minutes immediately after the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein at least 10000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein a reduction in infection by at least 10000-fold persists for 30 minutes immediately after the first 30 minutes of the treatment.

Optionally, the method comprises maintaining reduction of the infection by at least 100-fold for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the 25 subject to the programmed nuclease. Optionally, a reduction of the infection by at least 100-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease. Optionally, the method comprises maintaining reduction of the 30 infection by at least 1000-fold for at least 60 minutes (eg., at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease. Optionally, a reduction of the infection by at least 1000-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing 35 the subject to the programmed nuclease. Optionally, the method comprises maintaining reduction of the infection by at least 10000-fold for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease. Optionally, a reduction of the infec- 40 tion by at least 10000-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease.

Optionally, the method comprises reducing the infection at least 100-fold by the first 30 minutes (eg, by the first 15 45 minutes) of the treatment; and wherein reduction of the infection by at least 100-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease. Optionally, the method comprises reducing the infection at least 1000-fold 50 by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein reduction of the infection by at least 1000-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease. Optionally, the method comprises 55 reducing the infection at least 10000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein reduction of the infection by at least 10000-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed 60

Optionally, the method comprises reducing the infection at least 10000-fold by the first 15 minutes of the treatment; and wherein reduction of the infection by at least 10000-fold is maintained for at least 45 minutes after exposing the 65 subject to the programmed nuclease. This is exemplified below.

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In an example, the infection is durably treated, wherein a reduction of the infection by at least 100-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after commencement of the treatment. In an example, the infection is durably treated, wherein a reduction of the infection by at least 1000-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after commencement of the treatment. In an example, the infection is durably treated, wherein a reduction of the infection by at least 10000-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after commencement of the treatment.

Optionally, the infection is reduced at least 100,000-fold by the first 30 or 45 minutes of the treatment. Optionally, the infection is reduced at least 100,000-fold by the first 30 or 45 minutes of the treatment and the reduction is maintained until the  $60^{th}$  minute of the treatment.

Optionally, the infection is reduced at least 1000,000-fold by the first 30 or 45 minutes of the treatment. Optionally, the infection is reduced at least 1000,000-fold by the first 30 or 45 minutes of the treatment and the reduction is maintained until the  $60^{th}$  minute of the treatment.

Optionally, the infection is reduced at least 100-fold by the first 15 minutes of the treatment. Optionally, the infection is reduced at least 1000-fold by the first 15 minutes of the treatment. Optionally, the infection is reduced at least 100-fold by the first 15 minutes of the treatment and at least 1000-fold by the first 30 minutes of the treatment.

For example, the reduction is maintained for at least 15 further minutes, eg, the infection is reduced at least 100-fold or at least 1000-fold by the first 15 minutes of the treatment and the reduction is maintained from the 15-30<sup>th</sup> minute or 15-45<sup>th</sup> minute of the treatment

For example, the infection is reduced at least 100-fold or at least 1000-fold or at least 10000-fold by the first 15 minutes of the treatment in the first 15 minutes and the reduction is maintained for from the 15-30<sup>th</sup> minute or 15-45<sup>th</sup> minute of the treatment.

Optionally, the method comprises reducing the infection at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.

Optionally the method comprises reducing the infection at least 1000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.

Optionally the method comprises reducing the infection at least 10000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.

Optionally, the method comprises reducing the infection such that the reduction in infection persists for 30 minutes immediately after the first 30 minutes of the treatment, eg, the reduction may persist for at least 60 minutes after the first 30 minutes of the treatment. If the treatment is administered at time zero (T0), then the reduction in infection may be present at 60 minutes counted after T0, and indeed may persist after that 60 minutes. In FIGS. 1A, 5A and 6A, for example, reduction is seen at 60-180 minutes after T0. Optionally, the reduction in infection persists for at least 30 minutes after the first 30 minutes of the treatment.

In an example, the infection is reduced by at least 10, 20, 30, 40, 50, 60, 70, 80, 90, 91, 92, 93, 94, 95, 96, 97, 98 or 99%, eg, in the first 15 minutes of treatment. In an example, the infection is reduced by at least 10, 20, 30, 40, 50, 60, 70, 80, 90, 91, 92, 93, 94, 95, 96, 97, 98 or 99%, eg, in the first 30 minutes of treatment.

For determining killing or reduction in growth or proliferation of the target microbes, one can, for example, deter-

mine the difference in the number of microbes of the first species or strain in (i) a sample taken from the subject (eg, a blood, gut or leaf sample) immediately before commencement of the treatment and (ii) a sample (of the same type as the sample of (i), eg, a blood, gut or leaf sample respec- 5 tively) taken from the subject at 30 minutes of the treatment. For example, if the microbes are bacteria, the samples may be assessed for the difference in colony forming units (CFU)/ml sample, eg, when the samples have been plated on agar in respective petri dishes and incubated under identical conditions. Another example may use microscopic counting of microbes in samples, or other routine methods know to the skilled addressee.

In an example, at least 60, 70, 80, 90, 91, 92, 93, 94, 95, 96, 97, 98 or 99% killing of the microbes is achieved by the 15 first 30, 60, 90 or 120 minutes (eg, by the first 30 minutes; or by the first 120 minutes) of the treatment. For example, wherein the subject is a human or animal, the killing is determined comparing the prevalence (eg, by standard colony counting on an agar plate) of the microbes (eg. 20 bacteria) in a blood sample taken immediately before commencement of the treatment versus a sample taken after the first 15 or 30 minutes of the treatment. In an example, at least 80, 90, 91, 92, 93, 94, 95, 96, 97, 98 or 99% killing of the microbes is achieved by the first 0.5, 1 or 2 hours of the 25 treatment. In an example, at least 99% killing of the microbes is achieved by the first 30 minutes of the treatment. In an example, at least 99% killing of the microbes is achieved by the first 2 hours of the treatment. In an example, 100% killing is achieved. These are exemplified below. In an 30 embodiment, less than 100% of the microbes are killed.

Worked examples of killing in bacteria are shown below. Surprisingly, using a programmed nuclease to target the bacteria of choice, specific cutting resulted in rapid killing of or 10,000 fold killing) could be observed in very short spaces of time and surprisingly these were sustained for at least to an hour. Optionally, the infection is reduced at least 1000-fold by the first 15, 30 or 45 minutes of the treatment. Optionally, the infection is reduced at least 1000-fold by the 40 first 15, 30 or 45 minutes of the treatment and the reduction is maintained until the 60th, 120th or 180th minute of the treatment. Optionally, the infection is reduced at least 10,000-fold by the first 15, 30 or 45 minutes of the treatment. Optionally, the infection is reduced at least 10,000-fold by 45 the first 15, 30 or 45 minutes of the treatment and the reduction is maintained until the 60<sup>th</sup>, 120<sup>th</sup> or 180<sup>th</sup> minute of the treatment. See, for example, exemplification in FIG. 5A.

In an example, 100% killing is achieved by 24 hours after 50 commencement of the treatment.

In an example, the infection is reduced at least 1000-fold for 2 hours or more (eg, for 2-3 hours). Optionally also the infection is reduced by at least 1000-fold by the first 15 or 13 minutes of the treatment.

In an example, the infection is reduced at least 10,000fold for 2 hours or more (eg, for 2-3 hours). Optionally also the infection is reduced by at least 10,000-fold by the first 15 or 13 minutes of the treatment.

In an example, the infection is reduced by at least 90, 91, 60 92, 93, 94, 95, 96, 97, 98 or 99% for 1 hour; or for 1 hour or more; or for 2 hours or more (eg, for 2-3 hours). Optionally, the infection is reduced by at least 90% for 1 hour; or for 1 hour or more; or for 2 hours or more (eg, for 2-3 hours), and optionally by the first 30 minutes (eg, by the 65 first 15 minutes) of the treatment. Optionally, the infection is reduced by at least 90% for 1 hour or more, and by the first

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30 minutes (eg, by the first 15 minutes) of the treatment. Optionally, the infection is reduced by least 90, 91, 92, 93, 94, 95, 96, 97, 98 or 99% by the first 15 or 13 minutes of the treatment. Optionally, the infection is reduced by least 90, 91, 92, 93, 94, 95, 96, 97, 98 or 99% by the first 15 or 13 minutes of the treatment; and wherein the reduction is maintained for 1 hour or more (eg, for 2 hours or more; or for 3 hours or more; or for about 2 hours; or for 2 hours; or for about 3 hours; or for 3 hours). Exemplification below is provided, wherein the bacteria are E coli.

Optionally, the subject is a human or animal and the microbes are bacteria (eg, E coli or C dificile), wherein blood infection of the subject by the bacteria is reduced at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment. Optionally, the subject is a human or animal and the microbes are bacteria (eg, E coli or C dificile), wherein blood infection of the subject by the bacteria is reduced at least 1000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment. Optionally, the subject is a human or animal and the microbes are bacteria (eg. E coli or C dificile), wherein blood infection of the subject by the bacteria is reduced at least 10,00-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment. Optionally, the E coli are EHEC E coli.

Optionally, the programmed nuclease (eg., a Cas9 or Cas3) is capable of cutting a target site comprised by E. coli (EHEC) ATCC43888. Optionally, the programmed nuclease (eg, a Cas9 or Cas3) is capable of cutting a target site comprised by E. coli Nissle.

Optionally, the blood of the subject is infected with from  $10^7$  to  $10^{12}$  CFU/ml (eg, from  $10^{17}$  to  $10^{11}$ , from  $10^7$  to  $10^{10}$ , from 10<sup>7</sup> to 10<sup>9</sup> or from 10<sup>7</sup> to 10<sup>8</sup> CFU/ml) of the bacteria immediately before the treatment.

The worked example below shows improved survival the target bacteria—at least 3 or 4 logs of killing (ie, 1000- 35 using the method of the invention in an in vivo model. In an example, therefore, the method of the invention is for improving survival of the subject by treating acute microbial infection of a subject. In an example, the programmed nuclease herein is capable of carrying out the killing of bacteria of the first species or strain in Galleria mellonella larvae in vivo infection model.

> The nuclease may be, for example, a DNase (eg, a Cpf1, Cas9 or Cas3) or a RNase (eg, Cas13b).

> In an example, the nuclease is an isolated or recombinant nuclease. For example, the nuclease is a synthetic or nonnaturally occurring nuclease.

> In an example, the nuclease is ex vivo, eg, in vitro. In an example, the nucleic acid is ex vivo. In an example, the guide RNA or DNA encoding guide RNA(s) herein is ex vivo, eg, in vitro.

Optionally, the nuclease is a Cas nuclease (eg, a Cpf1, CasX, CasY, Cas13b, Cas 3 or 9), a meganuclease, a TALEN (Transcription activator-like effector nuclease) or zinc finger nuclease. In an example, the Cas is a *Streptococcus* (eg, 55 pyogenes or aureus) Cas9, Clostridium (eg, dificile), Salmonella (eg, typhimurium) or E coli Cas3. For example, the Cas is a spCas. In an example, the Cas9 is in combination with a tracrRNA or a DNA encoding a tracrRNA which is operable with the Cas. For example, the tracrRNA is of the same species as the Cas, eg, a Spyogenes tracrRNA or DNA encoding this.

In an example, the nuclease is a Cas 3 encoded by a nucleic acid comprising SEQ ID NO: 9 or a sequence that is at least 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. Optionally, also the bacteria are Clostridium (eg, C dificile) bacteria; or any Clostridium shown in Table 1. This is exemplified below.

In an example, the nuclease is a Cas 9 encoded by a nucleic acid comprising SEQ ID NO: 10 or a sequence that is at least 80, 85, 90, 95, 96, 97, 98 or 99% identical thereto. In an example, the nuclease is a Cas 9 comprising SEQ ID NO: 11 or a sequence that is at least 80, 85, 90, 95, 96, 97, 598 or 99% identical thereto. Optionally, also the bacteria are *Clostridium* (eg, *C dificile*) bacteria; or any *Clostridium* shown in Table 1. Optionally, also the bacteria are *E coli* (eg, EHEC). This is exemplified below.

Optionally, the method comprises administering to the 10 subject a RNA or a nucleic acid (eg, DNA) that encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject.

Optionally, the nuclease is administered simultaneously 15 or sequentially with the RNA or nucleic acid to the subject.

Optionally, subject comprises the nuclease prior to administration of the RNA or nucleic acid to the subject. For example, the nuclease is a Cas nuclease that is an endogenous Cas nuclease of bacterial cells of the first species or 20 strain that are comprised by the subject. Thus, in this example, the RNA or nucleic acid may be administered to the subject and introduced into the bacteria for programming endogenous Cas comprised by the bacteria, thereby forming programmed Cas nuclease that cuts the target site in the 25 genomes of the bacteria, whereby bacteria are killed or growth or proliferation of bacteria is reduced, thus treating or preventing the infection.

Optionally, a plurality of viruses (eg, phage or phagemids) are administered to the subject, wherein each virus comprises a copy (eg, one or more, eg, a plurality of copies) of the nucleic acid, wherein the viruses infect the microbes comprised by the subject to deliver thereto the nucleic acid. For example, viruses herein are phage or phagemids that infect (or are capable of infecting) the bacteria of the first 35 species or strain.

Optionally, the ratio of administered viruses:microbes comprised by the subject is from 10 to 150. For example, the microbes are bacteria and the ratio is from 10 to 100, ie, a multiplicity of infection (MOI) of from 1 to 100 (eg, wherein 40 the viruses are capable of replication, eg, are phage and not phagemid), eg, from 10 to 100. The ratio can be determined, for example, using a sample (eg, a blood or gut sample) from a human or animal subject immediately before the treatment and determining the number of microbes (eg, bacteria per ml 45 of blood or gut sample). The amount of viruses to be administered can then be worked out according to the determination using the sample.

Optionally, the microbes are bacteria. Alternatively, the microbes are archaea. Alternatively, the microbes are 50 viruses. Alternatively, the microbes are fungi. Alternatively, the microbes are algae. Alternatively, the microbes are protozoa.

In an example, the subject is a human and the infection is a nosocomial infection. In an example, the subject is a plant, 55 yeast, protist or amoeba.

Optionally, the subject is a human (eg, an adult, child, neonate, toddler, teenager, male or female) or animal (eg, a dog, cat, horse, cow, sheep, goat, salmon, chicken, turkey, pig, companion animal or livestock animal).

In an example, the subject is a human or animal and: Optionally, the infection is an infection of the lungs, abdomen or urinary tract. In an example, the subject is suffering from a urinary tract infection, lung infections, such as pneumonia, a kidney infection or an abdominal infection. In 65 an example, the subject is a surgery patient. In an example, the subject has

an infected wound (eg, a bacterially infected wound). In an example, the patient is suffering from AIDS or is infected by HIV. In an example, the subject is suffering from a cancer, such as a blood cancer, eg, leukaemia, eg, AML or CML or CLL or a lymphoma. In an example, the subject is a tissue or organ transplant patient, eg, a haematopoietic stem cell transplant or bone marrow transplant patient. In an example, the subject has a urinary or intravenous catheter. In an example, the subject is on mechanical ventilation. In an example, the subject has been receiving an immunosupressant. In an example, the subject is suffering from pneumonia. In an example, the subject is an intensive care unit (ICU) patient. In an example, the subject is an Acute respiratory distress syndrome (ARDS) patient. In an example, the subject is suffering from meningitis, an infection in pregnancy, a ruptured gallbladder (a gallbladder rupture is a medical condition where the gallbladder leaks or bursts. Ruptures are commonly caused by inflammation of the gallbladder), abortion with septic shock (abortion with septic shock can be an acute life-threatening illness), endometritis (endometritis is an inflammatory condition of the lining of the uterus, usually due to an infection), Acute Respiratory Distress Syndrome (Acute respiratory distress syndrome is a lung condition; it occurs when fluid fills up the air sacs in the lungs) or cellulitis.

The increase in average age of the population, more people with chronic diseases, on immunosuppressive drugs, and increase in the number of invasive procedures being performed has led to an increased rate of sepsis. Optionally, the subject has undergone surgery, is on an immunosuppressant medication and/or is suffering from a chronic disease.

Optionally, the subject is a human over 60, 65, 70, 75 or 80 years of age or is a paediatric patient. In an alternative, the subject is a paediatric patient (eg, a human baby or child) or adolescent. In an example, the method treats or prevents neonatal sepsis in the subject. In an example the subject is an immune-compromised human or animal, eg, suffering from an acute viral infection, such as HIV infection; or the subject is suffering from a cancer, eg, a blood cancer, such as a leukaemia; or the patient is a transplant patient, eg, that has received an organ, tissue or bone marrow transplant. In an example, the subject is a human or animal that is positive for gram negative bacterial lipopolysaccharide or lipid A. In an example, the subject is a human or animal that is positive for gram positive bacterial cell wall lipoteichoic acid.

Optionally, the method treats or prevents septicaemia and/or sepsis (eg, septic shock) in the subject.

SIRS (Systemic Inflammatory Response Syndrome) criteria has been used to define sepsis.

SIRS is the presence of two or more of the following: abnormal body temperature, heart rate, respiratory rate, or blood gas, and white blood cell count. Sepsis is, for example, SIRS in response to an infectious process. Severe sepsis is, for example, sepsis with sepsis-induced organ dysfunction or tissue hypoperfusion (manifesting as hypotension, elevated lactate, or decreased urine output). Septic shock is, for example, severe sepsis plus persistently low blood pressure, despite the administration of intravenous fluids.

In an embodiment, the method prevents or delays progression of end-organ dysfunction in the subject (when the subject is a human or animal).

Examples of end-organ dysfunction include the following:

Lungs: acute respiratory distress syndrome (ARDS) (PaO<sub>2</sub>/FiO<sub>2</sub><300)

Brain: encephalopathy symptoms including agitation, confusion, coma; causes may include ischemia, bleed-

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ing, formation of blood clots in small blood vessels, microabscesses, multifocal necrotizing leukoencephalopathy

Liver: disruption of protein synthetic function manifests acutely as progressive disruption of blood clotting due to an inability to synthesize clotting factors and disruption of metabolic functions leads to impaired bilirubin metabolism, resulting in elevated unconjugated serum bilirubin levels

Kidney: low urine output or no urine output, electrolyte abnormalities, or volume overload

Heart: systolic and diastolic heart failure, likely due to chemical signals that depress myocyte function, cellular damage, manifest as a troponin leak (although not necessarily ischemic in nature)

More specific definitions of end-organ dysfunction exist for SIRS in pediatrics.

Cardiovascular dysfunction (after fluid resuscitation with at least 40 ml/kg of crystalloid)

hypotension with blood pressure <5th percentile for age or systolic blood pressure <2 standard deviations below normal for age, or

vasopressor requirement, or

two of the following criteria:

unexplained metabolic acidosis with base deficit >5 mEq/l

lactic acidosis: serum lactate 2 times the upper limit of normal

oliguria (urine output<0.5 ml/kg/h)

prolonged capillary refill >5 seconds

core to peripheral temperature difference >3° C.

Respiratory dysfunction (in the absence of cyanotic heart disease or known chronic lung disease)

the ratio of the arterial partial-pressure of oxygen to the 35 from fraction of oxygen in the gases inspired (PaO<sub>2</sub>/FiO<sub>2</sub>) Sh <300 (the definition of acute lung injury), or

arterial partial-pressure of carbon dioxide (PaCO<sub>2</sub>)>65 torr (20 mmHg) over baseline PaCO<sub>2</sub> (evidence of hypercapnic respiratory failure), or

supplemental oxygen requirement of greater than FiO<sub>2</sub> 0.5 to maintain oxygen saturation≥92%

Neurologic dysfunction

Glasgow Coma Score (GCS)≤11, or

altered mental status with drop in GCS of 3 or more points 45 in a person with developmental delay/intellectual disability

Hematologic dysfunction

platelet count<80,000/mm³ or 50% drop from maximum in chronically thrombocytopenic, or

international normalized ratio (INR) >2

Disseminated intravascular coagulation

Kidney dysfunction

serum creatinine≥2 times the upper limit of normal for age or 2-fold increase in baseline creatinine in people with 55 chronic kidney disease

Liver dysfunction (only applicable to infants >1 month) total serum bilirubin ≥4 mg/dl, or

alanine aminotransferase (ALT)≥2 times the upper limit of normal

Table 2 sets out the criteria for a positive diagnosis of sepsis.

Optionally, the method reduces one or more symptoms in the patient selected from fever, low body temperature, rapid breathing, elevated heart rate, confusion, confusion, metabolic acidosis, respiratory alkalosis, low blood pressure, dysfunction of blood coagulation (such as blood clotting in 20

one or more organs, or bruising) and oedema. Optionally, the method reduces septic shock. Optionally, the sepsis is severe sepsis.

Optionally, at the start of the treatment, the subject (eg, a human) has a temperature of  $<36^{\circ}$  C. or  $>38^{\circ}$  C.; a heart rate of >90/min, a respiratory rate of >20 breaths/min or  $PaCO_2<4.3$  kPa; and white blood cell count of <4000/mm³ or >12,000/mm³.

Optionally, at the start of the treatment, the subject (eg, a human) has presence of two or more of the following: abnormal body temperature, abnormal heart rate, abnormal respiratory rate, abnormal blood gas and abnormal white blood cell count.

Optionally, the subject is a plant. In an example, the subject is a protist, eg, amoeba. Optionally in this example, the microbes are viruses (eg, large or gian viruses, eg, Mimiviruses). The nuclease, for example, is a Cas and is programmable using a guide RNA delivered by a virophage that infects the virus microbes

In an example the microbes are yeast, eg, Candida.

Preferably, the microbes are bacteria. Optionally, the bacteria are gram positive bacteria. Optionally, the bacteria are Staphylococcus, Streptococcus, Enterococcus, Legionella, Heamophilus, Ghonnorhea, Acinetobacter, Escherizohia, Klebsiella, Pseudomonas or Stenotrophomonas bacteria (eg, E coli (eg, EHEC E coli), C dificile, V cholera, Staphylococcus (eg, S aureus or MRSA), Streptococcus pyogenes, Acinetobacter baumannii, Legionella, Pseudomonas aeruginosa, Klebsiella pneumoniae bacteria).

Optionally, the first species is selected from the species in Table 1.

Optionally, the first species is enterohemorrhagic *E. coli* (EHEC), *E. coli* Serotype O157:H7 or Shiga-toxin producing *E. coli* (STEC)). In an example, the bacteria are selected from

Shiga toxin-producing *E. coli* (STEC) (STEC may also be referred to as Verocytotoxin-producing *E. coli* (VTEC); Enterohemorrhagic *E. coli* (EHEC) (this pathotype is the one most commonly heard about in the news in association with foodborne outbreaks);

Enterotoxigenic E. coli (ETEC);

Enteropathogenic E. coli (EPEC);

Enteroaggregative E. coli (EAEC);

Enteroinvasive E. coli (EIEC); and

Diffusely adherent E. coli (DAEC).

Enterohemorrhagic Escherichia coli (EHEC) serotype O157:H7 is a human pathogen responsible for outbreaks of bloody diarrhoea and haemolytic uremic syndrome (HUS) worldwide. Conventional antimicrobials trigger an SOS 50 response in EHEC that promotes the release of the potent Shiga toxin that is responsible for much of the morbidity and mortality associated with EHEC infection. Cattle are a natural reservoir of EHEC, and approximately 75% of EHEC outbreaks are linked to the consumption of contaminated bovine-derived products. EHEC causes disease in humans but is asymptomatic in adult ruminants. Characteristics of E. coli serotype O157:H7 (EHEC) infection includes abdominal cramps and bloody diarrhoea, as well as the life-threatening complication haemolytic uremic syndrome (HUS). Currently there is a need for a treatment for EHEC infections (Goldwater and Bettelheim, 2012). The use of conventional antibiotics exacerbates Shiga toxinmediated cytotoxicity. In an epidemiology study conducted by the Centers for Disease Control and Prevention, patients treated with antibiotics for EHEC enteritis had a higher risk of developing HUS (Slutsker et al., 1998). Additional studies support the contraindication of antibiotics in EHEC

infection; children on antibiotic therapy for hemorrhagic colitis associated with EHEC had an increased chance of developing HUS (Wong et al., 2000; Zimmerhackl, 2000; Safdar et al., 2002; Tarr et al., 2005). Conventional antibiotics promote Shiga toxin production by enhancing the 5 replication and expression of six genes that are encoded within a chromosomally integrated lambdoid prophage genome. The approach of the present invention relies on nuclease cutting. Stx induction also promotes phage-mediated lysis of the EHEC cell envelope, allowing for the release and dissemination of Shiga toxin into the environment (Karch et al., 1999; Matsushiro et al., 1999; Wagner et al., 2002). Thus, advantageously, the invention provides alternative means for treating EHEC in human and animal subjects. This is exemplified below with surprising results 15 on the speed and duration of anti-EHEC action produced by nuclease action (as opposed to conventional antibiotic action).

In an example, the subject (eg, a human) is suffering from or at risk of haemolytic uremic syndrome (HUS), eg, the  $^{20}$  subject is suffering from an  $E\ coli$  infection, such as an EHEC  $E\ coli$  infection.

An aspect of the invention provides: A plurality of viruses (eg, phage or phagemids for producing phage) for use with the nuclease of the invention in the method of treatment, 25 wherein each virus comprises a copy of a nucleic acid described herein, wherein the viruses are capable of infecting microbes comprised by the subject to deliver thereto the nucleic acid.

A aspect of the invention provides: A plurality of viruses 30 (eg, phage or phagemids for producing phage) for use with a programmable nuclease in a method of treating a microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by 35 the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the 40 target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated;

wherein each virus comprises a copy of a nucleic acid that encodes an RNA for expression of the RNA in the subject, 45 wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject;

wherein the viruses are capable of infecting microbes comprised by the subject to deliver thereto the nucleic acid.

Optionally, the method is for durable treatment, eg, as described herein; and/or optionally, the infection is acute infection.

Optionally, the method is for rapid treatment, eg, as described herein; and/or optionally, the infection is acute 55 infection.

Optionally, the nuclease is according to any nuclease of the invention herein. Optionally, the nucleic acid is according to any nucleic acid of the invention herein.

Optionally, the nuclease is according to any nuclease of 60 the invention herein. Optionally, the nucleic acid is according to any nucleic acid of the invention herein.

In an alternative, when the microbes are viruses, the plurality of viruses are phage that are capable of infecting host cells harbouring the microbes, wherein the nucleic 65 acids are introduced into the host cells for expression therein of the RNA. The RNA complexes with the nuclease in the

host cells to guide the nuclease to cut a target site of the microbes (ie, to cut viral RNA or DNA), thereby inactivating the viral microbes. For example, the microbes are viruses (eg, in amoeba; or in human or animal or plant cells) and the viruses of said plurality of viruses are capable of targeting the microbes, whereby the nuclease is programmed to cut the microbes (eg, in the amoeba or in said cells).

An aspect of the invention provides: A composition comprising a plurality of nucleic acids for programming the nuclease of the invention in the method of treatment, wherein each nucleic acid is a nucleic acid as defined herein.

An aspect of the invention provides: A composition comprising a plurality of nucleic acids for programming a programmable nuclease in a method of treating a microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and the nucleic acids wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is treated; wherein each nucleic acid encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject.

Optionally, the method is for durable treatment, eg, as described herein; and/or optionally, the infection is acute infection.

Optionally, the method is for rapid treatment, eg, as described herein; and/or optionally, the infection is acute infection.

Optionally, the nuclease is according to any nuclease of the invention herein. Optionally, each nucleic acid is according to any nucleic acid of the invention herein.

Optionally, the composition is a pharmaceutical composition comprising the nucleic acids and a pharmaceutically acceptable diluent, carrier or excipient. Optionally, the composition is for oral, intravenous, pulmonary, rectal, topical, buccal, ocular, intranasal, or subcutaneous administration to a human or animal subject. Optionally, the composition is a herbicide or pesticide or insecticide or nematodicide or aracnicide. Optionally, the composition is toxic to yeast. Optionally, the composition is toxic to giant viruses.

An aspect of the invention provides: A CRISPR/Cas system comprising a nuclease according to the invention for use in the method of treatment, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9 or any other Cas mentioned herein) and the system comprises one or more guide RNAs or DNA encoding one or more guide RNAs, wherein each guide RNA is capable of programming the Cas nuclease to cut a target site comprised by the genomes of the microbes.

In an example, each guide RNA mentioned herein is a single guide RNA (ie, a chimaeric guide RNA). In another example, each guide RNA comprises a crRNA that is hybridised to a tracrRNA.

In an example, a target site mentioned herein is comprised by an essential gene, virulence gene or antibiotic resistance gene of the bacteria. In an example, a target site mentioned herein is comprised by a multi-copy sequence (ie, a sequence that is present in more than one (eg, 2, 3, 4, 5, 6, 7, 8 or 9, or more) copies in each bacterial genome). For example, the target site is comprised by a ribosomal RNA gene. In an example, a target site mentioned herein is

comprised by a ribosomal RNA gene (eg, a 23S ribosomal RNA gene), a yapH gene; or a pks gene; or homologue or orthologue thereof.

Optionally, each guide RNA herein is capable of hybridizing to a protospacer sequence comprising the target site, 5 wherein the protospacer sequence is 15-45 nucleotides in length, eg, 15-25; 18-21; 20; or about 20 nucleotides in length. Optionally, each guide RNA herein comprises a spacer sequence that is 15-45 nucleotides in length, eg, 15-25; 18-21; 20; or about 20 nucleotides in length.

Optionally, each guide RNA herein is cognate to a 5'-NGG protospacer adjacent motif (PAM), eg, wherein the bacteria are *E coli*. Optionally, each guide RNA herein is cognate to a 5'-CCA or 5'-CCT protospacer adjacent motif (PAM), eg, wherein the bacteria are *C dificile*.

An aspect of the invention provides: A guide RNA or a DNA encoding a guide RNA for use in the system of the invention for use in the method of treating an acute microbial infection in the subject, eg, septicaemia or sepsis.

An aspect of the invention provides: A nucleic acid vector 20 comprising the guide RNA or DNA.

Optionally, the vector is a phage, phagemid, viriophage, virus, plasmid (eg, conjugative plasmid) or transposon. The example below shows almost complete killing can be achieved using a conjugative plasmid as the vector. Thus, in 25 an embodiment, each vector is a conjugative plasmid that is delivered from carrier bacteria eg, probiotic carrier bacteria for administration to the human or animal subject. In an example, the carrier bacteria are *Lactobacillus* (eg, *L reuteri*) or *E coli*. This is exemplified below and achieved complete 30 (100%) killing.

An aspect of the invention provides: An anti-sepsis or anti-septicaemia composition for administration to a human or animal for treating sepsis or septicaemia, the composition comprising a plurality of vectors, wherein each vector a 35 vector of the invention.

An aspect of the invention provides: A method of treating (eg, rapidly and/or durably treating) an acute microbial infection of a subject, wherein the method is as defined herein

An aspect of the invention provides: A method of treating (eg, rapidly and/or durably treating) an acute microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by 45 the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the 50 target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the subject is treated (eg, rapidly and/or durably treated).

An aspect of the invention provides: A method of treating (eg, rapidly and/or durably treating) an acute microbial 55 infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and a plurality of viruses wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the subject is treated (eg, rapidly and/or durably treated); wherein each virus comprises a copy of a

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nucleic acid that encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject; wherein the viruses are capable of infecting microbes comprised by the subject to deliver thereto the nucleic acid.

Optionally, the nuclease is according to any nuclease of the invention herein. Optionally, the nucleic acid is according to any nucleic acid of the invention herein.

An aspect of the invention provides: A method of treating (eg, rapidly and/or durably treating) an acute microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease and a plurality of nucleic acids wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the subject is treated (eg, rapidly and/or durably treated); wherein each virus comprises a copy of a nucleic acid that encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject; wherein each nucleic acid encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject.

Optionally, the nuclease is according to any nuclease of the invention herein. Optionally, each nucleic acid is according to any nucleic acid of the invention herein.

In an example, the invention is for medical or dental or opthalmic use (eg, for treating or preventing an infection in an organism or limiting spread of the infection in an organism)

In an example, the invention is for cosmetic use (eg, use in a cosmetic product, eg, make-up), or for hygiene use (eg, use in a hygiene product, eg, soap).

In an example, the vectors and/or nuclease prior to administration to the subject are comprised by a composition which is as any of the following (host here refers to the microbes of the first species or strain): In an example, the composition is a medical, opthalmic, dental or pharmaceutical composition (eg, comprised by an anti-host vaccine). In an example, the composition is an antimicrobial composition, eg, an antibiotic or antiviral, eg, a medicine, disinfectant or mouthwash. In an example, the composition is a cosmetic composition (eg, face or body make-up composition). In an example, the composition is a herbicide. In an example, the composition is a pesticide (eg, when the host is a Bacillus (eg, thuringiensis) host). In an example, the composition is a beverage (eg, beer, wine or alcoholic beverage) additive. In an example, the composition is a food additive (eg, where the host is an E coli, Salmonella, Listeria or Clostridium (eg, botulinum) host). In an example, the composition is a water additive. In an example, the composition is a additive for acquatic animal environments (eg, in a fish tank). In an example, the composition is an oil or petrochemical industry composition or comprised in such a composition (eg, when the host is a sulphate-reducing bacterium, eg, a Desulfovibrio host). In an example, the composition is a oil or petrochemical additive. In an example, the composition is a chemical additive. In an example, the composition is a disinfectant (eg, for sterilizing equipment

26 use. For example, it is for or used in agriculture, oil or petroleum industry, food or drink industry, clothing industry, packaging industry, electronics industry, computer industry, environmental industry, chemical industry, aeorspace industry, automotive industry, biotechnology industry, medical industry, healthcare industry, dentistry industry, energy industry, consumer products industry, pharmaceutical industry, mining industry, cleaning industry, forestry industry, fishing industry, leisure industry, recycling industry, cosmetics industry, plastics industry, pulp or paper industry, textile industry, clothing industry, leather or suede or animal hide industry, tobacco industry or steel industry. Host cells herein refers to the microbes of the first species or strain. Optionally, any host cell(s) herein is/are bacterial or archaeal cells. In an example, the cell(s) is/are in stationary phase. In an example, the cell(s) is/are in exponential phase. In an example, the cell(s) is/are in lag phase. In an example, the cell(s) is/are wild-type cells or naturallyoccurring cells, eg, comprised by a naturally-occurring micrombiome, eg, of a human, animal, plant, soil, water, sea,

waterway or environment. In an example, the cell(s) is/are

for human or animal use, eg, for surgical or medical use, or for baby feeding). In an example, the composition is a personal hygiene composition for human or animal use. In an example, the composition is a composition for environmental use, eg, for soil treatment or environmental decon- 5 tamination (eg, from sewage, or from oil, a petrochemical or a chemical, eg, when the host is a sulphate-reducing bacterium, eg, a Desulfovibrio host). In an example, the composition is a plant growth stimulator. In an example, the composition is a composition for use in oil, petrochemical, 10 metal or mineral extraction. In an example, the composition is a fabric treatment or additive. In an example, the composition is an animal hide, leather or suede treatment or additive. In an example, the composition is a dye additive. In an example, the composition is a beverage (eg, beer or 15 wine) brewing or fermentation additive (eg, when the host is a Lactobacillus host). In an example, the composition is a paper additive. In an example, the composition is an ink additive. In an example, the composition is a glue additive. In an example, the composition is an anti-human or animal 20 or plant parasitic composition. In an example, the composition is an air additive (eg, for air in or produced by air conditioning equipment, eg, where the host is a Legionella host). In an example, the composition is an anti-freeze additive (eg, where the host is a Legionella host). In an 25 example, the composition is an eyewash or opthalmic composition (eg, a contact lens fluid). In an example, the composition is comprised by a dairy food (eg, the composition is in or is a milk or milk product; eg, wherein the host is a Lactobacillus, Streptococcus, Lactococcus or Listeria 30 host). In an example, the composition is or is comprised by a domestic or industrial cleaning product (eg, where the host is an E coli, Salmonella, Listeria or Clostridium (eg, botulinum) host). In an example, the composition is comprised by a fuel. In an example, the composition is comprised by a 35 solvent (eg, other than water). In an example, the composition is a baking additive (eg., a food baking additive). In an example, the composition is a laboratory reagent (eg, for use in biotechnology or recombinant DNA or RNA technology). In an example, the composition is comprised by a fibre 40 retting agent. In an example, the composition is for use in a vitamin synthesis process. In an example, the composition is an anti-crop or plant spoiling composition (eg, when the host is a saprotrophic bacterium). In an example, the composition is an anti-corrosion compound, eg, for preventing or reduc- 45 ing metal corrosion (eg, when the host is a sulphate-reducing bacterium, eg, a Desulfovibrio host, eg for use in reducing or preventing corrosion of oil extraction, treatment or containment equipment; metal extraction, treatment or containment equipment; or mineral extraction, treatment or con- 50 tainment equipment). In an example, the composition is an agricultural or farming composition or comprised in such a composition. In an example, the composition is a silage additive. The invention provides a CRISPR array, gRNAencoding nucleotide sequence, vector or plurality of vectors 55 described herein for use in any of the compositions described in this paragraph or for use in any application described in this paragraph, eg, wherein the host cell is a bacterial or archaeal cell. The invention provides a method for any application described in this paragraph, wherein the 60 method comprises combining a CRISPR array, gRNA-encoding nucleotide sequence, vector or plurality of the invention with a host cell (eg, bacterial or archaeal cell). In an embodiment, the host cell is not present in or on a human (or human embryo) or animal.

artificially genetically modified. In an example, a plurality of vectors of the invention are introduced into a plurality of said host cells, wherein the host cells are comprised by a bacterial population, eg, ex vivo, in vivo or in vitro. In an example, the host cells are comprised by a microbiota population comprised by an organism or environment (eg, a waterway microbiota, water microbiota, human or animal gut microbiota, human or animal oral cavity microbiota, human or animal vaginal microbiota, human or animal skin or hair microbiota or human or animal armpit microbiota), the population comprising first bacteria that are symbiotic or commensal with the organism or environment and second bacteria comprising said host cells, wherein the host cells are detrimental (eg, pathogenic) to the organism or environment. In an embodiment, the population is ex vivo. In an example, the ratio of the first bacteria sub-population to the second bacteria sub-population is increased. In an example, the first bacteria are Bacteroides (eg, B fragalis and/or B thetaiotamicron) bacteria. Optionally, the Bacteroides comprises one, two, three or more Bacteroides species selected from caccae, capillosus, cellulosilyticus, coprocola, coprophilus, coprosuis, distasonis, dorei, eggerthii, faecis, finegoldii, fluxus, fragalis, intestinalis, melaninogenicus, nordii, oleiciplenus, oxalis, ovatus, pectinophilus, plebeius, stercoris, thetaiotaomicron, uniformis, vulgatus and xylanisolvens. For example, the Bacteroides is or comprises B thetaiotaomicron. For example, the Bacteroides is or comprises B fragalis.

Any aspect of the present invention is, for example, for an industrial or domestic use, or is used in a method for such

In an example, the host, first or second cells are any bacterial species disclosed in US20160333348, GB1609811.3, PCT/EP2017/063593 and all US equivalent applications. The disclosures of these species (including specifically, Table 1 of PCT/EP2017/063593), are incorporated herein in their entirety and for potential inclusion of one or more disclosures therein in one or more claims berein

In an example, the host cell(s) or bacterial population is harboured by a beverage or water (eg, a waterway or drinking water) for human consumption. In an example, the host cell(s) or said population is comprised by a composition (eg, a medicament (eg, bacterial gut transplant), beverage, mouthwash or foodstuff) for administration to a human or non-human animal for populating and rebalancing the gut or oral microbiota thereof (eg, wherein said use of the medicament is to treat or prevent a disease or condition in the human or animal). In an example, the host cell(s) or said

population are on a solid surface or comprised by a biofilm (eg, a gut biofilm or a biofilm on an industrial apparatus). In an example of the invention for in vitro treating an industrial or medical fluid, solid surface, apparatus or container (eg, for food, consumer goods, cosmetics, personal healthcare product, petroleum or oil production); or for treating a waterway, water, a beverage, a foodstuff or a cosmetic, wherein the host cell(s) are comprised by or on the fluid, surface, apparatus, container, waterway, water, beverage, foodstuff or cosmetic.

In an example, the invention provides a container for medical or nutritional use, wherein the container comprises the vectors for use in the method. For example, the container is a sterilised container, eg, an inhaler or connected to a syringe or IV needle.

In an example, the vectors or composition is for administration (or is administered) to the human or non-human animal subject by mucosal, gut, oral, intranasal, intrarectal, intravaginal, ocular or buccal administration.

Optionally, each host cell is of a strain or species found in 20 human microbiota, optionally wherein the host cells are mixed with cells of a different strain or species, wherein the different cells are *Enterobacteriaceae* or bacteria that are probiotic, commensal or symbiotic with humans (eg, in the human gut. In an example, the host cell is an *E coli* or 25 *Salmonella* cell.

The invention is optionally for altering the relative ratio of sub-populations of first and second bacteria in a mixed population of bacteria, eg, for altering human or animal microbiomes, such as for the alteration of the proportion of 30 Bacteroidetes (eg, *Bacteroides*, eg, *fragalis* and/or *thetaiotamicron*), Firmicutes and/or gram positive or negative bacteria in microbiota of a human.

In an example, the vectors or composition of the invention comprises a nucleotide sequence for expressing in the host 35 cell an endolysin for host cell lysis, optionally wherein the endolysin is a phage phi11, phage Twort, phage P68, phage phiWMY or phage K endolysin (eg, MV-L endolysin or P-27/HP endolysin).

In an example, the target site is comprised by a chromosome of each microbe host cell, eg, wherein the sequence is comprised by an antibiotic resistance gene, virulence gene or essential gene of the host cell. An example, provides the vectors of the invention in combination with an antibiotic agent (eg, a beta-lactam antibiotic), eg, wherein the vectors 45 target a protospacer sequence comprised by an antibiotic resistance gene comprised by host cell genome or episome (eg, a plasmid comprised by the host cells). In an example, the episome is a plasmid, transposon, mobile genetic element or viral sequence (eg, phage or prophage sequence).

In an example, the target is a chromosomal sequence, an endogenous host cell sequence, a wild-type host cell sequence, a non-viral chromosomal host cell sequence, not an exogenous sequence and/or a non-phage sequence (ie, one more or all of these), eg, the sequence is a a wild-type 55 host chromosomal cell sequence such as antibiotic resistance gene or essential gene sequence comprised by a host cell chromosome. In an example, the sequence is a host cell plasmid sequence, eg, an antibiotic resistance gene sequence.

Optionally, the nuclease is a Cas and the target site is comprised by a protospacer sequence that is a adjacent a NGG, NAG, NGA, NGC, NGGNG, NNGRRT or NNA-GAAW protospacer adjacent motif (PAM), eg, a AAAGAAA or TAAGAAA PAM (these sequences are written 5' to 3'). In an embodiment, the PAM is immediately adjacent the 3' end of the protospacer sequence. In an

example, the Cas is a *S aureus, S theromophilus* or *S pyogenes* Cas. In an example, the Cas is Cpf1 and/or the PAM is TTN or CTA. Optionally, the Cas is a Type I (eg, Type I-A, I-B, I-C, I-D, I-E, or I-F) CRISPR system Cas. Optionally, the Cas is a Type II CRISPR system Cas. Optionally, the Cas is a Type IIII CRISPR system Cas. Optionally, the Cas is a Type IV CRISPR system Cas. Optionally, the Cas is a Type V CRISPR system Cas. Optionally, the Cas is a Type VI CRISPR system Cas.

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Optionally, the nuclease is a Cas and each vector comprises a cognate CRISPR array that comprises multiple copies of the same spacer for targeting the target site. Optionally, there is provide a vector or plurality of vectors of the invention, wherein the vector(s) comprises a plurality of CRISPR arrays of said gRNA-encoding sequences for host cell protospacer sequence targeting, wherein the protospacers comprise the target site. Optionally, the or each vector comprises two, three or more of copies of nucleic acid sequences encoding crRNAs (eg, gRNAs), wherein the copies comprise the same spacer sequence for targeting a host cell target site (eg, a site comprised by a virulence, resistance or essential gene sequence).

In an example, at least two target sequences are modified by Cas, for example an antibiotic resistance gene and an essential gene. Multiple targeting in this way may be useful to reduce evolution of escape mutant host cells.

In an example, the Cas is a wild-type endogenous host cell Cas nuclease. In an example, target site cutting is carried out by a dsDNA Cas nuclease (eg, a Cas9, eg, a spCas9 or saCas9), whereby repair of the cut is by non-homologous end joining (NHEJ); alternatively the Cas is an exonuclease or Cas3

In an example, the array, gRNA-encoding sequence or vector is not in combination with a Cas endonuclease-encoding sequence that is naturally found in a cell together with repeat sequences of the array or gRNA-encoding sequence.

A tracrRNA sequence may be omitted from an array or vector of the invention, for example for Cas systems of a Type that does not use tracrRNA, or an endogenous tracrRNA may be used with the cRNA encoded by the vector.

In an example, the host target site is comprised by at least 5, 6, 7, 8, 9, 10, 20, 30 or 40 contiguous nucleotides.

In an example, the or each vector comprises an exogenous promoter functional for transcription of the crRNA or gRNA in the microbes.

Optionally, each vector is a plasmid, cosmid, virus, a virion, phage, phagemid or prophage. For example, the invention provides a plurality of bacteriophage comprising a plurality of vectors of the invention, eg, wherein the vectors are identical. In an example, the vector is a viral vector. Viral vectors have a particularly limited capacity for exogenous DNA insertion, thus virus packaging capacity needs to be considered. Room needs to be left for sequences encoding vital viral functions, such as for expressing coat proteins and polymerase. In an example, the vector is a phage vector or an AAV or lentiviral vector. Phage vectors are useful where the host is a bacterial cell. In an example, the vector is a virus capable of infecting an archaea host cell.

Optionally, vector components are comprised by a transposon that is capable of transfer into and/or between host cells. The transposon can be a transposon as described in US20160333348, GB1609811.3 and all US equivalent applications; the disclosures of these, including these specific transposon disclosures, are incorporated herein in its entirety and for potential inclusion of one or more disclosures therein in one or more claims herein.

In an example, the or each vector is provided by a nanoparticle or in liposomes.

In an example, the or each host cell (or first and/or second bacteria) is a gram positive cell. In an example, the or each host cell is an *Enterobacteriaceae*, eg, *Salmonella*, *Yersinia 5 pestis*, *Klebsiella*, *Shigella*, *Proteus*, *Enterobacter*, *Serratia*, or *Citrobacter* cells. Optionally, the or each cell is an *E coli* (eg, *E coli* K12) or *Salmonella* (eg, *S enteric* serovar *typhimurium*) cell. Optionally, the or each host cell (or first and/or second bacteria) is a gram negative cell.

Optionally, the host (or first and/or second bacteria) is a mycoplasma, chlamydiae, spirochete or mycobacterium. Optionally, the host (or first and/or second bacteria) is a Streptococcus (eg, pyogenes or thermophilus) host. Optionally, the host (or first and/or second bacteria) is a Staphy- 15 lococcus (eg, aureus, eg, MRSA) host. Optionally, the host (or first and/or second bacteria) is an E. coli (eg, O157: H7) host. Optionally, the host (or first and/or second bacteria) is a Pseudomonas (eg, aeruginosa) host. Optionally, the host (or first and/or second bacteria) is a *Vibro* (eg. *cholerae* (eg. 20 O139) or vulnificus) host. Optionally, the host (or first and/or second bacteria) is a Neisseria (eg, gonnorrhoeae or meningitidis) host. Optionally, the host (or first and/or second bacteria) is a Bordetella (eg, pertussis) host. Optionally, the host (or first and/or second bacteria) is a Haemophilus (eg. 25 influenzae) host. Optionally, the host (or first and/or second bacteria) is a Shigella (eg, dysenteriae) host. Optionally, the host (or first and/or second bacteria) is a Brucella (eg, abortus) host. Optionally, the host (or first and/or second bacteria) is a Francisella host. Optionally, the host (or first 30 and/or second bacteria) is a *Xanthomonas* host. Optionally, the host (or first and/or second bacteria) is a Agrobacterium host. Optionally, the host (or first and/or second bacteria) is a Erwinia host. Optionally, the host (or first and/or second bacteria) is a Legionella (eg, pneumophila) host. Optionally, 35 the host (or first and/or second bacteria) is a Listeria (eg, monocytogenes) host. Optionally, the host (or first and/or second bacteria) is a Campylobacter (eg, jejuni) host. Optionally, the host (or first and/or second bacteria) is a Yersinia (eg, pestis) host. Optionally, the host (or first and/or 40 second bacteria) is a Borelia (eg, burgdorferi) host. Optionally, the host (or first and/or second bacteria) is a Helicobacter (eg, pylori) host. Optionally, the host (or first and/or second bacteria) is a *Clostridium* (eg, dificile or botulinum) host. Optionally, the host (or first and/or second bacteria) is 45 a Erlichia (eg, chaffeensis) host. Optionally, the host (or first and/or second bacteria) is a Salmonella (eg, typhi or enterica, eg, serotype typhimurium, eg, DT 104) host. Optionally, the host (or first and/or second bacteria) is a Chlamydia (eg, pneumoniae) host. Optionally, the host (or 50 first and/or second bacteria) is a Parachlamydia host. Optionally, the host (or first and/or second bacteria) is a Corynebacterium (eg, amycolatum) host. Optionally, the host (or first and/or second bacteria) is a Klebsiella (eg, pneumoniae) host. Optionally, the host (or first and/or sec- 55 ond bacteria) is a Enterococcus (eg, faecalis or faecim, eg, linezolid-resistant) host. Optionally, the host (or first and/or second bacteria) is a Acinetobacter (eg, baumannii, eg, multiple drug resistant) host.

Optionally, the invention is for reducing the growth or 60 proliferation of host cell(s) in an environment (eg, soil, a composition comprising said host cells and yeast cells), human, animal or plant microbiome. This is useful, for example, when the microbiome is naturally-occurring.

Optionally, the nuclease is a Cas and the target is comprised by a protospacer sequence comprising at least 5, 6, 7, 8, 9 or 10 contiguous nucleotides immediately 3' of a

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cognate PAM in the genome of the host cell, wherein the PAM is selected from AWG, AAG, AGG, GAG and ATG. Non Medical. Ex Vivo & In Vitro Uses Etc

In certain configurations, the inventive observation of rapid and durable microbial killing and growth or proliferation inhibition using nuclease cutting finds application in subjects other than humans and animals (eg, to treat plants or yeast cultures), or for ex vivo or in vitro treatment of substrates, such as industrial surfaces, fluids and apparatus. Thus, the invention further provides the following Concepts. Any other feature herein of the invention, its configurations, aspects, embodiments, options and examples above and elsewhere herein are combinable mutatis mutandis with these Concepts (including for providing combinations of features in the claims herein).

A Concept provides: Use of a nuclease, plurality of viruses, system, guide RNA, DNA or vector of the invention, in the manufacture of a composition for carrying out a method of treatment as defined herein, wherein the subject is an organism other than a human or animal.

A Concept provides: Use of a nuclease, plurality of viruses, system, guide RNA, DNA or vector of the invention, in the manufacture of a composition for carrying out an ex vivo or in vitro a method of treatment of a microbial infection of a substrate, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the substrate, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the substrate is treated.

Herein, treatment of an infection of a substrate may mean the treatment of a bacterial population (eg, one or more colonies) on a surface of the substrate and/or incorporated in the material of the substrate. For example, the treatment may be the treatment to kill bacteria on the surface of an industrial apparatus or equipment (eg, medical equipment, such as a scalpel or medical device or tubing). In another example, the substrate is a fluid (eg, a liquid or a gas), such as a medical fluid or petroleum product in fluid form (eg, an oil or hydrocarbon fluid or liquid).

A Concept provides: Use of a programmable nuclease in the manufacture of a composition for carrying out an ex vivo method of treatment of a microbial infection of a substrate, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the substrate, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the substrate is treated.

Optionally the nuclease (eg, programmed nuclease) and/ or a nucleic acid that programs the nuclease to recognise and cut the target site is administered to the subject or substrate at a first time (T1) and at a second time (T2) wherein T2 is at least 1 hour after T1. T1 and T2 may be as defined herein.

Optionally, the infection is reduced at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment. Optionally, the infection is maintained by at least

100-fold for at least 60 minutes (eg, at least 120 minutes) after exposing the subject to the programmed nuclease.

Optionally, the reduction in infection persists for 30 minutes immediately after the first 30 minutes of the treatment.

Optionally, the method comprises administering to the subject or substrate a RNA or a nucleic acid that encodes an RNA for expression of the RNA in or on the subject or substrate, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes 10 comprised by the subject or substrate.

Optionally, the nuclease is administered simultaneously or sequentially with the RNA or nucleic acid to the subject or substrate.

Optionally, the subject or substrate comprises the nucle- 15 ase prior to administration of the RNA or nucleic acid.

Optionally, a plurality of viruses (eg, phage) are administered to the subject or substrate, wherein each virus comprises a copy of the nucleic acid, wherein the viruses infect the microbes comprised by the subject or substrate to deliver 20 thereto the nucleic acid.

Optionally, the ratio of administered viruses:microbes is from 10 to 150.

Optionally, the infection is reduced by at least 90% for 1 hour or more, optionally by the first 30 minutes (eg, by the 25 first 15 minutes) of the treatment.

Optionally, the infection is reduced at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein reduction of the infection by at least 100-fold is maintained for at least 60 minutes (eg, at least 30 120, 145 or 180 minutes) after exposing the subject or substrate to the programmed nuclease.

Optionally, the subject is a plant; or wherein the substrate is a metallic, plastic, concrete, stone, wood, glass or ceramic substrate. Optionally, the subject is a fluid (eg, a liquid or a 35 gas).

Optionally, the microbes are bacteria or archaea. Optionally, the bacteria are gram positive bacteria. Optionally, the bacteria are *Staphylococcus*, *Streptococcus*, *Enterococcus*, *Legionella*, *Heamophilus*, *Ghonnorhea*, *Acinetobacter*, 40 *Escherichia*, *Klebsiella*, *Pseudomonas* or *Stenotrophomonas* bacteria (eg, *E coli* (eg, EHEC *E coli*), *C dificile*, *V cholera*, *Staphylococcus* (eg, *S aureus* or MRSA), *Streptococcus pyogenes*, *Acinetobacter baumannii*, *Legionella*, *Pseudomonas* aeruginosa, *Klebsiella pneumoniae* bacteria).

Optionally, the nuclease is a Cas nuclease (eg, a Cas 3 or 9), a meganuclease, a TALEN (Transcription activator-like effector nuclease) or zinc finger nuclease.

Reference is made to WO2016177682, which discusses aspects of microbiologically influenced corrosion (MIC) or 50 biofouling of substrates and discloses methods for controlling MIC or biofouling of a substrate. The methods, nucleases, arrays, RNAs, vectors and viruses disclosed in that document can be employed in the present invention, for example for carrying out the method or use of the present invention and the disclosures of these parts and the substrates and bacteria disclosed in WO2016177682 are incorporated herein by reference for potentially providing disclosure of features possible to be used in one or more claims berein.

Optionally, the use of the present invention is for controlling microbiologically influenced corrosion (MIC) or biofouling of a substrate in an industrial or domestic system (eg, a system disclosed in WO2016177682, which disclosure is incorporated herein by reference). In an example, the 65 system comprises equipment (eg, for use in an industrial process) and the surface is a surface of said equipment. In an

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example, the biofouling comprises microbial biofilm and/or sludge formation, proliferation or maintenance. In an example, the microbes are sessile. In an example "controlling" comprises preventing, reducing or eliminating said MIC or biofouling, or reducing spread of said MIC or biofouling in the system. Cell growth or proliferation or maintenance is, for example, a characteristic of cell viability. Thus, in an example, the method reduces microbe proliferation and/or maintenance.

Optionally, the microbes are comprised by a microbial biofilm that is in contact with said substrate. Optionally, said surface and host cells are in contact with a fluid, such as an aqueous liquid (eg, sea water, fresh water, stored water or potable water).

Fresh water is naturally occurring water on the Earth's surface in ice sheets, ice caps, glaciers, icebergs, bogs, ponds, lakes, rivers and streams, and underground as groundwater in aquifers and underground streams. Fresh water is generally characterized by having low concentrations of dissolved salts and other total dissolved solids. The term specifically excludes sea water and brackish water, although it does include mineral-rich waters such as chalybeate springs. In an example said fresh water is any of these fresh water types. Potable water is water for human or animal (eg, livestock) consumption. In an example, the fluid is selected from industrial cooling water wherein the system is a cooling system; sewage water wherein the system is a sewage treatment or storage system; drinking water wherein the system is a drinking water processing, storage, transportation or delivery system; paper making water wherein the system is a paper manufacture or processing system; swimming pool water wherein the system is a swimming pool or swimming pool water treatment or storage system; fire extinguisher water wherein the system is a fire extinguishing system; or industrial process water in any pipe, tank, pit, pond or channel.

Optionally, the use is for controlling bacterial souring of a liquid in a reservoir or container), wherein the fluid comprises a population of first host cells of a first microbial species that mediates said biofouling, the method comprising

(i) contacting the population with a plurality of vectors that are capable of transforming or transducing the cells, each vector comprising a CRISPR array whereby CRISPR arrays are introduced into the host cells, wherein

- (a) each CRISPR array comprises one or more sequences for expression of a crRNA and a promoter for transcription of the sequence(s) in a host cell; and
- (b) each crRNA is capable of hybridising to a target sequence of a host cell to guide Cas (eg, a Cas nuclease) in the host cell to modify the target sequence (eg, to cut the target sequence); the target sequence being a gene sequence for mediating host cell viability; and

wherein the method comprises allowing expression of said cRNAs in the presence of Cas in host cells, thereby modifying target sequences in host cells, resulting in reduction of host cell viability and control of said biofouling.

In an example, the fluid is a liquid. In an example, the fluid is a gaseous fluid.

60 Systems:

An example system is selected from the group consisting of a:—

Petrochemical recovery, processing, storage or transportation system; hydrocarbon recovery, processing, storage or transportation system; crude oil recovery, processing, storage or transportation system; natural gas recovery, processing, storage or transportation system, (eg, an oil well, oil rig,

oil drilling equipment, oil pumping system, oil pipeline, gas rig, gas extraction equipment, gas pumping equipment, gas pipeline, oil tanker, gas tanker, oil storage equipment or gas storage equipment); Water processing or storage equipment; water reservoir (eg, potable water reservoir); Air or water 5 conditioning (eg, cooling or heating) equipment, eg, a coolant tube, condenser or heat exchanger; Medical or surgical equipment; Environmental (eg, soil, waterway or air) treatment equipment; Paper manufacturing or recycling equipment; Power plant, eg, a thermal or nuclear power plant; 10 Fuel (eg, hydrocarbon fuel, eg, petroleum, diesel or LPG) storage equipment; Mining or metallurgical, mineral or fuel recovery system, eg, a mine or mining equipment; Engineering system; Shipping equipment; Cargo or goods storage equipment (eg, a freight container); Food or beverage 15 manufacturing, processing or packaging equipment; Cleaning equipment (eg, laundry equipment, eg, a washing machine or dishwasher); Catering (eg, domestic or commercial catering) equipment; Farming equipment; Construction (eg, building, utilities infrastructure or road construction) 20 equipment; Aviation equipment; Aerospace equipment; Transportation equipment (eg, a motor vehicle (eg, a car, lorry or van); a railcar; an aircraft (eg, an aeroplane) or a marine or waterway vehicle (eg, a boat or ship, submarine or hovercraft)); Packaging equipment, eg, consumer goods 25 packaging equipment; or food or beverage packaging equipment; Electronics (eg, a computer or mobile phone or an electronics component thereof); or electronics manufacture or packaging equipment; Dentistry equipment; Industrial or domestic piping (eg, a sub-sea pipe) or storage vessel (eg, a 30 water tank or a fuel tank (eg, gasoline tank, eg, a gasoline tank of a vehicle)); Underground equipment; Building (eg, a dwelling or office or commercial premises or factory or power station); Roadway; Bridge; Agricultural equipment; Factory system; Crude oil or natural gas exploration equip- 35 ment; Office system; and a Household system.

In an example, the system is used in an industry or business selected from the group consisting of agriculture, oil or petroleum industry, food or drink industry, clothing industry, environmental industry, chemical industry, aerospace industry, automotive industry, biotechnology industry, medical industry, healthcare industry, dentistry industry, energy industry, consumer products industry, pharmaceutical industry, mining industry, cleaning industry, forestry 45 industry, fishing industry, leisure industry, recycling industry, cosmetics industry, plastics industry, pulp or paper industry, textile industry, clothing industry, leather or suede or animal hide industry, tobacco industry and steel industry. In an example, the surface or fluid to be treated is a surface 50 or fluid of equipment used in said selected industry. In an example, the system is used in the crude oil industry. In an example, the system is used in the natural gas industry. In an example, the system is used in the petroleum industry. In an example, the system is a sea container, platform or rig (eg, 55 oil or gas platform or rig for use at sea or at sea), ship or boat. In an embodiment, such a system is anchored at sea; eg, non-temporarily anchored at sea, eg, has been anchored at sea for 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24 or more months (eg, contiguous 60 months). In an embodiment, such a system is in the waters of a country or state; eg, non-temporarily at sea in such waters, eg, has been in waters of said country for 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24 or more months (eg, contiguous months).

In an example, the substrate surface to be treated comprises stainless steel, carbon steel, copper, nickel, brass, 34

aluminium, concrete, a plastic or wood. In an example, the substrate is a metal weld or join. In an example, the surface is a metallic (eg, steel or iron) or non-metallic (eg, plastic, concrete, asphalt, wood, rubber or stone) surface. In an example, the metal is an alloy (eg, stainless steel, brass or a nickel-, zinc-, copper-, nickel- or aluminium-alloy). In an example, the surface is a man-made polymer surface. In an example, the surface is a substrate coating. In an example, the substrate is in contact with soil, fresh water or sea water.

In an example, the fluid is potable water; a waterway; brackish water; or a liquid fuel, eg, gasoline or diesel (eg, for a car or motorized vehicle), LPG, kerosine, an alcohol (eg, ethanol, methanol or butanol), liquid hydrogen or liquid ammonia), in an example, the fuel is stored liquid fuel. In an example the fluid is an oil or non-aqueous liquid. In an example, the fluid is a liquid comprised by a waterway or body of water, eg, sea water, fresh water, potable water, a river, a stream, a pond, a lake, a reservoir, stored water (eg, in a water storage tank or cooling equipment), groundwater, well water, water in a rock formation, soil water or rainwater. In an example, the liquid is sea water. In an example, the substrate is in contact with a liquid mentioned in this paragraph. In an example, the fluid or liquid is selected from the group consisting of an oil, an aqueous solution, a hydraulic fracturing fluid, a fuel, carbon dioxide, a natural gas, an oil/water mixture, a fuel/water mixture, water containing salts, ocean or sea water, brackish water, sources of fresh water, lakes, rivers, stream, bogs, ponds, marshes, runoff from the thawing of snow or ice, springs, groundwater, aquifers, precipitation, any substance that is a liquid at ambient temperature (eg, at rtp) and is hydrophobic but soluble in organic solvents, hexanes, benzene, toluene, chloroform, diethyl ether, vegetable oils, petrochemical oils, crude oil, refined petrochemical products, volatile essential oils, fossil fuels, gasoline, mixtures of hydrocarbons, jet fuel, rocket fuel, biofuels. In an example the fluid is an oil/water mixture.

The terms "microbiologically influenced corrosion" or "MIC" as used herein, unless otherwise specified, refer to industry, packaging industry, electronics industry, computer 40 processes in which any element (substrate) of a system is structurally compromised due to the action of at least one member of a microbial population, eg, bacterial or archaeal population. The term "biofouling" as used herein, unless otherwise specified, refers to processes in which microorganisms (such as bacteria and/or archaea) accumulate on a substrate surface in contact with a fluid (eg, water or an aqueous liquid, or a hydrocarbon, or a petrochemical). Also included is the undesirable accumulation and proliferation of microorganisms (such as bacteria and/or archaea) in a fluid (eg, water or an aqueous liquid, or a hydrocarbon, or a petrochemical), ie, "souring" of the fluid. In an example, the bacteria are comprised by ship or boat ballast water and the bacteria are environmentally undesirable. The term "substrate" as used herein refers to any type of surface on which cells can attach and a biofilm can form and grow or on which biofouling (eg slime or sludge formation) can occur. The substrate may be an "industrial" substrate such as the surface of equipment in an petrochemical, fuel, crude oil or gas piping system, or a "non-industrial" (eg, domestic, eg, household or office) substrate such as a kitchen counter or a shower substrate or a garden substrate.

In an alternative, instead of a population of host bacterial cells, the population is a population of archaeal cells of a first species.

Optionally, said fluid is an aqueous liquid (eg, sea water, fresh water, stored water or potable water).

In an alternative, instead the microbes are algal cells.

Optionally, the microbes are sulphate reducing bacteria (SRB) cells (eg, Desulfovibrio or Desulfotomaculum cells). In an example, the cells are selected from the group consisting of Desulfotomaculum nigrificans, Desulfacinum infernum, Thermodesulfobacterium mobile, Thermodesulforhabdus norvegicus, Archaeoglobus fulgidus, Desulfomicrobium apsheronum, Desulfovibrio gabonensis, Desulfovibrio longus, Desulfovibrio vietnamensis, Desulfobacterium cetonicum, Desulfomaculum halophilum, Desulfobacter vibrioformis and Desulfotomaculum thermocisternum cells. 10 In an example, the population comprises a mixture of two or more of these cell species.

Optionally, the surface or fluid is comprised by a crude oil, gas or petrochemicals recovery, processing, storage or transportation equipment. Crude oil is one of the most 15 important energetic resources in the world. It is used as raw material in numerous industries, including the refinery-petrochemical industry, where crude oil is refined through various technological processes into consumer products such as gasoline, oils, paraffin oils, lubricants, asphalt, 20 domestic fuel oil, vaseline, and polymers. Oil-derived products are also commonly used in many other chemical processes. In an alternative, the fluid is a said consumer product or the surface is in contact with such a consumer product.

Optionally, the surface is in contact with sea water, a fracking liquid or liquid in a well; or wherein the fluid is sea water, a fracking liquid or liquid in a well.

Optionally, step (i) of the method comprises providing a population of microbial cells of a second species (second 30 host cells), the second cells comprising said vectors, wherein the vectors are capable of transfer from the second host cells to the first host cells; and combining the second host cells with the first host cells, whereby vectors are introduced into the first host cells. In an example, the second cell(s) are 35 environmentally-, industrially-, or domestically-acceptable in an environment (eg, in a water or soil environment) and the first host cell(s) are not acceptable in the environment.

Optionally, the first host cells are comprised by a mixture of microbial cells (eg, comprised by a microbial biofilm) 40 before contact with said vectors, wherein the mixture comprises cells of said second species.

Optionally, said second species is a species of *Bacillus* or nitrate-reducing bacteria or nitrate reducing sulfide oxidizing bacteria (NRB)

Optionally, the NRB is selected from the group consisting of *Campylobacter* sp., *Nitrobacter* sp., *Nitrosomonas* sp., *Thiomicrospira* sp., *Sulfurospirillum* sp., *Thauera* sp., *Paracoccus* sp., *Pseudomonas* sp., *Rhodobacter* sp. and *Desulfovibrio* sp; or comprises at least 2 of said species.

Optionally, the NRB is selected from the group consisting of Nitrobacter vulgaris, Nitrosomonas europea, Pseudomonas stutzeri, Pseudomonas aeruginosa, Paracoccus denitrificans, Sulfurospirillum deleyianum, andRhodobacter sphaeroides.

Optionally, the method comprises contacting the host cells of said first species with a biocide simultaneously or sequentially with said vectors. In an example, the vectors and biocide are provided pre-mixed in a composition that is contacted with the host cells.

Optionally, the biocide is selected from the group consisting of tetrakis hydroxymethyl phosphonium sulfate (THPS), glutaraldehyde, chlorine monoxide, chlorine dioxide, calcium hypochlorite, potassium hypochlorite, sodium hypochlorite, dibromonitriloproprionamide (DBNPA), methylene bis(thiocyanate) (MBT), 2-(thiocyanomethylthio) benzothiazole (TCMTB), bronopol, 2-bromo-2-nitro-

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1,3-propanediol (BNPD), tributyl tetradecyl phosphonium chloride (TTPC), taurinamide and derivatives thereof, phenols, quaternary ammonium salts, chlorine-containing agents, quinaldinium salts, lactones, organic dyes, thiosemicarbazones, quinones, carbamates, urea, salicylamide, carbanilide, guanide, amidines, imidazolines, acetic acid, benzoic acid, sorbic acid, propionic acid, boric acid, dehydroacetic acid, sulfurous acid, vanillic acid, p-hydroxybenzoate esters, isopropanol, propylene glycol, benzyl alcohol, chlorobutanol, phenylethyl alcohol, formaldehyde, iodine and solutions thereof, povidone-iodine, hexamethylenetetramine, noxythiolin, 1-(3-chloroallyl)-3,5,7-triazo-1azoniaadamantane chloride, taurolidine, taurultam, N-(5nitro-2-furfurylidene)-1-amino-hydantoin, 5-nitro-2furaldehyde semicarbazone, 3,4,4'-trichlorocarbanilide, 3,4', 5-tribromosalicylanilide, 3-trifluoromethyl-4,4'dichlorocarbanilide, 8-hydroxy quinoline, 1-cyclopropyl-6fluoro-1,4-dihydro-4-oxo-7-(1-piperazinyl)-3-

quinolinecarboxylic acid, 1,4-dihydro-1-ethyl-6-fluoro-4-oxo-7-(1-piperazinyl)-3-quinolinecarboxylic acid, hydrogen peroxide, peracetic acid, sodium oxychlorosene, parachlorometaxylenol, 2,4,4'-trichloro-2'-hydroxydiphenol, thymol, chlorhexidine, benzalkonium chloride, cetylpyridinium chloride, silver sulfadiazine, silver nitrate, bromine, ozone, isothiazolones, polyoxyethylene (dimethylimino) ethylene (dimethylimino) ethylene (dimethylimino) ethylene dichloride, 2-(tert-butylamino)-4-chloro-6-ethylamino-5'-triazine (terbutylazine), and combinations thereof. In an example the biocide is tetrakis hydroxymethyl phosphonium sulfate (THPS). In an example, the biocide is a quaternary ammonium compound.

Optionally, the system is used in an industry operation selected from the group consisting of mining; shipping; crude oil, gas or petrochemicals recovery or processing; hydraulic fracturing; air or water heating or cooling; potable water production, storage or delivery; transportation of hydrocarbons; and wastewater treatment.

Optionally, the surface is a surface of equipment used in said selected industry; or wherein the fluid is a fluid comprised by equipment used in said selected industry.

Optionally, the surface is a surface of kitchen, bathing or gardening equipment; or wherein the fluid is comprised by kitchen, bathing or gardening equipment. For example, the equipment is used in a domestic setting.

Optionally, the fluid is a potable liquid contained in a container (eg, water tank or bottle) and the surface is a surface of the container in contact with the liquid.

Optionally, each vector comprises a mobile genetic element (MGE), wherein the MGE comprises an origin of transfer (oriT) and a said CRISPR array; wherein the MGE is capable of transfer between a host cell of said first species and a further microbial host cell in said industrial or domestic system. For example, the further cell(s) are environmentally-, industrially-, or domestically-acceptable in an environment (eg, in a water or soil environment) and the first host cell(s) are not acceptable in the environment. Optionally, the oriT is functional in the first and further host cells.

Optionally, the first and further host cells are comprised by a biofilm of fluid in contact with said surface; or wherein said cells are comprised by said fluid.

Optionally, each MGE is or comprises an integrative and conjugative element (ICE); or wherein each vector is a phage that is capable of infecting host cells of said first species and each MGE is a phage nucleic acid that is capable of said transfer between the cells. Optionally, each ICE is a transposon, eg, a conjugative transposon. Optionally, each vector is a plasmid, optionally comprising an MGE as

described herein. Optionally, the sequences are comprised by a conjugative transposon of the first cell and/or further cell.

In an example, the method is a method of controlling microbiologically influenced corrosion (MIC) or biofouling of a substrate comprised by a crude oil, gas or petrochemicals recovery, processing, storage or transportation equipment (eg, a crude oil tanker, oil rig or oil drilling equipment), wherein a surface of the substrate is in contact with a population of first host cells, wherein the first host cells are sulphur- or sulphate-reducing bacteria (SRB), extracellular polymeric substance-producing bacteria (EPSB), acid-producing bacteria (APB), sulphur- or sulphide-oxidizing bacteria (SOB), iron-oxidising bacteria (JOB), manganese-oxidising bacteria (MOB), ammonia producing bacteria (AmPB) or acetate producing bacteria (AcPB) of a first species that mediates MIC or biofouling of the substrate, wherein the surface and cell population are in contact with a liquid selected from sea water, fresh water, a fracking 20 liquid or liquid in a well (eg, oil or natural gas well), the method comprising

- (i) contacting the cell population with vectors by mixing the liquid with a plurality of vectors that are capable of transforming or transducing first host cells, each vector comprising a CRISPR array whereby CRISPR arrays are introduced into the host cells, wherein
  - (a) each CRISPR array comprises one or more sequences for expression of a crRNA and a promoter for transcription of the sequence(s) in a host cell;
  - (b) each crRNA is capable of hybridising to a target sequence of a host cell to guide Cas (eg, a Cas nuclease, eg, a Cas9 or Cfp1) in the host cell to modify the target sequence (eg, to cut the target sequence); the target sequence being a gene sequence for mediating host cell viability;
  - (c) wherein each sequence of (a) comprises a sequence R1-S1-R1' for expression and production of the respective crRNA in a first host cell, wherein R1 is a first 40 CRISPR repeat, R1' is a second CRISPR repeat, and R1 or R1' is optional; and S1 is a first CRISPR spacer that comprises or consists of a nucleotide sequence that is 70, 75, 80, 85, 90 or 95% or more identical to a target sequence of a said first host cell and
- (ii) allowing expression of said cRNAs in the presence of Cas in host cells, thereby modifying target sequences in host cells, resulting in reduction of host cell viability and control of MIC or biofouling of said substrate. In an embodiment, both R1 and R1' are present.

In an example, the method is a method of controlling bacterial biofouling in ballast water of a ship or boat, wherein the water comprises a population of first host cells of a first microbial species that mediates said biofouling, the method comprising

- (i) contacting the population with a plurality of vectors that are capable of transforming or transducing the cells, each vector comprising a CRISPR array whereby CRISPR arrays are introduced into the host cells, wherein
  - (a) each CRISPR array comprises one or more sequences 60 for expression of a crRNA and a promoter for transcription of the sequence(s) in a host cell; and
  - (b) each crRNA is capable of hybridising to a target sequence of a host cell to guide Cas (eg, a Cas nuclease) in the host cell to modify the target sequence (eg, to cut 65 the target sequence); the target sequence being a gene sequence for mediating host cell viability; and

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(ii) allowing expression of said cRNAs in the presence of Cas in host cells, thereby modifying target sequences in host cells, resulting in reduction of host cell viability and control of said biofouling.

Optionally, the first host cells are *Vibrio cholerae*, *E coli* or *Enterococci* sp cells.

Optionally, step (i) comprises mixing the ballast water with the vectors, eg, in the hull of a ship or boat. Optionally, the ship or boat is a marine vehicle and the water is sea water. Optionally, instead of a ship or boat, the ballast water is comprised by a container or a drilling platform at sea, eg, an oil platform or oil rig. In an example, the ship, boat, container, platform or rig is anchored at sea (ie, not temporarily in its location).

In an example, the method is a method of discharging ballast water from a ship or boat, wherein the discharged ballast water comprises water treated by the method. Optionally, the water is discharged into a body of water, eg, a sea, ocean or waterway (eg, a river, canal, lake or reservoir) or into a container.

#### Paragraphs:

The invention provides the following Paragraphs, which are supported by the Examples below:—

- 1. A programmable Cas (eg, Cas3 or Cas9) nuclease for use in a method of treating E coli or C dificile infection of a subject, wherein the Cas nuclease is programmable with a guide RNA to cut a target site comprised by the genomes of E coli or C dificile bacteria that have infected the subject, whereby E coli or C dificile cells are killed, or growth or proliferation of the cells is reduced, the treatment method comprising exposing the subject to the Cas nuclease wherein the nuclease is programmed with guide RNA to cut the target site, whereby genomes of the E coli or C dificile bacteria comprised by the subject are cut and the infection of the subject is reduced by at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 2. A programmable Cas (eg, Cas3 or Cas9) nuclease (optionally according to paragraph 1) for use in a method of treating E coli or C dificile infection of a subject, wherein the Cas nuclease is programmable with a guide RNA to cut a target site comprised by the genomes of E coli or C dificile bacteria that have infected the subject, whereby E coli or C dificile cells are killed, or growth or proliferation of the cells is reduced, the treatment method comprising exposing the subject to the Cas nuclease wherein the nuclease is programmed with guide RNA to cut the target site, whereby genomes of the E coli or C dificile bacteria comprised by the subject are cut and the infection of the subject is reduced, wherein a reduction of the infection by at least 100-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease.
- 3. The nuclease of any preceding Paragraph, wherein at least 60% of the infection is reduced by 60 minutes after exposing the subject to the programmed nuclease.
- 4. The nuclease of any preceding Paragraph, wherein the nuclease (eg, programmed nuclease) and/or a nucleic acid encoding the guide RNA is administered to the subject at a first time (T1) and at a second time (T2) wherein T2 is at least 1 hour after T1.
- 5. The nuclease of any preceding Paragraph, wherein the method comprises reducing the infection such that the reduction in infection persists for 30 minutes immediately after the first 30 minutes of the treatment.

- 6. The nuclease of any preceding Paragraph, wherein the method comprises administering to the subject the RNA or a nucleic acid that encodes the RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to 5 cut the target site in microbes comprised by the subject.
- The nuclease of any preceding Paragraph, wherein the nuclease is administered simultaneously or sequentially with the RNA or nucleic acid encoding the RNA to the subject.
- The nuclease of Paragraph 7, wherein the subject comprises the nuclease prior to administration of the RNA or nucleic acid to the subject.
- 9. The nuclease of any preceding Paragraph, wherein a plurality of viruses (eg, phage) are administered to the 15 subject, wherein each virus comprises a copy of a nucleic acid encoding the RNA, wherein the viruses infect the microbes comprised by the subject to deliver thereto the nucleic acid.
- 10. The nuclease of Paragraph 9, wherein the ratio of 20 administered viruses:microbes comprised by the subject is from 10 to 150.
- 11. The nuclease according to any preceding Paragraph, wherein the subject is a human or animal, optionally wherein the subject is a human over 65 years of age or 25 is a paediatric patient.
- 12. The nuclease according to Paragraph 11, wherein the infection is an infection of the lungs, abdomen or urinary tract; or wherein the subject has undergone surgery, is on an immunosuppressant medication and/or 30 is suffering from a chronic disease.
- 13. The nuclease according to any preceding Paragraph, wherein the infection is reduced by at least 90% for 1 hour or more, optionally by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 14. The nuclease according to any preceding Paragraph, wherein the method comprises reducing the infection at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein reduction of the infection by at least 100-fold is maintained for at least 40 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease.
- 15. The nuclease according to any one of Paragraphs 11 to 14, wherein the method treats or prevents septicaemia and/or sepsis (eg, septic shock) in the subject.
- 16. The nuclease of Paragraph 16, wherein at the start of the treatment, the subject (eg, a human) has a temperature of <36° C. or >38° C.; a heart rate of >90/min, a respiratory rate of >20 breaths/min or PaCO<sub>2</sub><4.3 kPa; and white blood cell count of <4000/mm³ or >12,000/ 50 mm³
- 17. The nuclease of Paragraph 15 or 16, wherein at the start of the treatment, the subject (eg, a human) has presence of two or more of the following: abnormal body temperature, abnormal heart rate, abnormal respiratory rate, abnormal blood gas and abnormal white blood cell count.
- 18. The nuclease of any preceding Paragraph, wherein the subject is a human or animal and the microbes are bacteria (eg, *E coli* or *C dificile*), wherein blood infection of the subject by the bacteria is reduced at least 100- or 1000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 19. The nuclease of any one of Paragraphs 11 to 18, wherein the blood of the subject is infected with from 65 10<sup>7</sup> to 10<sup>12</sup> CFU/ml of the bacteria immediately before the treatment.

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- 20. The nuclease according to any one of Paragraphs 1 to 10, wherein the subject is a plant.
- 21. The nuclease according to any preceding Paragraph, wherein the bacteria are comprised by a microbiome.
- 22. The nuclease according to Paragraph 21, wherein the microbiome comprises *Lactobacillus* and/or *Streptococcus* bacteria.
- 23. The nuclease according to any preceding Paragraph, wherein the *E coli* are EHEC *E coli*.
- 24. The nuclease according to any preceding Paragraph, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9), a meganuclease, a TALEN (Transcription activator-like effector nuclease) or zinc finger nuclease.
- 25. A plurality of viruses (eg, phage or phagemids for producing phage) for use with the nuclease of any preceding Paragraph in the method of treatment, wherein each virus comprises a copy of a nucleic acid encoding the RNA, wherein the viruses are capable of infecting microbes comprised by the subject to deliver thereto the nucleic acid.
- 26. A composition comprising a plurality of nucleic acids for programming the nuclease of any one of Paragraphs 1 to 24 in the method of treatment, wherein each nucleic acid is a nucleic acid as defined in any one of Paragraphs 6 to 9.
- 27. A CRISPR/Cas system comprising a nuclease according to any preceding Paragraph for use in the method of treatment, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9) and the system comprises one or more guide RNAs or DNA encoding one or more guide RNAs, wherein each guide RNA is capable of programming the Cas nuclease to cut a target site comprised by the genomes of the microbes.
- 28. A guide RNA or a DNA encoding a guide RNA for use in the system of Paragraph 27 for use in the method of treating an acute microbial infection in the subject, eg, septicaemia or sepsis.
- A nucleic acid vector comprising the guide RNA or DNA recited in Paragraph 27 or 28.
- The vector of Paragraph 29 wherein the vector is a phage, phagemid, viriophage, virus, plasmid (eg, conjugative plasmid) or transposon.
- 31. An anti-sepsis or anti-septicaemia composition for administration to a human or animal for treating sepsis or septicaemia, the composition comprising a plurality of vectors, wherein each vector is according to Paragraph 29 or 30.
- 32. A method of treating an acute microbial infection of a subject, wherein the method is as defined by any preceding Paragraph.
- 33. Use of a nuclease, plurality of viruses, system, guide RNA, DNA or vector of any one of Paragraphs 1 to 25 and 27 to 30, in the manufacture of a composition for carrying out a method of treatment as defined by any preceding Paragraph, wherein the subject is an organism other than a human or animal.
- 34. Use of a nuclease, plurality of viruses, system, guide RNA, DNA or vector of any one of Paragraphs 1 to 25 and 27 to 30, in the manufacture of a composition for carrying out an ex vivo method of treatment of a microbial infection of a substrate, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the substrate, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method com-

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prising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the substrate is treated.

- 35. Use of a programmable nuclease in the manufacture of a composition for carrying out an ex vivo method of treatment of a microbial infection of a substrate, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the substrate, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the  $_{15}$ treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial infection of the substrate is treated.
- 36. The use of Paragraph 33, 34, 35, wherein the nuclease (eg, programmed nuclease) and/or a nucleic acid that programs the nuclease to recognise and cut the target site is administered to the subject or substrate at a first time (T1) and at a second time (T2) wherein T2 is at 25 least 1 hour after T1.
- 37. The use of any one of Paragraphs 33 to 36, wherein the infection is reduced at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 38. The use of any one of Paragraphs 33 to 37, wherein 30 the reduction of the infection is maintained by at least 100-fold for at least 60 minutes (eg, at least 120 minutes) after exposing the subject to the programmed
- 39. The use of any one of Paragraphs 33 to 38, wherein 35 the reduction in infection persists for 30 minutes immediately after the first 30 minutes of the treatment.
- 40. The use of any one of Paragraphs 33 to 39, wherein the method comprises administering to the subject or substrate a RNA or a nucleic acid that encodes an RNA 40 for expression of the RNA in or on the subject or substrate, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject or substrate.
- 41. The use of Paragraph 40, wherein the nuclease is 45 administered simultaneously or sequentially with the RNA or nucleic acid to the subject or substrate.
- 42. The use of Paragraph 40, wherein the subject or substrate comprises the nuclease prior to administration of the RNA or nucleic acid.
- 43. The use of any one of Paragraphs 40 to 42, wherein a plurality of viruses (eg, phage) are administered to the subject or substrate, wherein each virus comprises a copy of the nucleic acid, wherein the viruses infect the microbes comprised by the subject or substrate to 55 deliver thereto the nucleic acid.
- 44. The use of Paragraph 43, wherein the ratio of administered viruses:microbes is from 10 to 150.
- 45. The use of any one of Paragraphs 33 to 44, wherein the infection is reduced by at least 90% for 1 hour or 60 more, optionally by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 46. The use of any one of Paragraphs 44 to 45, wherein the infection is reduced at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; 65 and wherein reduction of the infection by at least 100-fold is maintained for at least 60 minutes (eg, at

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least 120, 145 or 180 minutes) after exposing the subject or substrate to the programmed nuclease.

- 47. The use of any one of Paragraphs 33 to 46, wherein the subject is a plant; or wherein the substrate is a metallic, plastic, concrete, stone, wood, glass or ceramic substrate.
- 48. The use of any one of Paragraphs 33 to 47, wherein the microbes are bacteria.
- 49. The use according to Paragraph 48, wherein the bacteria are gram positive bacteria.
- 50. The use according to Paragraph 48 or 49, wherein the bacteria are Staphylococcus, Streptococcus, Enterococcus, Legionella, Heamophilus, Ghonnorhea, Acinetobacter, Escherichia, Klebsiella, Pseudomonas or Stenotrophomonas bacteria (eg, E coli (eg, EHEC E coli), C dificile, V cholera, Staphylococcus (eg, S aureus or MRSA), Streptococcus pyogenes, Acinetobacter baumannii, Legionella, Pseudomonas aeruginosa, Klebsiella pneumoniae bacteria).
- 51. The use of any one of Paragraphs 33 to 50, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9), a meganuclease, a TALEN (Transcription activator-like effector nuclease) or zinc finger nuclease.

Treatment of Pathogenic Bacterial Infections

Infectious complications are a serious cause of morbidity and mortality in cancer patients, especially those with underlying haematological malignancies where autopsy studies demonstrate that approximately 60% of deaths are infection related. Although fewer data exist on infectious mortality in patients with solid organ tumours, approximately 50% of these patients are estimated to have an infection as either the primary or an associated cause of death ("Epidemiology of Infections in Cancer Patients", in "Infectious Complications in Cancer Patients", Springer International Publishing Switzerland (2014)). Bacterial infections dominate. These infectious complications remain a significant limitation of cancer treatment modalities.

The detrimental effects of classic antibiotic treatment with broad-spectrum antibiotics have been demonstrated in immune checkpoint inhibitor (ICI)-treated cancer patients. Routy et al investigated how the gut microbiome influences efficacy of PD-1-based immunotherapy against epithelial tumours (Routy et al Science 2018, 359, 91-97). In this work, the authors also analyzed datasets for infections/ antibiotic use in patients with advanced NSCLC (n=140), renal cell carcinoma (n=67), or urothelial carcinoma (n=42) who received antibody ICI against PD-1/PD-L1 interaction after one or several prior therapies. Among these patients, they were prescribed broad-spectrum antibiotics (betalactam+/- inhibitors, fluoroquinolones, or macrolides) within 2 months before, or 1 month after, the first administration of PD-1/PD-L1 mAb. Patients generally took antibiotic orally for common indications (dental, urinary, and pulmonary infections). The detrimental effect of treating infections in cancer patients undergoing ICI therapy with classical, broad-spectrum antibiotics was observed. See FIG. 8, which shows that the antibiotic treatment during ICI therapy has fatal outcomes: there was a medial overall survival of 21.9 months in the absence of antibiotic treatment, compared to an overall survival of 9.8 months with antibiotic treatment. So, the median overall survival in patients treated with classical antibiotics is <50% (or >12 months shorter) that of patients not receiving antibiotic treatment.

The work by Gopalakrishnan et al is another recent example lending support to the importance of a "healthy" microbiome in immuno-oncology therapy outcomes (Go-

palakrishnan et al, Science 2018, 359, 97-103). See FIGS. 9A and 9B. It was observed that the gut microbiome modulates the efficacy of anti-PD-1 inhibition in melanoma patients.

Several other studies add to the expanding evidence base 5 of the critical link between the microbiome and immunooncology outcomes:

"Microbiota: a key orchestrator of cancer therapy", Nat. Rev. Cancer 2017, 17, 271-285

Matson et al, Science 2018, 359, 104-108

L. Derosa et al Annals of Oncology 2018 (epub 30 Mar. 2018)

M. Vétizou et al Science. 2015, 350, 1079-84

Sivan et al Science 2015, 350, 1084-1089

Another report—claiming to be the first systematic review 15 of infection among patients receiving immune checkpoint blockade for cancer therapy—investigated serious infections in melanoma patients treated with immune checkpoint inhibitors (against CTLA-4, PD-1, and/or PD-L1) (M. Del Castillo et al Clin. Infect. Dis. 2016, 63, 1490-1493). Serious 20 infections were defined as infections requiring hospitalization or parenteral antimicrobials. Of 740 patients (898 courses of immune checkpoint blockade), serious infection developed in 54 patients (7.3%). Nine patients (17%) were deemed to have died of an infection. Total number of 25 infections was 58, as some patients developed >1 infection. The majority of infections were bacterial in origin (~80%; i.e., bacterial infections: 80% of 7.3%: 5.8% of patients). Pneumonia and bloodstream infections were the two dominating bacterial infection types.

Immune checkpoint-blocking drugs are associated with immune-related adverse effects (irAEs) related to the upregulated immune system. The complications are managed with immunosuppressive drugs, such as steroids (immunosuppression is a risk factor for subsequent opportunis- 35 tic infections). Of the 740 patients, 46% received steroids during the course of treatment. Risk of serious infections was 13.5% in the cohort receiving corticosteroids or infliximab (vs. 7.3% in the overall population).

In yet another report, the emerging concern of infectious 40 diseases in lung cancer patients receiving ICI therapy was investigated. Of 84 NSCLC patients receiving nivolumab (a PD-1 inhibitor), 20 patients (23.8%) developed an infectious disease. Bacterial infections accounted for 75% of infeccommon type of bacterial infection was pneumonia. See K. Fujita et al Eur. Resp. J. 2017, 50, OA1478.

The Gram-negative bacillus E. coli is one of the most common causes of bacteraemia in patients with cancer. The all-cause 30-day mortality rate for this pathogen is high 50 (~15%) (Y. E. Ha et al Int. J. Antimicr. Agen. 2013, 42, 403-409). Published estimates of 30-day all-cause mortality among E. coli bacteraemia patients (cancer/non-cancer) vary from around 10 to 35% (J. K. Abernethy et al Clin. Microbiol. Infect. 2015, 21, 251.e1-251.e8), clearly highlighting 55 the high burden associated with just this pathogen. Overall, causative pathogens in bacteraemia are primarily Gramnegative bacteria (65%), with E. coli (18.3%), P. aeruginosa (18.3%), and K. pneumoniae (17.3%) being the most common organisms encountered; the three pathogens together 60 account for 54% of the bacteraemia cases, or 85% of Gram-negative cases, according to a study investigating >100 bacteraemia cases in cancer patients (G. Samonis et al Support Care Cancer 2013, 21, 2521-2526). In-hospital mortality was 26.2% in this study. Comparable numbers can 65 be found elsewhere. For example, a study of neutropenic and non-neutropenic adult cancer patients with bloodstream

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infections investigated 399 cases of bloodstream infections in 344 cancer patients: The largest causative pathogen group was Gram-negative bacilli (45%). Of the clinical isolates, E. coli (35%) accounted for the most cases within Gramnegatives, followed by K. pneumoniae (20%) and P. aeruginosa (19%) (E. Velasco et al Eur. J. Clin. Microbiol. Infect. Dis. 2006, 25, 1-7). The three pathogens collectively account for 33% of the bacteraemia cases (or 74% of Gram-negative cases). The overall 30-day mortality rate was 32% in this study. Two other reports looked at causative agents of bloodstream infection in patients with solid tumours and also found Gram-negative bacteria to be the dominating pathogen type (47-55% of the infections, across several hundred patients) (M. Marin et al Medicine 2014, 93, 143-149; M. Anatoliotaki et al Infection 2004, 32, 65-71; see also C. Gudiol et al Virulence 2016, 7, 298-308). In the larger of the two studies (with more robust numbers for individual pathogens), the three main pathogens within the Gram-negative group were again E. coli (55%), P. aeruginosa (18%), and Klebsiella spp. (11%)—corresponding to 92% of the Gram-negative cases, or 51% of the 528 total cases of bloodstream infections studied.

The above data on specific causative infectious pathogens in cancer patients are summarized in Table 5 below.

Thus, available data on bloodstream infections in cancer indicate that Gram-negative pathogens are involved in 45-65% of the infection cases, with three key pathogens—*E*. coli, K pneumoniae, and P. aeruginosa—being the culprits in the vast majority of Gram-negative cases (73-92%).

The inventors, thus, formulated an oncologist's dilemma: Reduction in efficacy of the cancer therapy is likely due to the reduced microbiome diversity resulting from antibiotic therapy

At least 1/3 of patients on checkpoint inhibitors get serious and life-threatening infections

Not treating these infections could result in death from the infection (1-2 weeks)

Treatment with classic antibiotics leads to reduction in progression free survival after 4 years from >40% to around 10%.

The choice is to treat the immediate need of a potentially fatal infection (which must be addressed) at the risk of seriously undermining the cancer therapy.

The inventors realised, therefore, that there is a need for tions; i.e., bacterial infections in 18% of patients. Most 45 methods that can treat a bacterial pathogenic infection in a different way that minimizes compromise to the cancer therapy. The inventors realised that this need would also be useful in other therapy settings where the microbiome composition can modulate therapy outcomes, eg, in transplant settings.

Whilst not wishing to be bound by any particular theory, the inventors believe that alleviating the detrimental effect of traditional antibiotic therapy on overall survival in ICI patients using the invention may, in some embodiments, translate to as much as a doubling of overall survival (or >12 months). Capturing a treatment effect of several months in terms of median overall survival is a very substantial achievement in this space. In fact, an effect size of this order of magnitude is comparable to the outcomes reported for ICI trials (i.e., where benefits usually are measured in months, not years). Additionally, PD-1/PD-L1 drugs are projected to dominate the ICI market. In 2023, PD-1/PD-L1 are projected to account for 94% of \$46B USD global sales of ICIs (CTLA-4 blockers only account for 6%), source: "Landscape & Forecast Immune Checkpoint Inhibitors", Decision Resources, December 2017. Thus, a need for improving treatment using immune checkpoint inhibitors of PD-1 or

PD-L1 is particularly pressing in medicine, and we believe that the present invention finds particular benefit in this

In an example, the method removes the need to administer a classic antibiotic, such as a broad-spectrum antibiotic (or 5 any other one disclosed herein). In another example, the invention reduces the amount or dosing frequency of a classic antibiotic, such as a broad-spectrum antibiotic (or any other one disclosed herein) that is administered to the subject for treating the infection. For example, the subject 10 can be administered a low-dose broad-spectrum antibiotic (eg, 50, 40, 30, 20, 10% or less of a conventional dose) whilst the guided nuclease cutting is used, and thus treatment of the infection in this setting. The invention may be particularly beneficial for patients on immunosuppressants, 15 eg, for cancer patients, transplant patients or patients suffering from a viral infection (eg, HIV (human immunodeficiency virus), CMV (cytomegalovirus) or RSV (respiratory synctial virus) infection).

The term "broad-spectrum antibiotic" can refer to an 20 antibiotic that acts on the two major bacterial groups, gram-positive and gram-negative, or any antibiotic that acts against a wide range of disease-causing bacteria. These medications are used when a bacterial infection is suspected but the group of bacteria is unknown (also called empiric 25 therapy) or when infection with multiple groups of bacteria is suspected. Although powerful, broad-spectrum antibiotics pose specific risks, particularly the disruption of native, normal bacteria and the development of antimicrobial resistance. Examples of commonly used broad-spectrum antibi- 30 otics are: Aminoglycosides (except for streptomycin), Ampi-Amoxicillin, Amoxicillin, clavulanic cillin. (Augmentin), Carbapenems (e.g. imipenem), Piperacillin, tazobactam, Quinolones (e.g. ciprofloxacin), Tetracyclines, Chloramphenicol, Ticarcillin, Trimethoprim and sulfame- 35 thoxazole (Bactrim). In veterinary medicine, examples are co-amoxiclav, (eg, in small animals), penicillin, streptomycin, oxytetracycline and potentiated sulfonamides.

The invention, therefore, in one aspect provides the 40 following Clauses that are directed to the treatment of a pathogenic bacterial infection using a programmed nuclease.

1. A method for treating a pathogenic bacterial infection in a human or animal subject caused by bacteria (first prising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a programmable nuclease that is programmed to cut the target site, wherein the subject 50 is suffering from a further disease or condition other than the pathogenic bacterial infection and the method comprises administering a therapy to the subject for treating or preventing the further disease or condition, wherein the nuclease treats the infection and the 55 therapy is efficacious in the presence of the programmed nuclease to treat or prevent the disease or condition.

In an example, Clause 1 provides:-

A method for treating a pathogenic bacterial infection in 60 a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is pro- 65 grammed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the

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subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer.

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer;

Wherein

- (a) The immunotherapy comprises administering to the patient an anti-PD-1 antibody optionally selected from pembrolizumab (or KEYTRUDATM) and nivolumab (or OPDIVOTM); and
- (b) The cancer is selected from metastatic melanoma; renal cell carcinoma; bladder cancer; a solid tumour; non-small cell lung cancer (NSCLC); forehead and neck squamous cell carcinoma (HNSCC); Hodgkin's lymphoma; a cancer that overexpresses PD-L land no mutations in EGFR or in ALK; colorectal cancer and hepatocellular carcinoma; and
- (c) The first bacteria are selected from Pseudomonas aeruginosa, Klebsiella pneumoniae, E coli, Salmonella (eg, S typhimurium), Clostridium dificile, Staphylococcus (eg, S aureus or S epidermis), Streptococcus (eg, S viridans or S thermophilus), Pneumococcus and Enterococcus bacteria.

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious bacteria) of a first species or strain, the method com- 45 in the presence of the programmed nuclease to treat the cancer:

Wherein

- (a) The immunotherapy comprises administering to the patient an anti-PD-L1 antibody optionally selected from atezolimumab (or TECENTRIQTM), avelumab (or BAVENCIO<sup>TM</sup>) and durvalumab (or IMFINZI<sup>TM</sup>); and
- (b) The cancer is selected from metastatic melanoma; renal cell carcinoma; a solid tumour; non-small cell lung cancer (NSCLC); forehead and neck squamous cell carcinoma (HNSCC); Merkel cell carcinoma; Hodgkin's lymphoma; a cancer that overexpresses PD-L1 and no mutations in EGFR or in ALK; colorectal cancer and hepatocellular carcinoma; and
- (c) The first bacteria are selected from Pseudomonas aeruginosa, Klebsiella pneumoniae, E coli, Salmonella (eg, S typhimurium), Clostridium dificile, Staphylococcus (eg, S aureus or S epidermis), Streptococcus (eg, S viridans or S thermophilus), Pneumococcus and Enterococcus bacteria.

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing

first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer:

Wherein

- (a) The immunotherapy comprises administering to the patient an anti-CD52, antibody optionally alemtuzumab (or CAMPATH<sup>TM</sup>); and
- (b) The cancer is B-cell chronic lymphocytic leukemia (CLL); and
- (c) The first bacteria are selected from Pseudomonas aeruginosa, Klebsiella pneumoniae, E coli, Salmonella (eg, S typhimurium), Clostridium dificile, Staphylococcus (eg, S aureus or S epidermis), Streptococcus (eg, S viridans or S thermophilus), Pneumococcus and 20 Enterococcus bacteria.

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site 25 comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer:

Wherein

- (a) The immunotherapy comprises administering to the 35 patient an anti-CD20 antibody, optionally of atumumab (or ARZERRA<sup>TM</sup>) or rituximab (or RITUXAN<sup>TM</sup>); and
- (b) The cancer is B-cell chronic lymphocytic leukemia (CLL) (eg, refractory CLL) or non-Hodgkin lymphoma; and
- (c) The first bacteria are selected from *Pseudomonas* aeruginosa, Klebsiella pneumoniae, E coli, Salmonella (eg, S typhimurium), Clostridium dificile, Staphylococcus (eg, S aureus or S epidermis), Streptococcus (eg, S viridans or S thermophilus), Pneumococcus and 45 Enterococcus bacteria.

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site 50 comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer:

Wherein

- (a) The immunotherapy comprises administering to the 60 patient an anti-KIR antibody, optionally lirilumab; and
- (b) The cancer is optionally acute myeloid leukaemia or squamous cell carcinoma of the head and neck (SCCHN); and
- (c) The first bacteria are selected from *Pseudomonas* 65 aeruginosa, Klebsiella pneumoniae, E coli, Salmonella (eg, S typhimurium), Clostridium dificile, Staphylococ-

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cus (eg, S aureus or S epidermis), Streptococcus (eg, S viridans or S thermophilus), Pneumococcus and Enterococcus bacteria.

A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer:

Wherein

- (a) The immunotherapy comprises administering to the patient an anti-CD19 CAR-T optionally selected from axicabtagene ciloleucel (or YESCARTA<sup>TM</sup>) and tisagenlecleucel (or KYMRIAH<sup>TM</sup>); and
- (b) The cancer is selected from a B-cell lymphoma (eg, non-Hodgkin's lymphoma (NHL); diffuse large B-cell lymphoma (DLBCL); primary mediastinal large B-cell lymphoma; or high grade B-cell lymphoma); B-cell acute lymphoblastic leukaemia (ALL); or central nervous system lymphoma; and
- (c) The first bacteria are selected from *Pseudomonas* aeruginosa, Klebsiella pneumoniae, E coli, Salmonella (eg, S typhimurium), Clostridium dificile, Staphylococcus (eg, S aureus or S epidermis), Streptococcus (eg, S viridans or S thermophilus), Pneumococcus and Enterococcus bacteria.

Alternatively, the CAR-T is an anti-CD30, CD38 or CD22 CAR-T. In an example the cancer is large B-cell lymphoma after at least two other kinds of treatment failed. In an example the cancer is high grade B-cell lymphoma and DLBCL arising from follicular lymphoma. In an example the cancer is relapsing/remitting B cell acute lymphoblastic leukaemia. In an example the cancer is primary central nervous system lymphoma.

In an example, the nuclease treats the infection without causing reduction in efficacy of the therapy. In an embodiment, "without causing reduction in efficacy of the therapy" means the efficacy of the therapy compared to a reduction caused in patients by the administration of a broad-spectrum antibiotic (or an antibiotic disclosed herein) that kills a plurality of different species, wherein the plurality comprises the first species. In an embodiment, "without causing reduction in efficacy of the therapy" means the efficacy of the therapy is reduced by no more than 70, 80, 90 or 95% compared to administration of the therapy in the absence of treatment of the pathogenic bacterial infection (or compared to therapy as typically achieved in patients suffering from the disease or condition and receiving said therapy therefor). This may be assessed, for example, by determining the duration of progression-free survival of the subject or treatment of the disease or condition, or overall survival of the subject; and/or by determining a reduction in one or more symptoms of the disease or condition.

In an example, the infection is treated completely or substantially completely. In another example, the infection is reduced (eg, by at least 80, 90 or 95% as determined by a marker of the infection or a symptom thereof). A marker may, for example, be CFUs of bacteria of the first species or strain per ml of a blood sample taken from the patient after the method has been carried out, eg, within 24 hours of that method being carried out, eg, from 1-12 hours or 1-24 hours

after carrying out the method or from 1-12 hours or 1-24 hours after administering a RNA or DNA encoding the RNA to programme the nuclease in the subject. For example, the RNA is a guide RNA and the nuclease is Cas (eg, a Cas3 or a Cas9). The reduction may be compared to a sample taken from the subject immediately prior to the commencement of the method. Alternatively, the sample may be a stool, saliva or urine sample.

In an example, the invention increases overall survival rate in a human subject (compared to median overall survival rate in humans suffering from the same cancer and receiving the same cancer therapy treatment (eg, administration of the same immune checkpoint inhibitor, such as nivolumab, pembrolizumab or another antibody disclosed herein)). In an example any composition, or other product of the invention herein is provided for use in such method of treatment.

In an example, the method is practiced on a population of human subjects and the median overall survival rate for the 20 population is 120-250% (eg, 150-200%) of the median overall survival rate in humans suffering from the same cancer and receiving the same cancer therapy treatment (eg, administration of the same immune checkpoint inhibitor, such as nivolumab, pembrolizumab or another antibody 25 disclosed herein). In an example any composition, or other product of the invention herein is provided for use in such method of treatment.

A "pathogenic bacterial infection" is a health-threatening infection of the subject, for example, a life-threatening infection. In an embodiment, a pathogenic bacterial infection is an infection requiring hospitalization or parenteral antimicrobials. The infection may be an acute bacterial infection, such as a systemic infection or a localised infection. Bacterial pathogens often cause infection in specific areas of the body. Others are generalists. A pathogenic bacterial infection is contrasted with an infection of commensal bacteria, such as commensal gut bacteria; in this case the bacteria do not cause an immediate health- or life-threatening situation.

The infection (or symptom thereof) can be any of the following:—

Bacterial vaginosis: this is caused by bacteria that change the vaginal microbiota caused by an overgrowth of bacteria that crowd out the *Lactobacilli* species that maintain healthy vaginal microbial populations.

Bacterial meningitis: this is a bacterial inflammation of the meninges, that is, the protective membranes covering the brain and spinal cord.

Bacterial pneumonia: this is a bacterial infection of the lungs.

Urinary tract infection: this is predominantly caused by bacteria. Symptoms include the strong and frequent sensation or urge to urinate, pain during urination, and 55 urine that is cloudy. The main causal agent is *Escherichia coli*. Bacteria can ascend into the bladder or kidney and causing cystitis and nephritis.

Bacterial gastroenteritis: this is caused by enteric, pathogenic bacteria. These pathogenic species are usually 60 distinct from the usually harmless bacteria of the normal gut flora. But a different strain of the same species may be pathogenic.

Bacterial skin infections: these include:

Impetigo, which is a highly contagious bacterial skin 65 infection commonly seen in children. It is caused by *Staphylococcus aureus*, and *Streptococcus pyogenes*.

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Erysipelas, which is an acute *streptococcus* bacterial infection of the deeper skin layers that spreads via with lymphatic system.

Cellulitis, which is a diffuse inflammation of connective tissue with severe inflammation of dermal and subcutaneous layers of the skin. Cellulitis can be caused by normal skin flora or by contagious contact, and usually occurs through open skin, cuts, blisters, cracks in the skin, insect bites, animal bites, burns, surgical wounds, intravenous drug injection, or sites of intravenous catheter insertion. In most cases it is the skin on the face or lower legs that is affected, though cellulitis can occur in other tissues.

In an example, the first bacteria are *Streptococcus* and the patient is suffering from chest infection, cellulitis or tonsillitis. In an example, the first bacteria are *Enterococcus* and the patient is suffering from bladder infection or septicaemia. In an example, the first bacteria are *Pseudomonas aeruginosa* and the patient is suffering from diarrhoea. In an example, the first bacteria are *E coli* and the patient is suffering from diarrhoea.

- 2. The method of Clause 1, wherein the subject is a cancer patient and the therapy is a cancer therapy.
- 3. The method of Clause 2, wherein the therapy comprises administration of a haematopoietic stem cell transplant, chemotherapeutic agent, immune checkpoint inhibitor, immune checkpoint agonist or an immune cell (eg, T-cell and/or NK cell) enhancer; adoptive cell therapy (eg, CAR-T therapy); radiation or surgery.

In an example, the therapy is immunotherapy. Examples of suitable immunotherapy are administration of adoptive cell therapy (eg, CAR-T therapy), an immune checkpoint inhibitor, an immune checkpoint agonist or an immune cell (eg, T-cell and/or NK cell) enhancer. For example, administration of an anti-CTLA4, PD-1, PD-L1, PD-L2, LAG3, 0X40, CD28, BTLA, CD137, CD27, HVEM, KIR, TIM-3, VISTA, ICOS, GITR, TIGIT or SIRPa antibody, such as administration of an antibody selected from ipilimumab (or YERVOY<sup>TM</sup>), tremelimumab, nivolumab (or OPDIVO<sup>TM</sup>), pembrolizumab (or KEYTRUDA<sup>TM</sup>), pidilizumab, BMS-936559, durvalumab and atezolizumab, or a CAR-T therapy such as axicabtagene ciloleucel (Yescarta<sup>TM</sup>) or tisagenle-cleucel (Kymriah<sup>TM</sup>).

acterial vaginosis: this is caused by bacteria that change In an example, the immune enhancer comprises an interthe vaginal microbiota caused by an overgrowth of 45 leukin-2 (IL-2) or fragment or deletion mutant thereof.

In an example, the surgery comprises the removal of necrotic or cancerous tissue.

In an example, the chemotherapy comprises administration of a platinum-containing chemotherapy drug. In an example, the chemotherapy comprises administration of gefitinib.

In an example, the therapy comprises administering Cyclophosphamide, methotrexate and 5-fluorouracil (CMF); or doxorubicin and cyclophosphamide (AC); docetaxel, doxorubicin and cyclophosphamide (TAC); or doxorubicin, bleomycin, vinblastine and dacarbazine (ABVD); or mustine, vincristine, procarbazine and prednisolone (MOPP); cyclophosphamide, doxorubicin, vincristine and prednisolone (CHOP); bleomycin, etoposide and cisplatin (BEP); epirubicin, cisplatin and 5-fluorouracil (ECF); or epirubicin, cisplatin and capecitabine (ECX); methotrexate, vincristine, doxorubicin and cisplatin (MVAC); cyclophosphamide, doxorubicin and vincristine (CAV); or 5-fluorouracil, folinic acid and oxaliplatin (FOLFOX).

In an example, the cancer is breast cancer and the therapy comprises administering CMF or AC. In an example, the cancer is Hodgkin's lymphoma and the therapy comprises

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administering TAC, ABVD or MOPP. In an example, the cancer is Non-Hodgkin's lymphoma and the therapy comprises administering CHOP. In an example, the cancer is germ cell cancer and the therapy comprises administering BEP. In an example, the cancer is stomach cancer and the 5 therapy comprises administering ECF or ECX. In an example, the cancer is bladder cancer and the therapy comprises administering MVAC. In an example, the cancer is lung cancer and the therapy comprises administering CAV. In an example, the cancer is colorectal cancer and the 10 therapy comprises administering FOLFOX.

4. The method of Clause 3, wherein the therapy is an immune checkpoint inhibitor antibody.

Optionally the antibody is an anti-CTLA4, PD-1, PD-L1, PD-L2, LAG3, 0X40, CD28, BTLA, CD137, CD27, 15 HVEM, KIR, TIM-3, VISTA, ICOS, GITR, TIGIT or SIRPa antibody. In an example, the antibody is an anti-PD-1 antibody. In an example, the antibody is an anti-PD-L1 antibody. In an example, the antibody is an anti-CTLA4 antibody.

5. The method of Clause 3, wherein the therapy is administration of an antibody selected from ipilimumab (or YERVOYTM), tremelimumab, nivolumab (or OPDIVO<sup>TM</sup>), pembrolizumab (or KEYTRUDA<sup>TM</sup>), pidilizumab, BMS-936559, durvalumab and atezoli- 25

Optionally, the antibody (eg, anti-PD-L1 antibody) is administered with an anti-CTLA4 antibody (eg, ipilimumab or tremelimumab).

In an example, the an anti-PD-1 antibody herein is 30 selected from nivolumab, pembrolizumab, pidillizumab, OPDIVO®, KEYTRUDA®, AMP-514, REGN2810, CT-011, BMS 936559, MPDL3280A and AMP-224.

In an example, the an anti-CTLA4 antibody herein is selected from tremelimumab, YERVOY® and ipilimumab. 35 In an example the therapy is administration of an anti-KIR antibody, eg, lirilumab.

In an example, the checkpoint inhibitor is selected from an inhibitor of CTLA-4, PD-1, PD-L1, PD-L2, LAG-3, BTLA, B7H3, B7H4, TIM3, KIR, or A2aR. In certain 40 aspects, the immune checkpoint inhibitor is a human programmed cell death 1 (PD-1) axis-binding antagonist. In some aspects, the PD-1 axis-binding antagonist is selected from the group consisting of a PD-1 binding antagonist, a PD-L1-binding antagonist and a PD-L2-binding antagonist. 45 the subject or patient orally, by IV injection, by subcutane-In certain aspects, the PD-1 axis-binding antagonist is a PD-1-binding antagonist. In some aspects, the PD-1-binding antagonist inhibits the binding of PD-1 to PD-L1 and/or PD-L2.

In some embodiments, the immune checkpoint inhibitor is 50 a PD-L1 antagonist such as durvalumab, also known as MEDI4736, atezolizumab, also known as MPDL3280A, or avelumab, also known as MSB00010118C. In certain aspects, the immune checkpoint inhibitor is a PD-L2 antagonist such as rHIgM12B7. In some aspects, the immune 55 checkpoint inhibitor is a LAG-3 antagonist such as IMP321 or BMS-986016. The immune checkpoint inhibitor may be an adenosine A2a receptor (A2aR) antagonist such as PBF-

In some embodiments, the antibody described herein 60 (such as an anti-PD-1 antibody, an anti-PD-L1 antibody, or an anti-PD-L2 antibody) further comprises a human or murine constant region. In a still further aspect, the human constant region is selected from the group consisting of IgG1, IgG2, IgG2, IgG3, and IgG4. In a still further specific 65 aspect, the human constant region is IgG1. In a still further aspect, the murine constant region is selected from the group

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consisting of IgG1, IgG2A, IgG2B, and IgG3. In a still further specific aspect, the antibody has reduced or minimal effector function. In a still further specific aspect, the minimal effector function results from production in prokaryotic, CHO, Cos or HEK cells. In a still further specific aspect the minimal effector function results from an "effector-less Fc mutation" or aglycosylation.

For example, the therapy comprises a haemopoietic stem cell transplant, eg, a bone marrow transplant (such as when the patient is a cancer patient, eg, a blood cancer or leukaemia patient).

For example, the therapy comprises a stem cell transplant, a skin graft, or an organ transplant, eg, a heart, liver, kidney or lung transplant.

- 6. The method of Clause 1 or 2, wherein the therapy is a tissue, organ or cell transplant.
- 7. The method of any preceding Clause, wherein the treatment of the bacterial infection is carried out simultaneously with the administration of the therapy to the subject.
- 8. The method of any one of Clauses 1 to 6, wherein the treatment of the bacterial infection is carried out immediately before administering the therapy to the subject.

In an example, the treatment of the bacterial infection is carried out no more than 7, 6, 5, 4, 3, 2, or 1 day, or 24, 12, 6, 5, 4, 3, 2, 1 or 0.5 hours before the therapy of the further disease or condition. In an example, the treatment of the bacterial infection is carried out no more than 7, 6, 5, 4, 3, 2, or 1 day, or 24, 12, 6, 5, 4, 3, 2, 1 or 0.5 hours after the therapy of the further disease or condition.

The treatment of the infection and the administration of the therapy may be carried out simultaneously or sequentially.

- 9. The method of any one of Clauses 1 to 6, wherein the treatment of the bacterial infection is carried out immediately after administering the therapy to the subject.
- 10. The method of any preceding Clause, wherein the method comprises administering to the subject a RNA (eg, a gRNA) or a nucleic acid that encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in first bacteria comprised by the subject, thereby killing the first bacteria.

The RNA or nucleic acid is, for example, administered to ous injection or by inhalation.

11. The method of any preceding Clause, comprising administering a vector (eg, phage or plasmids) to the subject, wherein the vector encodes the programmable nuclease.

The nuclease is, for example, administered to the subject or patient orally, by IV injection, by subcutaneous injection or by inhalation.

- 12. The method of any one of Clauses 1 to 10, wherein the programmable nuclease is an endogenous nuclease (eg, Cas nuclease) of the first cells.
- 13. The method of any preceding Clause, wherein the efficacy of the therapy in the presence of the programmed nuclease is greater than the efficacy of the therapy in the presence of a broad-spectrum antibiotic.

In an example, the efficacy being greater is assessed by determining the duration of progression-free survival or treatment of the disease or condition; and/or by determining a reduction in one or more symptoms of the disease or condition. For example, this determination is compared to an analogous determination in a patient suffering from the disease or condition as well as the bacterial infection and

being treated with the therapy and the antibiotic (rather than the nuclease killing of first bacteria as per the invention).

- 14. The method of any preceding Clause, wherein the efficacy of the therapy in the presence of the programmed nuclease is greater than the efficacy of the 5 therapy in the presence of an antibiotic selected from methicillin, vancomycin, linezolid, daptomycin, quinupristin, dalfopristin; teicoplanin; cephalosporin; carbapenem; fluoroquinolone; aminoglycoside; colistin; erythromycin; clindamycin; beta-lactam; macrolide; 10 amoxicillin; azithromycin; penicillin; ceftriaxone; azithromycin; ciprofloxacin; isoniazid (INH); rifampicin (RMP); amikacin; kanamycin; capreomycin; trimethoprim; itrofurantoin; cefalexin; amoxicillin; metronidazole (MTZ); cefixime; tetracycline; and 15 meropenem.
- 15. The method of any preceding Clause, wherein the first bacteria is selected from (i) Staphylococcus aureus that is resistant to an antibiotic selected from methicillin, vancomycin, linezolid, daptomycin, quinupristin, dal- 20 fopristin and teicoplanin; (ii) Pseudomonas aeuroginosa that is resistant to an antibiotic selected from cephalosporins, carbapenems, fluoroquinolones, aminoglycosides and colistin; (iii) Klebsiella species that is resistant to carbapenem; (iv) Streptoccocus spe- 25 cies that is resistant to an antibiotic selected from erythromycin, clindamycin, beta-lactam, macrolide, amoxicillin, azithromycin and penicillin; (v) Salmonella species that is resistant to an antibiotic selected from ceftriaxone, azithromycin and ciprofloxacin; (vi) 30 Shigella species that is resistant to ciprofloxacin or azithromycin; (vii) Mycobacterium tuberculosis that is resistant to an antibiotic selected from Resistance to isoniazid (INH), rifampicin (RMP), fluoroquinolone, amikacin, kanamycin, capreomycin and azithromycin; 35 (viii) Enterococcus species that is resistant to vancomycin; (ix) Enterobacteriaceae species that is resistant to an antibiotic selected from cephalosporin and carbapenem; (x) E coli that is resistant to an antibiotic selected from trimethoprim, itrofurantoin, cefalexin 40 and amoxicillin; (xi) Clostridium species that is resistant to metronidazole (MTZ), fluoroquinolone or carbapenem; (xii) Neisseria gonnorrhoea that is resistant to an antibiotic selected from cefixime, ceftriaxone, azithromycin and tetracycline; (xiii) Acinetoebacter 45 baumannii that is resistant to an antibiotic selected from beta-lactam, meropenem and carbapenem; and (xiv) Campylobacter species that is resistant to ciprofloxacin or azithromycin.
- 16. The method of any preceding Clause, wherein the 50 treatment of the infection treats or prevents in the subject a condition selected from vaginosis, meningitis, pneumonia, urinary tract infection, cystitis, nephritis, gastroenteritis, a skin infection, impetigo, erysipelas, dental infection and cellulitis.
- 17. The method of any preceding Clause, wherein the treatment of the infection treats or prevents septicaemia or sepsis in the subject.

In an example, the infection is a bloodstream infection.

18. The method of any preceding Clause, wherein the further disease or condition is a cancer; autoimmune disease or condition; viral infection or GI tract disease or condition. In an example, the cancer is metastatic. In an example, the cancer is melanoma. In an example, the cancer is a solid tumour with mismatch repair deficiency or microsatellite instability. In an example, the cancer is NSCLC. In an example, the cancer is

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HNSCC. In an example, the cancer is Hodgkin's lymphoma. In an example, the cancer is urothelial cancer. In an example, the cancer is lung cancer. In an example, the cancer is head and neck cancer. In an example, the cancer is head cancer. In an example, the cancer is neck cancer.

In an example, the viral infection is a HIV, CMV or RSV infection.

- 19. The method of any preceding Clause, wherein the subject comprises bacteria (second bacteria) of one or more strains or species that are different to the first strain or species, wherein the genomes of the second bacteria do not comprise the target site, wherein the genomes of the second bacteria are not cut by the programmed nuclease in the subject, whereby second bacteria survive in the presence of the programmed nuclease in the patient; and wherein the therapy is efficacious in the presence of the second bacteria.
- 20. The method of Clause 19, wherein reduction in the second bacteria in patients (eg, in the gut microbiome) is associated with reduced efficacy of the therapy.

Optionally, the therapy is efficacious in the presence of the second bacteria in the gut of the subject.

Optionally, the first and/or second bacteria are present in the gut of the subject immediately prior to carrying out the method.

Optionally, the first and/or second bacteria are present in the blood of the subject immediately prior to carrying out the method

Optionally, the first bacteria are present in the blood of the subject and the second bacteria are present in the gut of the subject immediately prior to carrying out the method.

Optionally, the first bacteria are present in the gut of the subject and the second bacteria are present in the blood of the subject immediately prior to carrying out the method.

Optionally, first bacteria in the blood of the subject is killed.

Optionally, the bacteria are gram positive bacteria. Optionally, the bacteria are gram negative bacteria.

Optionally, the first and second bacteria are capable of being killed by the same antibiotic. Optionally, the method does not comprise administering the antibiotic to the subject. In an example, the antibiotic is selected from methicillin, vancomycin, linezolid, daptomycin, quinupristin, dalfopristin; teicoplanin; cephalosporin; carbapenem; fluoroquinolone; aminoglycoside; colistin; erythromycin; clindamycin; beta-lactam; macrolide; amoxicillin; azithromycin; penicillin; ceftriaxone; azithromycin; ciprofloxacin; isoniazid (INH); rifampicin (RMP); amikacin; kanamycin; capreomycin; trimethoprim; itrofurantoin; cefalexin; amoxicillin; metronidazole (MTZ); cefixime; tetracycline; and meropenem. In an example, the antibiotic is selected from Aminoglycosides, Ampicillin, Amoxicillin, Amoxicillin or 55 clavulanic acid, Carbapenems (e.g. imipenem), Piperacillin or tazobactam, Quinolones (e.g. ciprofloxacin), Tetracyclines, Chloramphenicol, Ticarcillin, Trimethoprim or sulfamethoxazole, penicillin, streptomycin, oxytetracycline and potentiated sulfonamides. In an example, the first bacteria are resistant to an antibiotic selected from Aminoglycosides, Ampicillin, Amoxicillin, Amoxicillin or clavulanic acid, Carbapenems (e.g. imipenem), Piperacillin or tazobactam, Quinolones (e.g. ciprofloxacin), Tetracyclines, Chloramphenicol, Ticarcillin, Trimethoprim or sulfamethoxazole, penicillin, streptomycin, oxytetracycline and potentiated sulfonamides. In an alternative, the antibiotic is selected from a beta-lactam, fluoroquinolone and macrolide.

Optionally, the first and second bacteria are bacteria of the same species, but are different strains of the species.

Optionally, the first and second bacteria are bacteria of the same genus, but are bacteria of different species of the genus.

Optionally, the first and second bacteria are bacteria of the same family, but are bacteria of different genera of the family.

Optionally, the first and second bacteria are gram positive bacteria

Optionally, the first and second bacteria are gram-negative bacteria.

Optionally, the therapy is efficacious in the presence of the second bacteria.

Optionally, reduction in the second bacteria in patients is associated with reduced efficacy of the therapy. Optionally, reduction in the second bacteria in patients reduces efficacy of the therapy.

Optionally, the presence of the second bacteria in patients 20 is associated with enhanced efficacy of the therapy. Optionally, the presence of the second bacteria in patients enhances efficacy of the therapy. For example, enhanced efficiency is efficiency compared to therapy in the absence or a reduced presence of the second bacteria, such as in the presence of 25 an antibiotic that kills the second bacteria.

In an example, the therapy is efficacious in the presence of the second bacteria, wherein the disease or condition (or a symptom thereof) is reduced in the subject by at least 20, 30, 40, 50, 60, 70, 80, 90 or 95%. In an example, the therapy 30 is efficacious in the presence of the second bacteria, wherein the progression of the disease or condition (or a symptom thereof) is reduced in the subject by at least 20, 30, 40, 50, 60, 70, 80, 90 or 95%. In an example, the therapy is efficacious in the presence of the second bacteria, wherein 35 disease-free progression of the disease or condition (or a symptom thereof) is reduced in the subject by at least 20, 30, 40, 50, 60, 70, 80, 90 or 95%. In an example, the therapy is efficacious in the presence of the second bacteria, wherein the duration of the disease or condition (or a symptom 40 thereof) is reduced in the subject by at least 20, 30, 40, 50, 60, 70, 80, 90 or 95%. In an example, the therapy is efficacious in the presence of the second bacteria, wherein the severity of the disease or condition (or a symptom thereof) is reduced in the subject by at least 20, 30, 40, 50, 45 60, 70, 80, 90 or 95%. In an example, the therapy is efficacious in the presence of the second bacteria, wherein the disease or condition (or a symptom thereof) is reduced for at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 21 or 28 days or for at least 1, 2, 3, 4 5, 6 or 12 months in the 50 patient by at least 20, 30, 40, 50, 60, 70, 80, 90 or 95%. In an example, the therapy is efficacious in the presence of the second bacteria, wherein the disease or condition (or a symptom thereof) is treated for at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 21 or 28 days or for at least 1, 2, 3, 4 55 5, 6 or 12 months in the patient by at least 20, 30, 40, 50, 60, 70, 80, 90 or 95%. In an example, the therapy is efficacious in the presence of the second bacteria, wherein the disease or condition (or a symptom thereof) is undetectable for at least 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 21 or 28 days 60 or for at least 1, 2, 3, 4 5, 6 or 12 months in the patient by at least 20, 30, 40, 50, 60, 70, 80, 90 or 95%.

21. The method of Clause 19 or 20, wherein the second bacteria are selected from the group consisting of Akkermansia, Alistipes, Bacteroides, Barnesiella, Bifidobacterium, Clostridium, Collinsella, Enterococcus, Fusobacterium, Lactobacillus, Propionibacterium,

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Ruminococcus, Segmented filamentous bacteria (SFB); Veillonella, Prevotella, Escherichia and Streptococcus bacteria.

In an example the second bacteria that produce short chain fatty acids (eg, butyrate-producing bacteria). In particular aspects, the species of bacteria produce butyrate. For example, the second bacteria are Clostridiales. The Clostridiales bacteria may be substantially or include bacteria in spore form. In particular aspects, the second bacteria are of the family Ruminococcaceae, Christensenellaceae, Clostridiaceae or Coriobacteriacease. In some embodiments, the Clostridiales (eg, Clostridium) bacteria comprise a first family and a second family. In some embodiments, the first family is selected from the group consisting of Ruminococcaceae, Christensenellaceae, Clostridiaceae and Coriobacteriacease, and the second family is not identical to the first family. In an example, the second bacteria are Faecalibacterium prausnitzii, Ruminococcus albus, Ruminococcus bromii, Ruminococcus callidus, Ruminococcus flavefaciens, Ruminococcus champanellensis, Ruminococcus faecis, Ruminococcus gauvreauii, Ruminococcus gnavus, Ruminococcus hansenii, Ruminococcus hydrogenotrophicus, Ruminococcus lactaris, Ruminococcus luti, Ruminococcus obeum, Ruminococcus palustris, Ruminococcus pasteurii, Ruminococcus productus, Ruminococcus schinkii, Ruminococcus torques, Subdoligranulum variabile, Butyrivibrio fibrisolvens, Roseburia intestinalis, Anaerostipes caccae, Blaufia obeum, Eubacterium nodatum or Eubacterium oxidoreducens. In particular aspects, the second bacteria are Faecalibacterium prausnitzii. In an example the second bacteria are Firmicutes.

In certain embodiments, the first bacteria are Bacteroidia or Prevotellaceae, eg, Bacteroidetes or *Bacteroides*.

In an embodiment, the treatment results in or maintains a microbiome (eg, gut and/or blood microbiome) of the subject, which is beneficial for the immune checkpoint inhibition or other therapy. In an example, the microbiome comprises a high relative abundance of one or more bacterial species from the phylum Firmicutes, class Clostridia, order Clostridiales, family Ruminococcaceae, genus Ruminococcus, genus Hydrogenoanaerobacterium, genus Faecalibacterium, phylum Actinobacteria, class Coriobacteriia, order Coriaobacteriales, family Coriobacteriaceae, domain Archaea, phylum Cyanobacteria, phylum Euryarchaeota or famly Christensenellaceae. Additionally or alternatively, the microbiome comprises a low relative abundance of bacteria from the genus Dialister, family Veillonellaceae, phylum Bacteroidetes, class Bacteroida, order Bacteroidales or family Prevotellaceae. Accordingly, a favorable microbial profile would have a higher relative abundance of one or more bacterial species from the phylum Firmicutes, class Clostridia, order Clostridiales, family Ruminococcaceae, genus Ruminococcus, genus Hydrogenoanaerobacterium, phylum Actinobacteria, class Coriobacteria, order Coriaobacteriales, family Coriobacteriaceae, domain Archaea, phylum Cyanobacteria, phylum Euryarchaeota or family Christensenellaceae, and/or has a decreased abundance of one or more bacterial species from genus Dialister, family Veillonellaceae, phylum Bacteroidetes, class Bacteroida, order Bacteroidales and/or family Prevotellaceae.

For example, the microbiome comprises a higher relative abundance of Firmicutes compared to Bacteroidetes, Bacteroida, Bacteroidales or Prevotellaceae. For example, the microbiome comprises a higher relative abundance of Firmicutes compared to Bacteroidetes, Bacteroida, Bacteroidales and Prevotellaceae.

Optionally, the second bacteria are selected from the group consisting of Akkermansia muciniphila; Alistipes shahii; Bacteroides fragilis; Bacteroides uniformis; Barnesiella intestinihominis; Bacteroides dorei; Bifidobacterium adolescentis; Bifidobacterium breve; Bifidobacterium longum; 5 Clostridium orbiscindens; Clostridium novyi; Clostridium perfringens; Collinsella aerofaciens; Enterococcus hirae; Fusobacterium nucleatum; Lactobacillus casei Shirota; L. casei AO47; Lactobacillus rhamnosus; Propionibacterium granulosum; Ruminococcus gnavus; Segmented filamentous bacteria (SFB); Veillonella; Lactobacilli; Bacteroides; Clostridia; Prevotella; E. coli Nissle; Lactobacillus plantarum; Lactobacillus delbrueckii (eg, subsp. Bulgaricus); Lactobacillus paracasei; Lactobacillus acidophilus; Bifidobacterium infantis; and Streptococcus salivarius (eg., subsp. 15 Thermophilus). See "The microbiome in cancer immunotherapy: Diagnostic tools and therapeutic strategies"; Laurence Zitvogel et al; Science 23 Mar. 2018: Vol. 359, Issue 6382, pp. 1366-1370; DOI: 10.1126/science.aar6918.

In an example, the second bacteria are commensal bacteria in humans.

In an example, the first bacteria are comprised by gut microbiota, skin microbiota, oral cavity microbiota, throat microbiota, hair microbiota, armpit microbiota, vaginal microbiota, rectal microbiota, anal microbiota, ocular microbiota, liver microbiota, tiver microbiota, kidney microbiota, genital microbiota, penile microbiota, scrotal microbiota, mammary gland microbiota, ear microbiota, urethra microbiota, labial microbiota, organ microbiota or dental microbiota.

In an example, the second bacteria are comprised by gut microbiota, skin microbiota, oral cavity microbiota, throat microbiota, hair microbiota, armpit microbiota, vaginal microbiota, rectal microbiota, anal microbiota, ocular microbiota, liver microbiota, tongue microbiota, lung microbiota, penile microbiota, scrotal microbiota, genital microbiota, penile microbiota, urethra microbiota, labial microbiota, organ microbiota or dental microbiota.

phage, phagemid plasm transposon.

The phage are capable of presence of a helper phage.

32. A pharmaceutical or nucleic acid vector (or a microbiota, organ microbiota or dental microbiota.

In an example, the first and/or second bacteria are blood- 40 borne bacteria.

22. The method of any preceding Clause, wherein the first bacteria are selected from the group consisting Staphylococcus, Streptococcus, Enterococcus, Helicobacter, Legionella, Heamophilus, Ghonnorhea, Acinetobacter, 45 Escherichia, Klebsiella, Pseudomonas or Stenotrophomonas bacteria.

H pylori has been implicated in gastric cancer and gastric ulcers. Thus, in an example, the first bacteria are H pylori and optionally the disease is a cancer, such as gastric cancer. 50 In an embodiment, the therapy is chemotherapy or therapy with an immune checkpoint inhibitor (eg, an antibody). In an example, the first bacteria are H pylori and the disease is gastric ulcer(s). In an embodiment, triple therapy for gastric ulcers is administered to the subject.

In an example, the first bacteria are Gram-negative bacteria and optionally the infection is a blood infection. In an example, the first bacteria are selected from *E. coli, P. aeruginosa* and *K pneumoniae*, and optionally the infection is a blood infection.

23. The method of Clause 22, wherein the first bacteria are selected from the group consisting of E coli (eg, EHEC E coli), C dificile, V cholera, Staphylococcus (eg, S aureus or MRSA), Streptococcus pyogenes, Helicobacter pylori, Acinetobacter baumannii, Legionella, 65 Pseudomonas aeruginosa and Klebsiella pneumoniae bacteria.

In an example, the subject has been administered an immunosuppressant drug, or is on a course of an immunosuppressant drug, eg, a steroid, such as a corticosteroid.

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- 24. A programmable nuclease for use in the method of any preceding Clause.
- 25. The method or nuclease of any preceding Clause, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9), a meganuclease, a TALEN (Transcription activator-like effector nuclease) or zinc finger nuclease.
- 26. A CRISPR/Cas system comprising a nuclease according to Clause 24 or 25 for use in the method of any one of Clauses 1 to 23, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9) and the system comprises one or more guide RNAs (gRNAs) or DNA encoding one or more guide RNAs, wherein each guide RNA is capable of programming the Cas nuclease to cut a target site comprised by the genomes of first bacteria.
- 27. A guide RNA or a DNA encoding a guide RNA for use in the system of Clause 26.
- 28. A guide RNA or a DNA encoding a guide RNA for use in the method of treating a pathogenic bacterial infection according to any one of Clauses 1 to 23, wherein the guide RNA is capable of programming the nuclease, wherein the nuclease is a Cas nuclease (eg, a Cas9, Cas3, Cas13, CasX, CasY or Cpf1 nuclease).
- 29. A nucleic acid vector comprising the guide RNA or DNA recited in any one of Clauses 26 to 28.
- 30. A nucleic acid vector encoding the nuclease of Clause 24 or 25 and optionally the guide RNA of Clause 29.
- 31. The vector of Clause 29 or 30 wherein the vector is a phage, phagemid plasmid (eg, conjugative plasmid) or transposon.

The phage are capable of infecting first bacteria and the phagemids are capable of producing such phage in the presence of a helper phage.

- 32. A pharmaceutical composition comprising a first nucleic acid vector (or a plurality thereof) encoding the nuclease of Clause 24 or 25 and a second nucleic acid vector (or a plurality thereof) encoding the guide RNA of Clause 29, the composition further comprising a pharmaceutically acceptable diluent, excipient or carrier.
- 33. A pharmaceutical composition comprising the CRISPR/Cas system of claim 26 and a pharmaceutically acceptable diluent, excipient or carrier.
- 34. A pharmaceutical composition comprising the vector of claim 31 and a pharmaceutically acceptable diluent, excipient or carrier.

Preventing a disease or condition herein may, for example, be reducing the risk of the disease or condition in the subject or patient.

In an alternative, instead of first bacteria, the infection is caused by first archaea and in this embodiment all of the features of the method and other configurations of the invention relating to killing first bacteria instead relate mutatis mutandis to killing first archaea.

In an embodiment, the method comprises carrying out the method of treating an acute microbial infection as described herein, and thus features of that method as described herein are combinable with the present method of treating a pathogenic bacterial infection (ie, where the pathogenic bacterial infection is the acute microbial infection in the first method). In an embodiment, the method comprises carrying out the method of durably treating a microbial infection as described herein, and thus features of that method as described herein are combinable with the present method of treating a pathogenic bacterial infection (ie, where the

pathogenic bacterial infection is the microbial infection in the first method). Any of the optional features of the first method herein may apply mutatis mutandis to the present method of treating a pathogenic bacterial infection. Aspects:—

Thus, the invention provides the following Aspects, which are optional features of Clauses above:—

- 1. The method of any one of Clauses 1-23, wherein the infection is reduced at least 100-fold by the first 30 minutes of carrying out step (b). Optionally, the infec- 10 tion is reduced at least 1000-fold by the first 30 minutes of carrying out step (b). Optionally, the reduction in infection persists for 30 minutes immediately after the first 30 minutes of carrying out step (b). For example, the reduction can be assessed by determining the dif- 15 ference in the number of bacteria of the first species or strain in (i) a sample taken from the subject (eg, a blood sample) immediately before commencement of the method and (ii) a sample (of the same type as the sample of (i), eg, a blood sample) taken from the 20 subject at 30 minutes of the treatment. For example, the samples may be assessed for the difference in colony forming units (CFU)/ml sample, eg, when the samples have been plated on agar in respective petri dishes and incubated under identical conditions. Another example 25 may use microscopic counting of bacteria in samples, or other routine methods know to the skilled addressee.
- 2. The method of any one of Clauses 1-23, wherein blood infection of the subject by the first bacteria is reduced at least 100- or 1000-fold by the first 30 minutes of 30 carrying out step (b).
- 3. The method of Aspect 2, wherein the blood is infected with from 10<sup>5</sup> to 10<sup>12</sup> (eg, 10<sup>7</sup> to 10<sup>12</sup>) CFU/ml of the first bacteria immediately before the treatment.
- 4. The method of any one of Clauses 1-23 or any 35 preceding Aspect, wherein the method comprises administering to the subject a nucleic acid (eg, a RNA) and nuclease, wherein the nucleic acid complexes with the nuclease to program the nuclease to cut the target site in the first bacteria comprised by the subject.
- The method of Aspect 4, wherein the nuclease is administered simultaneously or sequentially with the nucleic acid to the subject.
- The method of Aspect 4, wherein the subject comprises the nuclease prior to administration of the nucleic acid 45 to the subject.
- 7. The method of any one of Aspects 4 to 6, wherein a plurality phage are administered to the subject, wherein each phage comprises a copy of the nucleic acid, wherein the phage infect first bacteria comprised by the 50 subject to deliver thereto the nucleic acid.
- 8. The method of Aspect 7, wherein the ratio of administered phage: first bacteria comprised by the subject is from 10 to 150. For example, the ratio is from 10 to 100, ie, a multiplicity of infection (MOI) of from 10 to 55 100.

The ratio can be determined, for example, using a sample (eg, a blood or gut sample) from a human or animal subject immediately before the treatment and determining the number of bacteria per ml of blood or gut sample. The amount 60 of phage to be administered can then be worked out according to the determination using the sample.

- The method of any one of Clauses 1-23 or any preceding Aspect, wherein the infection is an infection of the lungs, brain, skin, abdomen or urinary tract.
- 10. The method of any one of Clauses 1-23 or any preceding Aspect, wherein the subject has undergone

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- surgery, is on an immunosuppressant medication, suffering from burns, suffering from diabetes, suffering from cancer or is suffering from a chronic disease.
- 11. The method of any one of Clauses 1-23 or any preceding Aspect, wherein the subject is a human over 65 years of age or is a paediatric patient.
- 12. The method of any one of Clauses 1-23 or any preceding Aspect, wherein the method treats or prevents sepsis in the subject.
- 13. The method of Clause 12, wherein at the start of the treatment, the subject (eg, a human) has a temperature of <36° C. or >38° C.; a heart rate of >90/min, a respiratory rate of >20 breaths/min or PaCO<sub>2</sub><4.3 kPa; and white blood cell count of <4000/mm³ or >12,000/mm³
- 14. The method of Clause 12 or 13, wherein at the start of the treatment, the subject (eg, a human) has presence of two or more of the following: abnormal body temperature, abnormal heart rate, abnormal respiratory rate, abnormal blood gas and abnormal white blood cell count

### Immune Checkpoint Modulation

Immune checkpoints of the invention either turn up a signal (e.g., co-stimulatory molecules) or turn down a signal. Inhibitory immune checkpoint molecules that may be targeted by immune checkpoint modulation in the invention include adenosine A2A receptor (A2AR), B7-H3 (also known as CD276), B and T lymphocyte attenuator (BTLA), cytotoxic T-lymphocyte-associated protein 4 (CTLA-4, also known as CD 152), indoleamine 2,3-dioxygenase (IDO), killer-cell immunoglobulin (KIR), lymphocyte activation gene-3 (LAG3), programmed death 1 (PD-1), T-cell immunoglobulin domain and mucin domain 3 (TIM-3) and V-domain Ig suppressor of T cell activation (VISTA). In particular, the immune checkpoint inhibitors target the PD-1 axis and/or CTLA-4.

The immune checkpoint inhibitors may be drugs such as small molecules, recombinant forms of ligand or receptors, or, antibodies, such as human antibodies (e.g., WO2015016718; Pardoll, Nat Rev Cancer, 12(4): 252-64, 2012; both incorporated herein by reference). Known inhibitors of the immune checkpoint proteins or analogs thereof may be used, in particular chimerised, humanised or human forms of antibodies may be used. As the skilled person will know, alternative and/or equivalent names may be in use for certain antibodies mentioned in the present disclosure. Such alternative and/or equivalent names are interchangeable in the context of the present invention. For example it is known that lambrolizumab is also known under the alternative and equivalent names MK-3475 and pembrolizumab.

It is contemplated that any of the immune checkpoint inhibitors that are known in the art to stimulate immune responses may be used. This includes inhibitors that directly or indirectly stimulate or enhance antigen-specific T-lymphocytes. These immune checkpoint inhibitors include, without limitation, agents targeting immune checkpoint proteins and pathways involving PD-L2, LAG3, BTLA, B7H4 and TIM3. For example, LAG3 inhibitors known in the art include soluble LAG3 (IMP321, or LAG3-Ig disclosed in WO2009044273) as well as mouse or humanized antibodies blocking human LAG3 (e.g., IMP701 disclosed in WO2008132601), or fully human antibodies blocking human LAG3 (such as disclosed in EP 2320940). Another example is provided by the use of blocking agents towards BTLA, including without limitation antibodies blocking human BTLA interaction with its ligand (such as 4C7 disclosed in WO2011014438). Yet another example is pro-

vided by the use of agents neutralizing B7H4 including without limitation antibodies to human B7H4 (disclosed in WO 2013025779, and in WO2013067492) or soluble recombinant forms of B7H4 (such as disclosed in US20120177645). Yet another example is provided by 5 agents neutralizing B7-H3, including without limitation antibodies neutralizing human B7-H3 (e.g. MGA271 disclosed as BRCA84D and derivatives in US 20120294796). Yet another example is provided by agents targeting TIM3, including without limitation antibodies targeting human 10 TIM3 (e.g. as disclosed in WO 2013006490 A2 or the anti-human TIM3, blocking antibody F38-2E2 disclosed by Jones et ah, J Exp Med. 2008; 205(12):2763-79). A. PD-1 Axis Antagonists

T cell dysfunction or anergy occurs concurrently with an 15 induced and sustained expression of the inhibitory receptor, programmed death 1 polypeptide (PD-1). Thus, therapeutic targeting of PD-1 and other molecules which signal through interactions with PD-1, such as programmed death ligand 1 (PD-L1) and programmed death ligand 2 (PD-L2) is pro- 20 vided herein. PD-L1 is overexpressed in many cancers and is often associated with poor prognosis (Okazaki T et ah, Intern. Immun 2007 19(7):813). Thus, improved methods of treating cancer by inhibiting the PD-L1/PD-1 interaction in combination with modulating the microbiome is provided 25 herein.

For example, PD-1 axis binding antagonists include a PD-1 binding antagonist, a PD-L1 binding antagonist and a PD-L2 binding antagonist. Alternative names for "PD-1" include CD279 and SLEB2. Alternative names for "PD-L1" 30 include B7-H1, B7-4, CD274, and B7-H. Alternative names for "PD-L2" include B7-DC, Btdc, and CD273. In some embodiments, PD-1, PD-L1, and PD-L2 are human PD-1, PD-L1 and PD-L2.

In some embodiments, the PD-1 binding antagonist is a 35 molecule that inhibits the binding of PD-1 to its ligand binding partners. In a specific aspect, the PD-1 ligand binding partners are PD-L1 and/or PD-L2. In another embodiment, a PD-L1 binding antagonist is a molecule that inhibits the binding of PD-L1 to its binding partners. In a 40 specific aspect, PD-L1 binding partners are PD-1 and/or B7-1. In another embodiment, the PD-L2 binding antagonist is a molecule that inhibits the binding of PD-L2 to its binding partners. In a specific aspect, a PD-L2 binding partner is PD-1. The antagonist may be an antibody, an 45 antigen binding fragment thereof, an immunoadhesin, a fusion protein, or oligopeptide. Exemplary antibodies are described in U.S. Pat. Nos. 8,735,553, 8,354,509, and 8,008, 449, all incorporated herein by reference. Other PD-1 axis antagonists for use in the methods provided herein are 50 known in the art such as described in U.S. Patent Application US20140294898, US2014022021, US20110008369, all incorporated herein by reference.

In some embodiments, the PD-1 binding antagonist is an antibody, or a chimeric antibody). In some embodiments, the anti-PD-1 antibody is selected from the group consisting of nivolumab, pembrolizumab, and CT-011. In some embodiments, the PD-1 binding antagonist is an immunoadhesin (e.g., an immunoadhesin comprising an extracellular or 60 PD-1 binding portion of PDL1 or PD-L2 fused to a constant region (e.g., an Fc region of an immunoglobulin sequence). In some embodiments, the PD-1 binding antagonist is AMP-224. Nivolumab, also known as MDX-1106-04, MDX-1106, ONO-4538, BMS-936558, and OPDIVO® is an anti-PD-1 65 antibody described in WO2006/121168. Pembrolizumab, also known as MK-3475, Merck 3475, lambrolizumab,

KEYTRUDA® and SCH-900475, is an anti-PD-1 antibody described in WO2009/114335. CT-011, also known as hBAT or hBAT-1, is an anti-PD-1 antibody described in WO2009/ 101611. AMP-224, also known as B7-DCIg, is a PD-L2-Fc fusion soluble receptor described in WO2010/027827 and WO2011/066342. Additional PD-1 binding antagonists include pidilizumab, also known as CT-011, MEDI0680, also known as AMP-514, and REGN2810.

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In some embodiments, the immune checkpoint inhibitor is a PD-L1 antagonist such as durvalumab, also known as MEDI4736, atezolizumab, also known as MPDL3280A, or avelumab, also known as MSB00010118C. In certain aspects, the immune checkpoint inhibitor is a PD-L2 antagonist such as rHIgM12B7. In some aspects, the immune checkpoint inhibitor is a LAG-3 antagonist such as, but not limited to, IMP321, and BMS-986016. The immune checkpoint inhibitor may be an adenosine A2a receptor (A2aR) antagonist such as PBF-509.

In some embodiments, any antibody described herein (such as an anti-PD-1 antibody, an anti-PD-L1 antibody, or an anti-PD-L2 antibody) further comprises a human or murine constant region. In a still further aspect, the human constant region is selected from the group consisting of IgG1, IgG2, IgG2, IgG3, and IgG4. In a still further specific aspect, the human constant region is IgG1. In a still further aspect, the murine constant region is selected from the group consisting of IgG1, IgG2A, IgG2B, and IgG3. In a still further specific aspect, the antibody has reduced or minimal effector function. In a still further specific aspect, the minimal effector function results from production in prokaryotic cells. In a still further specific aspect the minimal effector function results from an "effector-less Fc mutation" or aglycosylation. Glycosylation of antibodies is typically either N-linked or O-linked. N-linked refers to the attachment of the carbohydrate moiety to the side chain of an asparagine residue. The tripeptide sequences asparagine-Xserine and asparagine-X-threonine, where X is any amino acid except proline, are the recognition sequences for enzymatic attachment of the carbohydrate moiety to the asparagine side chain. Thus, the presence of either of these tripeptide sequences in a polypeptide creates a potential glycosylation site. O-linked glycosylation refers to the attachment of one of the sugars N-aceylgalactosamine, galactose, or xylose to a hydroxy amino acid, most commonly serine or threonine, although 5-hydroxyproline or 5-hydroxy lysine may also be used. Removal of glycosylation sites form an antibody is conveniently accomplished by altering the amino acid sequence such that one of the above-described tripeptide sequences (for N-linked glycosylation sites) is removed. The alteration may be made by substitution of an asparagine, serine or threonine residue within the glycosylation site another amino acid residue (e.g., glycine, alanine or a conservative substitution).

The antibody or antigen binding fragment thereof, may be anti-PD-1 antibody {e.g., a human antibody, a humanized 55 made using methods known in the art, for example, by a process comprising culturing a host cell containing nucleic acid encoding any of the previously described anti-PD-L1, anti-PD-1, or anti-PD-L2 antibodies or antigen-binding fragment in a form suitable for expression, under conditions suitable to produce such antibody or fragment, and recovering the antibody or fragment. B. CTLA-4

> Another immune checkpoint that can be targeted in the methods provided herein is the cytotoxic T-lymphocyteassociated protein 4 (CTLA-4), also known as CD152. The complete cDNA sequence of human CTLA-4 has the Genbank accession number L15006. CTLA-4 is found on the

surface of T cells and acts as an "off switch when bound to CD80 or CD86 on the surface of antigen-presenting cells. CTLA4 is a member of the immunoglobulin superfamily that is expressed on the surface of Helper T cells and transmits an inhibitory signal to T cells. CTLA4 is similar to 5 the T-cell co-stimulatory protein, CD28, and both molecules bind to CD80 and CD86, also called B7-1 and B7-2 respectively, on antigen-presenting cells. CTLA4 transmits an inhibitory signal to T cells, whereas CD28 transmits a stimulatory signal. Intracellular CTLA4 is also found in 10 regulatory T cells and may be important to their function. T cell activation through the T cell receptor and CD28 leads to increased expression of CTLA-4, an inhibitory receptor for B7 molecules.

In some embodiments, the immune checkpoint inhibitor is an anti-CTLA-4 antibody (e.g., a human antibody, a humanized antibody, or a chimeric antibody), an antigen binding fragment thereof, an immunoadhesin, a fusion protein, or oligopeptide.

Anti-human-CTLA-4 antibodies (or VH and/or VL 20 domains derived therefrom) suitable for use in the present methods can be generated using methods well known in the art. Alternatively, art recognized anti-CTLA-4 antibodies can be used. For example, the anti-CTLA-4 antibodies disclosed in: U.S. Pat. No. 8,119,129, WO 01/14424, WO 25 98/42752; WO 00/37504 (CP675, 206, also known as tremelimumab; formerly ticilimumab), U.S. Pat. No. 6,207,156; Hurwitz et al, 1998; can be used in the methods disclosed herein. The teachings of each of the aforementioned publications are hereby incorporated by reference. Antibodies that compete with any of these art-recognized antibodies for binding to CTLA-4 also can be used. For example, a humanized CTLA-4 antibody is described in International Patent Application No. WO2001014424, WO2000037504, and U.S. Pat. No. 8,017,114; all incorporated herein by 35

An exemplary anti-CTLA-4 antibody is ipilimumab (also known as 10D1, MDX-010, MDX-101, and Yervoy®) or antigen binding fragments and variants thereof (see, e.g., WO01/14424). In other embodiments, the antibody com- 40 prises the heavy and light chain CDRs or VRs of ipilimumab. Accordingly, in one embodiment, the antibody comprises the CDR1, CDR2, and CDR3 domains of the VH region of ipilimumab, and the CDR1, CDR2 and CDR3 domains of the VL region of ipilimumab. In another embodi- 45 ment, the antibody competes for binding with and/or binds to the same epitope on CTLA-4 as the above-mentioned antibodies. In another embodiment, the antibody has at least about 90% variable region amino acid sequence identity with the above-mentioned antibodies (e.g., at least about 50 90%, 95%, or 99% variable region identity with ipilimumab).

Other molecules for modulating CTLA-4 include soluble CTLA-4 ligands and receptors such as described in U.S. Pat. Nos. 5,844,905, 5,885,796 and International Patent Application Nos. WO1995001994 and WO1998042752; all incorporated herein by reference, and immunoadhesins such as described in U.S. Pat. No. 8,329,867, incorporated herein by reference.

### C. Killer Immunoglobulin-Like Receptor (KIR)

Another immune checkpoint inhibitor for use in the present invention is an anti-KIR antibody. Anti-human-KIR antibodies (or VH/VL domains derived therefrom) suitable for use in the present methods can be generated using methods well known in the art.

Alternatively, art recognized anti-KIR antibodies can be used. The anti-KIR antibody can be cross-reactive with

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multiple inhibitory KIR receptors and potentiates the cytotoxicity of NK cells bearing one or more of these receptors. For example, the anti-KIR antibody may bind to each of KIR2D2DL1, KIR2DL2, and KIR2DL3, and potentiate NK cell activity by reducing, neutralizing and/or reversing inhibition of NK cell cytotoxicity mediated by any or all of these KIRs. In some aspects, the anti-KIR antibody does not bind KIR2DS4 and/or KIR2DS3. For example, monoclonal antibodies 1-7F9 (also known as IPH2101), 14F1, 1-6F1 and 1-6F5, described in WO 2006/003179, the teachings of which are hereby incorporated by reference, can be used. Antibodies that compete with any of these art-recognized antibodies for binding to KIR also can be used. Additional art-recognized anti-KIR antibodies which can be used include, for example, those disclosed in WO 2005/003168, WO 2005/009465, WO 2006/072625, WO 2006/072626, WO 2007/042573, WO 2008/084106, WO 2010/065939, WO 2012/071411 and WO 2012/160448.

An exemplary anti-KIR antibody is lirilumab (also referred to as BMS-986015 or IPH2102). In other embodiments, the anti-KIR antibody comprises the heavy and light chain complementarity determining regions (CDRs) or variable regions (VRs) of lirilumab. Accordingly, in one embodiment, the antibody comprises the CDR1, CDR2, and CDR3 domains of the heavy chain variable (VH) region of lirilumab, and the CDR1, CDR2 and CDR3 domains of the light chain variable (VL) region of lirilumab. In another embodiment, the antibody has at least about 90% variable region amino acid sequence identity with lirilumab.

Examples of cancers contemplated for treatment include lung cancer, head and neck cancer, breast cancer, pancreatic cancer, prostate cancer, renal cancer, bone cancer, testicular cancer, cervical cancer, gastrointestinal cancer, lymphomas, pre-neoplastic lesions in the lung, colon cancer, melanoma, metastatic melanoma, basal-cell skin cancer, squamous-cell skin cancer, dermatofibrosarcoma protuberans, Merkel cell carcinoma, Kaposi's sarcoma, keratoacanthoma, spindle cell tumours, sebaceous carcinomas, microcystic adnexal carcinoma, Paget's disease of the breast, atypical fibroxanthoma, leiomyosarcoma, and angiosarcoma, Lentigo Maligna, Lentigo Maligna Melanoma, Superficial Spreading Melanoma, Nodular Melanoma, Acral Lentiginous Melanoma, Desmoplastic Melanoma, and bladder cancer.

In some embodiments, the subject has cancer that is resistant (has been demonstrated to be resistant) to one or more anti-cancer therapies. In some embodiments, resistance to anti-cancer therapy includes recurrence of cancer or refractory cancer. Recurrence may refer to the reappearance of cancer, in the original site or a new site, after treatment. In some embodiments, resistance to anti-cancer therapy includes progression of the cancer during treatment with the anti-cancer therapy. In some embodiments, the cancer is at early stage or at late stage. The subject may have a cancer that expresses (has been shown to express e.g., in a diagnostic test) PD-L1 biomarker. In some embodiments, the patient's cancer expresses low PD-L1 biomarker. In some embodiments, the patient's cancer expresses high PD-L1 biomarker. The PD-L1 biomarker can be detected in the sample using a method selected from the group consisting of FACS, Western blot, ELISA, immunoprecipitation, immunohistochemistry, immunofluorescence, radioimmunoassay, dot blotting, immunodetection methods, HPLC, surface plasmon resonance, optical spectroscopy, mass spectrometery, HPLC, qPCR, RT-qPCR, multiplex qPCR or RTqPCR, RNA-seq, microarray analysis, SAGE, MassARRAY technique, and FISH, and combinations thereof.

In some embodiments, the cancer has low levels of T cell infiltration. In some embodiments, the cancer has no detectable T cell infiltrate. In some embodiments, the cancer is a non-immunogenic cancer (e.g., non-immunogenic colorectal cancer and/or ovarian cancer).

For example, a therapeutically effective or sufficient amount of the immune checkpoint inhibitor, such as an antibody, is administered to a human will be in the range of about 0.01 to about 50 mg/kg of patient body weight whether by one or more administrations. In some embodi- 10 ments, the antibody used is about 0.01 to about 45 mg/kg, about 0.01 to about 40 mg/kg, about 0.01 to about 35 mg/kg, about 0.01 to about 30 mg/kg, about 0.01 to about 25 mg/kg, about 0.01 to about 20 mg/kg, about 0.01 to about 15 mg/kg, about 0.01 to about 10 mg/kg, about 0.01 to about 5 mg/kg, 15 or about 0.01 to about 1 mg/kg administered daily, for example. In some embodiments, the antibody is administered at 15 mg/kg. However, other dosage regimens may be useful. In one embodiment, an anti-PD-L1 antibody described herein is administered to a human at a dose of 20 about 100 mg, about 200 mg, about 300 mg, about 400 mg, about 500 mg, about 600 mg, about 700 mg, about 800 mg, about 900 mg, about 1000 mg, about 1100 mg, about 1200 mg, about 1300 mg or about 1400 mg on day 1 of 21-day cycles. The dose may be administered as a single dose or as 25 multiple doses (e.g., 2 or 3 doses), such as infusions. The progress of this therapy is easily monitored by conventional techniques.

### Anti-Cancer and Other Therapies

In some embodiments, the immune checkpoint inhibitor 30 may be administered in combination with at least one additional therapeutic. The additional therapy may be a cancer therapy such as radiation therapy, surgery, chemotherapy, gene therapy, DNA therapy, viral therapy, RNA therapy, immunotherapy, bone marrow transplantation, 35 nanotherapy, monoclonal antibody therapy, or a combination of the foregoing. The additional therapy may be in the form of adjuvant or neoadjuvant therapy.

In an example, the therapy of the cancer (whether with or without administration of an immune checkpoint inhibitor) 40 or any other disease (eg, viral infection or autoimmune disease) may radiation therapy, surgery, chemotherapy, gene therapy, DNA therapy, viral therapy, RNA therapy, immunotherapy, bone marrow transplantation, nanotherapy or monoclonal antibody therapy. The therapy may be a combination of the foregoing. An additional therapy may be administered

In some embodiments, the therapy (or the additional cancer therapy) is the administration of a small molecule enzymatic inhibitor or anti-metastatic agent. In some 50 embodiments, the additional therapy is the administration of side-effect limiting agents (e.g., agents intended to lessen the occurrence and/or severity of side effects of treatment, such as anti-nausea agents, etc.).

In some embodiments, the therapy (or the additional 55 cancer therapy) is radiation therapy. In some embodiments, the therapy (or the additional cancer therapy) is surgery. In some embodiments, the therapy (or the additional cancer therapy) is a combination of radiation therapy and surgery. In some embodiments, the therapy (or the additional cancer therapy) is gamma irradiation. In some embodiments, the therapy (or the additional cancer therapy) is therapy targeting PBK/AKT/mTOR pathway, HSP90 inhibitor, tubulin inhibitor, apoptosis inhibitor, and/or chemopreventative agent. The therapy (or the additional cancer therapy) may be 65 one or more of the chemotherapeutic agents known in the

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Administration of any compound or therapy of the present embodiments to a patient will follow general protocols for the administration of such compounds, taking into account the toxicity, if any, of the agents. Therefore, in some embodiments there is a step of monitoring toxicity that is attributable to combination therapy.

The therapy can comprise or consist of administration to the subject of any of the following:—

#### 1. Chemotherapy

A wide variety of chemotherapeutic agents may be used in accordance with the present embodiments. The term "chemotherapy" refers to the use of drugs to treat cancer. A "chemotherapeutic agent" is used to connote a compound or composition that is administered in the treatment of cancer. These agents or drugs are categorized by their mode of activity within a cell, for example, whether and at what stage they affect the cell cycle. Alternatively, an agent may be characterized based on its ability to directly cross-link DNA, to intercalate into DNA, or to induce chromosomal and mitotic aberrations by affecting nucleic acid synthesis.

Examples of chemotherapeutic agents include alkylating agents, such as thiotepa and cyclosphosphamide; alkyl sulfonates, such as busulfan, improsulfan, and piposulfan; aziridines, such as benzodopa, carboquone, meturedopa, and uredopa; ethylenimines and methylamelamines, including altretamine, triethylenemelamine, trietylenephosphoramide, triethiylenethiophosphoramide, and trimethylolomelamine; acetogenins (especially bullatacin and bullatacinone); a camptothecin (including the synthetic analogue topotecan); bryostatin; callystatin; CC-1065 (eg, its adozelesin, carzelesin and bizelesin synthetic analogues); cryptophycins (particularly cryptophycin 1 and cryptophycin 8); dolastatin; duocarmycin (eg, the synthetic analogues, KW-2189 and CB 1-TM1); eleutherobin; pancratistatin; a sarcodictyin; spongistatin; nitrogen mustards, such as chlorambucil, chlornaphazine, cholophosphamide, estramustine, ifosfamide, mechlorethamine, mechlorethamine oxide hydrochloride, melphalan, novembichin, phenesterine, prednimustine, trofosfamide, and uracil mustard; nitrosureas, such as carmustine, chlorozotocin, fotemustine, lomustine, nimustine, and ranimnustine; antibiotics, such as the enediyne antibiotics (e.g., calicheamicin, especially calicheamicin gammall and calicheamicin omegall); dynemicin, including dynemicin A; bisphosphonates, such as clodronate; an esperamicin; as well as neocarzinostatin chromophore and related chromoprotein enediyne antiobiotic chromophores, aclacinomy sins, actinomycin, authrarnycin, azaserine, bleomycins, cactinomycin, carabicin, carminomycin, carzinophilin, chromomycinis, dactinomycin, daunorubicin, detorubicin, 6-diazo-5-oxo-L-norleucine, doxorubicin morpholino-(eg, doxorubicin, cyanomorpholino-doxorubicin, 2-pyrrolinodoxorubicin and deoxydoxorubicin), epirubicin, esorubicin, idarubicin, marcellomycin, mitomycins, such as mitomycin C, mycophenolic acid, nogalarnycin, olivomycins, peplomycin, potfiromycin, puromycin, quelamycin, rodorubicin, streptonigrin, streptozocin, tubercidin, ubenimex, zinostatin, and zorubicin; anti-metabolites, such as methotrexate and 5-fluorouracil (5-FU); folic acid analogues, such as denopterin, pteropterin, and trimetrexate; purine analogs, such as fludarabine, 6-mercaptopurine, thiamiprine, and thioguanine; pyrimidine analogs, such as ancitabine, azacitidine, 6-azauridine, carmofur, cytarabine, dideoxyuridine, doxifluridine, enocitabine, and floxuridine; androgens, such as calusterone, dromostanolone propionate, epitiostanol, mepitiostane, and testolactone; anti-adrenals, such as mitotane and trilostane; folic acid replenisher, such as frolinic acid; aceglatone; aldophosphamide glycoside; aminolevulinic

acid; eniluracil; amsacrine; bestrabucil; bisantrene; edatraxate; defofamine; demecolcine; diaziquone; elformithine; elliptinium acetate; an epothilone; etoglucid; gallium nitrate; hydroxyurea; lentinan; lonidainine; maytansinoids, such as maytansine and ansamitocins; mitoguazone; mitoxantrone; mopidanmol; nitraerine; pentostatin; phenamet; pirarubicin; losoxantrone; podophyllinic acid; 2-ethylhydrazide; procarbazine; PSKpolysaccharide complex; razoxane; rhizoxin; sizofiran; spirogermanium; tenuazonic acid; triaziquone; 2,2',2"-trichlorotriethylamine; trichothecenes (eg, T-2 toxin, verracurin A, roridin A and anguidine); urethan; vindesine; dacarbazine; mannomustine; mitobronitol; mitolactol; pipobroman; gacytosine; arabinoside ("Ara-C"); cyclophosphamide; taxoids, e.g., paclitaxel and docetaxel gemcitabine; 15 6-thioguanine; mercaptopurine; platinum coordination complexes, such as cisplatin, oxaliplatin, and carboplatin; vinblastine; platinum; etoposide (VP-16); ifosfamide; mitoxantrone; vincristine; vinorelbine; novantrone; teniposide; edatrexate; daunomycin; aminopterin; xeloda; ibandronate; 20 irinotecan (e.g., CPT-11); topoisomerase inhibitor RFS 2000; difluorometlhylornithine (DMFO); retinoids, such as retinoic acid; capecitabine; carboplatin, procarbazine, plicomycin, gemcitabien, navelbine, farnesyl-protein tansferase inhibitors, transplatinum, and pharmaceutically acceptable 25 salts, acids, or derivatives of any of the above

### 2. Radiotherapy

Other factors that cause DNA damage and have been used extensively include what are commonly known as y-rays, X-rays, and/or the directed delivery of radioisotopes to 30 tumour cells. Other forms of DNA damaging factors are also contemplated, such as microwaves, proton beam irradiation (U.S. Pat. Nos. 5,760,395 and 4,870,287), and UV-irradiation. It is most likely that all of these factors affect a broad range of damage on DNA, on the precursors of DNA, on the 35 replication and repair of DNA, and on the assembly and maintenance of chromosomes. Dosage ranges for X-rays range from daily doses of 50 to 200 roentgens for prolonged periods of time (3 to 4 weeks), to single doses of 2000 to 6000 roentgens. Dosage ranges for radioisotopes vary 40 widely, and depend on the half-life of the isotope, the strength and type of radiation emitted, and the uptake by the neoplastic cells.

### 3. Immunotherapy

The skilled artisan will understand that immunotherapies 45 may be used in combination or in conjunction with the methods described herein. In the context of cancer treatment, immunotherapeutics generally rely on the use of immune effector cells and molecules to target and destroy cancer cells. Rituximab (RITUXAN®) is an example of an 50 immunotherapy. The immune effector may be, for example, an antibody specific for a marker on the surface of a tumour cell. The antibody alone may serve as an effector of therapy or it may recruit other cells to actually effect cell killing. The antibody also may be conjugated to a drug or toxin (chemo 55 therapeutic, radionuclide, ricin A chain, cholera toxin, pertussis toxin, etc.) and serve as a targeting agent. Alternatively, the effector may be a lymphocyte carrying a surface molecule that interacts, either directly or indirectly, with a tumour cell target. Various effector cells include cytotoxic T 60 cells and NK cells.

In an example, the immunotherapy comprises adoptive cell therapy, such as CAR-T administration, eg, anti-CD19 or CD20 CAR-T administration.

In an example, the immunotherapy comprises or consists 65 of administration of an IL-2 (eg, a truncated IL-2 or pegylated IL-2 or Fc-fused IL-2).

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Antibody-drug conjugates have emerged as a breakthrough approach to the development of cancer therapeutics. Antibody-drug conjugates (ADCs) comprise monoclonal antibodies (MAbs) that are covalently linked to cell-killing drugs. This approach combines the high specificity of MAbs against their antigen targets with highly potent cytotoxic drugs, resulting in "armed" MAbs that deliver the payload (drug) to tumour cells with enriched levels of the antigen. Targeted delivery of the drug also minimizes its exposure in normal tissues, resulting in decreased toxicity and improved therapeutic index. The approval of two ADC drugs, ADCE-TRIS® (brentuximab vedotin) in 2011 and KADCYLA® (trastuzumab emtansine or T-DM1) in 2013 by FDA validated the approach. There are currently more than 30 ADC drug candidates in various stages of clinical trials for cancer treatment. As antibody engineering and linker-payload optimization are becoming more and more mature, the discovery and development of new ADCs are increasingly dependent on the identification and validation of new targets that are suitable to this approach and the generation of targeting MAbs. Two criteria for ADC targets are upregulated/high levels of expression in tumour cells and robust internaliza-

In one aspect of immunotherapy, the tumour cell must bear some marker that is amenable to targeting, i.e., is not present on the majority of other cells. Many tumour markers exist and any of these may be suitable for targeting in the context of the present embodiments. Common tumour markers include CD20, carcinoembryonic antigen, tyrosinase (p97), gp68, TAG-72, HMFG, Sialyl Lewis Antigen, MucA, MucB, PLAP, laminin receptor, erb B, and pi 55. An alternative aspect of immunotherapy is to combine anticancer effects with immune stimulatory effects Immune stimulating molecules also exist including: cytokines, such as IL-2, IL-4, IL-12, GM-CSF, gamma-IFN, chemokines, such as MIP-1, MCP-1, IL-8, and growth factors, such as FLT3 ligand.

### 4. Surgery

The cancer or other disease or condition may be treated by surgery in the invention.

Approximately 60% of persons with cancer will undergo surgery of some type, which includes preventative, diagnostic or staging, curative, and palliative surgery. Curative surgery includes resection in which all or part of cancerous tissue is physically removed, excised, and/or destroyed and may be used in conjunction with other therapies, such as the treatment of the present embodiments, chemotherapy, radiotherapy, hormonal therapy, gene therapy, immunotherapy, and/or alternative therapies. Tumour resection refers to physical removal of at least part of a tumour. In addition to tumour resection, treatment by surgery includes laser surgery, cryosurgery, electro surgery, and microscopically-controlled surgery (Mohs' surgery).

Upon excision of part or all of cancerous cells, tissue, or tumour, a cavity may be formed in the body. Treatment may be accomplished by perfusion, direct injection, or local application of the area with an additional anti-cancer therapy. Such treatment may be repeated, for example, every 1, 2, 3, 4, 5, 6, or 7 days, or every 1, 2, 3, 4, and 5 weeks or every 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, or 12 months. These treatments may be of varying dosages as well.

### 5. Bacterial Transplants

In an embodiment, the therapy comprises administering to the subject a bacterial transplant, eg, a faecal microbial transplant, comprising defined bacteria. For example, the transplant is any composition disclosed in WO2018064165, the disclosure of which (especially the compositions therein)

are incorporated herein by reference in its entirety for possible application in the present invention. For example, the transplant is according to any of the following Paragraphs (tables nd sequence numbers referring to the tables and sequences in WO2018064165, which are explicitly 5 incorporated herein for possible use in the Claims):—

- 1. A composition comprising at least one isolated or purified population of bacteria belonging to one or more of the families Ruminococcaceae, Clostridiaceae, Lachnospiraceae, Micrococcaceae, and/or Veilonellaceae.
- 2. A composition comprising at least two isolated or purified populations of bacteria belonging to one or more of the families Ruminococcaceae, Clostridiaceae, Lachnospiraceae, Micrococcaceae, and/or Veilonellaceae.
- 3. The composition of Paragraph 1 or Paragraph 2, wherein each of the populations of bacteria is present in the composition at a concentration of at least 10<sup>3</sup> CFU.
- 4. The composition of Paragraph 1 or Paragraph 2, wherein the composition is a live bacterial product or a live biotherapeutic product.
- 5. The composition of Paragraph 1 or Paragraph 2, wherein the at least one isolated or purified population bacteria or the at least two isolated or purified populations of bacteria are provided as bacterial spores.
- 6. The composition of Paragraph 1 or Paragraph 2, wherein the at least one population of bacteria or the at least two isolated or purified populations of bacteria belong to Clostridiales Family XII and/or Clostridiales Family XIII.
- 7. The composition of Paragraph 1 or Paragraph 2, wherein 30 the at least one isolated or purified population bacteria or the at least two isolated or purified populations of bacteria belong to the family Ruminococcaceae and/or of the family Clostridiaceae.
- 8. The composition of Paragraph 1 or Paragraph 2, wherein 35 the population of bacteria belonging to the family Ruminococcaceae is further defined as a population of bacteria belonging to the genus *Ruminococcus*.
- 9. The composition of Paragraph 8, wherein the population of bacteria belonging to the genus *Ruminococcus* is further defined as a population of bacteria belonging to the species *Ruminococcus bromii.*404403, 1348, 253314, 258515, 33033, 1118061, 357276, 214851, 320502, 217731, 246787, 29371, 649764, 901, 29374, 33043, 39778, 682400, 871665, 160404, 745368, 408, 1584, 333367, 47246, 1096246, 53342, 438033,
- 10. The composition of Paragraph 1 or Paragraph 2, wherein the population of bacteria belonging to the family Ruminococcaceae is further defined as a population of bacteria 45 belonging to the genus *Faecalibacterium*.
- 11. The composition of Paragraph 10, wherein the population of bacteria belonging to the genus *Faecalibacterium* is further defined as a population of bacteria belonging to the species *Faecalibacterium prausnitzii*.
- 12. The composition of Paragraph 1 or Paragraph 2, wherein the population of bacteria belonging to the family Micrococcaceae is further defined as a population of bacteria belonging to the genus *Rothia*.
- 13. The composition of Paragraph 1 or Paragraph 2, wherein 55 the composition further comprises a population of bacteria belonging to the species *Porphyromonas pasteri*, the species *Clostridium hungatei*, the species *Phascolarctobacterium faecium*, the genus *Peptomphilus*, and/or the class Mollicutes.
- 14. The composition of Paragraph 1 or Paragraph 2, wherein the composition is essentially free of populations of bacteria belonging to the order Bacteroidales.
- 15. The composition of Paragraph 1 or Paragraph 2, wherein the at least one isolated or purified population bacteria or the 65 at least two isolated or purified populations of bacteria belongs to one or more of the species, subspecies or bacterial

70strains selected from the group consisting of the species in Table 1 with an enrichment index (ei) greater than 0.5.

16. The composition of Paragraph 1 or Paragraph 2, wherein the at least one isolated or purified population bacteria or the at least two isolated or purified populations of bacteria are selected from the group consisting of the species in Table 1 with an "ei" equal to 1.

17. The composition of Paragraph 1 or Paragraph 2, wherein the at least one isolated or purified population bacteria or the at least two isolated or purified populations of bacteria comprise a 16S ribosomal RNA (rRNA) nucleotide sequence that is at least 90% identical (eg, at least 91, 92, 93, 94, 95, 96, 97, 98 or 99% identical) to the 16S rRNA nucleotide sequence of bacteria identified by NCBI Taxonomy IDs selected from the group consisting of NCBI Taxonomy ID: 717959, 587, 758823, 649756, 44749, 671218, 1264, 1122135, 853, 484018, 46503, 54565, 290052, 216931, 575978, 433321, 1796646, 213810, 228924, 290054, 1509, 1462919, 29375, 337097, 1298596, 487174, 642492, 1735, 1297424, 742766, 46680, 132925, 411467, 1318465, 1852367, 1841857, 169679, 1175296, 259063, 172901, 39488, 57172, 28118, 166486, 28133, 1529, 694434, 1007096, 84030, 56774, 102148, 626947, 216933, 1348613, 1472417, 100176, 824, 1471761, 1297617, 288966, 1317125, 28197, 358743, 264639, 1265, 1335, 66219, 69473, 115117, 341220, 1732, 873513, 396504, 1796619, 45851, 2741, 105841, 86332, 1349822, 84037, 180311, 54291, 1217282, 762984, 1185412, 154046, 663278, 1543, 398512, 69825, 1841867, 1535, 1510, 84026, 1502, 1619234, 39497, 1544, 29343, 649762, 332095, 536633, 1033731, 574930, 742818, 177412, 1121308, 419208, 1673717, 55779, 28117, 626937, 180332, 1776382, 40519, 34062, 40518, 74426, 1216062, 293826, 850, 645466, 474960, 36835, 115544, 1515, 88431, 216932, 1417852, 39492, 1583, 420247, 118967, 169435, 37658, 138595, 31971, 100886, 1197717, 234908, 537007, 319644, 168384, 915173, 95159, 1816678, 626940, 501571, 1796620, 888727, 1147123, 376806, 1274356, 1267, 39495, 404403, 1348, 253314, 258515, 33033, 1118061, 357276, 29374, 33043, 39778, 682400, 871665, 160404, 745368, 408, 1584, 333367, 47246, 1096246, 53342, 438033, 351091, 1796622, 1776384, 817, 48256, 720554, 500632, 36849, 301302, 879970, 655811, 264463, 1532, 285, 995, 242750, 29539, 1432052, 622312, 1796636, 1337051, 328814, 28446, 1492, 820, 39496, 52786, 1549, 1796618, 582, 46507, 109327, 1531, 1382, 33039, 311460, 230143, 216935, 539, 35519, 1681, 328813, 214853, 89014, 1121115, 1585974, 29466, 1363, 292800, 270498, 214856,  $142877, \ 133926, \ 209880, \ 179628, \ 1121102, \ 105612,$ 1796615, 39777, 29353, 1579, 163665, 53443, 261299, 1302, 1150298, 938289, 358742, 471875, 938278, 1796613, 1118057, 1077144, 1737, 218205, 1121298, 684066, 433659, 52699, 204516, 706562, 253257, 328812, 1280, 147802, 58134, 1335613, 891, 585394, 1582, 235931, 308994, 1589, 1682.

1736, 28129, 178001, 551788, 2051, 856, 118562, 101070, 515619, 40215, 187979, 82979, 29363, 1776391, 1285191, 84112, 157688, 38304, 36850, 341694, 287, 75612, 818, 371674, 338188, 88164, 588581, 676965, 546271, 1236512, 178338, 862517, 157687, 158, 51048, 1583331, 529, 888745, 394340, 40545, 855, 553973, 938293, 93063, 708634, 179995, 1351, 476652, 1464038, 555088, 237576, 879566, 1852371, 742727, 1377, 35830, 997353, 218538, 83771, 1605, 28111, 131109, 46609, 690567, 46206, 155615, 51616, 40542, 203, 294, 1034346, 156456, 80866, 554406, 796942, 1002367, 29347, 796944, 61592, 487175,

1050201, 762948, 137732, 1211819, 1019, 272548, 1717, 384636, 216940, 2087, 45634, 466107, 1689, 47678, 575, 979627, 840, 1660, 1236517, 617123, 546, 28135, 82171, 483, 501496, 99656, 1379, 84032, 39483, 1107316, 584, 28124, 1033744, 657309, 536441, 76123, 1118060, 89152, 576122, 303, 1541, 507751, 515620, 38302, 53419, 726, 40324, 1796610, 988946, 1852370, 1017, 1168289, 76936, 94869, 1161098, 215580, 1125779, 327575, 549, 1450648 and 478.

18. The composition of Paragraph 1 or Paragraph 2, 10 wherein the at least one isolated or purified population of bacteria or the at least two isolated or purified populations of bacteria are a species, subspecies or bacterial strains comprising a 16S rRNA gene sequence at least 80% identical (eg, at least 85, 90, 95 or 98% 15 identical) to any one of the sequences of SEQ ID NOs: 1-876 in WO2018064165.

19. The composition of Paragraph 1 or Paragraph 2, wherein the at least one isolated or purified population bacteria or the at least two isolated or purified popu- 20 lations of bacteria belong to the species, subspecies or bacterial strains selected from the group consisting of Bacteroides coagulans, Clostridium aldenense, Clostridium aldrichii, Clostridium alkalicellulosi, Clostridium amvgdalinum, Clostridium asparagiforme, 25 Clostridium cellulosi, Clostridium citroniae, Clostridium clariflavum DSM 19732, Clostridium clostridioforme, Clostridium colinum, Clostridium fimetarium, Clostridium hiranonis, Clostridium hun-Clostridium hylemonae DSM 15053, 30 Clostridium indolis, Clostridium lactatifermentans, Clostridium leptum, Clostridium methylpentosum, Clostridium oroficum, Clostridium papyrosolvens DSM 2782, Clostridium populeti, Clostridium propionicum, Clostridium saccharolyticum, Clostridium 35 scindens, Clostridium sporosphaeroides, Clostridium Clostridium straminisolvens. stercorarium Clostridium sufflavum, Clostridium termifidis, Clostridium thermosuccino genes, Clostridium viride, Clostridium xylanolyticum, Desulfotomaculum guttoi- 40 deum, Eubacterium rectale ATCC 33656, Eubacterium dolichum, Eubacterium eligens ATCC 27750, Eubacterium hallii, Eubacterium infirmum, Eubacterium siraeum, Eubacterium tenue, Ruminococcus torques, Acetanaerobacterium elongatum, Acetatifactor muris, 45 Acetivibrio cellulolyticus, Acetivibrio ethanolgignens, Acholeplasma brassicae 0502, Acholeplasma parvum, Acholeplasma vituli, Acinetobacter junii, Actinobacillus porcinus, Actinomyces bowdenii, Actinomyces dentalis, Actinomyces odontolyticus, Acutalibacter muris, 50 Aerococcus viridans, Aeromicrobium fasfidiosum, Alistipes finegoldii, Alistipes obesi, Alistipes onderdonkii, Alistipes putredinis, Alistipes shahii, Alistipes shahii WAL 8301, Alistipes timonensis JC136, Alkalibacter Alkaliphilus metalliredigens 55 saccharofermentans, OYMF, Allisonella histaminiformans, Allobaculum stercoricanis DSM 13633, Alloprevotella cava, Alloprevotella tannerae, Anaerobacterium chartisolvens, Anaerobiospirillum thomasii, Anaerobium acetethylicum, Anaerococcus octavius NCTC 9810, Anaerococ- 60 cus provenciensis, Anaerococcus vaginalis ATCC 51170, Anaerocolumna jejuensis, Anaerofilum agile, Anaerofustis stercorihominis, Anaeroglobus geminatus, Anaeromassilibacillus senegalensis, Anaeroplasma abactoclasficum, Anaerorhabdus furcosa, Anaero- 65 mobilis, Anaerostipes butyraticus, sporobacter Anaerostipes caccae, Anaerostipes hadrus,

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Anaerotruncus colihominis, Anaerovorax odorimutans, Anoxybacillus rupiensis, Aquabacterium limnoficum, Arcobacter butzleri, Arthrospira platensis, Asaccharobacter celatus, Atopobium parvulum, Bacteroides caccae, Bacteroides caecimuris, Bacteroides cellulosilyticus, Bacteroides clarus YIT 12056, Bacteroides dorei, Bacteroides eggerthii, Bacteroides finegoldii, Bacteroides fragilis, Bacteroides gallinarum, Bacteroides massiliensis, Bacteroides oleiciplenus YIT 12058, Bacteroides plebeius DSM 17135, Bacteroides rodentium JCM 16496, Bacteroides thetaiotaomicron, Bacteroides uniformis, Bacteroides xylanisolvens XB1A, Bacteroides xylanolyticus, Barnesiella intestinihominis, Beduini massiliensis, Bifidobacterium bifidum, Bifidobacterium denfium, Bifidobacterium longum subsp. infantis, Blaufia caecimuris, Blaufia coccoides, Blaufia faecis, Blaufia glucerasea, Blaufia hansenii DSM 20583, Blaufia hydrogenotrophica, Blaufia luti, Blaufia luti DSM 14534, Blaufia wexlerae DSM 19850, Budvicia aquatica, Butyricicoccus pullicaecorum, Butyricimonas paravirosa, Butyrivibrio crossotus, Caldicoprobacter oshimai, Caloramator coolhaasii, Caloramator proteoclasficus, Caloramator quimbayensis, Campylobacter gracilis, Campylobacter rectus, Campylobacter ureolyticus DSM 20703, Capnocytophaga gingivalis, Capnocytophaga leadbetteri, Capnocytophaga sputigena, Casaltella massiliensis, Catabacter hongkongensis, Catenibacterium mitsuokai, Christensenella minuta, Christensenella timonensis, Chryseobacterium taklimakanense, Citrobacter freundii, Cloacibacillus porcorum, Clostridioides difficile ATCC 9689=DSM 1296, Clostridium amylolyticum, Clostridium bowmanii, Clostridium butvricum, Clostridium Clostridium cadaveris, colicanis, Clostridium gasigenes, Clostridium lentocellum DSM 5427, Clostridium oceanicum, Clostridium oryzae, Clostridium paraputrificum, Clostridium pascui, Clostridium perfringens, Clostridium Clostridium saccharobutylicum, Clostridium sporogenes, Clostridium ventriculi, Collinsella aerofaciens, Comamonas testosteroni, Coprobacter fasfidiosus NSB1, Coprococcus eutactus, Corvnebacterium diphtheriae, Corynebacterium durum, Corynebacterium mycetoides, Corynebacterium pyruviciproducens ATCC BAA-1742, Corynebacterium tuberculostearicum, Culturomica massiliensis, Cuneatibacter caecimuris, Defluviitalea saccharophila, Delftia acido-Desulfitobacterium chlororespirans, Desulfitobacterium metallireducens, Desulfosporosinus acididurans, Desulfotomaculum halophilum, Desulfotomaculum intricatum, Desulfotomaculum tondesulfuricans gense, Desulfovibrio desulfuricans, Desulfovibrio idahonensis, Desulfovibrio litoralis, Desulfovibrio piger, Desulfovibrio simplex, Desulfovibrio zosterae, Desulfuromonas acetoxidans, Dethiobacter alkaliphilus AHT Dethiosulfafibacter aminovorans, Dialister invisus, Dialister propionicifaciens, Dielma fasfidiosa, Dietzia alimentaria 72, Dorea longicatena, Dysgonomonas gadei ATCC BAA-286, Dysgonomonas mossii, Eggerthella lenta, Eikenella corrodens, Eisenbergiella tayi, Emergencia timonensis, Enorma massiliensis phi, Enterococcus faecalis, Enterorhabdus muris, Ethanoligenens harbinense YUAN-3, Eubacterium coprostanoligenes, Eubacterium limosum, Eubacterium oxi-Eubacterium sulci ATCC doreducens. Eubacterium uniforme, Eubacterium ventriosum,

Eubacterium xylanophilum, Extibacter muris, Ezakiella peruensis, Faecalibacterium prausnitzii, Faecalicoccus acidiformans, Faecalitalea cylindroides, Filifactor villosus, Flavonifr actor plautii, Flinfibacter butyricus, Frisingicoccus caecimuris, Fucophilus 5 fucoidanolyticus, Fusicatenibacter saccharivorans, Fusobacterium mortiferum, Fusobacterium nucleatum subsp. vincentii, Fusobacterium simiae, Fusobacterium varium, Garciella nitratireducens, Gemella haemolysans, Gemmiger formicilis, Gordonibacter urolithinfa- 10 ciens, Gracilibacter thermotolerans JW/YJL-S1, Granulicatella elegans, Guggenheimella bovis, Haemophilus haemolyticus, Helicobacter typhlonius, Hespellia stercorisuis, Holdemanella biformis, Holdemania massiliensis AP2. Howardella ureilytica, 15 Hungatella Hungatella hathewayi, Hydrogenoanaerobacterium saccharovorans, Ihubacter massiliensis, Intestinibacter bartlettii, Intestinimonas butyriciproducens, Irregularibacter muris, Kiloniella laminariae DSM 19542, Kroppenstedtia guangzhouensis, Lachno- 20 anaerobaculum orale. Lachnoanaerobaculum umeaense, Lachnoclostridium phytofermentans, Lactobacillus acidophilus, Lactobacillus algidus, Lactobacillus animalis, Lactobacillus casei, Lactobacillus delbrueckii, Lactobacillus fomicalis, Lactobacillus iners, 25 Lactobacillus pentosus, Lactobacillus rogosae, Lactococcus garvieae, Lactonifactor longoviformis, Leptotrichia buccalis, Leptotrichia hofstadii, Leptotrichia hongkongensis, Leptotrichia wadei, Leuconostoc inhae, Levylla massiliensis, Loriellopsis cavemicola, 30 Lutispora thermophila, Marinilabilia salmonicolor JCM 21150, Marvinbryanfia formatexigens, Mesoplasma photuris, Methanobrevibacter smithii ATCC 35061, Methanomassiliicoccus luminyensis BIO, Methylobacterium extorquens, Mitsuokellajalaludinii, 35 Mobilitalea sibirica, Mobiluncus curtisii, Mogibacterium pumilum, Mogibacterium timidum, Moorella glycerini, Moorella humiferrea, Moraxella nonliquefaciens, Moraxella osloensis, Morganella morganii, Moryella indoligenes, Muribaculum intestinale, Muri- 40 monas intestini, Natranaerovirga pectinivora, Neglecta timonensis, Neisseria cinerea, Neisseria oxalis, Nocardioides mesophilus, Novibacillus thermophilus, Ochroanthropi, Odoribacter splanchnicus, Olsenella profusa, Olsenella uli, Oribacterium asac- 45 charolyticum ACB7, Oribacterium sinus, Oscillibacter ruminantium GH1, Oscillibacter valericigenes, Oxobacter pfennigii, Pantoea agglomerans, Papillibacter cinnamivorans, Parabacteroides faecis, Parabacteroides goldsteinii, Parabacteroides gordonii, Para- 50 bacteroides merdae, Parasporobacterium paucivorans, Parasutterella excrementihominis, Parasutterella secunda, Parvimonas micra, Peptococcus niger, Peptoniphilus duerdenii ATCC BAA-1640, Peptoniphilus grossensis ph5, Peptoniphilus koenoeneniae, Pep- 55 toniphilus senegalensis JC140, Peptostreptococcus stomatis, Phascolarctobacterium succinatutens, Phocea massiliensis, Ponfibacter indicus, Porphyromonas bennonis, Porphyromonas endodontalis, Porphyromonas pasteri, Prevotella bergensis, Prevotella buccae 60 ATCC 33574, Prevotella denficola, Prevotella enoeca, Prevotella fusca JCM 17724, Prevotella loescheii, Prevotella nigrescens, Prevotella oris, Prevotella pollens ATCC 700821, Prevotella stercorea DSM 18206, P rev ote llamas silia timonensis, Propionispira arcuata, 65 Proteus mirabilis, Providencia rettgeri, Pseudobacteroides cellulosolvens ATCC 35603=DSM 2933,

Pseudobutyrivibrio ruminis, Pseudoflavonifr actor capillosus ATCC 29799, Pseudomonas aeruginosa, Pseudomonas fluorescens, Pseudomonas mandelii, Pseudomonas nitroreducens, Pseudomonas pufida, Raoultella ornithinolytica, Raoultella planficola, Raoultibacter massiliensis, Robinsoniella peoriensis, Romboutsia timonensis, Roseburia faecis, Roseburia hominis A2-183, Roseburia intestinalis, Roseburia inulinivorans DSM 16841, Rothia dentocariosa ATCC 17931, Ruminiclostridium thermocellum, Ruminococcus albus, Ruminococcus bromii, Ruminococcus callidus, Ruminococcus champanellensis 18P13=JCM 17042, Ruminococcus faecis JCM 15917, Ruminococcus flavefaciens, Ruminococcus gauvreauii, Ruminococcus lactaris ATCC 29176, Rummeliibacillus pyc-Saccharofermentans acetigenes, Scardovia wiggsiae, Schlegelella thermodepolymerans, Sedimenfibacter hongkongensis, Selenomonas sputigena ATCC 35185, Slackia exigua ATCC 700122, Slackia piriformis YIT 12062, Solitalea canadensis, Solobacterium moorei, Sphingomonas aquafilis, Spiroplasma alleghenense, Spiroplasma chinense, Spiroplasma chrysopicola, Spiroplasma culicicola, Spiroplasma lampyridicola, Sporobacter termifidis, Staphylococcus aureus, Stenotrophomonas maltophilia, Stomatobaculum longum, Streptococcus agalactiae ATCC 13813, Streptococcus cristatus, Streptococcus equinus, Streptococcus gordonii, Streptococcus lactarius, Streptococcus parauberis, Subdoligranulum variabile, Succinivibrio dextrinosolvens, Sutterella stercoricanis, Sutterella wadsworthensis, Syntrophococcus sucromutans, Syntrophomonas zehnderi OL-4, Terrisporobacter mayombei, Thermoleophilum album, Treponema denficola, Treponema socranskii, Tyzzerella nexilis DSM 1787, Vallitalea guaymasensis, Vallitalea pronyensis, Vampirovibrio chlorellavorus. Veillonella atypica. Veillonella denficariosi, Veillonella dispar, Veillonella parvula. Victivallis vadensis. Vulcanibacillus modesticaldus and Weissella confusa.

In an example, the transplant comprises or consists of SER-109 or SER-262 (and optionally the condition is a C dificile infection); VE202 or SER-287 (and optionally the disease is ulcerative colitis); SER-301 (and optionally the disease is IBD); SER-401 (and optionally the condition is a cancer; eg, wherein the therapy further comprises administration of an anti-PD-1 axis antibody, eg, an anti-PD-1 antibody); VE800 or SER-155 (and optionally the therapy further comprises the administration of a transplant, eg, a haematopoietic stem cell or solid organ transplant); EDP1066 or EDP1815 (and optionally the disease is an inflammatory condition, eg, colitis, Crohn's disease, asthma, rheumatoid arthritis (RA), psoriasis, dermatitis (eg, atopic dermatitis) or IBD); or EDP1503 (and the disease is a cancer, eg, colorectal cancer, renal cell carcinoma, melanoma or a PD-1 relapsed cancer). In an example, the therapy comprises the administration of SGM-1019, SG-2-0776 or EB8018 (and optionally the disease or condition is NASH or IBD or an inflammatory condition, eg, colitis, Crohn's disease, asthma, rheumatoid arthritis (RA), psoriasis and dermatitis (eg, atopic dermatitis). Those starting "VE" are developed by Vadanta Biosciences, SER are developed by Seres Therapeutics, EDP are developed by Evelo Biosciences, SG are developed by Second Genome and EB are developed by Enterome.

In an example, the disease or condition herein is an inflammatory condition, eg, colitis, Crohn's disease, asthma, rheumatoid arthritis (RA), psoriasis, dermatitis (eg, atopic dermatitis) or IBD.

### 6. Other Agents

It is contemplated that other agents may be used in combination with certain aspects of the present embodiments to improve the therapeutic efficacy of treatment. These additional agents include agents that affect the upregulation of cell surface receptors and GAP junctions, 10 cytostatic and differentiation agents, inhibitors of cell adhesion, agents that increase the sensitivity of the hyperproliferative cells to apoptotic inducers, or other biological agents. Increases in intercellular signalling by elevating the number of GAP junctions would increase the anti-hyperproliferative 15 effects on the neighbouring hyperproliferative cell population. In other embodiments, cytostatic or differentiation agents can be used in combination with certain aspects of the present embodiments to improve the anti-hyperproliferative efficacy of the treatments. Inhibitors of cell adhesion are 20 contemplated to improve the efficacy of the present embodiments. Examples of cell adhesion inhibitors are focal adhesion kinase (FAKs) inhibitors and Lovastatin. It is further contemplated that other agents that increase the sensitivity of a hyperproliferative cell to apoptosis, such as the antibody 25 c225, could be used in combination with certain aspects of the present embodiments to improve the treatment efficacy. Diseases and Conditions

Optionally, the disease or condition is selected from

- (a) A neurodegenerative disease or condition;
- (b) A brain disease or condition;
- (c) A CNS disease or condition;
- (d) Memory loss or impairment;
- (e) A heart or cardiovascular disease or condition, eg, heart attack, stroke or atrial fibrillation;
- (f) A liver disease or condition;
- (g) A kidney disease or condition, eg, chronic kidney disease (CKD);
- (h) A pancreas disease or condition;
- (i) A lung disease or condition, eg, cystic fibrosis or COPD; 40
- (i) A gastrointestinal disease or condition;
- (k) A throat or oral cavity disease or condition;
- (1) An ocular disease or condition;
- (m) A genital disease or condition, eg, a vaginal, labial, penile or scrotal disease or condition;
- (n) A sexually-transmissible disease or condition, eg, gonorrhea, HIV infection, syphilis or *Chlamydia* infection;
- (o) An ear disease or condition;
- (p) A skin disease or condition;
- (q) A heart disease or condition;
- (r) A nasal disease or condition
- (s) A haematological disease or condition, eg, anaemia, eg, anaemia of chronic disease or cancer;
- (t) A viral infection;
- (u) A pathogenic bacterial infection;
- (v) A cancer;
- (w) An autoimmune disease or condition, eg, SLE;
- (x) An inflammatory disease or condition, eg, rheumatoid arthritis, psoriasis, eczema, asthma, ulcerative colitis, colitis, Crohn's disease or IBD;
- (y) Autism;
- (z) ADHD;
- (aa) Bipolar disorder;
- (bb) ALS [Amyotrophic Lateral Sclerosis];
- (cc) Osteoarthritis;
- (dd) A congenital or development defect or condition;
- (ee) Miscarriage;

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- (ff) A blood clotting condition;
- (gg) Bronchitis;
- (hh) Dry or wet AMD;
- (ii) Neovascularisation (eg, of a tumour or in the eye);
- (jj) Common cold;
- (kk) Epilepsy;
- (11) Fibrosis, eg, liver or lung fibrosis;
- (mm) A fungal disease or condition, eg, thrush;
- (nn) A metabolic disease or condition, eg, obesity, anorexia, diabetes, Type I or Type II diabetes.
- (00) Ulcer(s), eg, gastric ulceration or skin ulceration;
- (pp) Dry skin;
- (qq) Sjogren's syndrome;
- (rr) Cytokine storm;
- 5 (ss) Deafness, hearing loss or impairment;
  - (tt) Slow or fast metabolism (ie, slower or faster than average for the weight, sex and age of the subject);
- (uu) Conception disorder, eg, infertility or low fertility;
- (vv) Jaundice:
- (ww) Skin rash:
- (xx) Kawasaki Disease;
- (yy) Lyme Disease;
- (zz) An allergy, eg, a nut, grass, pollen, dust mite, cat or dog fur or dander allergy;
- 5 (aaa) Malaria, typhoid fever, tuberculosis or cholera;
  - (bbb) Depression;
  - (ccc) Mental retardation;
  - (ddd) Microcephaly;
- (eee) Malnutrition;
- 30 (fff) Conjunctivitis;
  - (ggg) Pneumonia;
  - (hhh) Pulmonary embolism;
  - (iii) Pulmonary hypertension;
  - (jjj) A bone disorder;
- (kkk) Sepsis or septic shock;
  - (111) Sinusitus;
  - (mmm) Stress (eg, occupational stress);
- (nnn) Thalassaemia, anaemia, von Willebrand Disease, or haemophilia;
- 10 (000) Shingles or cold sore;
  - (ppp) Menstruation;
  - (qqq) Low sperm count.
  - Neurodegenerative or CNS Diseases or Conditions for Treatment or Prevention by the Method

In an example, the neurodegenerative or CNS disease or condition is selected from the group consisting of Alzheimer disease, geriopsychosis, Down syndrome, Parkinson's disease, Creutzfeldt Jakob disease, diabetic neuropathy, Parkinson syndrome, Huntington's disease, Machado-Joseph disease, amyotrophic lateral sclerosis, diabetic neuropathy, and Creutzfeldt Creutzfeldt-Jakob disease. For example, the disease is Alzheimer disease. For example, the disease is Parkinson syndrome.

In an example, wherein the method of the invention is practiced on a human or animal subject for treating a CNS or neurodegenerative disease or condition, the method causes downregulation of Treg cells in the subject, thereby promoting entry of systemic monocyte-derived macrophages and/or Treg cells across the choroid plexus into the brain of the subject, whereby the disease or condition (eg, Alzheimer's disease) is treated, prevented or progression thereof is reduced. In an embodiment the method causes an increase of IFN-gamma in the CNS system (eg, in the brain and/or CSF) of the subject. In an example, the method restores nerve fibre and/or reduces the progression of nerve fibre damage. In an example, the method restores nerve

myelin and//or reduces the progression of nerve myelin

damage. In an example, the method of the invention treats or prevents a disease or condition disclosed in WO2015136541 and/or the method can be used with any method disclosed in WO2015136541 (the disclosure of this document is incorporated by reference herein in its entirety, eg, for providing disclosure of such methods, diseases, conditions and potential therapeutic agents that can be administered to the subject for effecting treatment and/or prevention of CNS and neurodegenerative diseases and conditions, eg, agents such as immune checkpoint inhibitors, eg, anti-PD-1, anti-PD-L1, anti-TIM3 or other antibodies disclosed therein).

Cancers for Treatment or Prevention by the Method

Cancers that may be treated include tumours that are not vascularized, or not substantially vascularized, as well as vascularized tumours. The cancers may comprise non-solid tumours (such as haematological tumours, for example, leukaemias and lymphomas) or may comprise solid tumours. Types of cancers to be treated with the invention include, but are not limited to, carcinoma, blastoma, and sarcoma, and certain leukaemia or lymphoid malignancies, benign and malignant tumours, and malignancies e.g., sarcomas, carcinomas, and melanomas. Adult tumours/cancers and paediatric tumours/cancers are also included.

Haematologic cancers are cancers of the blood or bone 25 marrow. Examples of haematological (or haematogenous) cancers include leukaemias, including acute leukaemias (such as acute lymphocytic leukaemia, acute myelocytic leukaemia, acute myelogenous leukaemia and myeloblasts, promyeiocytic, myelomonocytic, monocytic and erythroleukaemia), chronic leukaemias (such as chronic myelocytic (granulocytic) leukaemia, chronic myelogenous leukaemia, and chronic lymphocytic leukaemia), polycythemia vera, lymphoma, Hodgkin's disease, non-Hodgkin's lymphoma (indolent and high grade forms), multiple myeloma, 35 Waldenstrom's macroglobulinemia, heavy chain disease, myeiodysplastic syndrome, hairy cell leukaemia and myelodysplasia.

Solid tumours are abnormal masses of tissue that usually do not contain cysts or liquid areas. Solid tumours can be 40 benign or malignant. Different types of solid tumours are named for the type of cells that form them (such as sarcomas, carcinomas, and lymphomas). Examples of solid tumours, such as sarcomas and carcinomas, include fibrosarcoma, myxosarcoma, liposarcoma, chondrosarcoma, 45 osteosarcoma, and other sarcomas, synovioma, mesothelioma, Ewing's tumour, leiomyosarcoma, rhabdomyosarcoma, colon carcinoma, lymphoid malignancy, pancreatic cancer, breast cancer, lung cancers, ovarian cancer, prostate cancer, hepatocellular carcinoma, squamous eel! carcinoma, 50 basal cell carcinoma, adenocarcinoma, sweat gland carcinoma, medullary thyroid carcinoma, papillary thyroid carcinoma, pheochromocytomas sebaceous gland carcinoma, papillary carcinoma, papillary adenocarcinomas, medullary carcinoma, bronchogenic carcinoma, renal cell carcinoma, 55 hepatoma, bile duct carcinoma, choriocarcinoma, Wilms' tumour, cervical cancer, testicular tumour, seminoma, bladder carcinoma, melanoma, and CNS tumours (such as a glioma (such as brainstem glioma and mixed gliomas), glioblastoma (also known as glioblastoma multiforme) 60 astrocytoma, CNS lymphoma, germinoma, medu!loblastoma, Schwannoma craniopharyogioma, ependymoma, pineaioma, hemangioblastoma, acoustic neuroma, oligodendroglioma, menangioma, neuroblastoma, retinoblastoma and brain metastases).

In an example, the cancer is a haematological cancer. In an example, the cancer is NSCLC. In an example, the cancer is renal cell carcinoma. In an example, the cancer is urothelial carcinoma. In an example, the cancer is melanoma. Autoimmune Diseases for Treatment or Prevention by the Method

Acute Disseminated Encephalomyelitis (ADEM)
Acute necrotizing hemorrhagic leukoencephalitis

Addison's disease

Agammaglobulinemia

Alopecia areata

Amyloidosis

Ankylosing spondylitis

Anti-GBM/Anti-TBM nephritis

Antiphospholipid syndrome (APS)

Autoimmune angioedema

Autoimmune aplastic anemia

Autoimmune dysautonomia

Autoimmune hepatitis

Autoimmune hyperlipidemia

Autoimmune immunodeficiency

Autoimmune inner ear disease (AIED)

Autoimmune myocarditis

Autoimmune oophoritis

Autoimmune pancreatitis

Autoimmune retinopathy

Autoimmune thrombocytopenic purpura (ATP)

Autoimmune thyroid disease

Autoimmune urticaria

Axonal & neuronal neuropathies

Balo disease

Behcet's disease

Bullous pemphigoid

Cardiomyopathy

Castleman disease

Celiac disease

Chagas disease

Chronic fatigue syndrome

Chronic inflammatory demyelinating polyneuropathy (CIDP)

Chronic recurrent multifocal ostomyelitis (CRMO)

Churg-Strauss syndrome

Cicatricial pemphigoid/benign mucosal pemphigoid

Crohn's disease

Cogans syndrome

Cold agglutinin disease

Congenital heart block Coxsackie myocarditis

CREST disease

Essential mixed cryoglobulinemia

Demyelinating neuropathies

Dermatitis herpetiformis

Dermatomyositis

Devic's disease (neuromyelitis optica)

Discoid lupus

Dressler's syndrome

Endometriosis

Eosinophilic esophagitis

Eosinophilic fasciitis

Erythema nodosum

Experimental allergic encephalomyelitis

Evans syndrome

Fibromyalgia

Fibrosing alveolitis

Giant cell arteritis (temporal arteritis)

Giant cell myocarditis

Glomerulonephritis

Goodpasture's syndrome

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Granulomatosis with Polyangiitis (GPA) (formerly called Wegener's Granulomatosis)

Graves' disease

Guillain-Barre syndrome
Hashimoto's encephalitis
Hashimoto's thyroiditis
Hemolytic anemia
Henoch-Schonlein purpura
Herpes gestationis

Hypogammaglobulinemia

Idiopathic thrombocytopenic purpura (ITP)

IgA nephropathy

IgG4-related sclerosing disease Immunoregulatory lipoproteins Inclusion body myositis Interstitial cystitis Juvenile arthritis

Juvenile diabetes (Type 1 diabetes)

Juvenile myositis Kawasaki syndrome Lambert-Eaton syndrome Leukocytoclastic vasculitis

Lichen planus Lichen sclerosus Ligneous conjunctivitis Linear IgA disease (LAD)

Lupus (SLE) Lyme disease, chronic Meniere's disease Microscopic polyangiitis

Mixed connective tissue disease (MCTD)

Mooren's ulcer

Mucha-Habermann disease Multiple sclerosis Myasthenia gravis Myositis

Narcolepsy

Neuromyelitis optica (Devic's)

Neutropenia

Ocular cicatricial pemphigoid

Optic neuritis

Palindromic rheumatism

PANDAS (Pediatric Autoimmune Neuropsychiatric Dis-

orders Associated with *Streptococcus*)
Paraneoplastic cerebellar degeneration
Paroxysmal nocturnal hemoglobinuria (PNH)

Parry Romberg syndrome Parsonnage-Turner syndrome Pars planitis (peripheral uveitis)

Pemphigus

Peripheral neuropathy Perivenous encephalomyelitis

Pernicious anemia POEMS syndrome Polyarteritis nodosa

Type I, II, & III autoimmune poly glandular syndromes

Polymyalgia rheumatica

Polymyositis

Postmyocardial infarction syndrome

Postpericardiotomy syndrome Progesterone dermatitis Primary biliary cirrhosis Primary sclerosing cholangitis

Psoriasis

Psoriatic arthritis

Idiopathic pulmonary fibrosis Pyoderma gangrenosum Pure red cell aplasia Raynauds phenomenon Reactive Arthritis

Reflex sympathetic dystrophy

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Reiter's syndrome
Relapsing polychondritis
Restless legs syndrome
Retroperitoneal fibrosis
Rheumatic fever

Rheumatoid arthritis

Sarcoidosis Schmidt syndrome

Scleritis Scleroderma

Sjogren's syndrome

Sperm & testicular autoimmunity

Stiff person syndrome

Subacute bacterial endocarditis (SBE)

Susac's syndrome Sympathetic ophthalmia Takayasu's arteritis

Temporal arteritis/Giant cell arteritis Thrombocytopenic purpura (TTP)

Tolosa-Hunt syndrome
Transverse myelitis
Type 1 diabetes
Ulcerative colitis

Undifferentiated connective tissue disease (UCTD)

0 Uveitis Vasculitis

Vesiculobullous dermatosis

Vitiligo

Wegener's granulomatosis (now termed Granulomatosis

with Polyangiitis (GPA). Inflammatory Diseases for Treatment or Prevention by the Method

Alzheimer's

ankylosing spondylitis

arthritis (osteoarthritis, rheumatoid arthritis (RA), psori-

atic arthritis)

atherosclerosis Crohn's disease

45 colitis dermatitis diverticulitis fibromyalgia hepatitis

irritable bowel syndrome (IBS) systemic lupus erythematous (SLE)

nephritis

Parkinson's disease ulcerative colitis.

55 Concepts:—

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The invention provides the following Concepts.

1. A programmable nuclease for use in a method of treating an acute microbial infection of a subject, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes

- comprised by the subject are cut and acute microbial infection of the subject is treated.
- 2. A programmable nuclease for use in a method of durably treating a microbial (eg, bacterial) infection of a subject, wherein the microbial infection is caused by 5 microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the subject, whereby microbes of the first species or strain are durably killed, or growth or proliferation of the 10 microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and microbial infection of the subject is durably 15 treated.
- 3. The nuclease of Concept 2, wherein the nuclease (eg, programmed nuclease) and/or a nucleic acid that programs the nuclease to recognise and cut the target site is administered to the subject at a first time (T1) and at 20 a second time (T2) wherein T2 is at least 1 hour after T1.
- 4. The nuclease of any preceding Concept, wherein the method comprises reducing the infection at least 100fold by the first 30 minutes (eg, by the first 15 minutes) 25 of the treatment.
- 5. The nuclease of any preceding Concept, wherein the method comprises maintaining reduction of the infection by at least 100-fold for at least 60 minutes (eg, at least 120 minutes) after exposing the subject to the 30 programmed nuclease.
- 6. The nuclease of any preceding Concept, wherein the method comprises reducing the infection such that the reduction in infection persists for 30 minutes immediately after the first 30 minutes of the treatment.
- 7. The nuclease of any preceding Concept, wherein the method comprises administering to the subject a RNA or a nucleic acid that encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the 40 target site in microbes comprised by the subject.
- 8. The nuclease of Concept 7, wherein the nuclease is administered simultaneously or sequentially with the RNA or nucleic acid to the subject.
- The nuclease of Concept 7, wherein the subject comprises the nuclease prior to administration of the RNA or nucleic acid to the subject.
- 10. The nuclease of any one of Concepts 7 to 9, wherein a plurality of viruses (eg, phage) are administered to the subject, wherein each virus comprises a copy of the 50 nucleic acid, wherein the viruses infect the microbes comprised by the subject to deliver thereto the nucleic acid.
- 11. The nuclease of Concept 10, wherein the ratio of administered viruses:microbes comprised by the subject is from 10 to 150.
- 12. The nuclease according to any preceding Concept, wherein the subject is a human or animal, optionally wherein the subject is a human over 65 years of age or is a paediatric patient.
- 13. The nuclease according to Concept 12, wherein the infection is an infection of the lungs, abdomen or urinary tract; or wherein the subject has undergone surgery, is on an immunosuppressant medication and/or is suffering from a chronic disease.
- 14. The nuclease according to any preceding Concept, wherein the infection is reduced by at least 90% for 1

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- hour or more, optionally by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 15. The nuclease according to any preceding Concept, wherein the method comprises reducing the infection at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein reduction of the infection by at least 100-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject to the programmed nuclease.
- 16. The nuclease according to any one of Concepts 12 to 15, wherein the method treats or prevents septicaemia and/or sepsis (eg, septic shock) in the subject.
- 17. The nuclease of Concept 16, wherein at the start of the treatment, the subject (eg, a human) has a temperature of <36° C. or >38° C.; a heart rate of >90/min, a respiratory rate of >20 breaths/min or PaCO<sub>2</sub><4.3 kPa; and white blood cell count of <4000/mm³ or >12,000/mm³.
- 18. The nuclease of Concept 16 or 17, wherein at the start of the treatment, the subject (eg, a human) has presence of two or more of the following: abnormal body temperature, abnormal heart rate, abnormal respiratory rate, abnormal blood gas and abnormal white blood cell count.
- 19. The nuclease of any preceding Concept, wherein the subject is a human or animal and the microbes are bacteria (eg, *E coli* or *C dificile*), wherein blood infection of the subject by the bacteria is reduced at least 100- or 1000-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 20. The nuclease of any one of Concepts 12 to 19, wherein the blood of the subject is infected with from  $10^7$  to  $10^{12}$  CFU/ml of the bacteria immediately before the treatment
- 21. The nuclease according to any one of Concepts 1 to 11, wherein the subject is a plant.
- The nuclease according to any preceding Concept, wherein the microbes are bacteria.
- 23. The nuclease according to Concept 22, wherein the bacteria are gram positive bacteria.
- 24. The nuclease according to Concept 22 or 23, wherein the bacteria are Staphylococcus, Streptococcus, Enterococcus, Legionella, Heamophilus, Ghonnorhea, Acinetobacter, Escherichia, Klebsiella, Pseudomonas or Stenotrophomonas bacteria (eg, E coli (eg, EHEC E coli), C dificile, V cholera, Staphylococcus (eg, S aureus or MRSA), Streptococcus pyogenes, Acinetobacter baumannii, Legionella, Pseudomonas aeruginosa, Klebsiella pneumoniae bacteria).
- 25. The nuclease according to any preceding Concept, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9), a meganuclease, a TALEN (Transcription activator-like effector nuclease) or zinc finger nuclease.
- 26. A plurality of viruses (eg, phage or phagemids for producing phage) for use with the nuclease of any preceding Concept in the method of treatment, wherein each virus comprises a copy of a nucleic acid as defined in any one of Concepts 7 to 9, wherein the viruses are capable of infecting microbes comprised by the subject to deliver thereto the nucleic acid.
- 27. A composition comprising a plurality of nucleic acids for programming the nuclease of any one of Concepts 1 to 25 in the method of treatment, wherein each nucleic acid is a nucleic acid as defined in any one of Concepts 7 to 9.
- 28. A CRISPR/Cas system comprising a nuclease according to any preceding Concept for use in the method of

treatment, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9) and the system comprises one or more guide RNAs or DNA encoding one or more guide RNAs, wherein each guide RNA is capable of programming the Cas nuclease to cut a target site comprised by the genomes of the microbes.

- 29. A guide RNA or a DNA encoding a guide RNA for use in the system of Concept 28 for use in the method of treating an acute microbial infection in the subject, eg, septicaemia or sepsis.
- A nucleic acid vector comprising the guide RNA or DNA recited in Concept 27 or 29.
- 31. The vector of Concept 30 wherein the vector is a phage, phagemid, viriophage, virus, plasmid (eg, conjugative plasmid) or transposon.
- 32. An anti-sepsis or anti-septicaemia composition for administration to a human or animal for treating sepsis or septicaemia, the composition comprising a plurality of vectors, wherein each vector is according to Concept 20 30 or 31.
- 33. A method of treating an acute microbial infection of a subject, wherein the method is as defined by any preceding Concept.
- 34. Use of a nuclease, plurality of viruses, system, guide 25 RNA, DNA or vector of any one of Concepts 1 to 26 and 28 to 30, in the manufacture of a composition for carrying out a method of treatment as defined by any preceding Concept, wherein the subject is an organism other than a human or animal.
- 35. Use of a nuclease, plurality of viruses, system, guide RNA, DNA or vector of any one of Concepts 1 to 26 and 28 to 30, in the manufacture of a composition for carrying out an ex vivo method of treatment of a microbial infection of a substrate, wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the substrate, whereby microbes of the first 40 species or strain are killed, or growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are 45 cut and acute microbial infection of the substrate is treated.
- 36. Use of a programmable nuclease in the manufacture of a composition for carrying out an ex vivo method of treatment of a microbial infection of a substrate, 50 wherein the microbial infection is caused by microbes of a first species or strain and the nuclease is programmable to cut a target site comprised by the genomes of microbes that have infected the substrate, whereby microbes of the first species or strain are killed, or 55 growth or proliferation of the microbes is reduced, the treatment method comprising exposing the subject to the nuclease wherein the nuclease is programmed to cut the target site, whereby genomes of the microbes comprised by the subject are cut and acute microbial 60 infection of the substrate is treated.
- 37. The use of Concept 34, 35 or 36, wherein the nuclease (eg, programmed nuclease) and/or a nucleic acid that programs the nuclease to recognise and cut the target site is administered to the subject or substrate at a first 65 time (T1) and at a second time (T2) wherein T2 is at least 1 hour after T1.

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- 38. The use of any one of Concepts 34 to 37, wherein the infection is reduced at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 39. The use of any one of Concepts 34 to 38, wherein the reduction of the infection is maintained by at least 100-fold for at least 60 minutes (eg, at least 120 minutes) after exposing the subject to the programmed nuclease.
- 40. The use of any one of Concepts 34 to 39, wherein the reduction in infection persists for 30 minutes immediately after the first 30 minutes of the treatment.
- 41. The use of any one of Concepts 34 to 40, wherein the method comprises administering to the subject or substrate a RNA or a nucleic acid that encodes an RNA for expression of the RNA in or on the subject or substrate, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in microbes comprised by the subject or substrate.
- 42. The use of Concept 41, wherein the nuclease is administered simultaneously or sequentially with the RNA or nucleic acid to the subject or substrate.
- 43. The use of Concept 41, wherein the subject or substrate comprises the nuclease prior to administration of the RNA or nucleic acid.
- 44. The use of any one of Concepts 41 to 43, wherein a plurality of viruses (eg, phage) are administered to the subject or substrate, wherein each virus comprises a copy of the nucleic acid, wherein the viruses infect the microbes comprised by the subject or substrate to deliver thereto the nucleic acid.
- 45. The use of Concept 44, wherein the ratio of administered viruses:microbes is from 10 to 150.
- 46. The use of any one of Concepts 34 to 45, wherein the infection is reduced by at least 90% for 1 hour or more, optionally by the first 30 minutes (eg, by the first 15 minutes) of the treatment.
- 47. The use of any one of Concepts 34 to 46, wherein the infection is reduced at least 100-fold by the first 30 minutes (eg, by the first 15 minutes) of the treatment; and wherein reduction of the infection by at least 100-fold is maintained for at least 60 minutes (eg, at least 120, 145 or 180 minutes) after exposing the subject or substrate to the programmed nuclease.
- 48. The use of any one of Concepts 34 to 47, wherein the subject is a plant; or wherein the substrate is a metallic, plastic, concrete, stone, wood, glass or ceramic substrate.
- 49. The use of any one of Concepts 34 to 48, wherein the microbes are bacteria.
- 50. The use according to Concept 49, wherein the bacteria are gram positive bacteria.
- 51. The use according to Concept 49 or 50, wherein the bacteria are Staphylococcus, Streptococcus, Enterococcus, Legionella, Heamophilus, Ghonnorhea, Acinetobacter, Escherichia, Klebsiella, Pseudomonas or Stenotrophomonas bacteria (eg, E coli (eg, EHEC E coli), C dificile, V cholera, Staphylococcus (eg, S aureus or MRSA), Streptococcus pyogenes, Acinetobacter baumannii, Legionella, Pseudomonas aeruginosa, Klebsiella pneumoniae bacteria).
- 52. The use of any one of Concepts 34 to 51, wherein the nuclease is a Cas nuclease (eg, a Cas 3 or 9), a meganuclease, a TALEN (Transcription activator-like effector nuclease) or zinc finger nuclease.

### Embodiments

1. A method for treating a pathogenic bacterial infection in a human or animal subject caused by bacteria (first

bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a programmable nuclease that is programmed to cut 5 the target site, wherein the subject is suffering from a further disease or condition other than the pathogenic bacterial infection and the method comprises administering a therapy to the subject for treating or preventing the further disease or condition, wherein the nuclease treats the infection and the therapy is efficacious in the presence of the programmed nuclease to treat or prevent the disease or condition.

- 2. The method of Embodiment 1, wherein the subject is a cancer patient and the therapy comprises administration 15 of a haematopoietic stem cell transplant, chemotherapeutic agent, immune checkpoint inhibitor, immune checkpoint agonist or an immune cell enhancer; adoptive cell therapy; radiation or surgery.
- 3. The method of Embodiment 2, wherein the therapy is an 20 immune checkpoint inhibitor antibody, or an antibody selected from ipilimumab (or YERVOY®), tremelimumab, nivolumab (or OPDIVO®), pembrolizumab (or KEYTRUDA®), pidilizumab, BMS-936559, durvalumab (or IMFINZI®) and atezolizumab (or TECENTRIQ®).
- 4. The method of Embodiment 1, wherein the therapy is a tissue, organ or cell transplant.
- 5. The method of Embodiment 1, wherein the treatment of the bacterial infection is carried out simultaneously with the administration of the therapy to the subject.
- 6. The method of Embodiment 1, wherein the treatment of the bacterial infection is carried out immediately before or after administering the therapy to the subject.
- 7. The method of Embodiment 1, wherein the method that encodes an RNA for expression of the RNA in the subject, wherein the RNA complexes with the nuclease to program the nuclease to cut the target site in first bacteria comprised by the subject, thereby killing the first bacteria.
- 8. The method of Embodiment 1, comprising administering 40 a nucleic acid vector to the subject, wherein the vector encodes the programmable nuclease.
- 9. The method of Embodiment 1, wherein the programmable nuclease is an endogenous nuclease of the first cells.
- the therapy in the presence of the programmed nuclease is greater than the efficacy of the therapy in the presence of a broad-spectrum antibiotic.
- 11. The method of Embodiment 1, wherein the efficacy of the therapy in the presence of the programmed nuclease is 50 greater than the efficacy of the therapy in the presence of an antibiotic selected from methicillin, vancomycin, linezolid, daptomycin, quinupristin, dalfopristin; teicoplanin; cephalosporin; carbapenem; fluoroquinolone; aminoglycoside; colistin; erythromycin; clindamycin; beta- 55 18. The method of Embodiment 1, wherein the first bacteria lactam; macrolide; amoxicillin; azithromycin; penicillin; ceftriaxone; azithromycin; ciprofloxacin; isoniazid (INH); rifampicin (RMP); amikacin; kanamycin; capreomycin; trimethoprim; itrofurantoin; cefalexin; amoxicillin; metronidazole (MTZ); cefixime; tetracycline; and 60 19. The method of Embodiment 1, wherein the nuclease is meropenem.
- 12. The method of Embodiment 1, wherein the first bacteria is selected from (i) Staphylococcus aureus that is resistant to an antibiotic selected from methicillin, vancomycin, linezolid, daptomycin, quinupristin, dalfopristin and tei- 65 coplanin; (ii) Pseudomonas aeuroginosa that is resistant to an antibiotic selected from cephalosporins, carbapen-

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ems, fluoroquinolones, aminoglycosides and colistin; (iii) Klebsiella species that is resistant to carbapenem; (iv) Streptoccocus species that is resistant to an antibiotic selected from erythromycin, clindamycin, beta-lactam, macrolide, amoxicillin, azithromycin and penicillin; (v) Salmonella species that is resistant to an antibiotic selected from ceftriaxone, azithromycin and ciprofloxacin; (vi) Shigella species that is resistant to ciprofloxacin or azithromycin; (vii) Mycobacterium tuberculosis that is resistant to an antibiotic selected from Resistance to isoniazid (INH), rifampicin (RMP), fluoroquinolone, amikacin, kanamycin, capreomycin and azithromycin; (viii) Enterococcus species that is resistant to vancomycin; (ix) Enterobacteriaceae species that is resistant to an antibiotic selected from cephalosporin and carbapenem; (x) E coli that is resistant to an antibiotic selected from trimethoprim, itrofurantoin, cefalexin and amoxicillin; (xi) Clostridium species that is resistant to metronidazole (MTZ), fluoroquinolone or carbapenem; (xii) Neisseria gonnorrhoea that is resistant to an antibiotic selected from cefixime, ceftriaxone, azithromycin and tetracycline; (xiii) Acinetoebacter baumannii that is resistant to an antibiotic selected from beta-lactam, meropenem and carbapenem; and (xiv) Campylobacter species that is resistant to ciprofloxacin or azithromycin.

- 13. The method of Embodiment 1, wherein the treatment of the infection treats or prevents in the subject a condition selected from vaginosis, meningitis, pneumonia, urinary tract infection, cystitis, nephritis, gastroenteritis, a skin infection, impetigo, erysipelas, cellulitis, septicaemia or sepsis in the subject.
- 14. The method of Embodiment 1, wherein the further disease or condition is a cancer; autoimmune disease or condition; or GI tract disease or condition.
- comprises administering to the subject a or a nucleic acid 35 15. The method of Embodiment 1, wherein the subject comprises bacteria (second bacteria) of one or more strains or species that are different to the first strain or species, wherein the genomes of the second bacteria do not comprise the target site, wherein the genomes of the second bacteria are not cut by the programmed nuclease in the subject, whereby second bacteria survive in the presence of the programmed nuclease in the patient; and wherein the therapy is efficacious in the presence of the second bacteria.
- 10. The method of Embodiment 1, wherein the efficacy of 45 16. The method of Embodiment 15, wherein reduction in the second bacteria in patients is associated with reduced efficacy of the therapy.
  - 17. The method of Embodiment 15, wherein the second bacteria are selected from the group consisting of Akkermansia, Alistipes, Bacteroides, Barnesiella, Bifidobacterium, Clostridium, Collinsella, Enterococcus, Fusobacterium, Lactobacillus, Propionibacterium, Ruminococcus, Segmented filamentous bacteria (SFB); Veillonella, Prevotella, Escherichia and Streptococcus bacteria.
  - are selected from the group consisting of E coli, C dificile, V cholera, Staphylococcus, Streptococcus pyogenes, Acinetobacter baumannii, Legionella, Pseudomonas aeruginosa and Klebsiella pneumoniae bacteria.
  - a Cas nuclease, a meganuclease, a Transcription activatorlike effector nuclease (TALEN) or zinc finger nuclease.
  - 20. A method for treating a pathogenic bacterial infection in a cancer patient caused by bacteria (first bacteria) of a first species or strain, the method comprising selectively killing first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria,

wherein the cutting is carried out using a Cas nuclease that is programmed by guide RNA to cut the target site, wherein the method comprises administering an immunotherapy to the subject for treating cancer in the patient, wherein the nuclease treats the infection and the immunotherapy is efficacious in the presence of the programmed nuclease to treat the cancer.

It will be understood that particular embodiments described herein are shown by way of illustration and not as limitations of the invention. The principal features of this invention can be employed in various embodiments without departing from the scope of the invention. Those skilled in the art will recognize, or be able to ascertain using no more than routine study, numerous equivalents to the specific 15 procedures described herein. Such equivalents are considered to be within the scope of this invention and are covered by the claims. All publications and patent applications mentioned in the specification are indicative of the level of skill of those skilled in the art to which this invention 20 pertains. All publications and patent applications and all US equivalent patent applications and patents are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and individually indicated to be incorporated by reference. The use of the word "a" or "an" when used in conjunction with the term "comprising" in the claims and/or the specification may mean "one," but it is also consistent with the meaning of "one or more," "at least one," and "one or more than one." 30 The use of the term "or" in the claims is used to mean "and/or" unless explicitly indicated to refer to alternatives only or the alternatives are mutually exclusive, although the disclosure supports a definition that refers to only alternatives and "and/or." Throughout this application, the term "about" is used to indicate that a value includes the inherent variation of error for the device, the method being employed to determine the value, or the variation that exists among the study subjects.

As used in this specification and claim(s), the words "comprising" (and any form of comprising, such as "comprise" and "comprises"), "having" (and any form of having, such as "have" and "has"), "including" (and any form of including, such as "includes" and "include") or "containing" 45 (and any form of containing, such as "contains" and "contain") are inclusive or open-ended and do not exclude additional, unrecited elements or method steps

The term "or combinations thereof" or similar as used herein refers to all permutations and combinations of the 50 listed items preceding the term. For example, "A, B, C, or combinations thereof is intended to include at least one of: A, B, C, AB, AC, BC, or ABC, and if order is important in a particular context, also BA, CA, CB, CBA, BCA, ACB, BAC, or CAB. Continuing with this example, expressly 55 included are combinations that contain repeats of one or more item or term, such as BB, AAA, MB, BBC, AAAB-CCCC, CBBAAA, CABABB, and so forth. The skilled artisan will understand that typically there is no limit on the number of items or terms in any combination, unless otherwise apparent from the context.

Any part of this disclosure may be read in combination with any other part of the disclosure, unless otherwise apparent from the context.

All of the compositions and/or methods disclosed and 65 claimed herein can be made and executed without undue experimentation in light of the present disclosure. While the

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compositions and methods of this invention have been described in terms of preferred embodiments, it will be apparent to those of skill in the art that variations may be applied to the compositions and/or methods and in the steps or in the sequence of steps of the method described herein without departing from the concept, spirit and scope of the invention. All such similar substitutes and modifications apparent to those skilled in the art are deemed to be within the spirit, scope and concept of the invention as defined by the appended claims.

The present invention is described in more detail in the following non limiting Examples.

### **EXAMPLES**

Precision Fast Bacteria Killing with Programmable Nucleases

The examples provide a method for fast and precision killing of *Escherichia coli* and *Clostridium dificile* strains. As a model programmable nuclease system, we used a CRISPR guided vector (CGV<sup>TM</sup>) system to specifically target enterohemorrhagic *E. coli* (EHEC) and probiotic *E. coli* Nissle.

## Example 1. Precision Killing of Target Strain Enterohemorrhagic *E. coli* (EHEC)

1.1. Design, Construction and Delivery of CRISPR Guided Vector (CGV) System Targeting *E. coli* (EHEC) ATCC43888.

The invention provides a CGV system to specifically target enterohemorrhagic E. coli (EHEC) ATCC43888 (a human fecal isolate obtained from the American Type Culture Collection). The CGV system comprises two vectors: (a) a vector containing a tracrRNA and the Cas9 protein from Streptococcus pyogenes (SpCas); (b) a vector containing a guide RNA (gRNA) that comprises a nucleotide sequence capable of hybridizing to a target sequence in the host cells to guide SpCas9 to the target sequence. To enable specific killing of E. coli (EHEC) ATCC43888, a particular sequence from the genome of this strain was chosen to target. Specifically, the sequence contains 20 nucleotides from the 23S ribosomal RNA gene from E. coli (EHEC) ATCC43888. Additionally, the 5'-NGG protospacer adjacent motif (PAM) was located adjacent to the selected target sequence. The selected target sequence in the 23S rRNA gene can be found in Table 3.

1.2 Characterization of the CGV System Targeting *E. coli* (EHEC) ATCC43888.

To establish the CGV system functionality in mediating sequence-specific killing in *E. coli* (EHEC) ATCC43888, the system was transformed into *E. coli* (EHEC) ATCC43888 cells. Overnight cultures were diluted 1:100 in fresh lysogeny broth (LB) and grown to mid-exponential phase OD600~0.6. The CRISPR system was induced by adding theophylline and arabinose (2 mM theophylline and 1% arabinose), and survival of the strain was followed over time by plating the cultures in serial dilutions every 15 minutes, for 1 h (FIG. 1B). CRISPR induction in *E. coli* (EHEC) surprisingly triggered a rapid killing of the cells, achieving 99.98% killing within 30 minutes of induction (FIG. 1A).

Example 2. In Vivo CRISPR Killing of Target Strain Enterohemorrhagic *E. coli* (EHEC) In *Galleria mellonella* Larvae In Vivo Infection Model

## 2.1. CRISPR Efficacy Against E. coli (EHEC) ATCC43888 Infections in Galleria mellonella

CRISPR killing of target strain *E. coli* (EHEC) ATCC43888 was tested in *G. mellonella* in vivo infection model. To this aim, *G. mellonella* larvae were delivered <sup>10</sup> injections of bacteria 10<sup>8</sup> CFU *E. coli* (EHEC) ATCC43888) behind the final left proleg. Approximately 1 h after the injection, CRISPR inducers (2 mM theophylline and 1% arabinose) were administered behind the final right proleg. Larvae were incubated at 37° C. and they were sacrificed <sup>15</sup> after 1 and 2 h after induction. As shown in FIGS. **2** and **3**, CRISPR induction killed 99% of the population after 2 h, as compared to the off-target control.

2.2. Survival Curves of *G. mellonella* Larvae Infected with Enterohemorrhagic *E. coli* (EHEC).

G. mellonella larvae were delivered injections of bacteria (8×10<sup>4</sup> CFU E. coli ATCC43888) behind the final left proleg. Approximately 1 h after the injection, CRISPR inducers (2 mM theophylline and 1% arabinose) were administered behind the final right proleg. Larvae were 25 incubated at 37° C. and survival was monitored for 115 h, with death indicated by lack of movement and unresponsiveness to touch. CRISPR killing of target strain E. coli (EHEC) ATCC43888 in G. mellonella larvae significantly improved survival of the larvae compared to the off-target 30 control (FIG. 4) (log-rank test, P<0.03).

### Example 3. Precision Killing of Target Strain Probiotic *E. coli* Nissle 1917

## 3.1. Design, Construction and Delivery of CRISPR Guided Vector (CGV) System Targeting *E. coli* Nissle 1917.

The invention provides a CGV system to specifically target *E. coli* Nissle 1917. The CGV system comprises two vectors: (a) a vector containing a tracrRNA and the Cas9 40 from *Streptococcus pyogenes* (SpCas); (b) a vector containing a guide RNA (gRNA) that comprises a nucleotide sequence capable of hybridizing to a target sequence in the host cells to guide SpCas9 to the target sequence. To enable specific killing of *E. coli* Nissle 1917, a specific sequence 45 from the genome of this strain was chosen to target. Specifically, the sequence contains 20 nucleotides from the pks gene from *E. coli* Nissle 1917. Additionally, the 5'-NGG protospacer adjacent motif (PAM) was located adjacent to the selected target sequence. The selected target sequence in 50 the pks gene can be found in Table 3.

Furthermore, a different genome target was selected to specifically kill *E. coli* Nissle 1917. The sequence contains 20 nucleotides from the yapH gene. Additionally, the 5'-NGG protospacer adjacent motif (PAM) was located 55 adjacent to the selected target sequence. The selected target sequence in yapH gene can be found in Table 3.

3.2. Construction and Delivery of CRISPR Guided Vectors (CGV) Targeting *E. coli* Nissle 1917

To establish CGVs functionality in mediating sequence-specific killing in *E. coli* Nissle 1917, the CGV system was transformed into *E. coli* Nissle 1917 cells. Overnight cultures were diluted 1:100 in fresh lysogeny broth (LB) and grown to mid-exponential phase OD600~0.6. The CRISPR system was induced by adding theophylline and arabinose (2 65 mM theophylline and 1% arabinose), and survival of the strain was followed over time by plating the cultures in serial

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dilutions every 15 minutes, for 3 h (FIGS. **5**B and **6**B). FIGS. **5**B and **6**A show CRISPR killing assay in *E. coli* Nissle 1917, targeting pks gene and yapH gene, respectively. In both cases, CRISPR induction triggers a rapid killing of *E. coli* Nissle 1917 cells, achieving 99.98% killing within only 15 minutes of induction.

### Example 4. In Vitro CRISPR Killing of Clostridium difficile by Conjugative Plasmid Vectors and Cas3

This experiment involves the precision killing of Clostridium difficile using a gRNA-encoding CRISPR array that is delivered from a probiotic carrier bacterial species by conjugative plasmids as vectors (which we call CRISPR guided vectors (CGV<sup>TM</sup>)). A carrier bacterium (E. coli donor strain containing the CRISPR guided vector (CGV<sup>TM</sup>)) was mated with Clostridium difficile which was killed upon delivery of the CGV<sup>TM</sup> containing the designed array. This CGV<sup>TM</sup> harnessed the endogenous Cas3 machinery of Clostridium difficile 630Δerm. A 100% killing of Clostridium difficile cells was achieved. Introduction

Clostridium difficile (C. difficile) is a spore-forming human opportunistic pathogen that can asymptomatically colonize the intestine of healthy individuals. The two main risk factors for contracting C. difficile-associated diseases, such as nosocomial diarrhea, are age and antibiotic treatment and can have fatal consequences. C. difficile 630 $\Delta$ erm, the subject of our study, is a well-characterized strain and it is widely used for the generation of mutant specimens. Study Objectives

Objective 1: Delivery of CGVs by Conjugation.

A CRISPR guided vector (CGV) containing an array to specifically target and kill *C. difficile* was designed and assembled. The same CGV lacking the array was assembled to use as a control for conjugation efficiency. Both CGVs were transformed into the carrier strain *Escherichia coli* CA434, which was used as a donor strain to conjugate the plasmid into our strain of interest *C. difficile* 630Δerm. Objective 2: Harnessing *Clostridium difficile* Endogenous Cas3 Machinery.

Upon transcription of the delivered CRISPR array in the recipient target strain C. difficile, the endogenous Cas3 was guided to cut its own DNA; leading to bacterial death. Objective 3: Eradication of  $Clostridium\ difficile\ 630\Delta erm$ .

Achievement of efficient killing of transconjugant *C. difficile* cells using designed CGVs.

Materials and Methods

Bacterial Strains and Growth Conditions

 $\it E.~coli$  strain CA434 was acquired from Chain Biotech. It was cultured on nutrient-rich media (2xYT) and grown overnight at 37° C. and 250 rpm. Medium was supplemented with 12.5  $\mu g/mL$  of thiamphenicol when required to maintain the CGVs.

Clostridium difficile  $630\Delta erm$  was grown on BHI agar supplemented with 5 g/L of yeast extract, 0.03% L-cystein, 250 ug/ml D-cycloserine and 8 ug/ml of cefoxitin (BHIS+CC). C. difficile was grown overnight in a Coy vinyl anaerobic cabinet in an atmosphere of 92% N<sub>2</sub>, 6% CO<sub>2</sub> and 2% H<sub>2</sub> at  $37^{\circ}$  C. The mating of the donor CA434 and C. difficile was grown on plain BHI agar to allow for growth of the donor strain. Thiamphenicol was added to BHIS+CC plates to a final concentration of  $12.5 \mu g/mL$  for selection of transconjugants after mating. All plates were dried for 1.5

hours and transferred, along with the broth version of this medium, to the anaerobic chamber at least 3 hours before

### **CGV Transfer Procedures**

Carrier cells of E. coli CA434 were obtained by elec- 5 troporation of either of our CGVs (control vector pMTL84151-FJ797649 and CRISPR vector pMTL84151cdCRISPR1). In order to do that, overnight cultures of E. coli CA434 were diluted 1:100 in fresh 2x YT medium without selection and grown to OD600~0.5. Then, they were 10 made electrocompetent by standard procedures (Sharan et al., 2009). Electrocompetent cells were transformed with either plasmid pMTL84151-FJ797649 or pMTL84151-cd-CRISPR1 and recovered in 2xYT for 1 h at 37° C. with shaking (250 rpm). Finally, they were plated on LB agar 15 supplemented with 12.5 µg/mL thiamphenicol for selection of transformants. Transformants were grown in liquid 2xYT supplemented with 12.5 μg/mL thiamphenicol at 37° C. and 250 rpm for mating with  $\tilde{C}$ . difficile. 1 ml of donor cells was centrifuged at 4000×g for 2 minutes, supernatant removed 20 and carefully washed with 400 µl of PBS. After a second centrifugation cycle the pellet was transferred to the anaerobic chamber for mating with C. difficile in BHI non-selective plates. C. difficile was prepared for mating following a modified protocol (Des Purdy et al., 2002). C. difficile 25 630∆erm was incubated overnight in selective BHIS+CC plates, from which, a scrape was inoculated overnight in 1 ml of non-selective BHI and incubated over night for mating. 200 µl of that culture was used to resuspend the pelleted donor cells and mixed culture was plated in 20 µl spots on top of non-selective BHI plates. The mating was incubated 24 h to allow for conjugation. After incubation, the whole plate was thoroughly scraped with a sterile inoculation loop, resuspended in BHI and serial dilutions were plated on BHI+CC plates to prevent growth of donor

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E. coli and on BHI+CC supplemented with thiamphenicol for additional selection of transconjugants. Single colonies were counted after 48 hours.

Replicates of BHI+CC+Thiamphenicol plates, selecting for C. difficile transconjugants carrying the control CGV, showed a consistent number of colonies resulting in about  $\sim$ 600-750 CFUs per mating experiment. For the mating of C. difficile with E. coli CA434 carrying the CGV with the CRISPR array the plates were empty, no colonies were observed. This translates into 100% killing of transconjugant C. difficile 630Δerm cells receiving the CRISPR array (see FIG. 7: Killing of transconjugant C. difficile 630Δerm).

### DISCUSSION AND CONCLUSIONS

The results of this experiment show that we could successfully conjugate CGVs containing the desired CRISPR arrays into C. difficile 630\Delta erm from an E. coli carrier bacterium. We could also successfully harness C. difficile endogenous Cas3 machinery for very efficient CRISPR killing.

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		TABLE 1					
	Example Bacteria Optionally, the bacteria are selected from this Table.						
Abiotrophia Abiotrophia defectiva	Acidocella Acidocella aminolytica	Actinomyces Actinomyces bovis	Alkalilimnicola Alkalilimnicola ehrlichii	Aquaspirillum Aquaspirillum polymorphum			
Acaricomes Acaricomes phytoseiuli Acetitomaculum Acetitomaculum ruminis Acetivibrio Acetivibrio cellulolyticus Acetivibrio ethanolgignens Acetivibrio multivorans	Acidocella facilis Acidomonas Acidomonas methanolica Acidothermus Acidothermus cellulolyticus Acidovorax Acidovorax anthurii Acidovorax caeni	Actinomyces denticolens Actinomyces europaeus Actinomyces georgiae Actinomyces gerencseriae Actinomyces hordeovulneris Actinomyces howellii Actinomyces hyovaginalis	Alkaliphilus Alkaliphilus oremlandii Alkaliphilus transvaalensis Allochromatium Allochromatium vinosum Alloiococcus Alloiococcus Alloiokutzneria	Aquaspirillum putridiconchylium Aquaspirillum serpens Aquimarina Aquimarina latercula Arcanobacterium Arcanobacterium haemolyticum			
Acetoanaerobium Acetoanaerobium noterae Acetobacter	Acidovorax cattleyae Acidovorax citrulli Acidovorax defluvii	Actinomyces israelii Actinomyces johnsonii Actinomyces meyeri	Allokutzneria albata  Altererythrobacter Altererythrobacter ishigakiensis	Arcanobacterium pyogenes Archangium Archangium gephyra			
Acetobacter aceti Acetobacter cerevisiae Acetobacter cibinongensis Acetobacter estunensis Acetobacter fabarum	Acidovorax delafieldii Acidovorax facilis Acidovorax konjaci Acidovorax temperans Acidovorax valerianellae	Actinomyces naeslundii Actinomyces neuii Actinomyces odontolyticus Actinomyces oris Actinomyces radingae	Altermonas Altermonas haloplanktis Altermonas macleodii Alysiella Alysiella crassa	Arcobacter Arcobacter butzleri Arcobacter cryaerophilus Arcobacter halophilus Arcobacter nitrofigilis			
Acetobacter ghanensis Acetobacter indonesiensis Acetobacter lovaniensis Acetobacter malorum Acetobacter nitrogenifigens	Acinetobacter Acinetobacter baumannii Acinetobacter baylyi Acinetobacter bouvetii Acinetobacter calcoaceticus	Actinomyces slackii Actinomyces turicensis Actinomyces viscosus Actinoplanes Actinoplanes auranticolor	Alysiella filiformis Aminobacter Aminobacter aganoensis Aminobacter aminovorans Aminobacter niigataensis	Arcobacter skirrowii Arhodomonas Arhodomonas aquaeolei Arsenophonus Arsenophonus nasoniae			
Acetobacter oeni Acetobacter orientalis Acetobacter orleanensis Acetobacter pasteurianus Acetobacter pornorurn	Acinetobacter gerneri Acinetobacter haemolyticus Acinetobacter johnsonii Acinetobacter junii Acinetobacter lwoffi	Actinoplanes brasiliensis Actinoplanes consettensis Actinoplanes deccanensis Actinoplanes derwentensis Actinoplanes digitatis	Aminobacterium Aminobacterium mobile Aminomonas Aminomonas paucivorans Ammoniphilus	Arthrobacter Arthrobacter agilis Arthrobacter albus Arthrobacter aurescens Arthrobacter chlorophenolicus			

## TABLE 1-continued

Example Bacteria
Optionally, the bacteria are selected from this Table.

Optionally, the bacteria are selected from this Table.						
Acetobacter senegalensis Acetobacter xylinus	Acinetobacter parvus Acinetobacter radioresistens	Actinoplanes durhamensis Actinoplanes ferrugineus	Ammoniphilus oxalaticus Ammoniphilus oxalivorans	Arthrobacter citreus Arthrobacter crystallopoietes		
Acetobacterium	Acinetobacter schindleri	Actinoplanes globisporus	Amphibacillus	Arthrobacter cumminsii		
Acetobacterium bakii	Acinetobacter soli	Actinoplanes humidus	Amphibacillus xylanus	Arthrobacter globiformis		
Acetobacterium carbinolicum	Acinetobacter tandoii	Actinoplanes italicus	Amphritea	Arthrobacter		
Acetobacterium dehalogenans	Acinetobacter tjernbergiae	Actinoplanes liguriensis	Amphritea balenae	histidinolovorans		
Acetobacterium fimetarium	Acinetobacter towneri	Actinoplanes lobatus	Amphritea japonica	Arthrobacter ilicis		
Acetobacterium malicum Acetobacterium paludosum	Acinetobacter ursingii Acinetobacter venetianus	Actinoplanes missouriensis Actinoplanes palleronii	Amycolatopsis Amycolatopsis alba	Arthrobacter luteus Arthrobacter		
Acetobacterium tundrae	Acrocarpospora	Actinoplanes philippinensis	Amycolatopsis albidoflavus	methylotrophus Arthrobacter mysorens		
Acetobacterium wieringae	Acrocarpospora corrugata	Actinoplanes rectilineatus	Amycolatopsis azurea	Arthrobacter nicotianae		
Acetobacterium woodii	Acrocarpospora	Actinoplanes regularis	Amycolatopsis coloradensis	Arthrobacter nicotinovorans		
Acetofilamentum	macrocephala	Actinoplanes	Amycolatopsis lurida	Arthrobacter oxydans		
Acetofilamentum rigidum	Acrocarpospora pleiomorpha	teichomyceticus	Amycolatopsis mediterranei	Arthrobacter pascens		
Ace to halo bium	Actibacter	Actinoplanes utahensis	Amycolatopsis rifamycinica	Arthrobacter		
Acetohalobium arabaticum	Actibacter sediminis	Actinopolyspora	Amycolatopsis rubida	phenanthrenivorans		
Acetomicrobium	Actinoalloteichus	Actinopolyspora halophila	Amycolatopsis sulphurea	Arthrobacter		
Acetomicrobium faecale Acetomicrobium flavidum	Actinoalloteichus cyanogriseus	Actinopolyspora mortivallis Actinosynnema	Amycolatopsis tolypomycina Anabaena	polychromogenes Atrhrobacter		
1 aatawawa	Antiunallatainhua	Antimonium and antimon	tuah asua sulindaisa	protophormiae		
Acetonema Acetonema longum	Actinoalloteichus hymeniacidonis	Actinosynnema mirum Actinotalea	Anabaena cylindrica Anabaena flos-aquae	Arthrobacter psychrolactophilus		
Acetothermus	Actinoalloteichus spitiensis	Actinotalea fermentans	Anabaena variabilis	Arthrobacter ramosus		
Acetothermus paucivorans	Actinobaccillus	Aerococcus	Anaeroarcus	Arthrobacter sulfonivorans		
Acholeplasma Acholeplasma axanthum	Actinobacillus capsulatus Actinobacillus delphinicola	Aerococcus sanguinicola Aerococcus urinae	Anaeroarcus burkinensis Anaerobaculum	Arthrobacter sulfureus Arthrobacter uratoxydans		
Acholeplasma brassicae	Actinobacillus hominis	Aerococcus urinaeequi	Anaerobaculum mobile	Arthrobacter ureafaciens		
Acholeplasma cavigenitalium	Actinobacillus indolicus	Aerococcus urinaehominis	Anaerobiospirillum	Arthrobacter viscosus		
Acholeplasma equifetale	Actinobacillus lignieresii	Aerococcus viridans	Anaerobiospirillum	Arthrobacter woluwensis		
Acholeplasma granularum	Actinobacillus minor	Aeromicrobium	succiniciproducens	Asaia		
Acholeplasma hippikon	Actinobacillus muris	Aeromicrobium erythreum	Anaerobiospirillum thomasii	Asaia bogorensis		
Acholeplasma laidlawii	Actinobacillus	Aeromonas	Anaerococcus	Asanoa		
Acholeplasma modicum Acholeplasma morum	pleuropneumoniae Actinobacillus porcinus	Aeromonas allosaccharophila	Anaerococcus hydrogenalis Anaerococcus lactolyticus	Asanoa ferruginea Asticcacaulis		
Acholeplasma multilocale	Actinobacillus rossii	Aeromonas bestiarum	Anaerococcus prevotii	Asticcacaulis biprosthecium		
Acholeplasma oculi	Actinobacillus scotiae	Aeromonas caviae	Anaerococcus tetradius	Asticcacaulis excentricus		
Acholeplasma palmae	Actinobacillus seminis	Aeromonas encheleia	Anaerococcus vaginalis	Atopobacter		
Acholeplasma parvum	Actinobacillus succinogenes	Aeromonas	Anaerofustis	Atopobacter phocae		
Acholeplasma pleciae	Actinobaccillus suis	enteropelogenes	Anaerofustis stercorihominis	Atopobium		
Acholeplasma vituli	Actinobacillus ureae	Aeromonas eucrenophila	Anaeromusa	Atopobium fossor		
Achromobacter	Actinobaculum	Aeromonas ichthiosmia	Anaeromusa acidaminophila	Atopobium minutum		
Achromobacter denitrificans	Actinobaculum massiliense	Aeromonas jandaei	Anaeromyxobacter	Atopobium parvulum		
Achromobacter insolitus	Actinobaculum schaalii	Aeromonas media	Anaeromyxobacter	Atopobium rimae		
Achromobacter piechaudii Achromobacter ruhlandii	Actinobaculum suis Actinomyces urinale	Aeromonas popoffii Aeromonas sobria	dehalogenans Anaerorhabdus	Atopobium vaginae Aureobacterium		
Achromobacter spanius	Actinocatenispora	Aeromonas veronii	Anaerorhabdus furcosa	Aureobacterium barkeri		
Acidaminobacter	Actinocatenispora rupis	Agrobacterium	Anaerosinus	Aurobacterium		
Acidaminobacter	Actinocatenispora	Agrobacterium	Anaerosinus glycerini	Aurobacterium		
	•	3	3,	liquefaciens		
hydrogenoformans	thailandica	gelatinovorum	Anaerovirgula	Avibacterium		
Acidaminococcus	Actinocatenispora sera	Agrococcus	Anaerovirgula multivorans	Avibacterium avium		
Acidaminococcus fermentans	Actinocorallia	Agrococcus citreus	Ancalomicrobium	Avibacterium gallinarum		
Acidaminococcus intestini	Actinocorallia aurantiaca	Agrococcus jenensis	Ancalomicrobium adetum	Avibacterium paragallinarum		
Acidicaldus	Actinocorallia aurea	Agromonas	Ancylobacter	Avibacterium volantium		
Acidicaldus organivorans	Actinocorallia cavernae	Agromonas oligotrophica	Ancylobacter aquaticus	Azoarcus		
Acidimicrobium	Actinocorallia glomerata	Agromyces	Aneurinibacillus	Azoarcus indigens		
Acidimicrobium ferrooxidans	Actinocorallia herbida	Agromyces fucosus	Aneurinibacillus aneurinilyticus	Azoarcus tolulyticus		
Acidiphilium	Actinocorallia libanotica	Agromyces hippuratus	Aneurinibacillus migulanus	Azoarcus toluvorans		
Acidiphilium acidophilum	Actinocorallia longicatena	Agromyces luteolus	Aneurinibacillus	Azohydromonas		
Acidiphilium angustum	Actinomadura	Agromyces mediolanus	thermoaerophilus	Azohydromonas		
				australica		
Acidiphilium cryptum	Actinomadura alba	Agromyces ramosus	Angiococcus	Azohydromonas lata		
Acidiphilium multivorum	Actinomadura atramentaria	Agromyces rhizospherae	Angiococcus disciformis	Azomonas		
Acidiphilium organovorum	Actinomadura	Akkermansia	Angulomicrobium	Azomonas agilis		
Acidiphilium rubrum	bangladeshensis	Akkermansia muciniphila	Angulomicrobium tetraedrale	Azomonas insignis		
Acidisoma	Actinomadura catellatispora	Albidiferax	Anoxybacillus	Azomonas		
				macrocytogenes		

phenazinium

### TABLE 1-continued

		TABLE 1-continued		
	Optionally,	Example Bacteria the bacteria are selected from t	his Table.	
Acidisoma sibiricum Acidisoma tundrae	Actinomadura chibensis Actinomadura chokoriensis	Albidiferax ferrireducens Albidovulum	Anoxybacillus pushchinoensis Aquabacterium	Azorhizobium Azorhizobium caulinodans
Acidisphaera Acidisphaera rubrifaciens Acidithiobacillus Acidithiobacillus albertensis Acidithiobacillus caldus	Actinomadura citrea Actinomadura coerulea Actinomadura echinospora Actinomadura fibrosa Actinomadura formosensis	Albidovulum inexpectatum Alcaligenes Alcaligenes denitrificans Alcaligenes faecalis Alcanivorax	Aquabacterium commune Aquabacterium parvum	Azorhizophilus Azorhizophilus paspali Azospirillum Azospirillum brasilense Azospirillum halopraeferens
Acidithiobacillus ferrooxidans Acidithiobacillus thiooxidans Acidobacterium Acidobacterium capsulatum	Actinomadura hibisca Actinomadura kijaniata Actinomadura latina Actinomadura livida	Alcanivorax borkumensis Alcanivorax jadensis Algicola Algicola bacteriolytica		Azospirillum irakense Azotobacter Azotobacter beijerinckii Azotobacter chroococcum
	Actinomadura luteofluorescens Actinomadura macra Actinomadura madurae Actinomadura oligospora Actinomadura rubrobrunea Actinomadura rugatobispora Actinomadura umbrina Actinomadura verrucosospora Actinomadura vinacea Actinomadura viridilutea Actinomadura viridilutea Actinomadura viridis	Alicyclobacillus Alicyclobacillus disulfidooxidans Alicyclobacillus sendaiensis Alicyclobacillus vulcanalis Alishewanella Alishewanella fetalis Alkalibacillus haloalkaliphilus		chrococcum Azotobacter nigricans Azotobacter salinestris Azotobacter vinelandii
Bacillus [see below] Bacteriovorax Bacteriovorax stolpii	Actinomadura yumaensis Bacteroides Bacteroides caccae Bacteroides coagulans Bacteroides eggerthii Bacteroides fragilis	Bibersteinia Bibersteinia trehalosi Bifidobacterium Bifidobacterium adolescentis Bifidobacterium angulatum	Borrelia Borrelia afzelii Borrelia americana Borrelia burgdorferi Borrelia carolinensis	Brevinema Brevinema andersonii Brevundimonas Brevundimonas alba Brevundimonas
	Bacteroides galacturonicus Bacteroides helcogenes	Bifidobacterium animalis Bifidobacterium asteroides	Borrelia coriaceae Borrelia garinii	aurantiaca Brevundimonas diminuta Brevundimonas
	Bacteroides ovatus	Bifidobacterium bifidum	Borrelia japonica	intermedia Brevundimonas
	Bacteroides pectinophilus	Bifidobacterium bourn	Bosea	subvibrioides Brevundimonas
	Bacteroides pyogenes	Bifidobacterium breve	Bosea minatitlanensis	vancanneytii Brevundimonas variabilis
	Bacteroides salyersiae	Bifidobacterium catenulatum	Bosea thiooxidans	Brevundimonas vesicularis
	Bacteroides stercoris Bacteroides suis	Bifidobacterium choerinum	Brachybacterium Brachybacterium	Brochothrix
	Bacteroides tectus	Bifidobacterium coryneforme Bifidobacterium cuniculi	alimentarium	Brochothrix campestris Brochothrix thermosphacta
	Bacteroides thetaiotaomicron Bacteroides uniformis	Bifidobacterium dentium Bifidobacterium gallicum	Brachybacterium faecium Brachybacterium	Brucella Brucella canis Brucella neotomae
	Bacteroides ureolyticus Bacteroides vulgatus Balnearium Balnearium lithotrophicum	Bifidobacterium gallinarum Bifidobacterium indicum Bifidobacterium longum Bifidobacterium	paraconglomeratum Brachybacterium rhamnosum Brachybacterium tyrofermentans	Bryobacter Bryobacter aggregatus Burkholderia
	Balneatrix Balneatrix alpica	magnumBifidobacterium merycicum	Brachyspira Brachyspira alvinipulli	Burkholderia ambifaria Burkholderia andropogonis
	Balneola Balneola vulgaris Barnesiella Barnesiella viscericola	Bifidobacterium minimum Bifidobacterium pseudocatenulatum Bifidobacterium	Brachyspira hyodysenteriae Brachyspira innocens Brachyspira murdochii Brachyspira pilosicoli	Burkholderia anthina Burkholderia caledonica Burkholderia caryophylli Burkholderia cenocepacia
	Bartonella Bartonella alsatica	pseudolongum Bifidobacterium pullorum	Bradyrhizobium Bradyrhizobium canariense	Burkholderia cepacia Burkholderia cocovenenans
	Bartonella bacilliformis Bartonella clarridgeiae Bartonella doshiae Bartonella elizabethae Bartonella grahamii Bartonella henselae Bartonella rochalimae	Bifidobacterium ruminantium Bifidobacterium saeculare Bifidobacterium subtile Bifidobacterium thermophilum Bilophila Bilophila wadsworthia	Bradyrhizobium elkanii Bradyrhizobium japonicum Bradyrhizobium liaoningense Brenneria Brenneria alni Brenneria nigrifluens Brenneria quercina	Burkholderia dolosa Burkholderia fungorum Burkholderia glathei Burkholderia glumae Burkholderia graminis Burkholderia kururiensis Burkholderia
	Bartonella vinsonii	Biostraticola	Brenneria quercina	multivorans Burkholderia phenazinium

B. firmus

B. flexus

B. foraminis

chitinolyticus

B. chondroitinus

B. choshinensis

### TABLE 1-continued

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## Example Bacteria Optionally, the bacteria are selected from this Table.

Bavariicoccus Biostraticola tofi Brenneria salicis Burkholderia plantarii Burkholderia pyrrocinia Bavariicoccus seileri Brevibacillus Bizionia Bdellovibrio Bizionia argentinensis Brevibacillus agri Burkholderia silvatlantica Bdellovibrio bacteriovorus Brevibacillus borstelensis Burkholderia stabilis Blastobacter Bdellovibrio exovorus Blastobacter capsulatus Brevibacillus brevis Burkholderia thailandensis Burkholderia tropica Beggiatoa Blastobacter denitrificans Brevibacillus centrosporus Beggiatoa alba Blastococcus Brevibacillus choshinensis Burkholderia unamae Beijerinckia Blastococcus aggregatus Brevibacillus invocatus Burkholderia vietnamiensis Beijerinckia derxii Blastococcus saxobsidens Brevibacillus laterosporus Buttiauxella Beijerinckia fluminensis Blastochloris Brevibacillus parabrevis Buttiauxella agrestis Beijerinckia indica Blastochloris viridis Brevibacillus reuszeri Buttiauxella brennerae Beijerinckia mobilis Blastomonas Brevibacterium Buttiauxella ferragutiae Blastomonas natatoria Belliella Brevibacterium abidum Buttiauxella gaviniae Belliella baltica Blastopirellula Brevibacterium album Buttiauxella izardii Blastopirellula marina Blautia Bellilinea Buttiauxella noackiae Brevibacterium aurantiacum Bellilinea caldifistulae Brevihacterium celere Buttiauxella warmboldiae Blautia coccoides Brevibacterium epidermidis Butvrivibrio Belnapia Butyrivibrio fibrisolvens Butyrivibrio hungatei Belnapia moabensis **Brevihacterium** Blautia hansenii Bergeriella frigoritolerans Blautia producta Bergeriella denitrificans Brevibacterium halotolerans Butyrivibrio Blautia wexlerae proteoclasticus Beutenhergia Bogoriella Rrevihacterium iodinum Bogoriella caseilytica Brevibacterium linens Beutenbergia cavernae Bordetella Brevibacterium lyticum Bordetella avium Brevibacterium mcbrellneri Bordetella bronchiseptica Brevihacterium otitidis Bordetella hinzii Brevibacterium oxydans Bordetella holmesii Brevibacterium paucivorans Bordetella parapertussis Brevibacterium stationis Bordetella pertussis Bordetella petrii Bordetella trematum Bacillus B. acidiceler B. aminovorans B. glucanolyticus B. taeanensis B. lautus B. acidicola B. amylolyticus B. gordonae B. tequilensis B. lehensis B. lentimorbus B. acidiproducens B. andreesenii B. gottheilii B. thermantarcticus B. acidocaldarius B. aneurinilyticus B. graminis B. thermoaerophilus B. lentus B. acidoterrestris B. anthracis B. halmapalus B. thermoamylovorans B. licheniformis B. aeolius B. aquimaris B. haloalkaliphilus B. thermocatenulatus B. ligniniphilus B. aerius B. arenosi B. halochares B. thermocloacae B. litoralis B. aerophilus B. arseniciselenatis B. halodenitrificans B. thermocopriae B. locisalis B. agaradhaerens B. arsenicus B. halodurans B. thermodenitrificans B. luciferensis B. aurantiacus B. halophilus B. thermoglucosidasius B. luteolus B. agri B. aidingensis B. arvi B. halosaccharovorans B. thermolactis B. luteus B. akibai B. aryabhattai B. hemicellulosilyticus B. thermoleovorans B. macauensis B. alcalophilus B. asahii B. hemicentroti B. thermophilus B. macerans B. algicola B. atrophaeus B. herbersteinensis B. thermoruber B. macquariensis B. alginolyticus B. axarquiensis B. horikoshii B. thermosphaericus B. macyae B. alkalidiazotrophicus B. azotofixans B. horneckiae B. thiaminolyticus B. malacitensis B. alkalinitrilicus B. azotoformans B. horti B. thioparans B. mannanilyticus B. huizhouensis B. marisflavi B. alkalisediminis B. badius B. thuringiensis B. alkalitelluris B. barbaricus B. humi B. tianshenii B. marismortui B. hwajinpoensis B. altitudinis B. bataviensis B. trypoxylicola B. marmarensis B. beijingensis B. idriensis B. tusciae B. alveayuensis B. massiliensis B. alvei B. benzoevorans B. indicus B. validus B. megaterium B. amyloliquefaciens B. beringensis B. infantis B. vallismortis B. mesonae B. berkelevi B. infernus B. vedderi B. methanolicus B. beveridgei a. subsp. amyloliquefaciens B. insolitus B. velezensis B. methylotrophicus B. a. subsp. plantarum B. bogoriensis B. invictae B. migulanus B. vietnamensis B. dipsosauri B. mojavensis B. boroniphilus B. iranensis R. vireti B. mucilaginosus
B. muralis B. drentensis B. borstelensis B. isabeliae B. vulcani B. wakoensis B. edaphicus B. brevis Migula B. isronensis B. butanolivorans B. weihenstephanensis B. ehimensis B. jeotgali B. murimartini B. kaustophilus B. eiseniae B. canaveralius B. xiamenensis B. mycoides B. kobensis B. kochii B. enclensis B. carboniphilus B. naganoensis B. xiaoxiensis B. cecembensis B. zhanjiangensis B. endophyticus B. nanhaiensis B. kokeshiiformis B. nanhaiisediminis B. endoradicis B. cellulosilyticus B. peoriae B. persepolensis B. farraginis B. centrosporus B. koreensis B. nealsonii B. fastidiosus B. cereus B. korlensis B. persicus B. neidei B. chagannorensis B. fengqiuensis B. kribbensis B. pervagus B. neizhouensis

B. krulwichiae

B. larvae

B. laevolacticus

B. plakortidis

B. polygoni

B. pocheonensis

B. niabensis

B. niacini

B. novalis

### TABLE 1-continued

		Examp	ole 1	Bacteria			
Optionally,	the	bacteria	are	selected	from	this	Table.

Optionally, the bacteria are selected from this Table.						
B. fordii B. formosus B. fortis B. finarioli B. finiculus B. fusiformis B. galactophilus B. galactosidilyticus B. galactosidilyticus B. gelatini B. gibsonii B. ginsengi B. ginsengihumi B. ginsengisoli B. globisporus (eg, B. g. subsp. Globisporus; or B. g. subsp. Marinus)  Caenimonas	B. chungangensis B. cibi B. circulans B. clarkii B. clausii B. coagulans B. coahuilensis B. cohnii B. composti B. corposti B. cytloheptanicus B. cytotoxicus B. daliensis B. decisifrondis B. decolorationis B. deserti	B. laterosporus B. salexigens B. saliphilus B. schegelii B. sediminis B. seleniarsenatis B. selenitireducens B. seohaeanensis B. shacketenii B. siamensis B. sincheensis B. shackletonii B. siamensis B. silvestris B. simplex B. siralis B. smithii B. soli B. soli B. solisalsi B. songklensis B. sonorensis B. sonorensis B. sporothermodurans B. stearothermophilus B. stratosphericus B. subterraneus B. subterraneus B. subtilis (eg. B. s. subsp. Spizizeni; or B.	B. polymyxa B. popilliae B. pseudalcalophilus B. pseudomycoides B. psychrodurans B. psychrosaccharolyticus B. psychrosaccharolyticus B. psychrotolerans B. pulyifaciens B. pulyifaciens B. purgationiresistens B. pycnus B. qingdaonensis B. qingdaonensis B. qingshengii B. reuszeri B. rhizosphaerae B. rigui B. ruris B. safensis B. salarius  Catenuloplanes	B. oceanisediminis B. odysseyi B. okhensis B. okhensis B. oleronius B. oryzaecorticis B. oshimensis B. pabuli B. pakistanensis B. pallidus B. pallidus B. panacisoli B. panaciterrae B. panaciterrae B. paraflexus B. paraflexus B. paragoniensis		
Caenimonas koreensis Caldalkalibacillus Caldalkalibacillus Caldanaerobacter Caldanaerobacter subterraneus Caldanaerobius Caldanaerobius Caldanaerobius polysaccharolyticus Caldanaerobius zeae Caldanaerobius zeae Caldanaerobius acetigignens Caldanaerovirga acetigignens Caldicellulosiruptor Caldicellulosiruptor bescii Caldicellulosiruptor kristjanssonii Caldicellulosiruptor owensensis	Campylobacter coli Campylobacter concisus Campylobacter curvus Campylobacter fetus Campylobacter fetus Campylobacter helveticus Campylobacter hominis Campylobacter hominis Campylobacter lari Campylobacter lari Campylobacter mucosalis Campylobacter rectus Campylobacter rectus Campylobacter sputorum Campylobacter sputorum Campylobacter sputorum Campylobacter upsaliensis Capnocytophaga Capnocytophaga canimorsus Capnocytophaga cynodegmi Capnocytophaga gingivalis Capnocytophaga granulosa Capnocytophaga granulosa Capnocytophaga achacea Capnocytophaga sputigena	Cardiobacterium hominis Carnimonas Carnimonas nigrificans Carnobacterium Carnobacterium alterfunditum Carnobacterium divergens Carnobacterium funditum	Catenuloplanes atrovinosus Catenuloplanes castaneus Catenuloplanes crispus Catenuloplanes indicus Catenuloplanes indicus Catenuloplanes nepalensis Catenuloplanes niger Chryseobacterium Chryseobacterium balustinum Citrobacter C. amalonaticus C. braakii C. diversus C. farmeri  C. freundii C. gillenii C. koseri C. murliniae C. pasteurii <sup>[1]</sup> C. rodentium C. sedlakii C. werkmanii C. youngae Clostridium (see below) Coccochloris Cocynebacterium flavescens Corynebacterium variabile	Curtobacterium albidum Curtobacterium citreus		

### Clostridium

Clostridium absonum, Clostridium aceticum, Clostridium acetireducens, Clostridium acetobutylicum, Clostridium acidisoli, Clostridium aciditolerans, Clostridium aciditolerans, Clostridium aerotolerans, Clostridium aerotolerans, Clostridium algidacarni, Clostridium algidacylanolyticum, Clostridium algifaecis, Clostridium algoriphilum, Clostridium alkalicellulosi, Clostridium aminophilum, Clostridium aminophilum, Clostridium aminophilum, Clostridium aminophilum, Clostridium aminophilum, Clostridium apparagiforme, Clostridium aurantibutyricum, Clostridium autoethanogenum, Clostridium baratii, Clostridium barkeri, Clostridium barlettii, Clostridium beijerinckii, Clostridium bipermentans, Clostridium bolteae, Clostridium bornimense, Clostridium botulinum, Clostridium bowmanii, Clostridium bryantii, Clostridium butyricum, Clostridium cadaveris, Clostridium caenicola, Clostridium caminithermale, Clostridium carboxidivorans, Clostridium carnis, Clostridium cavendishii, Clostridium cellulogicum, Clostridium cellulogicum, Clostridium cellulogicum, Clostridium cellulogis, Clostridium cellulogis, Clostridium cellulogis, Clostridium cellulogis, Clostridium clariflavum, Clostridium clostridium coccoides, Clostridium clostridium colletant, Clostridium disporicum, Clostridium darakei, Clostridium durum, Clostridium estertheticum estertheticum estertheticum estertheticum laramiense, Clostridium fallax, Clostridium felsineum,

### TABLE 1-continued

## Example Bacteria Optionally, the bacteria are selected from this Table.

Clostridium fervidum, Clostridium fimetarium, Clostridium formicaceticum, Clostridium frigidicarnis, Clostridium frigoris, Clostridium ganghwense, Clostridium gasigenes, Clostridium ghonii, Clostridium glycolicum, Clostridium glycyrrhizinilyticum, Clostridium grantii, Clostridium haemolyticum, Clostridium halophilum, Clostridium hastiforme, Clostridium hathewayi, Clostridium herbivorans, Clostridium hiranonis, Clostridium histolyticum, Clostridium homopropionicum, Clostridium huakuii, Clostridium hungatei, Clostridium hydrogeniformans, Clostridium hydroxybenzoicum, Clostridium hylemonae, Clostridium jejuense, Clostridium indolis, Clostridium innocuum, Clostridium intestinale, Clostridium irregulare, Clostridium isatidis, Clostridium josui, Clostridium kluyveri, Clostridium lactatifermentans, Clostridium lacusfryxellense, Clostridium laramiense, Clostridium lavalense, Clostridium lentocellum, Clostridium lentoputrescens, Clostridium leptum, Clostridium limosum, Clostridium litorale, Clostridium lituseburense, Clostridium ljungdahlii, Clostridium lortetii, Clostridium lundense, Clostridium magnum, Clostridium malenominatum, Clostridium mangenotii, Clostridium mayombei, Clostridium methoxybenzovorans, Clostridium methylpentosum, Clostridium neopropionicum, Clostridium nexile, Clostridium nitrophenolicum, Clostridium novyi, Clostridium oceanicum, Clostridium orbiscindens, Clostridium oroticum, Clostridium oxalicum, Clostridium papyrosolvens, Clostridium paradoxum, Clostridium paraperfringens (Alias: C. welchii), Clostridium paraputrificum, Clostridium pascui, Clostridium pasteurianum, Clostridium peptidivorans, Clostridium perenne, Clostridium perfringens, Clostridium pfennigii, Clostridium phytofermentans, Clostridium piliforme, Clostridium polysaccharolyticum, Clostridium populeti, Clostridium propionicum, Clostridium proteoclasticum, Clo Clostridium putrefaciens, Clostridium putrificum, Clostridium quercicolum, Clostridium quinii, Clostridium ramosum, Clostridium rectum, Clostridium roseum, Clostridium saccharobutylicum, Clostridium saccharogumia, Clostridium saccharolyticum, Clostridium saccharoperbutylacetonicum, Clostridium sardiniense, Clostridium sartagoforme, Clostridium scatologenes, Clostridium schirmacherense, Clostridium scindens, Clostridium septicum, Clostridium sordellii, Clostridium sphenoides, Clostridium spiroforme, Clostridium sporogenes, Clostridium sporosphaeroides, Clostridium stercorarium, Clostridium stercorarium leptospartum, Clostridium stercorarium stercorarium, Clostridium stercorarium thermolacticum, Clostridium sticklandii, Clostridium straminisolvens, Clostridium subterminale, Clostridium sufflavum, Clostridium sulfidigenes, Clostridium symbiosum, Clostridium tagluense, Clostridium tepidiprofundi, Clostridium termitidis, Clostridium tertium, Clostridium tetani, Clostridium tetanomorphum, Clostridium thermaceticum, Clostridium thermautotrophicum, Clostridium thermoalcaliphilum, Clostridium thermobutyricum, Clostridium thermocellum, Clostridium thermocopriae, Clostridium thermohydrosulfuricum, Clostridium thermolacticum, Clostridium thermopalmarium, Clostridium thermopapyrolyticum, Clostridium thermosaccharolyticum, Clostridium thermosuccinogenes, Clostridium thermosulfurigenes, Clostridium thiosulfatireducens, Clostridium tyrobutyricum, Clostridium uliginosum, Clostridium ultunense, Clostridium villosum, Clostridium vincentii, Clostridium viride, Clostridium xylanolyticum, Clostridium xylanovorans Dactylosporangium Deinococcus Delftia Echinicola Dactylosporangium Deinococcus aerius Delftia acidovorans Echinicola pacifica aurantiacum Dactylosporangium fulvum Deinococcus apachensis Desulfovibrio Echinicola vietnamensis Dactylosporangium Deinococcus aquaticus Desulfovibrio desulfuricans matsuzakiense Dactylosporangium roseum Deinococcus aquatilis Diplococcus Dactylosporangium Deinococcus caeni Diplococcus pneumoniae thailandense Dactylosporangium vinaceum Deinococcus radiodurans Deinococcus radiophilus Enterobacter Enterobacter kobei Faecalibacterium Flavobacterium E. aerogenes E. ludwigii Faecalibacterium prausnitzii Flavobacterium antarcticum E. amnigenus E. mori Flavobacterium aquatile Fangia E. agglomerans E. nimipressuralis Fangia hongkongensis Flavobacterium aquidurense E. arachidis E. oryzae Fastidiosipila Flavobacterium balustinum E. asburiae E. pulveris Fastidiosipila sanguinis Flavobacterium croceum E. cancerogenous E. pyrinus Fusobacterium Flavobacterium cucumis E. radicincitans Fusobacterium nucleatum Flavobacterium daejeonense E. cloacae E. taylorae Flavobacterium defluvii E. cowanii E. dissolvens E. turicensis Flavobacterium degerlachei E. gergoviae E. sakazakii Enterobacter soli Flavobacterium E. helveticus Enterococcus denitrificans E. hormaechei Enterococcus durans Flavobacterium filum E. intermedius Enterococcus faecalis Flavobacterium flevense Enterococcus faecium Flavobacterium frigidarium Erwinia Flavobacterium mizutaii Erwinia hapontici Flavobacterium Escherichia okeanokoites Escherichia coli Gaetbulibacter Haemophilus Ideonella Janibacter Gaetbulibacter Haemophilus aegyptius Ideonella azotifigens Janibacter anophelis saemankumensis Haemophilus aphrophilus Janibacter corallicola Gallibacterium Idiomarina Haemophilus felis Gallibacterium anatis Idiomarina abvssalis Janibacter limosus Haemophilus gallinarum Gallicola Idiomarina baltica Janibacter melonis Haemophilus haemolyticus Gallicola barnesae Idiomarina fontislapidosi Janibacter terrae Haemophilus influenzae Garciella Idiomarina loihiensis Jannaschia Haemophilus paracuniculus Garciella nitratireducens Idiomarina ramblicola Jannaschia cystaugens Geobacillus Haemophilus Jannaschia helgolandensis Idiomarina seosinensis parahaemolyticus Geobacillus Haemophilus parainfluenzae Idiomarina zobellii Jannaschia pohangensis thermoglucosidasius Geobacillus Haemophilus Ignatzschineria Jannaschia rubra stearothermophilus paraphrohaemolyticus Geobacter Ignatzschineria larvae Janthinobacterium Geobacter bemidjiensis Haemophilus parasuis Ignavigranum Janthinobacterium Geobacter bremensis Haemophilus pittmaniae Ignavigranum ruoffiae agaricidamnosum Geobacter chapellei Hafnia Ilumatobacter Janthinobacterium lividum Geobacter grbiciae Hafnia alvei Ilumatobacter fluminis

Ilyobacter

Geobacter hydrogenophilus

Hahella

Jejuia pallidilutea

### TABLE 1-continued

	Ontionally	Example Bacteria	his Table	
Geobacter lovleyi	Optionally,  Hahella ganghwensis	The bacteria are selected from the		
Geobacter metallireducens Geobacter pelophilus Geobacter pickeringii	Halalkalibacillus Halalkalibacillus halophilus Helicobacter	Ilyobacter delafieldii Ilyobacter insuetus Ilyobacter polytropus Ilyobacter tartaricus	Jeotgalibacillus Jeotgalibacillus alimentarius Jeotgalicoccus	
Geobacter sulfurreducens Geodermatophilus Geodermatophilus obscurus Gluconacetobacter Gluconacetobacter xylinus Gordonia	Helicobacter pylori		Jeotgalicoccus halotolerans	
Gordonia rubripertincta Kaistia	Labedella	Listeria ivanovii	Micrococcus	Nesterenkonia
Kaistia adipata	Labedella gwakjiensis	L. marthii	Micrococcus luteus	Nesterenkonia holobia
Kaistia soli	Labrenzia	L. monocytogenes	Micrococcus lylae	Nocardia
Kangiella Vanciella assimanina	Labrenzia aggregata Labrenzia alba	L. newyorkensis L. riparia	Moraxella Moraxella bovis	Nocardia argentinensis Nocardia corallina
Kangiella aquimarina Kangiella koreensis	Labrenzia alexandrii	L. riparia L. rocourtiae	Moraxella nonliquefaciens	Nocardia coranina Nocardia
Kerstersia	Labrenzia marina	L. seeligeri	Moraxella osloensis	otitidiscaviarum
Kerstersia gyiorum	Labrys	L. weihenstephanensis	Nakamurella	
Kiloniella	Labrys methylaminiphilus	L. welshimeri	Nakamurella multipartita	
Kiloniella laminariae	Labrys miyagiensis	Listonella	Nannocystis	
Klebsiella V. znamelomatia	Labrys monachus	Listonella anguillarum	Nannocystis pusilla Natranaerobius	
K. granulomatis K. oxytoca	Labrys okinawensis Labrys portucalensis	Macrococcus Macrococcus bovicus	Natranaerobius Natranaerobius	
K. pneumoniae	Lactobacillus	Marinobacter	thermophilus	
K. terrigena	[see below]	Marinobacter algicola	Natranaerobius trueperi	
K. variicola	Laceyella	Marinobacter bryozoorum	Naxibacter	
Kluyvera	Laceyella putida	Marinobacter flavimaris	Naxibacter alkalitolerans	
Kluyvera ascorbata Kocuria	Lechevalieria Lechevalieria	Meiothermus Meiothermus ruber	Neisseria Neisseria cinerea	
Nocuna	aerocolonigenes	Meioinermus ruber	iveisseria cinerea	
Kocuria roasea	Legionella	Methylophilus	Neisseria denitrificans	
Kocuria varians	[see below]	Methylophilus methylotrophus		
Kurthia	Listeria	Microbacterium	Neisseria lactamica	
Kurthia zopfii	L. aquatica	Microbacterium	Neisseria mucosa	
	L. booriae L. cornellensis	ammoniaphilum Microbacterium arborescens	Neisseria sicca Neisseria subflava	
	L. fleischmannii	Microbacterium liquefaciens	Neptunomonas	
	L. floridensis	Microbacterium oxydans	Neptunomonas japonica	
	L. grandensis			
	L. grayi			
Lactobacillus	L. innocua			
L. acetotolerans	L. catenaformis	L. mali	L. parakefiri	L. sakei
L. acidifarinae	L. ceti	L. manihotivorans	L. paralimentarius	L. salivarius
L. acidipiscis	L. coleohominis	L. mindensis	L. paraplantarum	L. sanfranciscensis
L. acidophilus Lactobacillus agilis	L. collinoides L. composti	L. mucosae L. murinus	L. pentosus L. perolens	L. satsumensis L. secaliphilus
L. algidus	L. concavus	L. nagelii	L. plantarum	L. sharpeae
L. alimentarius	L. coryniformis	L. namurensis	L. pontis	L. siliginis
L. amylolyticus	L. crispatus	L. nantensis	L. protectus	L. spicheri
L. amylophilus	L. crustorum	L. oligofermentans	L. psittaci	L. suebicus
L. amylotrophicus	L. curvatus	L. oris	L. rennini	L. thailandensis
L. amylovorus	L. delbrueckii subsp. bulgaricus	L. panis	L. reuteri	L. ultunensis
L. animalis	L. delbrueckii subsp.	L. pantheris	L. rhamnosus	L. vaccinostercus
L. antri	delbrueckii	L. parabrevis	L. rimae	L. vaginalis
L. apodemi	L. delbrueckii subsp. lactis	L. parabuchneri	L. rogosae	L. versmoldensis
L. aviarius	L. dextrinicus	L. paracasei	L. rossiae	L. vini
L. bifermentans	L. diolivorans	L. paracollinoides	L. ruminis	L. vitulinus
L. brevis L. buchneri	L. equi L. equigenerosi	L. parafarraginis	L. saerimneri L. jensenii	L. zeae
L. camelliae	L. equigenerosi L. farraginis	L. homohiochii L. iners	L. johnsonii	L. zymae L. gastricus
L. casei	L. farciminis	L. ingluviei	L. kalixensis	L. ghanensis
L. kitasatonis	L. fermentum	L. intestinalis	L. kefiranofaciens	L. graminis
L. kunkeei	L. fornicalis	L. fuchuensis	L. kefiri	L. hammesii
L. leichmannii	L. fructivorans	L. gallinarum	L. kimchii	L. hamsteri
L. lindneri	L. frumenti	L. gasseri	L. helveticus	L. harbinensis
L. malefermentans Legionella			L. hilgardii	L. hayakitensis
Legionella adelaidensis	Legionella drancourtii	Candidatus Legionella jeonii	Legionella quinlivanii	
Legionella anisa	Legionella dresdenensis	Legionella jordanis	Legionella rowbothamii	
Legionella beliardensis	Legionella drozanskii	Legionella lansingensis	Legionella rubrilucens	
Legionella birminghamensis	Legionella dumoffii	Legionella londiniensis	Legionella sainthelensi	
Legionella bozemanae	Legionella erythra	Legionella longbeachae	Legionella santicrucis	
Legionella brunensis	Legionella fairfieldensis	Legionella lytica	Legionella shakespearei	
Legionella busanensis	Legionella fallonii	Legionella maceachernii	Legionella spiritensis	

### TABLE 1-continued

## Example Bacteria Optionally, the bacteria are selected from this Table.

Legionella cardiaca Legionella cherrii Legionella cincinnatiensis Legionella clemsonensis Legionella donaldsonii Legionella feeleii
Legionella geestiana
Legionella genomospecies
Legionella gormanii
Legionella gratiana
Legionella gresilensis
Legionella hackeliae
Legionella impletisoli
Legionella israelensis
Legionella jamestowniensis

Legionella massiliensis
Legionella micdadei
Legionella mornovica
Legionella moravica
Legionella nagasakiensis
Legionella nautarum
Legionella norrlandica
Legionella oakridgensis
Legionella parisiensis
Legionella pittsburghensis
Legionella preumophila
Legionella quateirensis
Prevotella

Legionella steelei
Legionella steigerwaltii
Legionella taurinensis
Legionella tucsonensis
Legionella tunisiensis
Legionella wadsworthii
Legionella waltersii
Legionella worsleiensis
Legionella yabuuchiae

Oceanibulbus
Oceanibulbus indolifex
Oceanicaulis
Oceanicaulis alexandrii
Oceanicola
Oceanicola batsensis
Oceanicola granulosus
Oceanicola nanhaiensis
Oceanimonas
Oceanimonas baumannii
Oceaniserpentilla
Oceaniserpentilla haliotis

Paenibacillus
Paenibacillus thiaminolyticus
Pantoea
Pantoea agglomerans
Paracoccus
Paracoccus alcaliphilus
Paucimonas
Paucimonas lemoignei
Pectobacterium
Pectobacterium atrosepticum
Pectobacterium atrosepticum
Pectobacterium
Pectobacterium
Pectobacterium
Pectobacterium
Pectobacterium
Pectobacterium carticida
Pectobacterium carnegieana

Prevotella albensis Prevotella amnii Prevotella bergensis Prevotella bivia Prevotella brevis Prevotella buccae Prevotella buccais Prevotella copri Prevotella dentalis Prevotella dentalis Quadrisphaera
Quadrisphaera granulorum
Quatrionicoccus
Quatrioni coccus
australiensis
Quinella
Quinella ovalis
Ralstonia
Ralstonia eutropha
Ralstonia insidiosa
Ralstonia mannitolilytica
Ralstonia pickettii

Oceanisphaera
Oceanisphaera donghaensis
Oceanisphaera litoralis
Oceanithermus
Oceanithermus desulfurans
Oceanithermus profundus
Oceanobacillus
Oceanobacillus caeni
Oceanospirillum
Oceanospirillum

Pectobacterium atrosepticum
Pectobacterium
betavasculorum
Pectobacterium carticida
Pectobacterium carticida
Pectobacterium carotovorum
Pectobacterium chrysanthemi
Pectobacterium rhapontici
Pectobacterium wasabiae
Planococcus
Planococcus citreus
Planomicrobium
Planomicrobium okeanokoites
Plesiomonas
Plesiomonas shigelloides
Proteus vulgaris

Prevotella disiens Prevotella histicola Prevotella intermedia Prevotella maculosa Prevotella marshii Prevotella melaninogenica Prevotella micans Prevotella multiformis Prevotella nigrescens Prevotella oralis Prevotella oris Prevotella oulorum Prevotella pallens Prevotella salivae Prevotella stercorea Prevotella tannerae Prevotella timonensis Prevotella veroralis Providencia Providencia stuartii Pseudomonas Pseudomonas aeruginosa Pseudomonas alcaligenes Pseudomonas anguillispetica

Pseudomonas fluorescens Pseudoalteromonas haloplanktis Pseudomonas mendocina Pseudomonas pseudoalcaligenes Pseudomonas putida Pseudomonas tutzeri Pseudomonas syringae Pseudomobacter Ralstonia pseudosolanacearum Ralstonia syzygii Ralstonia solanacearum Ramlibacter Ramlibacter henchirensis Ramlibacter tataouinensis Raoultella Raoultella ornithinolytica Raoultella planticola Raoultella terrigena Rathayibacter Rathayibacter caricis Rathayibacter festucae Rathayibacter iranicus Rathayibacter rathayi Rathayibacter toxicus Rathayibacter tritici Rhodobacter Rhodobacter sphaeroides Ruegeria Ruegeria gelatinovorans

Saccharococcus
Saccharococcus thermophilus
Saccharomonospora azurea
Saccharomonospora cyanea
Saccharomonospora viridis
Saccharophagus
Saccharophagus degradans
Saccharopolyspora
Saccharopolyspora erythraea
Saccharopolyspora gregorii
Saccharopolyspora hirsuta
Saccharopolyspora hordei
Saccharopolyspora
Saccharopolyspora
Rectivirgula

Sagittula
Sagittula stellata
Salegentibacter
Salegentibacter salegens
Salimicrobium
Salimicrobium album
Salinibacter ruber
Salinicoccus
Salinicoccus alkaliphilus
Salinicoccus roseus
Salinispora
Salinispora arenicola

Psychrobacter faecalis Psychrobacter phenylpyruvicus Sanguibacter Sanguibacter keddieii Sanguibacter suarezii Saprospira Saprospira grandis Sarcina Sarcina maxima Sarcina ventriculi Sebaldella Sebaldella termitidis Serratia Serratia fonticola Serratia marcescens Sphaerotilus

Stenotrophomonas
Stenotrophomonas
maltophilia
Streptococcus
[also see below]
Streptomyces
Streptomyces
achromogenes
Streptomyces cesalbus
Streptomyces cescaepitosus
Streptomyces cesdiastaticus
Streptomyces cesexfoliatus
Streptomyces fimbriatus
Streptomyces fradiae

Tatlockia
Tatlockia maceachemii
Tatlockia micdadei
Tenacibaculum
Tenacibaculum
Tenacibaculum discolor
Tenacibaculum
gallaicum
Tenacibaculum
tenacibaculum
Tenacibaculum
lutimaris
Tenacibaculum
mesophilum
Tenacibaculum

### TABLE 1-continued

## Example Bacteria Optionally, the bacteria are selected from this Table.

Saccharopolyspora spinosa Saccharopolyspora taberi Saccharothrix australiensis Saccharothrix coeruleofusca Saccharothrix espanaensis Saccharothrix longispora Saccharothrix mutabilis Saccharothrix syringae Saccharothrix tangerinus Saccharothrix texasensis Salinispora tropica Salinivibrio Salinivibrio costicola Salmonella Salmonella bongori Salmonella enterica Salmonella subterranea Salmonella typhi

Sphaerotilus natans Sphingobacterium Sphingobacterium multivorum Staphylococcus [see below] Streptomyces fulvissimus Streptomyces griseoruber Streptomyces griseus Streptomyces lavendulae Streptomyces phaeochromogenes Streptomyces thermodiastaticus Streptomyces tubercidicus

S. schleiferi

S. sciuri

S. simiae

S. simulans

S. succinus

S. vitulinus

S. warneri

S. xylosus

S. stepanovicii

skagerrakense
Tepidanaerobacter
Tepidanaerobacter
syntrophicus
Tepidibacter
Tepidibacter
formicigenes
Tepidibacter thalassicus
Thermus
Thermus aquaticus
Thermus filiformis
Thermus thermophilus

Staphylococcus

S. arlettae
S. agnetis
S. aureus
S. aureus
S. capitis
S. caprae
S. carnosus
S. caseolyticus
S. chromogenes
S. condimenti
S. delphini
S. devriesei
S. epidermidis

Streptococcus Streptococcus agalactiae Streptococcus anginosus Streptococcus bovis Streptococcus canis Streptococcus constellatus Streptococcus downei Streptococcus dysgalactiae Streptococcus equines Streptococcus faecalis Streptococcus ferus Uliginosibacterium Uliginosibacterium gangwonense Ulvibacter Ulvibacter litoralis Umezawaea Umezawaea tangerina Undibacterium Undibacterium pigrum Ureaplasma Ureaplasma urealyticum Ureibacillus Ureibacillus composti Ureibacillus suwonensis Ureibacillus terrenus Ureibacillus thermophilus Ureibacillus thermosphaericus

S. equorum
S. felis
S. fleurettii
S. gallinarum
S. haemolyticus
S. hominis
S. hyicus
S. intermedius
S. kloosii
S. leei
S. lentus
S. lugdunensis
S. lutrae

Streptococcus infantarius

Streptococcus intermedius

Streptococcus lactarius

Streptococcus milleri

Streptococcus mutans

Streptococcus tigurinus

Vagococcus carniphilus

Vagococcus elongatus

Vagococcus fessus

Vagococcus lutrae

Vagococcus fluvialis

Vagococcus salmoninarum

Variovorax boronicumulans

Veillonella montpellierensis Veillonella parvula

Venenivibrio stagnispumantis

Verminephrobacter eiseniae

Verrucomicrobium spinosum

Variovorax dokdonensis

Variovorax paradoxus

Variovorax soli Veillonella

Veillonella atypica

Veillonella caviae

Veillonella criceti

Veillonella dispar

Veillonella ratti

Venenivibrio

Veillonella rodentium

Verminephrobacter

Verrucomicrobium

Streptococcus oralis

Vagococcus

Variovorax

Streptococcus mitis

Streptococcus iniae

S. lyticans S. massiliensis S. microti
S. muscae
S. nepalensis
S. pasteuri
S. petrasii
S. pettenkoferi
S. piscifermentans
S. pseudintermedius
S. pseudolugdunensis
S. pulvereri
S. rostri
S. saccharolyticus
S. saprophyticus

Streptococcus orisratti
Streptococcus parasanguinis
Streptococcus peroris
Streptococcus pneumoniae
Streptococcus
pseudopneumoniae
Streptococcus pyogenes
Streptococcus ratti
Streptococcus ratti

Vibrio Vibrio aerogenes

Vibrio albensis

Vibrio aestuarianus

Vibrio alginolyticus Vibrio campbellii Vibrio cholerae Vibrio cincinnatiensis Vibrio coralliilyticus Vibrio cyclitrophicus Vibrio diazotrophicus Vibrio fluvialis Vibrio furnissii Vibrio gazogenes Vibrio halioticoli Vibrio harveyi Vibrio ichthyoenteri Vibrio mediterranei Vibrio metschnikovii Vibrio mytili Vibrio natriegens Vibrio navarrensis Vibrio nereis Vibrio nigripulchritudo Vibrio ordalii Vibrio orientalis Vibrio parahaemolyticus Vibrio pectenicida Vibrio penaeicida Vibrio proteolyticus Vibrio shilonii Vibrio splendidus Vibrio tubiashii Vibrio vulnificus

Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus suis Streptococcus uberis Streptococcus viridans Streptococcus viridans Streptococcus zooepidemicus

Virgibacillus Virgibacillus

halodenitrificans Virgibacillus pantothenticus Weissella Weissella cibaria Weissella confusa Weissella halotolerans Weissella hellenica Weissella kandleri Weissella koreensis Weissella minor Weissella paramesenteroides Weissella soli Weissella thailandensis Weissella viridescens Williamsia Williamsia marianensis Williamsia maris Williamsia serinedens Winogradskyella Winogradskyella thalassocola Wolbachia Wolbachia persica Wolinella Wolinella succinogenes Zobellia Zobellia galactanivorans Zobellia uliginosa Zoogloea Zoogloea ramigera Zoogloea resiniphila

Xanthobacter Xanthobacter agilis

Xanthobacter aminoxidans Xanthobacter autotrophicus Xanthobacter flavus Xanthobacter tagetidis Xanthobacter viscosus Xanthomonas Xanthomonas albilineans Xanthomonas alfalfae Xanthomonas arboricola Xanthomonas axonopodis Xanthomonas campestris Xanthomonas citri Xanthomonas codiaei Xanthomonas cucurbitae Xanthomonas euvesicatoria Xanthomonas fragariae Xanthomonas fuscans Xanthomonas gardneri Xanthomonas hortorum Xanthomonas hvacinthi Xanthomonas perforans Xanthomonas phaseoli Xanthomonas pisi Xanthomonas populi Xanthomonas theicola Xanthomonas translucens

### TABLE 1-continued

Example Bacteria Optionally, the bacteria are selected from this Table.						
Xenophilus Xenophilus azovorans Xenorhabdus beddingii Xenorhabdus bovienii Xenorhabdus cabanillasii Xenorhabdus doucetiae Xenorhabdus griffiniae Xenorhabdus kominickii Xenorhabdus koppenhoeferi Xenorhabdus nematophila Xenorhabdus poinarii Xylanibacter Xylanibacter oryzae	Yangia Yangia pacifica Yaniella flava Yaniella flava Yaniella halotolerans Yeosuana Yeosuana aromativorans Yersinia Yersinia aldovae Yersinia bercovieri Yersinia enterocolitica Yersinia entomophaga Yersinia frederiksenii Yersinia intermedia Yersinia kristensenii	Yersinia mollaretii Yersinia philomiragia Yersinia pestis Yersinia roseudotuberculosis Yersinia rouckeri Yokenella Yokenella regensburgei Yonghaparkia Yonghaparkia alkaliphila Zavarzinia compransoris	Zooshikella Zooshikella ganghwensis Zunongwangia Zunongwangia profunda Zymobacter Zymobacter palmae Zymomonas Zymomonas mobilis Zymophilus paucivorans Zymophilus raffinosivorans	Xanthomonas vesicatoria Xylella fastidiosa Xylophilus ampelinus Zobellella Zobellella denitrifican Zobellella taiwanensis Zeaxanthinibacter zeaxanthinibacter enoshimensis Zhihengliuella halotolerans Xylanibacterium Xylanibacterium ulmi		

TABLE 2

TABLE 2			. 25	TABLE 3-continued			
Finding Temperature Heart rate	>90/min	° F.) or >38° C. (100.4° F.)	30	Sequences of genomic targets to selective  The 3-bp PAM are shown in bold ty  Expected SpCas9-gRNA cleavage sites at  (3 bp upstream of the		d type (ie, tgg, agg and cgg).	
Respiratory rate WBC	<4 × 10 <sup>9</sup> /L (<	CO2 <32 mmHg (4.3 kPa) 4000/mm <sup>3</sup> ), >12 × 10 <sup>9</sup> /L ), or 10% bands	•	Target strain	Target gene	Genomic target sequence	
TABLE 3  Sequences of genomic targets to selectively CRISPR-kill <i>E. coli</i> strains.  The 3-bp PAM are shown in bold type (ie, tgg, agg and cgg).  Expected SpCas9-gRNA cleavage sites are indicated by vertical arrows (3 bp upstream of the PAM).		35	E. coli Nissle 1917	уарН	caccegcattacetttatgcagg (SEQ ID NO: 2)		
Target strain	Target gene	Genomic target sequence		E. coli Nissle 1917	pks	1	
E. coli (EHEC) ATCC43888	23S ribosomal RNA	taggtgaagtccctcgcgga <b>tgg</b> (SEQ ID NO: 1)	45			gcgcgggtgtggttgtgctt <b>cgg</b> (SEQ ID NO: 3)	

TABLE 4

Seq	uences of	genomic targets & Cas to selectively CRISPR-kill C dificile
	(a) Overvi	lew of design spacers in pMTL84151 - cdCRISPR1 targeting  C. difficile
Gene	Name PAM	target
tcdA	TA2 CCA	ACACCTTAACCCAGCCATAGAGTCTGATAATAACTTC (SEQ ID NO: 4)
tcdB	TB2 CCA	GACTTATTTGAGTCTATAGAGAAACCTAGTTCAGTAA (SEQ ID NO: 5)
rRNA- 1	TRR1 CCT	TCGACGACTTCTTCCAAAAGGTTAGATAATCGGCTTC (SEQ ID NO: 6)
rRNA- 2	TRR2 CCA	$ \begin{array}{ll} {\tt GTACAGGATGGACCCGCGTCTGATTAGCTAGTTGGTA} & ({\tt SEQ\ ID} \\ {\tt NO:\ 7)} \end{array} $

gyrA TAR1 CCA TCCTCATGGAGATACTGCTGTTTATTATGCTATGGTA (SEQ ID NO: 8)

(b) Cas3 sequence (SEQ ID NO: 9) Sequence

TCAAATAAATTGGTCTATTTCATTACTTATAAGCACACCTTTACCAAATATCTGTT TTGTATGCTTATTTTCGTATATATCATATTTATATAAAAGTATTTTTAAATCTTCAA GACCTTTCACCTGTATGTCAATTACATTTTGTTTAGCTTTATATATTTGGTAAATTTA AATTGTCTATCATCATTACTATATGCATTTATAAGCTCTTGTCCCAACTCTTCATAA TCTTTGATTAAAGTTTCTTCAATTTTATTATATATATCCTCTGGAATTACTGTATAT CCTTCAATATTTCTAAGTATATTTTGTGCATCTTTAGAACCTAAACTATATGGCGTT  ${\tt ATAGTATCTAAAATATTTAAAGCACTGGTAAATCTCTTTTCAAAAGCTGTACCTTC}$ CTTCACACATTCTTTTCCATTTATAAAAGTTTTCAGTAGCTCAAGTCCTTTTTCTAC ATTTGGGCTATTTCTTCATACTCACGACTTCTATAACATCTACCAAATCGTTGAA ATAGACTGTCAAGTGTTGAATTTTCTGTATGAAGCTCATCAAAATCAATATCAAG GGATGCTTCCACTAATTGTGTAGTAATCCAAATACCATTACTATCACTATCTGCAA ATTCTTTTATATATTTTTCTAATTTTGCTCTATCTTCTTGTATATACATAGAATGTA AAAGATTTAAGTTGACATCTATTCCTTTTGGCTTAACTATTTCTTCTATTAACTCAT CACTTTGTATTATTTTACCTAAGTTTTCATCTATTGAATTTTCTACAATAGACACAC AATGTCTTATTTTTCTGTATTACATGTCAATTCAGCTAAGTTACTATTCATTACAC CTCTTTTTTTAATTCATCTATATATATGGTTGGCATAGTAGCTGTCATTATCATAA ATCTGCCACCTATCTTATGTATCATTTCTATACCTTTTACCAATACAGCTGCTATTT  $\tt CTGGTGAATATGCTTGTATCTCATCTATTACTACTTTTGAATATGCTAATGTTGAGT$ ACACTTTTCATACCCTCTATACAAAAAAGGAAATTTAAATATTTGGTCTATTGTA  ${\tt GAAAATGTCAGTTTGCAAGATAATAACTTTGCTAAATCTACAATCTCACTTGAATT}$  ${\tt TTCTTGATTACTTTCTTGGATAATCTATTGCTGTTGAATGTAACAAACCTAAGA}$ ATGTATCATTTGATTCTCCAACTCCAACTATATTTTTTTGCTCTATCAAATAATGCAT  ${\tt AAGGCAGTTTCTGTTTTTCCCATTCCTGTAGGTGCAATCAGTATTATATTCTTATTT}$ CTATTAGATTTAGCAAATGATTGAGCCTCTCTCAAACTACCAAACTCTTTCATTAA ATAATTTTCTGTCTGTTCTCCTATATTTATAACATTATTGCATTCCACAACTTCATG AGCAGAAGCACTGTGGTCTAATCTATGTAGTATTCCTTTTAACATAATATATAAAT TATAGTACTTGTGATTTTTATCTATTCTTTTTTTCTACACTTTGTAGATATACTTTACT CAATTTTTCTGTTTTTATTGGATATCTAACTTTAAATTCATGCTGTAGTTCATAAAC TTTATTTATAAATCTTCATCTAATATTTTTTTGTATTAAAATTTTTAAAATCTTTATCT ATAAAGATATCTCTTTCATGATGATATACAATAACTTGATTTAATACTGCTCTAAG TTCTTTATTTTTTCCTGTCTATATAACTATAATCAATAAATGCAGGAGAAAGAT TATCCATTCTGCTTTTTATTAACTCTTGAAACGGTGAAAATGCTTTTCCAATATCAT GAAATTCTATAACAAAGTCAAGTAATTGCCAAAATATCTCCTCTTCTAGAAAATCT AAGCTATTTATATTTTTCCATAACTTTCTCTTAATACATTCATTTGTTTTAAAAGT TCATCAGTATGTTCTCTAAGTGTTTCCACTGGATTAGATTTAGCATATAACAT

Sequences of Cas9 used to selectively kill E coli. SEQ ID NO: 10 (Cas9 nucleotide sequence)  $\tt ATGGATAAGAAATACTCAATAGGCTTAGATATCGGCACAAATAGCGTCGGATGGGCGGT$ GATCACTGATGAATATAAGGTTCCGTCTAAAAAGTTCAAGGTTCTGGGAAATACAGACC  $\tt GCCACAGTATCAAAAAAATCTTATAGGGGCTCTTTTATTTGACAGTGGAGAGACAGCG$ GAAGCGACTCGTCTCAAACGGACAGCTCGTAGAAGGTATACACGTCGGAAGAATCGTAT TCGACTTGAAGAGTCTTTTTTGGTGGAAGAAGACAAGAAGCATGAACGTCATCCTATTTT AAAAAATTGGTAGATTCTACTGATAAAGCGGATTTGCGCTTAATCTATTTGGCCTTAGC GCATATGATTAAGTTTCGTGGTCATTTTTTGATTGAGGGAGATTTAAATCCTGATAATAGT AGACGATTAGAAAATCTCATTGCTCAGCTCCCCGGTGAGAAGAAAATGGCTTATTTGG GAATCTCATTGCTTTGTCATTGGGTTTGACCCCTAATTTTAAATCAAATTTTGATTTGGCA GAAGATGCTAAATTACAGCTTTCAAAAGATACTTACGATGATGATTTAGATAATTTATTG GCGCAAATTGGAGATCAATATGCTGATTTGTTTTTTGGCAGCTAAGAATTTATCAGATGCT ATTTTACTTTCAGATATCCTAAGAGTAAATACTGAAATAACTAAGGCTCCCCTATCAGCT TCAATGATTAAACGCTACGATGAACATCATCAAGACTTGACTCTTTTAAAAGCTTTAGTT TGCAGGTTATATTGATGGGGGAGCTAGCCAAGAAGAATTTTATAAATTTATCAAACCAAT TTTAGAAAAATGGATGGTACTGAGGAATTATTGGTGAAACTAAATCGTGAAGATTTGCT GCGCAAGCAACGGACCTTTGACAACGGCTCTATTCCCCATCAAATTCACTTGGGTGAGCT GCATGCTATTTTGAGAAGACAAGAAGACTTTTATCCATTTTTAAAAGACAATCGTGAGAA  ${\tt GATTGAAAAAATCTTGACTTTTCGAATTCCTTATTATGTTGGTCCATTGGCGCGTGGCAAT}$ AGTCGTTTTGCATGGATGACTCGGAAGTCTGAAGAAACAATTACCCCATGGAATTTTGAA  ${\tt GAAGTTGTCGATAAAGGTGCTTCAGCTCAATCATTTATTGAACGCATGACAAACTTTGAT}$  ${\tt AAAAATCTTCCAAATGAAAAAGTACTACCAAAACATAGTTTGCTTTATGAGTATTTTACG}$ GTTTATAACGAATTGACAAAGGTCAAATATGTTACTGAAGGAATGCGAAAACCAGCATT  ${\tt TCTTTCAGGTGAACAGAAGAAGCCATTGTTGATTTACTCTTCAAAACAAATCGAAAAGT}$ 

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TABLE 4-continued

Sequences of genomic targets & Cas to selectively CRISPR-kill C dificile

AACCGTTAAGCAATTAAAAGAAGATTATTTCAAAAAAATAGAATGTTTTGATAGTGTTTTG AATTATTAAAGATAAAGATTTTTTGGATAATGAAGAAAATGAAGATATCTTAGAGGATA  $\tt TTGTTTTAACATTGACCTTATTTGAAGATAGGGAGATGATTGAGGAAAGACTTAAAACAT$  $\tt ATGCTCACCTCTTTGATGATAAGGTGATGAAACAGCTTAAACGTCGCCGTTATACTGGTT$ GGGGACGTTTGTCTCGAAAATTGATTAATGGTATTAGGGATAAGCAATCTGGCAAAACA ATATTAGATTTTTTGAAATCAGATGGTTTTGCCAATCGCAATTTTATGCAGCTGATCCATG ATGATAGTTTGACATTTAAAGAAGACATTCAAAAAGCACAAGTGTCTGGACAAGGCGAT AGTTTACATGAACATATTGCAAATTTAGCTGGTAGCCCTGCTATTAAAAAAGGTATTTTA CAGACTGTAAAAGTTGTTGATGAATTGGTCAAAGTAATGGGGCGGCATAAGCCAGAAAA TATCGTTATTGAAATGGCACGTGAAAATCAGACAACTCAAAAGGGCCAGAAAAATTCGC GAGAGCGTATGAAACGAATCGAAGAAGGTATCAAAGAATTAGGAAGTCAGATTCTTAAA GAGCATCCTGTTGAAAATACTCAATTGCAAAATGAAAAGCTCTATCTCTATTATCTCCAA AATGGAAGAGACATGTATGTGGACCAAGAATTAGATATTAATCGTTTAAGTGATTATGAT  $\tt GTCGATCACATTGTTCCACAAAGTTTCCTTAAAGACGATTCAATAGACAATAAGGTCTTA$ ACGCGTTCTGATAAAAATCGTGGTAAATCGGATAACGTTCCAAGTGAAGAAGTAGTCAA AAAGATGAAAAACTATTGGAGACAACTTCTAAACGCCAAGTTAATCACTCAACGTAAGT  $\tt TTGATAATTTAACGAAAGCTGAACGTGGAGGTTTGAGTGAACTTGATAAAGCTGGTTTTA$  ${\tt TCAAACGCCAATTGGTTGAAACTCGCCAAATCACTAAGCATGTGGCACAAATTTTGGATA}$ GTCGCATGAATACTAAATACGATGAAAATGATAAACTTATTCGAGAGGTTAAAGTGATT ACCTTAAAATCTAAATTAGTTTCTGACTTCCGAAAAGATTTCCAATTCTATAAAGTACGT  ${\tt GAGATTAACAATTACCATCATGCCCATGATGCGTATCTAAATGCCGTCGTTGGAACTGCT}$  $\tt TTGATTAAGAAATATCCAAAACTTGAATCGGAGTTTGTCTATGGTGATTATAAAGTTTAT$ GATGTTCGTAAAATGATTGCTAAGTCTGAGCAAGAAATAGGCAAAGCAACCGCAAAATA  $\tt TTTCTTTTACTCTAATATCATGAACTTCTTCAAAACAGAAATTACACTTGCAAATGGAGA$ GATTCGCAAACGCCCTCTAATCGAAACTAATGGGGAAACTGGAGAAATTGTCTGGGATA AAGGGCGAGATTTTGCCACAGTGCGCAAAGTATTGTCCATGCCCCAAGTCAATATTGTCA AGAAAACAGAAGTACAGACAGGCGGATTCTCCAAGGAGTCAATTTTACCAAAAAGAAAT TCGGACAAGCTTATTGCTCGTAAAAAAGACTGGGATCCAAAAAAATATGGTGGTTTTGAT  ${\tt AGTCCAACGGTAGCTTATTCAGTCCTAGTGGTTGCTAAGGTGGAAAAAGGGAAATCGAA}$ GAAGTTAAAATCCGTTAAAGAGTTACTAGGGATCACAATTATGGAAAGAAGTTCCTTTG AAAAAAATCCGATTGACTTTTTAGAAGCTAAAGGATATAAGGAAGTTAAAAAAAGACTTA ATCATTAAACTACCTAAATATAGTCTTTTTGAGTTAGAAAACGGTCGTAAACGGATGCTG  $\tt GCTAGTGCCGGAGAATTACAAAAAGGAAATGAGCTGGCTCTGCCAAGCAAATATGTGAA$ TTTTTTATATTTAGCTAGTCATTATGAAAAGTTGAAGGGTAGTCCAGAAGATAACGAACA AAAACAATTGTTTGTGGAGCAGCATAAGCATTATTTAGATGAGATTATTTGAGCAAATCAG  $\tt GTTGACGAATCTTGGAGCTCCCGCTGCTTTTAAATATTTTGATACAACAATTGATCGTAA$ ACGATATACGTCTACAAAAGAAGTTTTAGATGCCACTCTTATCCATCAATCCATCACTGG TCTTTATGAAACACGCATTGATTTGAGTCAGCTAGGAGGTGACTGA

SEQ ID NO: 11 (Cas9 amino acid sequence)  $\verb|MDKKYSIGLDIGTNSVGWAVITDEYKVPSKKFKVLGNTDRHSIKKNLIGALLFDSGETAEAT|$  $\verb|RLKRTARRRYTRKNRICYLQEIFSNEMAKVDDSFFHRLEESFLVEEDKKHERHPIFGNIVDE|$ VAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDLNPDNSDVDKLFIQL  $\verb"VQTYNQLFEENPINASGVDAKAILSARLSKSRRLENLIAQLPGEKKNGLFGNLIALSLGLTPNF"$ KSNFDLAEDAKLOLSKDTYDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKA PLSASMIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKP ILEKMDGTEELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFLKDNREKIEK ILTFRIPYYVGPLARGNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNLPNE KVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVTVKQLKE DYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIVLTLTLFEDRE MIEERLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDFLKSDGFANRN FMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKGILQTVKVVDELVKVMGRH KPENIVIEMARENQTTQKGQKNSRERMKRIEEGIKELGSQILKEHPVENTQLQNEKLYLYYLQ  $\tt NGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLTRSDKNRGKSDNVPSEEVVKK$ MKNYWROLLNAKLI TORKFDNLTKAERGGLSELDKAGFIKROLVETROITKHVAOILDSRM NTKYDENDKLIREVKVITLKSKLVSDFRKDFOFYKVREINNYHHAHDAYLNAVVGTALIKKY PKLESEFVYGDYKVYDVRKMIAKSEOEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIET NGETGEIVWDKGRDFATVRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDW  ${\tt DPKKYGGFDSPTVAYSVLVVAKVEKGKSKKLKSVKELLGITIMERSSFEKNPIDFLEAKGYKE}$ VKKDLIIKLPKYSLFELENGRKRMLASAGELOKGNELALPSKYVNFLYLASHYEKLKGSPED NEQKQLFVEQHKHYLDEIIEQISEFSKRVILADANLDKVLSAYNKHRDKPIREQAENIIHLFTL TNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQSITGLYETRIDLSQLGGD

**115** TABLE 5

# 116 TABLE 5-continued

Publication	Gram-neg. bacteria as causative infectious pathogen in cancer patients	E. coli/K. pneumoniae/ P. aeruginosa as causative pathogens in bacteraemia in cancer patients [fraction of Gram-neg. cases]	5	Publication	Gram-neg. bacteria as causative infectious pathogen in cancer patients	E. coli/K. pneumoniae/ P. aeruginosa as causative pathogens in bacteraemia in cancer patients [fraction of Gram-neg. cases]
Samonis et al	65%	54% [85%]		Marin et al Anatoliotaki et al	55% 47%	51% [92%] 34% [73%]
Velasco et al	45%	33% [74%]				[]

### SEQUENCE LISTING

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His	Leu	Phe	Asp	Asp 645	Lys	Val	Met	Lys	Gln 650	Leu	Lys	Arg	Arg	Arg 655	Tyr
Thr	Gly	Trp	Gly 660	Arg	Leu	Ser	Arg	Lys 665	Leu	Ile	Asn	Gly	Ile 670	Arg	Asp
Lys	Gln	Ser 675	Gly	Lys	Thr	Ile	Leu 680	Asp	Phe	Leu	Lys	Ser 685	Asp	Gly	Phe
Ala	Asn 690	Arg	Asn	Phe	Met	Gln 695	Leu	Ile	His	Asp	Asp 700	Ser	Leu	Thr	Phe
Lys 705	Glu	Asp	Ile	Gln	Lys 710	Ala	Gln	Val	Ser	Gly 715	Gln	Gly	Asp	Ser	Leu 720
His	Glu	His	Ile	Ala 725	Asn	Leu	Ala	Gly	Ser 730	Pro	Ala	Ile	Lys	Lys 735	Gly
Ile	Leu	Gln	Thr 740	Val	rys	Val	Val	Asp 745	Glu	Leu	Val	rya	Val 750	Met	Gly
Arg	His	Lys 755	Pro	Glu	Asn	Ile	Val 760	Ile	Glu	Met	Ala	Arg 765	Glu	Asn	Gln
Thr	Thr 770	Gln	Lys	Gly	Gln	Lys 775	Asn	Ser	Arg	Glu	Arg 780	Met	ГЛа	Arg	Ile
Glu 785	Glu	Gly	Ile	ГÀа	Glu 790	Leu	Gly	Ser	Gln	Ile 795	Leu	Lys	Glu	His	Pro 800
Val	Glu	Asn	Thr	Gln 805	Leu	Gln	Asn	Glu	Lys 810	Leu	Tyr	Leu	Tyr	Tyr 815	Leu
Gln	Asn	Gly	Arg 820	Asp	Met	Tyr	Val	Asp 825	Gln	Glu	Leu	Asp	Ile 830	Asn	Arg
Leu	Ser	Asp 835	Tyr	Asp	Val	Asp	His 840	Ile	Val	Pro	Gln	Ser 845	Phe	Leu	TÀa
Asp	Asp 850	Ser	Ile	Asp	Asn	Lув 855	Val	Leu	Thr	Arg	Ser 860	Asp	Lys	Asn	Arg
Gly 865	Lys	Ser	Asp	Asn	Val 870	Pro	Ser	Glu	Glu	Val 875	Val	Lys	Lys	Met	880 TÀs
Asn	Tyr	Trp	Arg	Gln 885	Leu	Leu	Asn	Ala	Lys 890	Leu	Ile	Thr	Gln	Arg 895	ГЛа
Phe	Asp	Asn	Leu 900	Thr	Lys	Ala	Glu	Arg 905	Gly	Gly	Leu	Ser	Glu 910	Leu	Asp
Lys		Gly 915	Phe	Ile	Lys		Gln 920		Val	Glu		Arg 925	Gln	Ile	Thr
Lys	His 930	Val	Ala	Gln	Ile	Leu 935	Asp	Ser	Arg	Met	Asn 940	Thr	ГЛа	Tyr	Asp
Glu 945	Asn	Asp	ГЛа	Leu	Ile 950	Arg	Glu	Val	ГЛа	Val 955	Ile	Thr	Leu	TÀa	Ser 960
Lys	Leu	Val	Ser	Asp 965	Phe	Arg	ГÀа	Asp	Phe 970	Gln	Phe	Tyr	ГЛа	Val 975	Arg
Glu	Ile	Asn	Asn 980	Tyr	His	His	Ala	His 985	Asp	Ala	Tyr	Leu	Asn 990	Ala	Val
Val	Gly	Thr 995	Ala	Leu	Ile	Lys	Lys 1000		Pro	Lys	Leu	Glu 1005		Glu	Phe
Val	Tyr 1010		Asp	Tyr	Lys	Val 1015		Asp	Val	Arg	Lys 1020	Met	Ile	Ala	Lys
Ser 1025		Gln	Glu	Ile	Gly 1030		Ala	Thr	Ala	Lys 1035		Phe	Phe		Ser .040

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Asn	Ile	Met	Asn	Phe 104!		ГÀа	Thr	Glu	Ile 1050		Leu	Ala	Asn	Gly 1055	
Ile	Arg	Lys	Arg 1060		Leu	Ile	Glu	Thr 106		Gly	Glu	Thr	Gly 1070		Ile
Val	Trp	Asp 1075	_	Gly	Arg	Asp	Phe	Ala O	Thr	Val	Arg	1089		Leu	Ser
Met	Pro 1090		Val	Asn	Ile	Val 1095	•	Lys	Thr	Glu	Val		Thr	Gly	Gly
Phe 1105		Lys	Glu	Ser	Ile 1110		Pro	Lys		Asn 1115		Asp	Lys		Ile L120
Ala	Arg	Lys	Lys	Asp 112		Asp	Pro	Lys	Lys 1130		Gly	Gly	Phe	Asp 1135	
Pro	Thr	Val	Ala 1140	_	Ser	Val	Leu	Val 114		Ala	ГÀа	Val	Glu 1150	_	Gly
Lys	Ser	Lys 1155	_	Leu	Lys	Ser	Val 1160	Lys	Glu	Leu	Leu	Gly 116		Thr	Ile
Met	Glu 1170		Ser	Ser	Phe	Glu 1175		Asn	Pro	Ile	Asp		Leu	Glu	Ala
Lys 1185		Tyr	Lys	Glu	Val 1190		Lys	Asp	Leu	Ile 119		Lys	Leu		Lys 1200
Tyr	Ser	Leu	Phe	Glu 120!		Glu	Asn	Gly	Arg 1210		Arg	Met	Leu	Ala 121	
Ala	Gly	Glu	Leu 1220		Lys	Gly	Asn	Glu 122!		Ala	Leu	Pro	Ser 1230	_	Tyr
Val	Asn	Phe 1235		Tyr	Leu	Ala	Ser 124	His O	Tyr	Glu	Lys	Leu 124	-	Gly	Ser
Pro	Glu 1250		Asn	Glu	Gln	Lys 1255		Leu	Phe	Val	Glu 126		His	Lys	His
Tyr 1265		Asp	Glu	Ile	Ile 1270		Gln	Ile	Ser	Glu 1279		Ser	Lys	_	Val 1280
Ile	Leu	Ala	Asp	Ala 128		Leu	Asp	Lys	Val 1290		Ser	Ala	Tyr	Asn 1295	_
His	Arg	Asp	Lys 1300		Ile	Arg	Glu	Gln 130		Glu	Asn	Ile	Ile 1310		Leu
Phe	Thr	Leu 1315		Asn	Leu	Gly	Ala 1320	Pro	Ala	Ala	Phe	Lys 1325		Phe	Asp
Thr	Thr 1330		Asp	Arg	Lys	Arg 1335		Thr	Ser	Thr	Lys 134		Val	Leu	Asp
Ala 1345		Leu	Ile	His	Gln 135		Ile	Thr	Gly	Leu 1355		Glu	Thr		Ile 1360
Asp	Leu	Ser	Gln	Leu 136!	_	Gly	Asp								

The invention claimed is:

1. A method for treating an acute pathogenic bacterial infection in a subject caused by first bacteria that are *E. coli*, *Pseudomonas aeruginosa* or *Klebsiella* bacteria, the method comprising selectively killing the first bacteria comprised by the subject by cutting a target site comprised by the genomes of the first bacteria, wherein the cutting is carried out using a Cas nuclease that is programmed by a guide RNA to cut the target site, wherein the subject has a cancer, wherein the method is carried out no more than 7 days before or after administration of an immunotherapy to the subject, wherein 65 the programmed Cas nuclease treats the infection and the immunotherapy treats the cancer;

55 wherein:

- (a) the immunotherapy comprises administering to the subject an anti-PD-1/PD-L1 axis antibody; and
- (b) the cancer is selected from melanoma; renal cell carcinoma; bladder cancer; a solid tumour; non-small cell lung cancer (NSCLC); forehead and neck squamous cell carcinoma (HNSCC); Hodgkin's lymphoma; a cancer that overexpresses PD-L1 and has no mutations in EGFR or in ALK; colorectal cancer and hepatocellular carcinoma.
- **2**. The method of claim **1**, wherein the first bacteria are *E. coli*

- **3**. The method of claim **2**, wherein the first bacteria are enterohemorrhagic *E. coli* (EHEC).
- **4**. The method of claim **1**, wherein the anti-PD-1/PD-L1 axis antibody is selected from the group consisting of pembrolizumab, nivolumab, atezolimumab, avelumab and 5 durvalumab.
- 5. The method of claim 1, wherein the treatment of the bacterial infection is carried out simultaneously with the administration of the immunotherapy to the subject.
- **6**. The method of claim **1**, wherein the treatment of the 10 bacterial infection is carried out immediately before or after administering the immunotherapy to the subject.
- 7. The method of claim 1, wherein the method comprises administering to the subject a nucleic acid vector comprising the guide RNA or a DNA encoding the guide RNA.
- **8**. The method of claim **7**, comprising administering a second nucleic acid vector to the subject, wherein the second vector encodes the Cas nuclease.
- **9**. The method of claim **7**, wherein the Cas nuclease is an endogenous Cas nuclease of the first bacteria.
- **10**. The method of claim **7**, wherein the vector is a phage, phagemid, viriophage, virus, plasmid, or transposon.
- 11. The method of claim 7, wherein the vector is a conjugative plasmid.
- 12. The method of claim 11, wherein the conjugative 25 plasmid is delivered from carrier bacteria.
- 13. The method of claim 12, wherein the carrier bacteria are *Lactobacillus* bacteria.

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- 14. The method of claim 1, wherein the Cas nuclease is Cas9.
- 15. The method of claim 1, wherein the Cas nuclease is Cas3.
- **16**. The method of claim **8**, wherein the Cas nuclease is Cas9.
- 17. The method of claim 16, wherein the second vector further comprises a tracrRNA or a DNA encoding a tracrRNA.
- 18. The method of claim 1, wherein the treatment of the bacterial infection treats in the subject a disease or condition selected from the group consisting of vaginosis, meningitis, pneumonia, urinary tract infection, cystitis, nephritis, gastroenteritis, a skin infection, impetigo, erysipelas, cellulitis, a dental infection, septicaemia and sepsis, or reduces the risk of the disease or condition in the subject, wherein the disease or condition is mediated by the first bacteria.
- 19. The method of claim 1, wherein the guide RNA does not program a Cas nuclease to cut the genome of second bacteria in the subject, wherein the second bacteria are selected from the group consisting of Akkermansia, Alistipes, Bacteroides, Barnesiella, Bifidobacterium, Clostridium, Collinsella, Enterococcus, Fusobacterium, Lactobacillus, Propionibacterium, Ruminococcus, Segmented filamentous bacteria (SFB); Veillonella, Prevotella, Escherichia and Streptococcus bacteria.

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