Abstract:
The invention relates to methods for the screening, identification and/or application of microorganisms of use in imparting beneficial properties to plants. In one embodiment, the method involves identifying one or more plant markers associated with a beneficial property, subjecting one or more plants to a growth medium in the presence of a set of microorganisms, selecting one or more plants on the basis that the one or more plant markers is present, acquiring one or more microorganisms associated with the selected plant(s) and repeating the process one or more times.
Screening methods for the selection of microorganisms capable of imparting a beneficial property to a plant

FIELD
The present invention relates to methods for the screening, identification and/or application of microorganisms of use in imparting beneficial properties to plants.

BACKGROUND
Geography, environmental conditions, disease and attack by insects are major factors influencing the ability to grow and cultivate different species of plant. Such factors can have a significant downstream economic and social impact on communities around the world. There would be benefit in identifying products and methods which might impart beneficial properties to a plant species to allow it to grow in a variety of geographical locations, in different weather conditions, to survive disease and to be resistant to attack by insects, for example.

There is an ever increasing demand for plants having other desirable characteristics such as improved quality, increased or decreased levels of certain compounds, improved or different taste, smell, colour or other physical or chemical properties.

Selective breeding techniques have been used to this end. Selective breeding relies principally on genetic diversity in a starting population coupled with selection to achieve a plant cultivar with characteristics beneficial for human use. As the available, unused genetic diversity of cultivatable plant species has diminished, the potential for improvement has decreased. This situation has stimulated the growth of plant genetic modification in which genes from closely related species are introgressed into the new cultivar to provide a new genetic base for imparting desirable traits into new cultivars. However, this process is extremely costly, slow, limited in its scope and fraught with regulatory difficulties. Few commercial successes have eventuated from over two decades of large-scale investment into this technology.

Despite many decades of successful scientific research into the conventional breeding of highly-productive crops and into development of transgenic crops, relatively little research effort has been directed at development of plant traits via other means.
The importance of providing "good" soil with a rich microbial diversity via composts, complex biomaterial fertilisers e.g. blood and bone, to plants to ensure their healthy growth has been understood by home gardeners' and producers of organic foods. However, the inventors have recognised that the complexity of the plant-microorganism associations that underpin the observable benefits is poorly understood. Benefits to plant growth and health in such soils are often microbially-mediated through improved nutrient availability. This may be the result of solubilisation of minerals from the soil biomass itself, or from colonization of the plants with microorganisms in endophytic, epiphytic or rhizospheric associations leading to nitrogen fixation, resistance to pests and diseases through direct microbial competition within the plant, or the elicitation of plant defence reactions. The science community has produced literature on the diverse mechanisms of endophytic, epiphytic and rhizospheric plant microorganism associations, largely in relation to crop plants and their soils. The nature of some associations is known, encompassing the genetic basis of plant-induced metabolite production by specific organisms and the reverse influence of the microbe on gene expression in the plant (e.g. Neotyphodium spp), and increases in plant growth following microbial application to certain crop plants or seeds has been documented. However, despite the potential of microorganisms to improve plant growth, commercial success is limited to a relatively small range of specific microbial applications e.g. Rhizobium spp. to legume seeds, or the use of products resulting from "uncontrolled" microbial fermentations e.g. compost teas, seaweed fermentations, fish waste fermentations etc.

There are many specific strains of potentially beneficial microorganisms for association with specific plant cultivars, making the task of finding an appropriate strain(s) for any particular crop a very onerous procedure. Current means primarily focus on the application of microorganisms singly or in limited combinations. Such microorganisms are likely to have been selected for specific potential properties based on their identity. It would be useful if there was no requirement for knowledge of microbial identity for success.

Bibliographic details of the publications referred to herein are collected at the end of the description.
OBJECT
It is an object of the present invention to provide a method for the selection of one or more microorganism and/or composition which is of use in imparting one or more beneficial properties to a plant which overcomes or ameliorates at least one of the disadvantages of known methods.

It is an alternative object of the invention to provide a method for identifying new plant markers for breeding plants with beneficial traits.

It is an alternative object of the invention to provide a method for identifying new microorganism markers which may identify the microorganism as having a trait which is of use in imparting one or more beneficial properties to a plant.

Alternatively it is an object of the invention to provide a method and/or system for assisting in the improvement of one or more plants.

Alternatively, it is an object to at least provide the public with a useful choice.

STATEMENT OF INVENTION
In a first broad aspect of the present invention there is provided a method for the selection of one or more microorganisms capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:

a) identifying one or more plant marker associated with one or more beneficial property;

b) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to a growth medium in the presence of a first set of one or more microorganisms;

c) selecting one or more plant following step b) on the basis that the one or more plant marker is present;

d) acquiring a second set of one or more microorganisms associated with said one or more plant selected in step c);

e) repeating steps a) to d) and/or steps b) to d) one or more times, wherein the second set of one or more microorganisms acquired in step d) is used as the first set of microorganisms in step b) of any successive repeat.
In a second broad aspect of the present invention there is provided a method for the selection of one or more microorganisms capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:

a) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to a growth medium in the presence of a first set of one or more microorganisms;

b) selecting one or more plant following step a);

c) acquiring a second set of one or more microorganisms associated with said one or more plant selected in step b);

d) repeating steps a) to c) one or more times, wherein the second set of one or more microorganisms acquired in step c) is used as the first set of microorganisms in step a) of any successive repeat;

wherein the method also comprises the step of identifying one or more plant marker and/or one or more microorganism marker associated with the beneficial property after step b) and/or after step c).

In one embodiment of the first or second aspect, the second set of one or more microorganisms are isolated from said one or more plant in step d (first aspect) or step c (second aspect).

In one embodiment of the first or second aspect, where two or more microorganisms are acquired in step d) (first aspect), or step c) (second aspect), the method further comprises the steps of separating the two or more microorganisms into individual isolates, selecting two or more individual isolates, and then combining the selected two or more isolates. The combined isolates may then be used in any successive repeats of the method or any part thereof. In addition, two or more methods or parts thereof may be performed separately and the one or more microorganisms acquired in the first aspect and/or the second aspect of the method combined. In one embodiment, the combined microorganisms are used in any successive rounds of the method or part thereof.

In another embodiment of the first or second aspect, the method further comprises repeating steps a) to d) or steps b) to d) (first aspect) or steps a) to c) (second aspect) one or more times, wherein where two or more microorganisms are acquired in step d) (first
aspect) or step c) (second aspect), the two or more microorganisms are separated into individual isolates, two or more individual isolates are selected and then combined, and the combined isolates are used in step b) (first aspect) or step a) (second aspect) of any successive repeat. Accordingly, where reference is made to using the one or more microorganisms acquired in step d) (first aspect) or step c) (second aspect) of the method, it should be taken to include using the combined isolates of this embodiment of the invention.

In another embodiment, two or more methods of the invention may be performed separately and the one or more microorganisms acquired in step d) (first aspect) or step c) (second aspect) of each separate method combined. In one embodiment, the combined microorganisms are used in step b) (first aspect) or step a) (second aspect) when performing a further method of the invention.

In one embodiment of the first or second aspects, one or more selective pressure is applied in step b) (first aspect) or step a) (second aspect).

In one embodiment, the selective pressure is biotic and includes but is not limited to exposure to one or more organisms that are detrimental to the plant. In one embodiment, the organisms include fungi, bacteria, viruses, insects, mites and nematodes.

In another embodiment, the selective pressure is abiotic. Abiotic selective pressures include, but are not limited to, exposure to or changes in the level of salt concentration, temperature, pH, water, minerals, organic nutrients, inorganic nutrients, organic toxins, inorganic toxins, and metals.

In one embodiment, the selective pressure is applied during substantially the whole time during which the one or more plant is subjected to the growth medium and one or more microorganisms. In one embodiment, the selective pressure is applied during substantially the whole growth period of the one or more plant. Alternatively, the selective pressure is applied at a discrete time point.
In one embodiment, one or more selective pressure applied in successive repeats is different. In another embodiment, the selective pressure(s) applied in successive repeats is the same.

In one embodiment, the method further comprises the following step which is conducted prior to step a) (first or second aspect) or step b) (first aspect): subjecting the one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to a growth medium in the presence of one or more microorganisms, and after a desired period, isolating one or more microorganisms associated with said one or more plant. In a preferred embodiment, the one or more microorganisms acquired from the one or more plant, are used in the process. In one embodiment, this step may be conducted two or more times.

In one embodiment of the first aspect or second aspect, the methods are combined, in any order or combinations. Such a method may be repeated any number of times.

In one embodiment of the first or second aspect, the first set and/or second set of one or more microorganisms are selected from the microorganisms detailed herein after.

In one embodiment of the first or second aspect, the growth medium is selected from the growth media detailed herein after.

In one embodiment of the first or second aspect the growth medium is selective for a specific marker of interest.

In one embodiment of the first or second aspect the marker in each iterative repeat of the methods of the invention could be the same or different. Two or more markers may be assessed in each iteration of the method.

In another embodiment of the first or second aspect, one or more plant may be selected on the basis of the presence of one or more marker in combination with one or more other selection criterion.
In one embodiment of the first or second aspect, the one or more selection criterion includes one or more phenotypic trait. In one preferred embodiment, the one or more phenotypic trait is a desirable phenotypic trait. In one embodiment, the phenotypic trait is one of those detailed herein after. In one embodiment, the one or more selection criterion is one or more genotypic trait. In one preferred embodiment, the one or more genotypic trait is a desirable genotypic trait. In one embodiment, the one or more selection criterion is a combination of one or more genotypic and one or more phenotypic traits. In one embodiment, different selection criteria may be used in each iteration of a method of the invention.

In one embodiment of the first or second aspect, the step of subjecting one or more plant to a growth medium involves growing or multiplying the plant.

In one embodiment of the first or second aspect, two or more plants are subjected to a growth medium in the presence of one or more microorganisms. In other embodiments, to 20 plants are subjected to a growth medium in the presence of the first set of one or more microorganisms. In other embodiments, 20 or more, 100 or more, 300 or more, 500 or more, or 1000 or more plants are subjected to a growth medium in the presence of the first set of one or more microorganisms.

In one embodiment of the first or second aspect, the second set of one or more microorganisms are acquired from the root, stem and/or foliar (including reproductive) tissue of the one or more plants selected. Alternatively, the second set of one or more microorganisms are acquired from whole plant tissue of the one or more plants selected. In another embodiment, the plant tissues may be surface sterilised and then one or more microorganisms acquired from any tissue of the one or more plants. This embodiment allows for the targeted selection of endophytic microorganisms. In another embodiment, the one or more microorganisms may be acquired from the growth medium surrounding selected plants.

In another embodiment of the first or second aspect, the second set of one or more microorganisms are used in any successive repeat of the method in crude form.
In one embodiment of the first or second aspect, the one or more microorganisms are acquired any time after germination of the one or more plants.

In one embodiment, the methods of the first or second aspect of the invention may also be useful in identifying and/or selecting one or more endophytic microorganisms capable of imparting one or more beneficial property to a plant.

In one embodiment, plant material (including for example seeds, seedlings, cuttings, and/or propagules thereof) may be used as the source of microorganisms for step a). In a preferred embodiment, the plant material used as a source for microorganisms in step a) is seed material. Preferably, the plant material is surface sterilised.

In another embodiment, the methods of the first or second aspect of the invention may be useful in identifying and/or selecting one or more unculturable microorganisms capable of imparting one or more beneficial property to a plant. In this embodiment, plant material (including for example seeds, seedlings, cuttings, and/or propagules thereof) may be used as the source of microorganisms for step a). In a preferred embodiment, the plant material used as a source for microorganisms in step a) is explant material (for example, plant cuttings). Preferably, the plant material is surface sterilised.

In a third broad aspect, there is provided a method for assisting in the improvement of one or more plants, comprising arranging for the evaluation of said plant(s) in the presence of one or more microorganisms and/or compositions. The method preferably comprises at least the steps of a method of the first, second, eighth (and/or related) and/or ninth (and/or related) aspects of the invention.

According to one embodiment, the plant(s) are for growing in a first region. The microorganism(s) may or may not (or at least to a significant extent) be present in the first region.

"Region" and "first region" are to be interpreted broadly as meaning one or more areas of land. The land areas may be defined by geographical / political / private land boundaries or by land areas having similar properties such as climate, soil properties, presence of a particular pest etc.
Preferably, the evaluation is performed in a second region in which the microorganism(s) are present, but this is in no way essential. Microorganisms may be obtained from other sources including microorganism depositaries and artificially associated with plant material and/or soil. Furthermore, while plant(s) may be cultivated in essentially a conventional manner but, in one particular embodiment in a region having microorganisms not normally associated with the plant(s), at least in the first region, artificial growing environments may alternatively be used as would be appreciated by those skilled in the art. Thus, possible beneficial microorganism/plant relationships may be identified that would not necessarily normally be utilised.

Preferably, the step of arranging comprises arranging for one or more of:

- receipt or transmission of an identity of one or more plants or plant types to be evaluated;
- receipt or transmission of plant material from one or more plants or plant types to be evaluated;
- identification and/or selection of the microorganism(s) and/or composition(s);
- acquisition of the microorganism(s) and/or composition(s); and
- associating the microorganism(s) and/or composition(s) with the plant material.

Preferably, the method comprises evaluating (or arranging for said evaluation of) said plant(s) in the presence of said microorganism(s) and/or composition(s).

The step of evaluating preferably comprises performing one or more of the steps of a method described herein, in particular embodiments a method of the first aspect, second aspect, eighth (and/or related) aspect or ninth (and/or related) aspect of the invention.

The various steps identified above may be performed by a single entity although it is preferred that at least two parties are involved, a first which makes a request and a second which actions the request. Note that various agents may act for one or both parties and that varying levels of automation may be used. For example, in response to a particular request the microorganism(s) may be selected by a processor querying a database based on known
microorganism associations for that or similar plant(s) with little or no input required from an operator.

Furthermore, the evaluation may be performed by the requesting party and/or in the first region. Performing the evaluation in the first region better ensures that the evaluation is accurate and that no unforeseen environmental factors that may impact on the plant(s) or the microorganism(s) are not considered.

Following the evaluation or during the course thereof, the method preferably further comprises one or more of:

- receiving or sending one or more microorganisms (or at least the identity thereof) and/or composition(s) to the first region, including in combination with plant material; and

- growing said plant(s) or other plants (preferably having similar properties) in the first region in the presence of said microorganism(s) and/or composition(s).

The method of the third aspect may be embodied by a first party:

- identifying a need for an improvement in a plant(s);
- sending the identity thereof and/or relevant plant material to a second party together with any relevant information, and
- receiving plant material and/or one or more microorganisms and/or the identities thereof and/or composition(s).

The step of receiving is preferably performed following or as a result of an assessment of plant/microorganism and/or plant/composition associations. Preferably, the assessment is made using a method as described herein, in particular embodiments a method of the first aspect, second aspect, the eighth (and/or related) aspect or the ninth (and/or related) aspect.

The method of the third aspect may additionally or alternatively be embodied by a second party:
- receiving an identity of a plant(s) and/or relevant plant material from a first party together with any relevant information, and
- sending plant material and/or one or more microorganism(s) and/or the identities thereof and/or composition(s) to the first party.

The step of sending is preferably performed following or as a result of an assessment of plant/microorganism and/or plant/composition associations. Preferably, the assessment is made using a method described herein, in particular embodiments a method of the second aspect, the eighth (and/or related) aspect or the ninth (and/or related) aspect.

According to a fourth aspect, there is provided a system for implementing the method of the third aspect.

The system of the fourth aspect preferably includes one or more of:

- means for receiving or transmitting an identity of one or more plants or plant types to be evaluated;
- means for receiving or transmitting plant material from one or more plants or plant types to be evaluated;
- means for identifying and/or selecting microorganism(s) and/or composition(s);
- means for acquiring the microorganism(s) and/or composition(s);
- means for associating the microorganism(s) and/or composition(s) with the plant material;
- means for evaluating said plant(s) in the presence of said microorganism(s) and/or composition(s);
- means for receiving or sending one or more microorganisms (or at least the identity thereof) and/or composition(s) to the first region, including in combination with plant material; and
- means for growing said plant(s) or other plants (preferably having similar properties) in the first region in the presence of said microorganism(s) and/or composition(s).

Means known to those skilled in the art may be used to provide the functionality required in the system of the third aspect. For example, conventional communication means,
including the internet, may be used to convey the identities of plants / microorganisms; conventional carrier means may be used to convey the plant material / microorganisms/ composition(s); conventional means and processes may be used to associate a microorganism and/or composition with plant material and conventional means for evaluating said plant(s) and / or the plant / microorganism and/or plant/composition associations may be used.

According to a preferred embodiment, the system of the invention is embodied by a facility configured to transmit request(s) for an improvement in a plant(s) and subsequently to receive plant material and / or one or more microorganisms and / or the identities thereof, preferably following or as a result of an assessment of plant / microorganism associations. Preferably, the assessment is made using a method described herein, in particular embodiments a method of the first aspect, the second aspect, the eighth (and/or related) aspect, or the ninth (and/or related) aspect.

The system of the fourth aspect may additionally or alternatively be embodied by a facility configured to receive an identity of a plant(s) and / or relevant plant material from together with any relevant information; and send plant material and / or one or more microorganisms and / or the identities thereof and/or composition(s), preferably following or as a result of an assessment of plant / microorganism or plant/composition associations. Preferably, the assessment is made using a method described herein, in particular embodiments a method of the first aspect, the second aspect, the eighth (and/or related) aspect or the ninth (and/or related) aspect.

Accordingly to a fifth broad aspect of the invention, there is provided a microorganism acquired, selected or isolated by a method as hereinbefore described. In one embodiment, the microorganism is an endophyte. In one embodiment, the microorganism is unculturable.

In a sixth broad aspect of the invention, there is provided a method for the production of a composition to support plant growth, quality and/or health, or a composition to suppress or inhibit the growth, quality and/or health of a plant the method comprising the steps of a method herein before described and the additional step of combining the one or more microorganisms selected by the method with one or more additional ingredients.
In a seventh broad aspect of the invention, there is provided a composition comprising one or more microorganism of the fifth broad aspect or as prepared by a method of the sixth broad aspect.

In an eighth broad aspect of the invention there is provided a method for the selection of a composition which is capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:

a) culturing one or more microorganism selected by a method of the first aspect and/or the second aspect in one or more media to provide one or more culture;

b) separating the one or more microorganism from the one or more media in the one or more culture after a period of time to provide one or more composition substantially free of microorganisms;

c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition of step b);

d) selecting one or more composition from step c) if it is observed to impart one or more beneficial property to the one or more plants.

In an aspect of the invention related to (but distinct from) the eighth broad aspect of the invention there is provided a method for the selection of a composition which is capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:

a) culturing one or more microorganisms selected by a method of the first aspect and/or the second aspect of the invention in one or more media to provide one or more culture;

b) inactivating the one or more culture of step a) to provide one or more composition containing one or more inactivated microorganisms;

c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition of step b);

d) selecting one or more composition from step c) if it is observed to impart one or more beneficial property to the one or more plants.

In a ninth broad aspect of the invention there is provided a method for the selection of one or more microorganisms which are capable of producing a composition which is capable of
imparting one or more beneficial property to a plant, the method comprising at least the
steps of:

a) culturing one or more microorganism selected by a method of the first aspect
and/or the second aspect in one or more media to provide one or more culture;
b) separating the one or more microorganism from the one or more media in the
one or more culture from step a) after a period of time to provide one or more
composition substantially free of microorganisms;
c) subjecting one or more plant (including for example seeds, seedlings, cuttings,
and/or propagules thereof) to the one or more composition from step b);
d) selecting the one or more microorganisms associated with one or more
composition observed to impart one or more beneficial property to the one or more
plants.

In an aspect related to (but distinct from) the ninth broad aspect of the invention there is
provided a method for the selection of one or more microorganisms which are capable of
producing a composition which is capable of imparting one or more beneficial property to
a plant, the method comprising at least the steps of:

a) culturing one or more microorganism in one or more media to provide one or
more culture;
b) separating the one or more microorganism from the one or more media in one or
more culture after a period of time to provide one or more composition
substantially free of microorganisms;
c) subjecting one or more plant (including for example seeds, seedlings, cuttings,
and/or propagules thereof) to the one or more composition of step b);
d) selecting the one or more microorganisms associated with (or in other words
used to produce the) one or more composition observed to impart one or more
beneficial property to the one or more plants; and,
e) using the one or more microorganisms selected in step d) of a method of the first
aspect of the invention or step c) of the second aspect of the invention.

In a related aspect, step b) of the method of the ninth (and/or related) aspect could be
substituted with the step of b) inactivating the one or more culture of step a) to provide one
or more composition containing one or more inactivated microorganisms, and then using
this composition in step c) of the process.
It should be appreciated that the methods of the first, second, eighth (and/or related) and ninth (and/or related) aspects may be combined in any combination, including the methods being run concurrently or sequentially in any number of iterations, with compositions and/or microorganisms selected, acquired or isolated from the methods being used individually or combined and used in iterative rounds of any one of the methods. By way of example, a method of the eighth (and/or related) aspect may be performed and a composition selected. The selection of a composition indicates that the one or more microorganism separated from the media in step b) is desirable for imparting beneficial properties to the one or more plant (as the one or more microorganism is capable of producing a selected composition). The one or more microorganism may then be used in another round of a method of the first aspect, second aspect, eighth (and/or related) aspect or ninth (and/or related) aspect. Alternatively, the combination of methods could be run in reverse. This could be repeated any number of times in any order and combination.

Accordingly, the invention provides for the use of one or more microorganism, composition or plant acquired, selected or isolated by a method of the invention in any other method of the invention.

In a tenth broad aspect of the invention there is provided a composition obtained as a result of the methods of the eighth (and/or related) or ninth (and/or related) broad aspects of the invention.

In an eleventh broad aspect of the invention there is provided a combination of two or more microorganisms acquired, selected or isolated by a method as herein before described.

In another aspect, the invention provides the use of one or more composition and/or microorganism acquired, selected or isolated by a method of the invention for imparting one or more beneficial property to one or more plant.

It should be appreciated that methods of the invention may also involve applying steps a) to e) (first aspect) or steps a) to d) (second aspect) of a method of the invention to two or more different species of plant so as to identify combinations of microorganisms that may impart a positive benefit to one species and a negative benefit to another species.
simultaneously. For example, one may wish to identify a group of microorganisms that may simultaneously improve the growth and survival of a food crop and suppress or inhibit the growth of a competing crop or weed. This may be achieved by using two or more different plant species or running separate methods on different species and at appropriate points combining the microorganisms acquired in those methods and conducting further iterations.

The invention also provides plants selected in a method of the invention.

The invention also provides the use of a method of the invention in a plant breeding programme, and a plant breeding programme comprising conducting a method of the invention.

In another broad aspect, the invention provides a method for identifying one or more plant marker that is associated with one or more beneficial trait the method comprising at least the step of conducting a method of the first or second aspect of the invention and identifying the one or more plant marker.

In another broad aspect, the invention provides a method for identifying one or more microorganism marker that is associated with one or more beneficial plant trait the method comprising at least the step of conducting a method of the first or second aspect of the invention and identifying the one or more microorganism marker.

In one embodiment, there is provided a method for identifying a combination of one or more plant marker and one or more microorganism marker comprising at least the step of conducting a method of the first or second aspect of the invention and identifying the combination of one or more plant marker and one or more microorganism marker.

The invention may also be said broadly to consist in the parts, elements and features referred to or indicated in the specification of the application, individually or collectively, in any or all combinations of two or more of said parts, elements or features, and where specific integers are mentioned herein which have known equivalents in the art to which the invention relates, such blown equivalents are deemed to be incorporated herein as if individually set forth.
FIGURES
These and other aspects of the present invention, which should be considered in all its novel aspects, will become apparent from the following description, which is given by way of example only, with reference to the accompanying figures, in which:

Figure 1: shows a system according to an embodiment of the invention;
Figure 2: shows the process flow of a method of an embodiment of the invention.

PREFERRED EMBODIMENT(S)
The following is a description of the preferred forms of the present invention given in general terms. The invention will be further elucidated from the Examples provided hereafter.

The inventor(s) have found that one can readily identify microorganisms capable of imparting one or more beneficial property to one or more plants through use of a method of the invention. The method is broadly based on the presence of variability (such as genetic variability, or variability in the phenotype, for example) in the plants and microbial populations used. The inventors have identified that this variability can be used to support a directed process of selection of one or more microorganisms of use to a plant and for identifying particular plant/microbe combinations which are of benefit for a particular purpose, and which may never have been recognised using conventional techniques.

The methods of the invention may be used as a part of a plant breeding programme. The methods may allow for, or at least assist with, the selection of plants which have a particular genotype/phenotype which is influenced by the microbial flora, in addition to identifying microorganisms and/or compositions that are capable of imparting one or more property to one or more plants.

The methods of the invention may be useful for improving the efficiency of crop breeding programs through the use of directed selection of crop-associated microbes that influence phenotypic traits under the control of quantitative trait loci (QTLs). The methods may indirectly manipulate the expression of crop QTLs that control the heritable variability of the traits and physiological mechanisms underlying desirable traits such as biomass
compartmentalization, abiotic stress tolerance, resistance to pest and diseases and nutrient assimilation.

Methods of the invention may be used to assist in improving plants by identifying microorganisms that optimise the expression of plant markers.

As used herein a plant "marker" is intended to mean any phenotypic or genotypic trait. In certain embodiments it includes genetic elements, including quantitative trait loci (QTL), any genetic variation including but not limited to polymorphisms (SNPs, RFLPs), mutations and allelic variation, the presence or absence or level of expression of one or more gene, the presence or absence or level of production of one or more compounds by a plant, including one or more metabolites.

As used herein a microorganism "marker" is intended to mean any phenotypic or genotypic trait. In certain embodiments it includes genetic elements, any genetic variation including but not limited to polymorphisms (SNPs, RFLPs), mutations and allelic variation, the presence or absence or level of expression of one or more gene, the presence or absence or level of production of one or more compounds by a microorganism, including one or more metabolites.

In one aspect the invention relates to a method for identifying one or more microorganisms capable of imparting one or more beneficial property to a plant. It should be appreciated that as referred to herein a "beneficial property to a plant" should be interpreted broadly to mean any property which is beneficial for any particular purpose including properties which may be beneficial to human beings, other animals, the environment, a habitat, an ecosystem, the economy, of commercial benefit, or of any other benefit to any entity or system. Accordingly, the term should be taken to include properties which may suppress, decrease or block one or more characteristic of a plant, including suppressing, decreasing or inhibiting the growth or growth rate of a plant. The invention may be described herein, by way of example only, in terms of identifying positive benefits to one or more plants or improving plants. However, it should be appreciated that the invention is equally applicable to identifying negative benefits that can be conferred to plants.
Such beneficial properties include, but are not limited to, for example: improved growth, health and/or survival characteristics, resistance to pests and/or diseases, tolerance to growth in different geographical locations and/or different environmental biological and/or physical conditions, suitability or quality of a plant for a particular purpose, structure, colour, chemical composition or profile, taste, smell, improved quality. By way of example, the invention may allow for the identification of microorganisms which allow a plant to grow in a variety of different temperatures (including extreme temperatures), pH, salt concentrations, mineral concentrations, in the presence of toxins, and/or to respond to a greater extent to the presence of organic and/or inorganic fertilisers. In other embodiments, beneficial properties include, but are not limited to, for example; decreasing, suppressing or inhibiting the growth of a plant identified to be a weed; constraining the height and width of a plant to a desirable ornamental size; limiting the height of plants used in ground cover applications such as motorway and roadside banks and erosion control projects; slowing the growth of plants used in turf applications such as lawns, bowling greens and golf courses to reduce the necessity of mowing; reducing ratio of foliage/flowers in ornamental flowering shrubs; regulate production of and/or response to plant pheromones (resulting in increased tannin production in surrounding plant community and decreased appeal to foraging species).

As used herein, "improved" should be taken broadly to encompass improvement of a characteristic of a plant which may already exist in a plant or plants prior to application of the invention, or the presence of a characteristic which did not exist in a plant or plants prior to application of the invention. By way of example, "improved" growth should be taken to include growth of a plant where the plant was not previously known to grow under the relevant conditions.

As used herein, "inhibiting and suppressing" and like terms should be taken broadly and should not be construed to require complete inhibition or suppression, although this may be desired in some embodiments.

To assist in describing the invention, the terms a "first set of one or more microorganisms" and a "second set of one or more microorganisms" may be used herein to distinguish the set or group of microorganism(s) applied, for example, in step b) of the first aspect and step a) of the second aspect and, the set or group of microorganism(s) acquired in a method
of the invention, for example in step d) of the first aspect and step c) of the second aspect.
In certain embodiments, the sets of microorganisms will be distinct; for example, the
second set may be a subset of the first set, as a result of combining the first set with the
plant and then selecting one or more plant based on one or more selection criterion.

However, it should be appreciated that this may not always be the case and accordingly,
the use of this terminology should not be construed in such a limited manner.

In certain embodiments, methods of the invention relate to selecting one or more
microorganisms which are capable of imparting one or more beneficial property to a plant.
As is further described herein after, such microorganism(s) may be contained within a
plant, on a plant, and/or within the plant rhizosphere. Accordingly, where reference is
made herein to acquiring a second set of one or more microorganisms "from" a plant,
unless the context requires otherwise, it should be taken to include reference to acquiring a
second set of microorganisms contained within a plant, on a plant and/or within the plant
rhizosphere. For ease of reference, the wording "associated with" may be used
synonymously to refer to microorganism(s) contained within a plant, on a plant and/or
within the plant rhizosphere.

A method of the invention broadly comprises at least the steps of growing one or more
plant in a growth medium in the presence of a first set of one or more microorganisms,
selecting one or more plant following the growth step and, acquiring a second set of one or
more microorganisms associated with said one or more of the selected plants. The one or
more plants, growth medium and one or more microorganisms may be provided separately
and combined in any appropriate order. In one embodiment the methods of the invention
also include the step of identification of one or more plant marker associated with one or
more beneficial property (as described in the first aspect of the invention hereinbefore). In
another embodiment, the methods of the invention require that one or more plant marker
and/or one or more microorganism marker associated with the beneficial property are
identified (as described in the second aspect of the invention hereinbefore). In particular,
the invention provides an iterative method in which the relevant method steps are repeated
one or more times, wherein the one or more microorganisms acquired in the method are
used in any successive repeat or cycle of the method. In one embodiment, the relevant
steps of the method are repeated once. In another embodiment, the relevant steps of the
method are repeated twice. In another embodiment, the relevant steps of the method are
repeated three times. In another embodiment, the relevant steps of the method are repeated at least until a desired beneficial property is observed.

In another aspect, the invention comprises a step of applying a selective pressure. The selective pressure is applied at the step in the method comprising subjecting one or more plant to a growth medium in the presence of a first set of one or more microorganisms. In particular, the invention of this aspect provides an iterative method in which the relevant method steps are repeated one or more times, wherein the one or more microorganisms acquired in the method are used in any successive repeat or cycle of the method. In one embodiment, the relevant steps of the method are repeated once. In another embodiment, the relevant steps of the method are repeated twice. In another embodiment, the relevant steps of the method are repeated three times. In another embodiment, the relevant steps of the method are repeated at least until a desired beneficial property is observed. Whilst the same selective pressure(s) may be applied in each iteration of the method, different selective pressures may be applied in each iteration. In addition, the method may comprise a further step (or steps) which are conducted prior to the first step of a method of the first or second aspect and include subjecting the one or more plant to a growth medium in the presence of one or more microorganisms but without a selective pressure. After a desired period, one or more microorganisms may be acquired from said one or more plant and can be used in any successive repeat of the process. It should also be appreciated, that successive rounds of this step may be conducted, with the microorganisms acquired being used in any subsequent round of the process, before embarking on the steps of the processes of the first aspect or second aspect as described hereinbefore. Further, the inventors envisage an iterative method in which the relevant steps (for example as described in the first and second aspects herein before) are repeated one or more times, but interspersed with the step of subjecting the plants to a growth medium and one or more microorganisms and isolating the microorganisms, without applying a selective pressure.

It will be appreciated that after a desired number of repeats of steps a) to d) of a method of the first aspect of the invention or steps a) to c) of a method of the second aspect of the invention, or any combination thereof, the method may conclude with the acquisition of a set of one or more microorganisms from step d) of the first aspect or step c) of the second aspect (as the case may be) of any such method.
It should be appreciated that the methods do not require the identification of the microorganisms in the population acquired in the method (for example in step d) of the first aspect or step c) of the second aspect) nor do they require a determination of the beneficial properties of individual microorganisms or combinations of microorganisms acquired. However, evaluation, identification and/or a determination of the beneficial properties could be conducted if desired. For example, it may be preferred in some cases to isolate and identify the microbes in the final step of a method of the invention to determine their safety for commercial use and to satisfy regulatory requirements. In such cases, genetic and/or phenotypic analyses may be conducted.

In one embodiment of methods of the invention, step a) is conducted using at least two plants. In other embodiments 10 to 20 plants are used. In yet other embodiments, 20 or more, 50 or more, 100 or more, 300 or more, 500 or more or 1000 or more plants are used. As noted hereinbefore, where two or more plants are used in a particular method of the invention, they need not be the same variety or species. For example, in one embodiment, it may be desirable to select microorganisms that can impart a positive benefit to one plant variety or species and a negative benefit to another plant variety or species.

In one embodiment, where two or more microorganisms are acquired (for example in step d) of the first aspect or in step c) of the second aspect), the method may further comprise the steps of separating the two or more microorganisms into individual isolates, selecting two or more individual isolates, and then combining the selected two or more isolates. This may result in the set of microorganisms acquired at the conclusion of a method of the invention. However, in one embodiment, the combined isolates may then be used in step b) of the first aspect or step a) of the second aspect of successive rounds of the method. By way of example, from two, three, four, five, six, seven, eight, nine or ten individual isolates may be combined. The inventors envisage an iterative method in which steps a) to d) and/or steps b) to d) (first aspect) or steps a) to c) (second aspect) are repeated one or more times, utilising these additional steps of separating, selecting and combining with each repeat of the method, or interspersed or otherwise combined with a method in which individual isolates are not selected and combined.

It is expected that these combinations will detect previously unknown, desirable property promoting (such as plant growth), synergistic interactions between previously acquired
microbes. Using the iterative steps of the methods will drive the starting population of two or more microorganisms toward microbes that interact with the plant to impart a desired property or characteristic. In other words, the process will allow for enrichment of suitable microorganisms within the plant microbiome.

Selection of individual isolates may occur on the basis of any appropriate selection criteria. For example, it may be random, it may be based on the beneficial property or properties observed by performing a method of the invention or, where information about the identity of the microorganism is known, it may be on the basis that the microorganism has previously been recognised to have a particular beneficial property.

In addition, two or more methods of the invention may be performed separately or in parallel and the microorganisms that result from each method combined into a single composition. For example, two separate methods may be performed, one to identify microorganisms capable of imparting one or more first beneficial property, and a second to identify microorganisms capable of imparting one or more second beneficial property. The separate methods may be directed to identifying microorganisms having the same beneficial property or having distinct beneficial properties. Where a method employing a selective pressure is used, the selective pressure in the separate methods may be the same or different. Similarly, the microorganisms and plants used in the separate methods may be the same or different. If further optimisation of the microorganisms is desired, the single composition of microorganisms may be applied to one or more further rounds of a method of the invention. Alternatively, the single composition of microorganisms may be used, as desired, to confer the relevant properties to plant crops, without further optimisation. Combining two or more methods of the invention in this way allows for the selection and combination of microorganisms which may ordinarily be separated by time and/or space in a particular environment.

In certain embodiments of the invention, the methods may comprise growing or propagating one or more plants selected in step c) (first aspect) or step b) (second aspect), to grow the population of the second set of one or more microorganisms associated with the selected one or more plants, either at the conclusion of a method of the invention, or prior to using the second set of one or more microorganisms in step b) or step a) of any successive repeat of the method. If the one or more plants (with associated
microorganisms) are grown or propagated at the conclusion of a method of the invention they may then be used or sold in that form. Alternatively, one or more microorganisms may be isolated from the one or more plants, or one or more plant tissue and/or one or more plant part with associated microorganisms may be used as a crude source of the one or more microorganisms in any successive repeat of the invention, or for any other purpose at the conclusion of the method. In one embodiment, the seeds (with associated microorganisms) of one or more plant that has been grown or propagated may be obtained and used as a source of the one or more microorganisms in any successive repeat of the method. Alternatively, if obtained at the conclusion of a method of the invention, the seeds and associated microbes may be sold or used for any other purpose.

In particular embodiments, the one or more microorganisms selected or acquired by the present methods can be formulated as a composition, as described herein after. Such compositions may, for example, be utilised to support plant growth, quality and/or health, or to suppress or inhibit plant growth, quality or health. In one embodiment, a composition is utilized as a seed coating for commercially important agricultural crops, for example.

In certain embodiments, the compositions comprise one or more microorganisms combined with one or more additional ingredients. In certain embodiments, the one or more microorganisms can be combined with known actives available in the agricultural space, such as: pesticide, herbicide, bactericide, fungicide, insecticide, virucide, miticide, nemataicide, acaricide, plant growth regulator, rodenticide, or anti-algae agent. Further, the one or more microorganisms can be combined with known fertilizers.

In some embodiments, when the one or more microorganisms is combined with an active chemical agent one witnesses an additive effect on a plant phenotypic trait of interest.

In some embodiments, when the one or more microorganisms is combined with a fertilizer one witnesses an additive effect on a plant phenotypic trait of interest.

In other embodiments, when the one or more microorganisms is combined with an active chemical agent one witnesses a synergistic effect. The synergistic effect obtained by the taught methods can be quantified according to Colby's formula \( (E) = X + Y - (X \times Y / 100) \).

*See* Colby, R. S., "Calculating Synergistic and Antagonistic Responses of Herbicide
Combinations", 1967 Weeds, vol. 15, pp. 20-22, incorporated herein by reference in its entirety. Thus, by "synergistic" is intended a component which, by virtue of its presence, increases the desired effect by more than an additive amount. The one or more microorganisms selected or acquired by the methods disclosed herein can synergistically increase the effectiveness of agricultural active compounds and also agricultural auxiliary compounds.

In other embodiments, when the one or more microorganisms is combined with a fertilizer one witnesses a synergistic effect.

The composition comprising the one or more microorganisms can be formulated with certain auxiliaries in order to improve the activity of a known active agricultural compound. This has the advantage that the amounts of active ingredient in the formulation may be reduced while maintaining the efficacy of the active compound, thus allowing costs to be kept as low as possible and any official regulations to be followed. In individual cases, it may also be possible to widen the spectum of action of the active compound since plants, where the treatment with a particular active ingredient without addition was insufficiently successful, can indeed be treated successfully by the addition of certain auxiliaries along with the one or more microorganisms selected or acquired by the methods disclosed herein. Moreover, the performance of the active may be increased in individual cases by a suitable formulation when the environmental conditions are not favorable.

Such auxiliaries that can be used in a composition comprising an active agricultural compound and one or more microorganism selected or acquired according to the disclosed methods can be an adjuvant. Frequently, adjuvants take the form of surface-active or salt-like compounds. Depending on their mode of action, they can roughly be classified as modifiers, activators, fertilizers, pH buffers, and the like. Modifiers affect the wetting, sticking, and spreading properties of a formulation. Activators break up the waxy cuticle of the plant and improve the penetration of the active ingredient into the cuticle, both short-term (over minutes) and long-term (over hours). Fertilizers such as ammonium sulfate, ammonium nitrate or urea improve the absorption and solubility of the active ingredient and may reduce the antagonistic behavior of active ingredients. pH buffers are conventionally used for bringing the formulation to an optimal pH.
Further methods and aspects of the invention are described herein after.

Microorganisms
As used herein the term "microorganism" should be taken broadly. It includes but is not limited to the two prokaryotic domains, Bacteria and Archaea, as well as eukaryotic fungi and protists. By way of example, the microorganisms may include Proteobacteria (such as Pseudomonas, Enterobacter, Stenotrophomonas, Burkholderia, Rhizobium, Herbaspirillum, Pantoea, Serratia, Rahnella, Azospirilium, Azorhizobium, Azotobacter; Duganella, Delftia, Bradyrhizobium, Sinorhizobium and Halomonas), Firmicutes (such as Bacillus, Paenibacillus, Lactobacillus, Mycoplasma, and Acetobacterium), Actinobacteria (such as Streptomyces, Rhodococcus, Microbacterium, and Curtobacterium), and the fungi Ascomycota (such as Trichoderma, Ampelomyces, Coniothyrium, Paecolomyces, Penicillium, Cladosporium, Hypocrea, Beauveria, Metarhizium, Verticullium, Cordyceps, Pichea, and Candida, Basidiomycota (such as Coprinus, Corticium, and Agaricus) and Oomycota (such as Pythium, Mucor, and Mortierella).

In a particularly preferred embodiment, the microorganism is an endophyte or an epiphyte or a microorganism inhabiting the plant rhizosphere. In one embodiment, the microorganism is a seed-borne endophyte.

In certain embodiments, the microorganism is unculturable. This should be taken to mean that the microorganism is not known to be culturable or is difficult to culture using methods known to one skilled in the art.

Microorganisms of use in the methods of the present invention (for example, as the first set of one or more microorganisms) may be collected or obtained from any source or contained within and/or associated with material collected from any source.

In one embodiment, the first set of one or more microorganisms are obtained from any general terrestrial environment, including its soils, plants, fungi, animals (including invertebrates) and other biota, including the sediments, water and biota of lakes and rivers; from the marine environment, its biota and sediments (for example sea water, marine muds, marine plants, marine invertebrates (for example sponges), marine vertebrates (for example, fish)); the terrestrial and marine geosphere (regolith and rock, for example
crushed subterranean rocks, sand and clays); the cryosphere and its meltwater; the atmosphere (for example, filtered aerial dusts, cloud and rain droplets); urban, industrial and other man-made environments (for example, accumulated organic and mineral matter on concrete, roadside gutters, roof surfaces, road surfaces).

In another embodiment the first set of one or more microorganisms are collected from a source likely to favour the selection of appropriate microorganisms. By way of example, the source may be a particular environment in which it is desirable for other plants to grow, or which is thought to be associated with terroir. In another example, the source may be a plant having one or more desirable traits, for example a plant which naturally grows in a particular environment or under certain conditions of interest. By way of example, a certain plant may naturally grow in sandy soil or sand of high salinity, or under extreme temperatures, or with little water, or it may be resistant to certain pests or disease present in the environment, and it may be desirable for a commercial crop to be grown in such conditions, particularly if they are, for example, the only conditions available in a particular geographic location. By way of further example, the microorganisms may be collected from commercial crops grown in such environments, or more specifically from individual crop plants best displaying a trait of interest amongst a crop grown in any specific environment, for example the fastest-growing plants amongst a crop grown in saline-limiting soils, or the least damaged plants in crops exposed to severe insect damage or disease epidemic, or plants having desired quantities of certain metabolites and other compounds, including fibre content, oil content, and the like, or plants displaying desirable colours, taste or smell. The microorganisms may be collected from a plant of interest or any material occurring in the environment of interest, including fungi and other animal and plant biota, soil, water, sediments, and other elements of the environment as referred to previously.

In certain embodiments, the microorganisms are sourced from previously performed methods of the invention (for example, the microorganisms acquired in step d) (first aspect) or step c) (second aspect) of the method), including combinations of individual isolates separated from the second set of microorganisms acquired in step d) (first aspect) or step c) (second aspect) or combinations of microorganisms resulting from two or more separately performed methods of the invention.
While the invention obviates the need for pre-existing knowledge about a microorganism's desirable properties with respect to a particular plant species, in one embodiment a microorganism or a combination of microorganisms of use in the methods of the invention may be selected from a pre-existing collection of individual microbial species or strains based on some knowledge of their likely or predicted benefit to a plant. For example, the microorganism may be predicted to: improve nitrogen fixation; release phosphate from the soil organic matter; release phosphate from the inorganic forms of phosphate (e.g. rock phosphate); "fix carbon" in the root microsphere; live in the rhizosphere of the plant thereby assisting the plant in absorbing nutrients from the surrounding soil and then providing these more readily to the plant; increase the number of nodules on the plant roots and thereby increase the number of symbiotic nitrogen fixing bacteria (e.g. *Rhizobium* species) per plant and the amount of nitrogen fixed by the plant; elicit plant defensive responses such as ISR (induced systemic resistance) or SAR (systemic acquired resistance) which help the plant resist the invasion and spread of pathogenic microorganisms; compete with microorganisms deleterious to plant growth or health by antagonism, or competitive utilisation of resources such as nutrients or space; change the colour of one or more part of the plant, or change the chemical profile of the plant, its smell, taste or one or more other quality.

In one embodiment a microorganism or combination of microorganisms (the first set of one or more microorganisms) is selected from a pre-existing collection of individual microbial species or strains that provides no knowledge of their likely or predicted benefit to a plant. For example, a collection of unidentified microorganisms isolated from plant tissues without any knowledge of their ability to improve plant growth or health, or a collection of microorganisms collected to explore their potential for producing compounds that could lead to the development of pharmaceutical drugs.

In one embodiment, the microorganisms are acquired from the source material (for example, soil, rock, water, air, dust, plant or other organism) in which they naturally reside. The microorganisms may be provided in any appropriate form, having regard to its intended use in the methods of the invention. However, by way of example only, the microorganisms may be provided as an aqueous suspension, gel, homogenate, granule, powder, slurry, live organism or dried material. The microorganisms may be isolated in substantially pure or mixed cultures. They may be concentrated, diluted or provided in the
natural concentrations in which they are found in the source material. For example, microorganisms from saline sediments may be isolated for use in this invention by suspending the sediment in fresh water and allowing the sediment to fall to the bottom. The water containing the bulk of the microorganisms may be removed by decantation after a suitable period of settling and either applied directly to the plant growth medium, or concentrated by filtering or centrifugation, diluted to an appropriate concentration and applied to the plant growth medium with the bulk of the salt removed. By way of further example, microorganisms from mineralized or toxic sources may be similarly treated to recover the microbes for application to the plant growth material to minimise the potential for damage to the plant.

In another embodiment, the microorganisms (including the first set of one or more microorganisms and/or the second set of one or more microorganisms) are used in a crude form, in which they are not isolated from the source material in which they naturally reside. For example, the microorganisms are provided in combination with the source material in which they reside; for example, as soil, or the roots, seed or foliage of a plant. In this embodiment, the source material may include one or more species of microorganisms.

It is preferred that a mixed population of microorganisms is used in the methods of the invention.

In embodiments of the invention where the microorganisms are isolated from a source material (for example, the material in which they naturally reside), any one or a combination of a number of standard techniques which will be readily known to skilled persons may be used. However, by way of example, these in general employ processes by which a solid or liquid culture of a single microorganism can be obtained in a substantially pure form, usually by physical separation on the surface of a solid microbial growth medium or by volumetric dilutive isolation into a liquid microbial growth medium. These processes may include isolation from dry material, liquid suspension, slurries or homogenates in which the material is spread in a thin layer over an appropriate solid gel growth medium, or serial dilutions of the material made into a sterile medium and inoculated into liquid or solid culture media.
Whilst not essential, in one embodiment, the material containing the microorganisms may be pre-treated prior to the isolation process in order to either multiply all microorganisms in the material, or select portions of the microbial population, either by enriching the material with microbial nutrients (for example, nitrates, sugars, or vegetable, microbial or animal extracts), or by applying a means of ensuring the selective survival of only a portion of the microbial diversity within the material (for example, by pasteurising the sample at 60°C-80°C for 10-20 minutes to select for microorganisms resistant to heat exposure (for example, bacilli), or by exposing the sample to low concentrations of an organic solvent or sterilant (for example, 25% ethanol for 10 minutes) to enhance the survival of actinomycetes and spore-forming or solvent-resistant microorganisms). Microorganisms can then be isolated from the enriched materials or materials treated for selective survival, as above.

In a preferred embodiment of the invention endophytic or epiphytic microorganisms are isolated from plant material. Any number of standard techniques known in the art may be used and the microorganisms may be isolated from any appropriate tissue in the plant, including for example root, stem and leaves, and plant reproductive tissues. By way of example, conventional methods for isolation from plants typically include the sterile excision of the plant material of interest (e.g. root or stem lengths, leaves), surface sterilisation with an appropriate solution (e.g. 2% sodium hypochlorite), after which the plant material is placed on nutrient medium for microbial growth (see, for example, Strobel G and Daisy B (2003) Bioprospecting for microbial endophytes and their natural products. Microbiology and Molecular Biology Reviews 67 (4): 491-502; Zinniel DK et al. (2002) Isolation and characterisation of endophytic colonising bacteria from agronomic crops and prairie plants. Applied and Environmental Microbiology 68 (5): 2198-2208. In one preferred embodiment of the invention, the microorganisms are isolated from root tissue. Further methodology for isolating microorganisms from plant material are detailed herein after.

As used herein, "isolate", "isolated" and like terms should be taken broadly. These terms are intended to mean that the one or more microorganism(s) has been separated at least partially from at least one of the materials with which it is associated in a particular environment (for example soil, water, plant tissue). "Isolate", "isolated" and like terms should not be taken to indicate the extent to which the microorganism(s) has been purified.
As used herein, "individual isolates" should be taken to mean a composition or culture comprising a predominance of a single genera, species or strain of microorganism, following separation from one or more other microorganisms. The phrase should not be taken to indicate the extent to which the microorganism has been isolated or purified. However, "individual isolates" preferably comprise substantially only one genus, species or strain of microorganism.

In one embodiment, the microbial population is exposed (prior to the method or at any stage of the method) to a selective pressure to enhance the probability that the eventually-selected plants will have microbial assemblages likely to have desired properties. For example, exposure of the microorganisms to pasteurisation before their addition to a plant growth medium (preferably sterile) is likely to enhance the probability that the plants selected for a desired trait will be associated with spore-forming microbes that can more easily survive in adverse conditions, in commercial storage, or if applied to seed as a coating, in an adverse environment.

It should be appreciated that the second set of microorganisms acquired in a method of the invention may be isolated from a plant or plant material, surface or growth media associated with a selected plant using any appropriate techniques known in the art, including but not limited to those techniques described herein before. However, in certain embodiments, as mentioned herein before, the microorganism(s) may be used in crude form and need not be isolated from a plant or a media. For example, plant material or growth media which includes the microorganisms identified to be of benefit to a selected plant may be obtained and used as a crude source of microorganisms for the next round of the method or as a crude source of microorganisms at the conclusion of the method. For example, whole plant material could be obtained and optionally processed, such as mulched or crushed. Alternatively, individual tissues or parts of selected plants (such as leaves, stems, roots, and seeds) may be separated from the plant and optionally processed, such as mulched or crushed. In certain embodiments, one or more part of a plant which is associated with the second set of one or more microorganisms may be removed from one or more selected plants and, where any successive repeat of the method is to be conducted, grafted on to one or more plant used in step b) of the first aspect or step a) of the second aspect.
The methods of the invention may be described herein in terms of the second set of one or more microorganisms being isolated from its source material. However, unless the context requires otherwise, this should also be taken to include reference to the use of microorganisms in crude form in which they have not been isolated from the source material.

Plants
Any number of a variety of different plants, including mosses and lichens and algae, may be used in the methods of the invention. In preferred embodiments, the plants have economic, social and/or environmental value. For example, the plants may include those of use: as food crops; as fibre crops; as oil crops; in the forestry industry; in the pulp and paper industry; as a feedstock for biofuel production; and/or, as ornamental plants. In other embodiments, the plants may be economically, socially and/or environmentally undesirable, such as weeds. The following is a list of non-limiting examples of the types of plants the methods of the invention may be applied to:

Food crops:
- Cereals (maize, rice, wheat, barley, sorghum, millet, oats, rye, triticale, buckwheat);
- leafy vegetables (brassicaceous plants such as cabbages, broccoli, bok choy, rocket; salad greens such as spinach, cress, lettuce);
- fruiting and flowering vegetables (e.g. avocado, sweet com, artichokes, curcubits e.g. squash, cucumbers, melons, courgettes, pumpkins; solononaceous vegetables/fruits e.g. tomatoes, eggplant, capsicums);
- podded vegetables (groundnuts, peanuts, peas, soybeans, beans, lentils, chickpea, okra);
- bulbed and stem vegetables (asparagus, celery, Allium crops e.g garlic, onions, leeks);
- roots and tuberous vegetables (carrots, beet, bamboo shoots, cassava, yams, ginger, Jerusalem artichoke, parsnips, radishes, potatoes, sweet potatoes, taro, turnip, wasabi);
- sugar crops including sugar beet (Beta vulgaris), sugar cane (Saccharum officinarum);
- **crops grown for the production of non-alcoholic beverages and stimulants**
  (coffee, black, herbal and green teas, cocoa, tobacco);
- **fruit crops** such as true berry fruits (e.g. kiwifruit, grape, currants, gooseberry, guava, feijoa, pomegranate), citrus fruits (e.g. oranges, lemons, limes, grapefruit), epigynous fruits (e.g. bananas, cranberries, blueberries), aggregate fruit (blackberry, raspberry, boysenberry), multiple fruits (e.g. pineapple, fig), stone fruit crops (e.g. apricot, peach, cherry, plum), pip-fruit (e.g. apples, pears) and others such as strawberries, sunflower seeds;
- **culinary and medicinal herbs** e.g. rosemary, basil, bay laurel, coriander, mint, dill, *Hypericum*, foxglove, aloe vera, rosehips);
- **crop plants producing spices** e.g. black pepper, cumin, cinnamon, nutmeg, ginger, cloves, saffron, cardamom, mace, paprika, masalas, star anise;
- **crops grown for the production of nuts** e.g. almonds and walnuts, Brazil nut, cashew nuts, coconuts, chestnut, macadamia nut, pistachio nuts; peanuts, pecan nuts;
- **crops grown for production of beers, wines and other alcoholic beverages** e.g. grapes, hops;
- **oilseed crops** e.g. soybean, peanuts, cotton, olives, sunflower, sesame, lupin species and brassicaeous crops (e.g. canola/oilseed rape); and,
- **edible fune** e.g. white mushrooms, Shiitake and oyster mushrooms;

**Plants used in pastoral agriculture:**
- **legumes**: *Trifolium* species, *Medicago* species, and *Lotus* species; White clover (*T. repens*); Red clover (*T. pratense*); Caucasian clover (*T. ambigum*); subterranean clover (*T. subterraneum*); Alfalfa/Lucerne (*Medicago sativum*); annual medic; barrel medic; black medic; Sainfoin (*Onobrychis viciifolia*); Birdfoot trefoil (*Lotus corniculatus*); Greater Birdfoot trefoil (*Lotus pedunculatus*);
- **seed legumes/pulses** including Peas (*Pisum sativum*), Common bean (*Phaseolus vulgaris*), Broad beans (*Vicia faba*), Mung bean (*Vigna radiata*), Cowpea (*Vigna unguiculata*), Chick pea (*Cicer arietum*), Lupins (*Lupinus* species);
- **Cereals** including Maize/corn (*Zea mays*), Sorghum (*Sorghum spp.*), Millet (*Panicum miliaceum, P. sumatrense*), Rice (*Oryza sativa indica, Oryza sativa*...
japonica), Wheat (Triticum sativa), Barley (Hordeum vulgare), Rye (Secale cereale), Triticale (Triticum X Secale), Oats (Avena sativa);

- **Forage and Amenity grasses**: Temperate grasses such as *Lolium* species; Festuca species; *Agrostis* spp., Perennial ryegrass (*Lolium perenne*); hybrid ryegrass (*Lolium hybridum*); annual ryegrass (*Lolium multiflorum*), tall fescue (*Festuca arundinacea*); meadow fescue (*Festuca pratensis*); red fescue (*Festuca rubra*); *Festuca ovina*; Festuloliuims (*LoHum* X *Festuca* crosses); Cocksfoot (*Dactylis glomerata*); Kentucky bluegrass *Poa pratensis*; *Poa palustris*; *Poa nemoralis*; *Poa trivialis*; *Poa compresa*; Bromus species; Phalaris (Phleum species); *Arrhenatherum elatius*; *Agropyron* species; *Avena strigosa*; Setaria italic;

- **Tropical grasses** such as: *Phalaris* species; *Brachiaria* species; *Eragrostis* species; *Panicum* species; Bahai grass (*Paspalum notatum*); *Brachypodium* species; and,

- Grasses used for biofuel production such as Switchgrass (*Panicum virgatum*) and *Miscanthus* species;

**Fibre crops**:
- cotton, hemp, jute, coconut, sisal, flax (*Linum* spp.), New Zealand flax (*Phonnium* spp.); plantation and natural forest species harvested for paper and engineered wood fibre products such as coniferous and broadleafed forest species;

**Tree and shrub species used in plantation forestry and bio-fuel crops**:
- Pine (*Pinus* species); Fir (*Pseudotsuga* species); Spruce (*Picea* species); Cypress (*Cupressus* species); Wattle (*Acacia* species); Alder (*Alnus* species); Oak species (*Quercus* species); Redwood (*Sequoiadendron* species); willow (*Salix* species); birch (*Betula* species); Cedar (*Cedrus* species); Ash (*Fraxinus* species); Larch (*Larix* species); *Eucalyptus* species; Bamboo (*Bambuseae* species) and Poplars (*Populus* species).

**Plants grown for conversion to energy, biofuels or industrial products by extractive, biological, physical or biochemical treatment**:
- Oil-producing plants such as oil palm, jatropha, soybean, cotton, linseed;
Latex-producing plants such as the Para Rubber tree, *Hevea brasiliensis* and the Panama Rubber Tree *Castillo elastica*;

- plants used as direct or indirect feedstocks for the production of biofuels i.e. after chemical, physical (e.g. thermal or catalytic) or biochemical (e.g. enzymatic pre-treatment) or biological (e.g. microbial fermentation) transformation during the production of biofuels, industrial solvents or chemical products e.g. ethanol or butanol, propane diols, or other fuel or industrial material including sugar crops (e.g. beet, sugar cane), starch-producing crops (e.g. C3 and C4 cereal crops and tuberous crops), cellulosic crops such as forest trees (e.g. Pines, Eucalypts) and Graminaceous and Poaceous plants such as bamboo, switch grass, miscanthus;

- crops used in energy, biofuel or industrial chemical production via gasification and/or microbial or catalytic conversion of the gas to biofuels or other industrial raw materials such as solvents or plastics, with or without the production of biochar (e.g. biomass crops such as coniferous, eucalypt, tropical or broadleaf forest trees, graminaceous and poaceous crops such as bamboo, switch grass, miscanthus, sugar cane, or hemp or softwoods such as poplars, willows; and,

- biomass crops used in the production of biochar;

**Crops producing natural products useful for the pharmaceutical, agricultural, nutraceutical and cosmeceutical industries:**

- crops producing pharmaceutical precursors or compounds or nutraceutical and cosmeceutical compounds and materials for example, star anise (silikimic acid), Japanese knotweed (resveratrol), kiwifruit (soluble fibre, proteolytic enzymes);

**Floricultural Ornamental and Amenity plants grown for their aesthetic or environmental properties:**

- Flowers such as roses, tulips, chrysanthemums;

- Ornamental shrubs such as Buxus, Hebe, Rosa, Rhododendron, Hedera

- Amenity plants such as Platanus, Choisya, Escallonia, Euphorbia, Carex

- Mosses such as sphagnum moss

**Plants grown for bioremediation:**
- Helianthus, Brassica, Salix, Populus, Eucalyptus

It should be appreciated that a plant may be provided in the form of a seed, seedling, cutting, propagule, or any other plant material or tissue capable of growing. In one embodiment the seed may surface-sterilised with a material such as sodium hypochlorite or mercuric chloride to remove surface-contaminating microorganisms. In one embodiment, the propagule is grown in axenic culture before being placed in the plant growth medium, for example as sterile plantlets in tissue culture.

**Growth Medium**

The term "growth medium" as used herein, should be taken broadly to mean any medium which is suitable to support growth of a plant. By way of example, the media may be natural or artificial including, but not limited to, soil, potting mixes, bark, vermiculite, hydroponic solutions alone and applied to solid plant support systems, and tissue culture gels. It should be appreciated that the media may be used alone or in combination with one or more other media. It may also be used with or without the addition of exogenous nutrients and physical support systems for roots and foliage.

In one embodiment, the growth medium is a naturally occurring medium such as soil, sand, mud, clay, humus, regolith, rock, or water. In another embodiment, the growth medium is artificial. Such an artificial growth medium may be constructed to mimic the conditions of a naturally occurring medium, however, this is not necessary. Artificial growth media can be made from one or more of any number and combination of materials including sand, minerals, glass, rock, water, metals, salts, nutrients, water. In one embodiment, the growth medium is sterile. In another embodiment, the growth medium is not sterile.

The medium may be amended or enriched with additional compounds or components, for example, a component which may assist in the interaction and/or selection of specific groups of microorganisms with the plant and each other. For example, antibiotics (such as penicillin) or sterilants (for example, quaternary ammonium salts and oxidizing agents) could be present and/or the physical conditions (such as salinity, plant nutrients (for example organic and inorganic minerals (such as phosphorus, nitrogenous salts, ammonia, potassium and micronutrients such as cobalt and magnesium), pH, and/or temperature) could be amended.
In certain embodiments of the invention, the growth medium may be pre-treated to assist in
the survival and/or selection of certain microorganisms. For example, the medium may be
pre-treated by incubating in an enrichment media to encourage the multiplication of
endogenous microbes that may be present therein. By way of further example, the medium
may be pre-treated by incubating in a selective medium to encourage the multiplication of
specific groups of microorganisms. A further example includes the growth medium being
pre-treated to exclude a specific element of the microbial assemblage therein; for example
pasteurization (to remove spore-forming bacteria and fungi) or treatment with organic
solvents such as various alcohols to remove microorganisms sensitive to these materials
but allow the survival of actinomycetes and spore-forming bacteria, for example.

Methods for pre-treating or enriching may be informed by culture independent microbial
community profiling techniques that provide information on the identity of microbes or
groups of microbes present. These methods may include, but are not limited to,
sequencing techniques including high throughput sequencing and phylogenetic analysis, or
microarray-based screening of nucleic acids coding for components of rRNA operons or
other taxonomically informative loci.

20 Growth Conditions
In accordance with the methods of the invention one or more plant is subjected to one or
more microorganism and a growth medium. The plant is preferably grown or allowed to
multiply in the presence of the one or more microorganism(s) and growth medium. The
microorganism(s) may be present in the growth medium naturally without the addition of
further microorganisms, for example in a natural soil. The growth medium, plant and
microorganisms may be combined or exposed to one another in any appropriate order. In
one embodiment, the plant, seed, seedling, cutting, propagule or the like is planted or sown
into the growth medium which has been previously inoculated with the one or more
microorganisms. Alternatively, the one or more microorganisms may be applied to the
plant, seed, seedling, cutting, propagule or the like which is then planted or sown into the
growth medium (which may or may not contain further microorganisms). In another
embodiment, the plant, seed, seedling, cutting, propagule or the like is first planted or sown
into the growth medium, allowed to grow, and at a later time the one or more
microorganisms are applied to the plant, seed, seedling, cutting, propagule or the like and/or the growth medium itself is inoculated with the one or more microorganisms.

The microorganisms may be applied to the plant, seedling, cutting, propagule or the like and/or the growth medium using any appropriate techniques known in the art. However, by way of example, in one embodiment, the one or more microorganisms are applied to the plant, seedling, cutting, propagule or the like by spraying or dusting. In another embodiment, the microorganisms are applied directly to seeds (for example as a coating) prior to sowing. In a further embodiment, the microorganisms or spores from microorganisms are formulated into granules and are applied alongside seeds during sowing. In another embodiment, microorganisms may be inoculated into a plant by cutting the roots or stems and exposing the plant surface to the microorganisms by spraying, dipping or otherwise applying a liquid microbial suspension, or gel, or powder. In another embodiment the microorganism(s) may be injected directly into foliar or root tissue, or otherwise inoculated directly into or onto a foliar or root cut, or else into an excised embryo, or radicle or coleoptile. These inoculated plants may then be further exposed to a growth media containing further microorganisms, however, this is not necessary. In certain embodiments, the microorganisms are applied to the plant, seedling, cutting, propagule or the like and/or growth medium in association with plant material (for example, plant material with which the microorganisms are associated).

In other embodiments, particularly where the microorganisms are unculturable, the microorganisms may be transferred to a plant by any one or a combination of grafting, insertion of explants, aspiration, electroporation, wounding, root pruning, induction of stomatal opening, or any physical, chemical or biological treatment that provides the opportunity for microbes to enter plant cells or the intercellular space. Persons of skill in the art may readily appreciate a number of alternative techniques that may be used.

It should be appreciated that such techniques are equally applicable to application of the first set of one or more microorganisms and the second set of microorganisms when used in step a) of a successive repeat of the method.

In one embodiment the microorganisms infiltrate parts of the plant such as the roots, stems, leaves and/or reproductive plant parts (become endophytic), and/or grow upon the surface
of roots, stems, leaves and/or reproductive plant parts (become epiphytic) and/or grow in
the plant rhizosphere. In one preferred embodiment microorganism(s) form a symbiotic
relationship with the plant.

The growth conditions used may be varied depending on the species of plant, as will be
appreciated by persons skilled in the art. However, by way of example, for clover, in a
growth room one would typically grow plants in a soil containing approximately 1/3rd
organic matter in the form of peat, 1/3rd compost, and 1/3rd screened pumice,
supplemented by fertilisers typically containing nitrates, phosphates, potassium and
magnesium salts and micronutrients and at a pH of between 6 and 7. The plants may be
grown at a temperature between 22-24°C in an 16:8 period of daylight:darkness, and
watered automatically.

Selective Pressure

In certain aspects and embodiments of the invention, at a desired time during the period
within which the plant is subjected to one or more microorganism and a growth medium, a
selective pressure is applied. The selective pressure may be any biotic or abiotic factor or
element which may have an impact on the health, growth, and/or survival of a particular
plant, including environmental conditions and elements which plants may be exposed to in
their natural environment or a commercial situation.

Examples of biotic selective pressures include but are not limited to organisms that are
detrimental to the plant, for example, fungi, bacteria, viruses, insects, mites, nematodes,
animals.

Abiotic selective pressures include for example any chemical and physical factors in the
environment; for example, water availability, soil mineral composition, salt, temperature,
alterations in light spectrum (e.g. increased UV light), pH, organic and inorganic toxins
(for example, exposure to or changes in the level of toxins), metals, organic nutrients,
inorganic nutrients, air quality, atmospheric gas composition, air flow, rain fall, and hail.

For example, the plant/microorganisms maybe exposed to a change in or extreme salt
concentrations, temperature, pH, higher than normal levels of atmospheric gases such as
CO₂, water levels (including drought conditions or flood conditions), low nitrogen levels,
provision of phosphorus in a form only available to the plant after microbial degradation, exposure to or changes in the level of toxins in the environment, soils with nearly toxic levels of certain minerals such as alurainates, or high winds.

In one embodiment the selective pressure is applied directly to the plant, the microorganisms and/or the growth medium. In another embodiment the selective pressure is applied indirectly to the plant, the microorganisms and/or the growth medium, via the surrounding environment; for example, a gaseous toxin in the air or a flying insect.

The selective pressure may be applied at any time, preferably during the time the plant is subjected to the one or more microorganism and growth medium. In one embodiment, the selective pressure is applied during for substantially the whole time during which a plant is growing and/or multiplying. In another embodiment, the selective pressure is applied at a discrete time point during growth and/or multiplication. By way of example, the selective pressure may be applied at different growth phases of the one or more plants which simulate a potential stress on the plant that might occur in a natural or commercial setting. For example, the inventor has observed that some pests attack plants only at specific stages of the plant's life. In addition, the inventor has observed that different populations of potentially beneficial microorganisms can associate with plants at different points in the plant's life. Simulating a pest attack on the plant at the relevant time point, may allow for the identification and isolation of microorganisms which may protect the plant from attack at that particular life stage. It should also be appreciated that the selective pressure may be present in the growth medium or in the general environment at the time the plant, seed, seedling, cutting, propagule or the like is planted or sown.

In one embodiment, the microbial population is exposed (prior to the method or at any stage of the method) to a selective pressure to enhance the probability that the eventually-selected plants will have microbial assemblages likely to have desired properties. For example, exposure of the microorganisms to pasteurisation before their addition to a plant growth medium (preferably sterile) is likely to enhance the probability that the plants selected for a desired trait will be associated with spore-forming microbes that can more easily survive in adverse conditions, in commercial storage, or if applied to seed as a coating, in an adverse environment. Another example is provided herein after in Example 11IB, in which microorganisms were subjected to a media which allowed for selection of
phosphate solubilising microbes. Such a step of applying a selective pressure to the microbial population may be referred to herein as an "enrichment step".

The plants may be grown and subjected to the selective pressure for any appropriate length of time before they are selected and harvested. By way of example only, the plants and any microorganisms associated with them may be selected and harvested at any time during the growth period of a plant, in one embodiment, any time after germination of the plant. In a preferred embodiment, the plants are grown or allowed to multiply for a period which allows one to distinguish between plants having desirable phenotypic features and those that do not. By way of general example wheat may be selected for improvements in the speed of foliar growth say after one month, but equally may be selected for superior grain yield on maturity of the seed head. The length of time a plant is grown depends on the timing required to express the plant trait that is desired to be improved by the invention, or the time required to express a trait correlated with the desired trait. For example, in the case of winter wheat varieties, mainly sown in the Northern Hemisphere, it may be important to select plants that display early tillering after exposure of seed to a growth medium containing microorganisms under conditions of light and temperature similar to those experienced by winter wheat seed in the Northern Hemisphere, since early tillering is a trait related to winter survival, growth and eventual grain yield in the summer.

Or, a tree species may be selected for improved growth and health at 4-6 months as these traits are related to the health and growth rate and size of trees of 10 years later, an impractical period product development using this invention.

It should be appreciated that the methods of the invention may involve applying two or more selective pressures simultaneously or successively in step b).

Selection
Typically, following growth of the one or more plants in the presence of one or more microorganisms, and in certain embodiments following exposure to a selective pressure, one or more plant is selected based on one or more selection criterion. In one embodiment the plants are selected on the basis of one or more phenotypic traits. Skilled persons will readily appreciate that such traits include any observable characteristic of the plant, including for example growth rate, height, weight, colour, taste, smell, changes in the production of one or more compounds by the plant (including for example, metabolites,
proteins, drugs, carbohydrates, oils, and any other compounds). Selecting plants based on genotypic information is also envisaged (for example, including the pattern of plant gene expression in response to the microorganisms, genotype, presence of genetic markers). It should be appreciated that in certain embodiments, plants may be selected based on the absence, suppression or inhibition of a certain feature or trait (such as an undesirable feature or trait) as opposed to the presence of a certain feature or trait (such as a desirable feature or trait).

Where the presence of one or more genetic marker is assessed, the one or more marker may already be known and/or associated with a particular characteristic of a plant; for example, a marker or markers may be associated with an increased growth rate or metabolite profile. This information could be used in combination with assessment based on other characteristics in a method of the invention to select for a combination of different plant characteristics that may be desirable. Such techniques may be used to identify novel QTLs which link desirable plant traits with a specific microbial flora - for example matching plant genotype to the microbiome type.

By way of example, plants may be selected based on growth rate, size (including but not limited to weight, height, leaf size, stem size, branching pattern, or the size of any part of the plant), general health, survival, tolerance to adverse physical environments and/or any other characteristic, as described herein before. Further non-limiting examples include selecting plants based on: speed of seed germination; quantity of biomass produced; increased root, and/or leaf/shoot growth that leads to an increased yield (herbage or grain or fibre or oil) or biomass production; effects on plant growth that results in an increased seed yield for a crop, which may be particularly relevant in cereal crops such as wheat, barley, oats, rye, maize, rice, sorghum, oilseed crops such as soybean, canola, cotton, sunflower, and seed legumes such as peas, beans; effects on plant growth that result in an increased oil yield, which may be particularly relevant in oil seed crops such as soybean, canola, cotton, jatropha and sunflower; effects on plant growth that result in an increased fibre yield (e.g. in cotton, flax and linseed) or for effects that result in an increased tuber yield in crops such as potatoes and sugar beet; effects on plant growth that result in an increased digestibility of the biomass which may be particularly relevant in forage crops such as forage legumes (alfalfa, clovers, medics), forage grasses (Lolium species; Festuca species; Paspalum species; Brachiaria species; Eragrostis species), forage crops grown for
silage such as maize and forage cereals (wheat, barley, oats); effects on plant growth which result in an increased fruit yield which may be particularly relevant to pip fruit trees (such as apples, pears, etc), berry fruits (such as strawberries, raspberries, cranberries), stone fruit (such as nectarines, apricots), and citrus fruit, grapes, figs, nut trees; effects on plant growth that lead to an increased resistance or tolerance disease including fungal, viral or bacterial diseases or to pests such as insects, mites or nematodes in which damage is measured by decreased foliar symptoms such as the incidence of bacterial or fungal lesions, or area of damaged foliage or reduction in the numbers of nematode cysts or galls on plant roots, or improvements in plant yield in the presence of such plant pests and diseases; effects on plant growth that lead to increased metabolite yields, for example in plants grown for pharmaceutical, nutraceutical or cosmeceutical purposes which may be particularly relevant for plants such as star anise grown for the production of shikimic acid critical for the production of anti-influenza drug oseltamivir, or the production of Japanese knotweed for the extraction of resveratrol, or the production of soluble fibre and dietary enzyme products from kiwifruit, or for example increased yields of "condensed tannins" or other metabolites useful for inhibiting the production of greenhouse gases such as methane in grazing animals; effects on plant growth that lead to improved aesthetic appeal which may be particularly important in plants grown for their form, colour or taste, for example the colour intensity and form of ornamental flowers, the taste of fruit or vegetable, or the taste of wine from grapevines treated with microorganisms; and, effects on plant growth that lead to improved concentrations of toxic compounds taken up or detoxified by plants grown for the purposes of bioremediation.

Selection of plants based on phenotypic or genotypic information maybe performed using techniques such as, but not limited to: high through-put screening of chemical components of plant origin, sequencing techniques including high through-put sequencing of genetic material, differential display techniques (including DDRT-PCR, and DD-PCR), nucleic acid microarray techniques, RNA-seq (Whole Transcriptome Shotgun Sequencing), qRT-PCR (quantitative real time PCR).

In certain embodiments of the invention, selection for a combination of plant traits may be desired; for example, in the embodiments of the invention which involve repeating the basic method steps and applying different selective pressures with each iteration of the method. This can be achieved in a number of ways, for example, one embodiment, multiple rounds
of iterative improvement for one trait, e.g. superior growth in nematode infested soils, are maintained until an acceptable level of nematode resistance is attained. Similar, but completely separate rounds of selection are undertaken to identify microorganisms that can confer at least different desirable traits, for example simply for improved growth or other characteristics, or improved growth resulting from improved microbial soil phosphate utilisation, or improved growth resulting from increased tolerance to sucking insect pests. Such separate rounds of selection may be performed using an iterative or stacking approach or a combination of separate methods could be used, with the microorganisms that result from those separate rounds or methods being combined into a single composition. At this point the microorganism(s) could be developed into a product containing combinations of separately-fermented microorganisms each shown to improve a different plant attribute. In a further embodiment, the separately selected sets of microorganisms may be combined in sets of two or more and used in further methods of the invention, h in another embodiment, the separately selected sets of microorganisms may be separated into individual isolates and then individual isolates combined in sets of two or more and used in further methods of the invention. In one embodiment, the combined microorganisms are applied to the plant and/or growth medium for the application of two or more selection pressures in the same iterative cycle. For example, in one combination, microorganisms able to improve plant growth in a medium containing low-levels of plant-available phosphorus are combined with microorganisms able to enhance plant growth in soils infested with plant parasitic nematodes. The combined microorganisms are then added to a plant growth medium with low levels of available phosphorus in which the plants are grown for a suitable period, nematodes applied and the plants are further grown until nematode damage can be expressed. The degree of nematode root damage and plant biomass is assessed non-destructively and microbes are isolated from the best-performing plants for use in a succeeding iteration. Similar iterative rounds may be continued until an acceptable level of plant growth is attained under both selective pressures. This approach will aid the selection of microbes that synergistically improve plant performance i.e. improve plant growth and nematode resistance to a degree better than that achieved if the microorganisms are applied simply as a combination of two separately-selected sets. The same approach could be used in methods of the invention which do not employ the application of one or more selective pressure. For example, the microorganisms acquired from separate runs of a method of the invention, each shown to be associated with different desirable plant characteristics (by way of non-limiting example, sugar or fibre content and
growth rate, or flower colour and height) could be combined into a single composition and applied to one or more plants in a further round of the method for selection of plants having both characteristics or yet a further characteristic.

5 Harvesting

Following selection, one or more plants are harvested and plant tissues may be examined to detect microorganisms forming associations with the plants (for example, endophytic, epiphytic or rhizospheric associations).

The techniques described in this section may be used in acquiring a second set of microorganisms at the conclusion of a method of the invention or for use in any successive repeat of the methods of the invention.

The one or more microorganisms may be isolated from any appropriate tissue of the plants selected; for example, whole plant, foliar tissue, stem tissue, root tissue, and/or seeds. In a preferred embodiment, the microorganisms are isolated from the root tissue, stem or foliar tissues and/or seeds of the one or more plants selected.

In certain embodiments, the microorganisms may be acquired in crude form, in which they are not isolated from the source material in which they reside (such as plant tissue or growth media).

Where isolation of the microorganisms occurs, they may be isolated from the plants using any appropriate methods known in the art. However, by way of example, methods for isolating endophytic microbes may include the sterile excision of the plant material of interest (e.g. root, stem lengths, seed), surface sterilisation with an appropriate solution (e.g. 2% sodium hypochlorite), after which the plant material is placed on nutrient medium for microbial outgrowth, especially filamentous fungi. Alternatively, the surface-sterilised plant material can be crushed in a sterile liquid (usually water) and the liquid suspension, including small pieces of the crushed plant material spread over the surface of a suitable solid agar medium, or media, which may or may not be selective (e.g. contain only phytic acid as a source of phosphorus). This approach is especially useful for bacteria and yeasts which form isolated colonies and can be picked off individually to separate plates of nutrient medium, and further purified to a single species by well-known methods.
Alternatively, the plant root or foliage samples may not be surface sterilised but only washed gently thus including surface-dwelling epiphytic microorganisms in the isolation process, or the epiphytic microbes can be isolated separately, by imprinting and lifting off pieces of plant roots, stem of leaves on to the surface of an agar medium and then isolating individual colonies as above. This approach is especially useful for bacteria and yeasts, for example. Alternatively, the roots may be processed without washing off small quantities of soil attached to the roots, thus including microbes that colonise the plant rhizosphere. Otherwise, soil adhering to the roots can be removed, diluted and spread out onto agar of suitable selective and non-selective media to isolate individual colonies of rhizospheric microbes. Further exemplary methodology can be found in: Strobel G and Daisy B (2003) Bioprospecting for microbial endophytes and their natural products. Microbiology and Molecular Biology Reviews 67 (4): 491-502; Zinniel DK et al. (2002) Isolation and characterisation of endophytic colonising bacteria from agronomic crops and prairie plants. Applied and Environmental Microbiology 68 (5): 2198-2208; Manual of Environmental Microbiology, Hurst et al., ASM Press, Washington DC.

Methods for isolation may be informed by culture independent community profiling techniques that provide information on the identity and activity of microbes present in a given sample. These methods may include, but are not limited to, sequencing techniques including high throughput sequencing and phylogenetic analysis, or microarray-based screening of nucleic acids coding for components of rRNA operons or other taxonomically informative loci.

Embodiments of the invention where two or more microorganism are isolated from plant material and then separated into individual isolates, any appropriate methodology for separating one or more microorganism from each other may be used. However, by way of example, microbial extracts prepared from plant material could be spread on agar plates, grown at an appropriate temperature for a suitable period of time and the resulting microbial colonies subsequently selected and grown in an appropriate media (for example, streaked onto fresh plates or grown in a liquid medium). The colonies may be selected based on morphology or any other appropriate selection criteria as will be understood in the art. By way of further example, selective media could be used.
The one or more microorganisms may be harvested (including in isolated or crude form) from the plants (including the rhizosphere as described herein before) at any appropriate time point. In one embodiment they are harvested at any time after germination of the plant. For example, they can be isolated from the period shortly after germination (where survival in the first few days after germination is an issue, for example with bacterial and fungal root and collar rots), then at any stage after that, depending on the timing required for a plant to grow in order to evidence a discriminatory benefit that enables it's selection from the plant population (for example, to discriminate say the top 10 of 200 plants).

The inventor(s) has observed that different microorganisms may associate with a plant at different stages of the plant's life. Accordingly, harvesting a plant at different time points may result in selection of a different population of microorganisms. Such microorganisms may be of particular benefit in improving plant condition, survival and growth at critical times during its life: by way of example, as mentioned herein before, a plant may be susceptible to attack by nematodes at discrete time points during its life and the invention may be used to identify and isolate a population of microorganisms which may increase resistance to such attack at that particular life stage.

In another embodiment of the invention, in the case of microorganisms that form an association with a plant that allows vertical transmission from one generation or propagule to the next (for example seed-endophytic or -epiphytic associations, or endophytic and epiphytic associations with plants/propagules multiplied vegetatively) the microorganisms may not be isolated from the plant(s). At the conclusion of a method of the invention a target or selected plant itself may be multiplied by seed or vegetatively (along with the associated microorganisms) to confer the benefit(s) to "daughter" plants of the next generation or multiplicative phase. Similarly, where a successive repeat of the method is desired, plant material (whole plant, plant tissue, part of the plant) comprising the set of one or more microorganisms can be used in step b) (first aspect) or step a) (second aspect) of the successive repeat.

Stac Jans
The inventor(s) envisage advantages being obtained by stacking selective pressures in repeated rounds of the method of the invention. This may allow for acquiring a population of microorganisms that may assist a plant in surviving in a number of different
environmental conditions, resisting a number of different diseases and attack by a number
of different organisms, for example.

Similarly, the inventor(s) envisage advantages being obtained by stacking the means of
selection (or the selection criteria) of plants in repeated rounds of the method of the
invention. This may allow for the acquiring a population of microorganisms that may
assist a plant in having a number of different desirable traits, for example.

One could also stack both selective pressures and selection criteria in methods of the
invention.

In one embodiment of the invention the one or more microorganisms acquired from the
one or more plants selected following exposure to a selective pressure, as previously
described, is used in a second round or cycle of the method; i.e. the microorganisms from
the selected plants are provided, along with one or more plants and a growth medium, a
selective pressure is applied, plants are selected at a desired time and microorganisms are
isolated from the selected plants. The microorganisms acquired from the second round of
the method may then be used in a subsequent round, and so on and so on.

In one embodiment, the selective pressure applied in each repeat of the method is different.
For example, in the first round the pressure may be a particular soil pH and in the second
round the pressure may be nematode attack. However, in other embodiments of the
invention, the selective pressure applied in each round may be the same. It could also be
the same but applied at differing intensities with each round. For example, in the first
round the selective pressure may be a particular concentration of salt present in the soil. In
the second round, the selective pressure may be a higher concentration of salt present in
the soil. In one embodiment, the selective pressure is increased in successive rounds in a
pattern that may be linear, stepped or curvilinear. For example in round 1 of an iterative
selective process wheat plus microorganisms may be exposed to 100 mM NaCl, in the
second to 110 mM salt, in the third to 120 mM salt, thus increasing the selective pressure
on the plants as adaptation occurs via improved plant/microorganism associations.
Alternatively, it may be advantageous to maintain a selective pressure of 120mM for
several rounds to allow for a slower adjustment in the microbial population balance
underlying improvements in the ability of wheat to grow productively in a higher salt environment.

In one embodiment, a selective pressure may be separated disjunctively from a specific step of the iterative process, particularly the first round of an iterative cycle. For example in round one the selective pressure may not be applied at all. But after the microorganisms have been isolated from the selected plants after exposure for a relevant period to a growth medium and microorganisms in round 1, they are applied to the plant growth medium along with the plant, seed, seedling, cutting, propagule or the like for round 2. After an appropriate time a selective pressure is applied in round 2 and in successive rounds. This type of selection may be especially relevant for selection factors that severely diminish the plant tissue that is the target of the selection. For example nematodes are especially destructive of root tissue and it may be advantageous to allow particular microbes to multiply to high levels on, in, or around the roots in round 1 to allow high concentrations of microorganisms from the roots of plants selected in round 1 to be applied to the growth medium in round 2.

Where selection criteria are stacked, the one or more microorganisms acquired from the one or more plants selected, as previously described, is used in a second round or cycle of the method, where a different selection criterion is used. For example, in the first round, one or more plants may have been selected based on biomass. In the second round, one or more plants may be selected based on production of a particular compound. The microorganisms from the second round of the method may then be used in a subsequent round, and so on and so on. Any number of different selection criteria may be employed in successive rounds of the method, as desired or appropriate.

In one embodiment, the selection criteria applied in each repeat of the method is different. However, in other embodiments of the invention, the selection criteria applied in each round may be the same. It could also be the same but applied at differing intensities with each round. For example, the selection criteria may be fibre levels and level of fibre required for a plant to be selected may increase with successive rounds of the method. The selective criteria may increase or decrease in successive rounds in a pattern that may be linear, stepped or curvilinear.
It should be appreciated that the methods of the first and second aspects of the invention maybe combined such that they alternate. For example, a method of the first aspect is performed and the microorganisms acquired in that method used in step a) of a method of the second aspect of the invention. The microorganisms acquired in step c) of a method of the second aspect of the invention may then be used in step b) of a method of the first aspect. This may be repeated any number of times as necessary or desired. It should be appreciated that the methods may be combined in any order and in any combination or permutation. For example, they need not alternate on a one by one basis. Any number of repeats of one a method of one aspect may be performed before switching to one or more round of a method of the other aspect. The methods of the first and second aspect may also be run independently and then the microorganisms acquired in the methods combined and used in a further round of one or both methods, and so on.

It should also be appreciated that in certain embodiments of the invention, where one or more microorganism(s) forms an endophytic or epiphytic relationship with a plant that allows vertical transmission from one generation or propagule to the next the microorganisms need not be isolated from the plant(s). At the conclusion of a method of the invention a target or selected plant itself may be multiplied by seed or vegetatively (along with the associated microorganisms) to confer the benefit(s) to "daughter" plants of the next generation or multiplicative phase. Similarly, where a successive repeat of the method is desired, plant material (whole plant, plant tissue, part of the plant) comprising the set of one or more microorganisms can be used in step b) (first aspect) or step a) (second aspect) of the successive repeat.

It should further be appreciated that two or more selective pressures and/or two or more selection criterion may be applied with each iteration of a method of the invention.

**Identification of genetic markers**

In one embodiment, the methods of the invention involve identifying one or more plant marker associated with one or more beneficial property in a plant (for example, the first aspect of the invention) prior to subjecting one or more plant to a growth media in the presence of a first set of one or more microorganisms. In another embodiment, the methods of the invention involve identifying one or more plant marker and/or one or more
microorganism marker associated with one or more beneficial property in a plant (for example, the second aspect of the invention).

In the second embodiment, the step of identifying one or more marker may occur after step b) and/or after step c) and prior to step a) of any successive repeat of the method. It may also occur after step c) at the conclusion of a method of the invention.

In one embodiment, the one or more marker may already be known and known to be associated with a beneficial property in a plant. In this way, it may be possible to identify the presence of a particular beneficial property in the plant, prior to its physical manifestation, for example. In other embodiments, the one or more marker may not have previously been known and the methods of the invention allow for the identification of novel markers or novel marker/beneficial property correlations or associations. Once any new marker/beneficial property correlations or associations are identified, they may be used in any successive repeat of a method of the invention to select desirable plants.

So, the invention also provides methods for identifying one or more plant marker that is associated with one or more beneficial trait and/or one or more microorganism marker that is associated with one or more beneficial plant trait.

These methods will generally involve conducting a method of the invention and screening for markers and/or correlations between one or more marker and the presence of one or more beneficial property in the plant.

Markers may be screened and identified using any standard methodologies known in the art to which the invention relates. However, by way of example, well known methods for detection and analysis of the following types of markers may be used: single nucleotide polymorphism (SNP), simple sequence repeats (SSRs or microsatellites), RFLPs, and transposable element positions. Similarly, correlations between the presence or absence of one or more marker and one or more beneficial property may be identified using any standard methodologies known in the art.
Microorganisms and compositions containing same
In addition to the methods described herein before, the invention relates to microorganisms obtained by such methods and compositions comprising such microorganisms. In its simplest form, a composition comprising one or more microorganisms includes a culture of living microorganism, or microorganisms in a live but inactive state(s), including frozen, lyophilised or dried cultures. However, the compositions may comprise other ingredients, as discussed below.

The invention should also be understood to comprise methods for the production of a composition to support plant growth, quality and/or health, or a composition to suppress or inhibit plant growth, quality and/or health, the method comprising the steps of a method herein before described and the additional step of combining the one or more microorganisms with one or more additional ingredients.

A "composition to support plant growth, health, and/or quality" should be taken broadly to include compositions which may assist the growth, general health and/or survival of a plant, the condition of a plant, or assist in the maintaining or promoting any desired characteristic, quality, and/or trait. It should be taken to include maintaining or altering the production of one or more metabolite or other compound by a plant as well altering gene expression and the like. The phrase should not be taken to imply that the composition is able to support plant growth, quality and/or health on its own. However, in one embodiment the compositions are suitable for this purpose. Exemplary compositions of this aspect of the invention include but are not limited to plant growth media, plant mineral supplements and micronutrients, composts, fertilisers, potting mixes, insecticides, fungicides, media to protect against infection or infestation of pests and diseases, tissue culture media, see coatings, hydroponic media, compositions that impart tolerance to drought or abiotic stress such as metal toxicity, compositions that modify soil pH.

A "composition to inhibit or suppress plant growth, health, and/or quality" should be taken broadly to include compositions which may assist in suppressing or inhibiting one or more characteristic or quality and/or trait of a plant, including its growth, general health and/or survival. It should be taken to include maintaining or altering the production of one or more metabolite or other compounds by a plant as well altering gene expression and the like. The phrase should not be taken to imply that the composition is able to suppress or
inhibit plant growth, quality and/or health on its own. However, in one embodiment the compositions are suitable for this purpose. Exemplary compositions of this aspect of the invention include but are not limited to plant growth suppression media, weed killer, fertilisers, potting mixes, plant mineral supplements and micronutrients, composts, mixes, insecticides, fungicides, tissue culture media, seed coatings, hydroponic media, compositions that impart tolerance to drought or abiotic stress such as metal toxicity, compositions that modify soil pH.

Skilled persons will readily appreciate the types of additional ingredients that may be combined with the one or more microorganisms, having regard to the nature of the composition that is to be made, the microorganisms to be used, and/or the method of delivery of the composition to a plant or its environment. However, by way of example, the ingredients may include liquid and/or solid carriers, microbial preservatives, microbial activators that induce specific metabolic activities, additives to prolong microbial life (such as gels and clays), wettable powders, granulated eariners, soil, sand, agents known to be of benefit to microbial survival and the growth and general health of a plant, peat, organic matter, organic and inorganic fillers, other microorganisms, wetting agents, organic and inorganic nutrients, and minerals.

Such compositions can be made using standard methodology having regard to the nature of the ingredients to be used.

Compositions developed from the methods of the invention may be applied to a plant by any number of methods known to those skilled in the art. These include for example: sprays; dusts; granules; seed-coating; seed spraying or dusting upon application; germinating the seed in a bed containing suitable concentrations of the composition prior to germination and planting out of the seedlings; prills or granules applied next to the seed or plant during sowing or planting, or applied to an existing crop through a process such as direct drilling; application to plant cuttings or other vegetative propagules by dipping the cut surface or the propagule into liquid or powdered microbial substrate prior to planting; application to the soil as a "soil treatment" in the form of a spray, dust, granules or composted composition that may or may not be applied with plant fertilisers prior to or after sowing or planting of the crop; application to a hydroponic growth medium; inoculation into plant tissues under axenic conditions via injection of compositions or
otherwise inoculated via a cut in such tissues, for the subsequent establishment of an
dendyrophytic relationship with the plant that extends to the seed, or propagative tissues, such
that the plant can be multiplied via conventional agronomic practice, along with the
dendyrophytic microbe providing a benefit(s) to the plant.

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**Method of producing alternative compositions**

When microorganisms are cultured they may produce one or more metabolites and which
are passed into the media in which they reside. Such metabolites may confer beneficial
properties to plants.

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Accordingly, the invention also provides a method of selecting or producing a composition
capable of imparting one or more beneficial property to a plant, for example to support
plant growth, quality and/or health, or for example to suppress or inhibit growth, quality
and/or health of a plant, or to identify microorganisms that are capable of producing such a
composition. In one embodiment, the composition is substantially free of microorganisms.

In one embodiment, the method is for the selection of a composition and comprises at least
the steps of:

a) culturing one or more microorganism selected by a method of the first aspect
and/or a method of the second aspect in one or more media to provide one or more
culture;

b) separating the one or more microorganism from the one or more media in the
one or more culture after a period of time to provide one or more composition
substantially free of microorganisms;

c) subjecting one or more plant (including for example seeds, seedlings, cuttings,
and/or propagules thereof) to the one or more composition of step b);

d) selecting one or more composition from step c) if it is observed to impart one or
more beneficial property to the one or more plants.

30 In another embodiment, the method is for the selection of a composition and comprises at
least the steps of:

a) culturing one or more microorganisms selected by a method of the first aspect
and/or the second aspect of the invention in one or more media to provide one or
more culture;
b) inactivating the one or more culture of step a) to provide one or more composition containing one or more inactivated microorganisms;

c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition of step b);

d) selecting one or more composition from step c) if it is observed to impart one or more beneficial property to the one or more plants.

In another embodiment, the method is for the selection of one or more microorganisms which are capable of producing a composition and comprises at least the steps of:

- a) culturing one or more microorganism selected by a method of the first aspect and/or the second aspect in one or more media to provide one or more culture;
- b) separating the one or more microorganism from the one or more media in the one or more culture from step a) after a period of time to provide one or more composition substantially free of microorganisms;
- c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition from step b);
- d) selecting the one or more microorganisms associated with (or in other words used to produce the) one or more composition observed to impart one or more beneficial property to the one or more plants.

In another embodiment the method is for the selection of one or more microorganisms which are capable of producing a composition and comprises at least the steps of:

- a) culturing one or more microorganism in one or more media to provide one or more culture;
- b) separating the one or more microorganism from the one or more media in one or more culture after a period of time to provide one or more composition substantially free of microorganisms;
- c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition of step b);
- d) selecting the one or more microorganisms associated with (or in other words used to produce the) one or more composition observed to impart one or more beneficial property to the one or more plants; and,
- e) using the one or more microorganisms selected in a method of the first, second, eighth (and/or related) and/or ninth (and/or related) aspects of the invention.
In another embodiment, step b) of the method of the previous two paragraphs could be substituted with the step of b) inactivating the one or more culture of step a) to provide one or more composition containing one or more inactivated microorganisms, and then using this composition in step c) of the process instead of the composition.

As used herein "inactivating" the one or more culture and "inactivated microorganisms" and like terms should be taken broadly to mean that the microorganisms are substantially inactivated, killed, fixed or otherwise destroyed. The term should not be taken to mean that all microorganisms are inactivated, killed or destroyed, however, this may be preferable. In one embodiment, the microorganisms are inactivated, fixed, killed or destroyed to the extent that self-sustained replication is no longer measurable using techniques known to one skilled in the art.

The microorganisms can be inactivated, killed or destroyed using any appropriate techniques known in the art. However, by way of example, one may use chemical agents and/or physical means to do so. In one embodiment, the cells are lysed. In another embodiment, cells are fixed by chemical means, so as to render the organisms non-viable, but retaining their structural integrity.

As used herein, a "composition substantially free of microorganisms" should be taken broadly and not be construed to mean that no microorganisms are present, although this may be preferred.

In certain embodiments of these methods, microorganisms are cultured in two or more (preferably a large number, for example, from at least approximately 10 to up to approximately 1000) mixed cultures using media that can support the growth of a wide variety of microorganisms. Any appropriate media known in the art may be used. However, by way of example, growth media may include, TSB (tryptic soy broth), Luria-Bertani (LB) broth, or R2A broth. In another embodiment, selective or enrichment media which are able to support the growth of microorganisms with an array of separate but desirable properties may be used. By way of example, the enrichment media referred to elsewhere herein may be used.
The microorganisms may be cultured in the media for any desired period. Following culture, the microorganisms are separated from the media and stored for later use. A separate composition also results. One or more plants in a suitable growth medium are then subjected to the composition (using any known methodology, or methodology as described herein before). After a period of time, growth of plants is assessed and plants selected (as described herein before, for example). Plants are preferably selected on the basis of size. However, other selection criteria as referred to herein may be used.

In one embodiment, the microorganism(s) producing the subset of compositions associated with the selected plants are recovered from storage. Two or more separate cultures of the microorganisms may then be mixed together and separated into two or more sub-cultures grown in two or more different media.

This process can be repeated iteratively as many times as is deemed efficacious, with progressive steps refining down to fewer media and a narrower diversity of microorganisms until a desirable effect on the growth plants is achieved with a mixture of microbes that can be identified, grown and stored indefinitely as a standard starting inoculum for the production the composition.

Compositions of this aspect of the invention may be used or formulated on their own or combined with one or more additional ingredients.

It should be appreciated that the general methodology described herein before may be applicable to this aspect of the invention, including but not limited to growth media, plants, microorganisms, selective pressures, timing, iterative processing, and combinations thereof.

**Additional Methodology**

The following methodology may be applied to a method of the invention for identifying one or more microorganisms as herein before described.

Figure 1 shows a system 10 according to an embodiment of the invention. System 10 includes requestors 11, request processor 12, growing facility 13, database or library 14 and depository 15.
Figure 2 provides a flow chart illustrating a method 20 according to an embodiment of the invention. The steps shown in Figure 2 will be described with reference to the system 10 shown in Figure 1.

This aspect of the invention is described in terms of identifying one or more microorganism that may impart one or more desired properties to one or more plants, with particular reference to the first and/or second aspects of the invention. However, it should be appreciated that it is equally applicable to the identification of one or more compositions that may impart one or more desired property to one or more plant, or one or more microorganism that produces a composition that may impart one or more desired property to one or more plant, as herein before described, and summarised in the and eighth (and/or related) and ninth (and/or related) aspects of the invention. Accordingly, unless the context requires otherwise, when describing the embodiments of the invention in this section of the specification, reference to the first aspect and/or second aspect of the invention should be taken to also include reference to the first, second, eighth (and/or related) and/or ninth (and/or related) aspects of the invention, and reference to one or more microorganism should be taken to include reference to one or more composition.

The method begins at step 21 with a requestor 11 identifying a plant (or a class or group of plants). Reasons why particular plants or types of plants maybe identified will be apparent to those skilled in the art. However, by way of example, it may have been found that a plant noted in general for having a high growth rate is growing at lower rates or not at all, there may simply be a desire to improve on existing growth rates or there may be a desire to introduce a plant to a different climate / environment / geographical region. The invention is not limited to conferring improvements to particular plant(s) and may be used to inhibit growth or otherwise adversely affect the plant(s).

At step 22, the requestor 11 sends the plant and / or the identity thereof to a request processor 12. The requestor 11 may provide further relevant information such as why or what properties they are seeking to improve. While only one request processor 12 is shown, it will be appreciated that more than one may be provided in the system 10.
Where a requestor 11 identifies a class or group of plants, more than one plant variety may be evaluated. Alternatively or additionally, selection of a one or more plant variety may be made elsewhere within the system 10 based on the group or class identified, including following evaluation of different varieties including using different microorganisms in accordance with methods of the invention.

Requests may conveniently be received over the internet via a web browser, although the invention is not limited thereto. Use of a web browser may additionally or alternatively be used to enable a requestor 11 to view reports on the progress being made in response to their request. For example, measures of growth may be provided.

At step 23, the request processor 12 receives and processes the request, essentially by initiating the performance of the method for the selection of one or more microorganism according to the first aspect of the invention. Note that the request processor 12 may or may not actively perform the method of the first aspect, or may only perform parts thereof. According to particular embodiments, the request processor 12 may act as an intermediary or agent between the requestor 11 and the parties able to perform the method of the first aspect. Also, different arrangements may be made in response to different requests. For example, for one request, the environment around the request processor 12 may be suitable for evaluating a particular plant but unsuitable for another, requiring the assistance of a third party facility. This could be due to a desire to test in a particular soil type, altitude or climate. Other factors will also be apparent although it is appreciated that "artificial" environments may be used. Furthermore, varying degrees of user interaction may take place at the request processor 12. According to one embodiment, a computer processor selects parameters or conditions for a study based on data input by a requestor 11. As will be appreciated, providing a structured information request may help to effect this, and where necessary, reference may be made to databases including database 14.

At step 24, parameters of the evaluation process are selected. For example, reference may be made to database 14 for microorganisms that may provide the desired improvement in the plant(s). While little data to date has been provided in the art on microorganisms having beneficial associations with particular plant varieties, this will be improved upon through ongoing operation of the methods of the invention and stored in database 14. Other parameters such as plant type(s) and environmental conditions may also be selected.
At step 25, the request (or portions thereof) and evaluation parameters are sent to growing facility 13 which may obtain suitable microorganisms from depository 15. These may or may not have been previously identified. While only one growing facility 13 and one depository 15 are shown, it will be appreciated that the invention is not so limited. Furthermore, any two or more of request processor 12, growing facility 13, database 14 and depository 15 may be co-located and/or under the same control.

At step 26, a selection process is performed, preferably according to the selection method of the first aspect.

At step 27, a response is sent to the request. A response may be sent to the requestor 11 and/or to a third party and preferably includes at least one of at least a subset of the results generated at step 26, identification of plant(s), plant(s), identification of microorganism(s), microorganism(s), or plant(s) provided in association with microorganisms, namely those that have been shown to provide benefits at step 26.

At step 28, database 14 may be updated with results of the selection process of step 26. This step may be performed prior to step 27, including periodically or at other various stages which the selection process is conducted. Preferably, at least details of new beneficial associations between plant(s) and microorganisms are recorded. It will be appreciated that incompatible or less beneficial associations will also preferably be recorded, thereby over time building a knowledge framework of plants and microorganisms.

It will be appreciated that one or more of the steps of Figure 2 may be omitted or repeated. For example, growing facility 13 may generate results at step 26 and in response thereto, one or more of steps 21 to 26 may be repeated.

Thus, the invention provides means and methods to improve plant(s) (or growth or other characteristics thereof). This is achieved by enabling a requestor 11 in a first geographical region (e.g. country) or otherwise defined environment (e.g. by parameters or characteristics affecting growing conditions such as such soil salinity or acidity) to access microbiological biodiversity not present or of limited presence in the first region for the
purposes of plant improvement in the first or another region. The other region may be or in a foreign country but may be otherwise defined by characteristics of that environment that affect a plant rather than being defined by political boundaries. Consequently, the invention may enable a requestor to obtain the beneficial effects of a particular microorganism(s) on a particular plant(s) in a first region, even though such microorganism(s) may not be present or are of limited presence in the first region.

An example implementation of the invention is provided below.

1. A company in say New Zealand (home company), enters into a contractual relationship with a second, say overseas, company (overseas company).

2. The overseas company agrees to send seeds, cuttings or other plant propagules (foreign cultivar) to the home company from plant cultivars adapted to the environment(s) in its own, or other foreign countries, in order to gain access to elements of New Zealand's terrestrial and marine microbial biodiversity that are able to form beneficial plant-microorganism associations with the foreign cultivar.

3. The nature of the benefit may encompass increased plant productivity, for example through any one or more of but not limited to: increased root or foliar mass, or through an increase in efficiency in nutrient utilisation through nitrogen fixation by diazatrophs such as Klebsiella or Rhizobium, or through release of plant nutrients from the soil, such as phosphates released soil through the production of microbial phytases, or through improved resistance to attack from pests and diseases spanning a broad range of nematodes, insects, microbial and virus diseases, or through improvements in the ability of the plant to resist adverse environmental conditions such as drought, salinity, extreme temperatures, toxic soil minerals, or through improvements in plant phenotype for example date of flowering, or changes in physical form e.g. colour frequency of root or foliar branching, or changes in chemical profile including compounds associated with taste, smell or properties which make the plant suitable for a particular purpose.

4. In New Zealand, the home company identifies which indigenous microorganisms can form an association with the foreign plant by exposing the seed to the microorganisms, with or without knowledge of their likely effects on the plant, by the method of germinating the seed and growing the plant in a growing material that ensures contact of the plant during its growth with indigenous microorganisms
via seed coating, direct inoculation into the seed or germinating seedling and/or contamination of the growing medium. The invention is not limited to this arrangement or methodology. For example, it may be apparent that microorganisms present in soil other than in New Zealand may provide benefits and testing may be conducted in such regions in addition to or instead of New Zealand. Also, artificial environments may be created. Referring to the immediately prior example, this may be achieved by obtaining soil and/or microorganisms from such regions and conducting the tests in say New Zealand.

As will be apparent, such embodiments may include provision for artificial control of climatic conditions among other parameters. Thus, the invention is not limited to conducting testing in a region based on its indigenous microorganisms - the microorganisms may be artificially introduced so as to conduct the testing elsewhere than in the microorganisms' natural environment.

5. The period of growth and the physical conditions under which they take place may vary widely according to plant species and specific plant improvement traits, including based on parameters desired or specified by the overseas company. After the relevant period of plant growth the nature of possible plant-microorganisms associations may be determined by microbiological assessment to determine whether microorganisms have formed an endophytic, epiphytic or rhizospheric association with the foreign crop. One or more of the previous steps may be repeated as required until a desired relationship is found.

6. Where such association(s) are demonstrated the microorganisms form a collection of (say New Zealand indigenous) microorganisms able to associate with the (say foreign) crop or plant.

7. In one embodiment of the invention, microbial isolates of the collection may, for example, be coated on to seeds, inoculated into seeds or seedlings, or inoculated into a growing medium that may or may not be sterile.

8. After a suitable period the plants are assessed for improved root and foliar growth or other desired characteristics and/or they may be exposed to environmental stressors designed to identify the plant-microorganism associations most able to provide benefit to the plant in the manner desired by the overseas company.

9. Examples of such stressors or selection criteria are provided in 3 above and elsewhere herein, and where identical pests, diseases or other parameters of the second, overseas environment are not present in the home or test region (i.e., New Zealand).
Zealand in the example), similar microbial diseases, nematode and insect pests or other parameters most similar to those in the overseas environment and that maybe considered acceptable to the overseas company may be selected. As mentioned in 4 above, the invention also includes introducing foreign material or creating otherwise artificial conditions in the home or test region.

10. The steps involving growing one or more plant in the presence of one or more microorganism, selecting one or more plants with desired characteristics, and acquiring the microorganism(s) forming an association with the plant will be repeated one or more time.

11. One or more steps involving identifying one or more plant and/or microorganism marker in accordance with the invention is performed at the appropriate time (such as as described in the first or second aspect described herein before).

12. Elite microorganisms providing commercially-significant benefit to the growth of the foreign cultivar are identified by this process and may be shipped to the overseas company for further testing and selection in the foreign environment.

13. In a further embodiment the overseas company will agree that microorganisms found on, or in, the seed, cuttings or propagules of the foreign cultivar will be added to the collection of the home company to enlarge the collection for use both on that cultivar or on other foreign cultivars received for similar testing from other companies.

In an alternative embodiment, the microbial isolates able to form plant-microorganism associations with the foreign cultivar i.e., the collection, are sent to the second company for testing and selection, such that items 7-12 above are performed by and/or in the grounds of the second company. This may be performed by or under the control of the first company.

As a further alternative, rather than identifying and using predetermined microorganism(s) of a collection, the home company may simply expose the seed to indigenous microorganisms, with or without knowledge of their likely effects on the plant, for example by germinating the seed and growing the plant in a growing material that ensures contact of the plant during its growth with indigenous microorganisms via seed coating, direct inoculation into the seed or germinating seedling and/or contamination of the growing medium or otherwise. As will be apparent, the home company may additionally
or alternatively arrange for similar testing in other regions, where the same or different microorganisms may be present. The period of growth and the physical conditions under which they take place may vary widely according to plant species and specific plant traits desired by the overseas company. After a period of plant growth the nature of possible plant-microorganism associations may be determined in a similar manner to that described above.

EXAMPLES
The invention is now further described by the following non-limiting examples.

Use of process to optimise expression of Quantitative Trait Loci (QTLs) for beneficial plant phenotypes

The expression of plant QTLs can often vary depending on environmental factors, including the composition and activity of local microbial communities. This invention uses the directed process of selection to generate plants with microbially-optimised QTL expression for enhanced phenotype stability and efficacy. Methods of the first 2 examples involve 3 key processes

- **Microbe capture:** plants are exposed to a diverse collection of microbes generating a pool of material enriched for microbes capable of associating with the plant species of interest. Extracts containing the microbes are prepared and used as an inoculum in the next step.
- **Selection:** Iterative rounds of microbial inoculation, plant growth and selection based on some phenotypic trait. Plants/seeds are initially inoculated using extracts from microbe capture round. Further rounds use extracts prepared from the plants grown in the preceding round. Growth conditions may include, but do not necessarily require some form of biotic or abiotic stress
- **Microbe isolation:** Microbes are isolated from plants exhibiting the desired phenotypic trait and are used as leads for commercial product development.

Example 1: Identification of microorganisms that optimise expression of Quantitative trait loci (QTL) associated with specific phenotypes.
**Step 1: Identify loci of interest.** Specific QTL and associated genotypic and/or phenotypic expression markers are identified within the plant of choice. For example, but not limited to, QTL may be identified that code for aluminium tolerance in wheat.

5 **Step 2: Microbe capture:** Untreated seeds from a selected cultivar or cultivars are planted in a wide variety of soils in pots. Soils may include additional amendments comprising pure cultures of microorganisms, mixtures of microorganisms or materials containing microorganisms that derived from other sources and may include GM microbes. Alternatively, untreated seeds are planted in a wide variety of soils that are selective for the QTL of interest (eg elevated Al concentrations for Al tolerance QTL).

10 **Step 3: Preparation of inoculum:** After a suitable period of growth, the plants are removed from or washed out of the soil, and the microorganisms isolated from roots and stems/foliage, either as individual isolates in pure culture, or as mixed populations e.g. as a microbial suspension from an aqueous root crush and/or a stem/foliar crush.

15 **Step 4: First round of selection.** The microorganisms are added to a plant growth medium into which untreated seeds are then planted. Alternatively, the microorganism(s) are mixed into a suitable seed coating material e.g. a gel, and coated onto seeds before being planted into a similar plant growth medium. Alternatively, the seeds are gminated and then exposed to the microorganisms for a short period of say 1 to 24 hours (to maximise the chance that the microbes may form an endophytic or epiphytic association with the germinating plant) and then planted into a similar growth medium. In each of these cases the growing medium may be initially sterile, although this is not essential and further microorganisms applied to the growth medium and/or plant.

The growth medium and or growth conditions are designed to induce the expression of the QTL, for example elevated Al levels for induction of Al tolerance QTL. The growth medium may be a specific soil type or may be a range of different soil types. After an appropriate period of growth, say two weeks (but at any desirable time point between germination and seed-harvesting), the plants are screened for expression of the QTL. Quantitative expression may be determined using methods that quantify genotypic markers or products thereof including qPCR of mRNA, microarray analysis of gene expression, high throughput sequencing of transcripts, proteome or metabolome analyses.
Step 5: Iterative rounds of selection. The best plants are selected as determined by the highest level of QTL expression. Selection criteria may or may not include other factors such as expression of other desirable phenotypic traits such as growth of herbage and/or roots determined by dry weight, or image analysis or similar, and/or disease symptoms.

The root and foliar microorganisms from selected plants are isolated and made ready to be added to the seeds and/or the growing medium as individual or combined suspensions, as in step 3.

The entire process from step 3 to the end of step 5 can then be repeated iteratively, with or without increasing selective pressure for QTL expression.

Step 6: Isolation. A number of iterations to the point at which it is deemed that the expression of QTL is stabilised, and the best phenotype is achieved, the microbes in the best-performing plants are isolated. These microbial strains are then used individually or in a mixture to develop a commercial product that optimises the expression of QTLs and associated phenotypes.

Example 2: Identification of microbially-induced Quantitative trait loci (QTLs) associated with beneficial plant traits.

Step 1: Microbe capture: Untreated seeds from a selected cultivar are planted in a wide variety of soils. Soils may include additional amendments comprising pure cultures of microorganisms, mixtures of microorganisms or materials containing microorganisms that derive from other sources and may include GM microbes. Alternatively, untreated seeds are planted in a wide variety of soils that are selective for the QTLs of interest (e.g. elevated Al concentrations for Al tolerance QTL).

Step 2: Preparation of inoculum: After a suitable period of growth, the plants are removed from or washed out of the soil, and the microorganisms isolated from roots and stems/foliage, either as individual isolates in pure culture, or as mixed populations e.g. as a microbial suspension from an aqueous root crush and/or a stem/foliar crush.

Step 3: Selection. The microorganisms are added to a plant growth medium into which untreated seeds are then planted. Alternatively, the microorganism(s) are mixed into a
suitable seed coating material e.g. a gel, and coated onto seeds before being planted into a similar plant growth medium. Alternatively, the seeds are geminated and then exposed to the microorganisms for a short period of say 1 to 24 hours (to maximise the chance that the microbes may form an endophytic or epiphytic association with the germinating plant) and then planted into a similar growth medium. In each of these cases the growing medium may be initially sterile, although this is not essential and further microorganisms applied to the growth medium and/or plant.

The growth medium and or growth conditions may be designed to induce a response to a specific biotic or abiotic stress and may be a specific soil type or may be a range of different soil types. After an appropriate period of growth, (but at any desirable time point between germination and seed-harvesting), the plants are screened for expression of the desired phenotype (eg increased foliar weight) and the best plants are selected. Selection criteria may include a single factor or a combination of factors. The root and foliar microorganisms from selected plants are isolated and made ready to be added to the seeds and/or the growing medium as individual or combined suspensions, as in step 2.

The entire process from step 2 to the end of step 3 can then be repeated iteratively, with or without increasing selective pressure for a specific phenotype. All rounds include a control comprising plants in normal soil to which a microbial extract has not added.

Step 4: Microbial isolation and assessment of plant gene expression. After a number of iterations to the point at which it is deemed that the best phenotype is achieved, the microbes in the best-performing plants are isolated and the gene expression of the plant is assessed, relative to controls in which no microbial selection has been applied. Patterns of expression may be determined using methods that quantify genotypic markers or products thereof including qPCR of mRNA, microarray analysis of gene expression, high throughput sequencing of transcripts, proteome or metabolome analyses. Conventional and high throughput genotyping methods (e.g. SNP analysis) may also be included to assess the expression patterns in the context of genotypic variation.

Isolation of microbes may be informed through use of methods that enable culture independent identification of the constituents of the plant microbiome. Such methods may include, but are not limited to, high throughput sequencing or microarray analysis of rRNA genes.
Step 5: Elaboration of microbially-induced gene expression patterns that associate with beneficial plant traits. The microbial strains isolated in step 4 may be used individually or in a mixture to further elaborate the effects of microbial interaction on plant gene expression under different growth conditions and at different time points over the life of the plant. Pre-conditioning of microbes prior to application to the plant (e.g. through growth on different media or in the presence of biotic or abiotic stress) can also be assessed for effects on plant gene expression, enabling further optimisation of microbially induced traits.

Once QTLs have been identified that associate with microbially-induced beneficial plant traits, markers can be developed for use in crop breeding using marker-assisted selection (MAS). QTL may also be used as targets for genetic modifications that enable expression of the phenotype in the absence of a microbial inducer.

Example 3: Use of process to select seed-borne endophytes conveying a beneficial crop trait

Forage grasses expressing beneficial traits such as insect-resistance and improved tolerance to both biotic and abiotic stressors via strains of the seed-borne fungus *Neotyphodium* ¾?. have been widely adopted by faixaers in New Zealand and elsewhere. It would be desirable to extend the benefits of traits similar to those expressed by this seed-borne fungus and other similar species in the fungal family, to a broader range seed-borne endophytic microbes thereby providing access to a much wider range of beneficial crop traits.

Step 1. Untreated ryegrass seeds are planted in a wide variety of soils in small pots. Soils may include additional amendments comprising pure cultures of microorganisms, mixtures of microorganisms or materials containing microorganisms that derived from other sources.

Step 2. After a suitable period of growth, the plants are washed out of the soil, surface sterilised with a combination of ethanol and sodium hypochlorite or other methods known to people skilled in the art, and the endophytic microorganisms (endophytes) isolated from internal tissues of roots and stems/foliage and seeds, either as individual isolates in pure culture, or as mixed populations e.g. as a microbial suspension from an aqueous root crush and/or a stem/foliar crush.
Step 3. The endophytic microorganisms are then added to a plant growth medium into which pre-germinated surface-sterilised ryegrass seeds are planted (seeds checked for sterility by germinating on nutrient agar plates). Alternatively, the microorganism(s) are mixed into a suitable seed coating material e.g. a gel, and coated onto surface-sterilised seeds before being planted into a similar plant medium. Alternatively, the surface-sterilised seeds are gminated on nutrient agar plates, checked for sterility and then exposed to the microorganisms for a short period (usually between 1 - 24 hours to maximise the chance that the microbes may form an endophytic or epiphytic association with the germinating plant) and then planted into a similar growth medium. In each of these cases the growing medium may be initially sterile, although this is not essential and further microorganisms may be applied to the growth medium and/or plant.

Step 4. After a period of suitable growth, e.g. 4 - 6 weeks, plants are assessed for expression of the desired phenotype. Phenotypes may include relative resistance to abiotic selection pressures such as relative growth in nutrient deficient conditions e.g. nitrogen or phosphorus, growth in a medium high in salinity or deficient in water. The selection pressures could also be biotic such as exposure to insect pests, plant parasitic nematodes, or plant diseases. Alternatively no selection may be applied and the plants selected merely on a phenotypic attribute such as improved colour, plant form, metabolite expression, or the like.

Step 5. Selected plants are permitted to grow onward to the point of seed set. At this stage a subset of seeds from each plant may be screened for endophyte carnegie using culture dependent or independent methods. The remaining seeds from plants yielding positive results in the screen are germinated and planted without microbial addition in a further round of selection to enrich for endophyte carriage and the ability to transmit the desired phenotype as described in steps 3 - 5.

Alternatively, endophytic microbes may be acquired from a subset of seeds from each plant either as isolates from surface sterilised seeds or as explants, or as a microbial suspension prepared, for example, by crushing the surface sterilised seed in aqueous solution. Isolates and preparations are used as an inoculum for plants arising from surface sterilised seeds as described in step 3.

In a further variation of the method, the selection for seed transmission of the trait may take place in the following generation by surface sterilising a subset of seeds (with or without prior screening) from the selected plants of the prior generation and allowing them to germinate and grow on for the period at which point phenotypic screening is conducted.
as generally described in steps 3 and 4 (i.e. prior to seed set). Plants exhibiting the desired phenotype in this generation (i.e. by seed transmission), are selected and either tissue explants are prepared, and/or microbes isolated from plant tissues, and/or crude microbial suspensions made by crushing the surface foliage or roots in an aqueous solution. One or a combination of these preparations are used as an inoculum for further iterative rounds of growth and selection and seed harvest, as described in steps 3-5. Alternatively, the remaining seeds of plants exhibiting the desired seed-borne trait may be germinated and planted without microbial addition in a further round of selection to enrich for endophyte carriage and the ability to transmit the desired phenotype as described in steps 3-5. Step 6). At the end of successive rounds of this iterative process, as determined by the generation of a desired seed-borne phenotype, the best seed lines are selected for commercial assessment and cultivar development.

In this example, one or more QTL of a plant and/or microorganism which is associated with one or more beneficial trait is identified prior to the process. Alternatively one or more such QTL is identified following acquisition of one or more microorganism or at the time of selection of one or more plant.

The invention has been described herein, with reference to certain preferred embodiments, in order to enable the reader to practice the invention without undue experimentation. However, a person having ordinary skill in the art will readily recognise that many of the components and parameters maybe varied or modified to a certain extent or substituted for known equivalents without departing from the scope of the invention. It should be appreciated that such modifications and equivalents are herein incorporated as if individually set forth. In addition, titles, headings, or the like are provided to enhance the reader's comprehension of this document, and should not be read as limiting the scope of the present invention.

The entire disclosures of all applications, patents and publications, cited above and below, if any, are hereby incorporated by reference. However, the reference to any applications, patents and publications in this specification is not, and should not be taken as, an acknowledgment or any form of suggestion that they constitute valid prior art or form part of the common general knowledge in any country in the world.
Throughout this specification and any claims which follow, unless the context requires otherwise, the words "comprise", "comprising" and the like, are to be construed in an inclusive sense as opposed to an exclusive sense, that is to say, in the sense of "including, but not limited to".
BIBLIOGRAPHY
WHAT WE CLAIM IS:

1. A method for the selection of one or more microorganisms capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:
   a) identifying one or more plant marker associated with one or more beneficial property;
   b) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to a growth medium in the presence of a first set of one or more microorganisms;
   c) selecting one or more plant following step b) on the basis that the one or more plant marker is present;
   d) acquiring a second set of one or more microorganisms associated with said one or more plant selected in step c);
   e) repeating steps a) to d) and/or steps b) to d) one or more times, wherein the second set of one or more microorganisms acquired in step d) is used as the first set of microorganisms in step b) of any successive repeat.

2. A method according to claim 1, wherein one or more selective pressure is applied in step b).

3. A method for the selection of one or more microorganisms capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:
   a) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to a growth medium in the presence of a first set of one or more microorganisms;
   b) selecting one or more plant following step a);
   c) acquiring a second set of one or more microorganisms associated with said one or more plant selected in step b);
   d) repeating steps a) to c) one or more times, wherein the second set of one or more microorganisms acquired in step c) is used as the first set of microorganisms in step a) of any successive repeat; wherein the method also comprises the step of identifying one or more plant marker and/or one or more microorganism marker associated with the beneficial property after step b) and/or after step c).
4. A method according to claim 3, wherein one or more selective pressure is applied in step a).

5. A method according to claim 1 or 2, wherein the growth medium is selective for a specific marker of interest.

6. A method according to claim 1 or 3, wherein one or more plant may be selected on the basis of the presence of one or more marker in combination with one or more other selection criterion.

7. A method for the production of a composition to support plant growth, quality and/or health, or a composition to suppress or inhibit the growth, quality and/or health of a plant the method comprising the steps of a method according to claim 1 or 3 and the additional step of combining the one or more microorganisms selected by the method with one or more additional ingredients.

8. A method for the selection of a composition which is capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:
   a) culturing one or more microorganisms selected by a method of claim 1 and/or claim 3 in one or more media to provide one or more culture;
   b) separating the one or more microorganisms from the one or more media in the one or more culture after a period of time to provide one or more composition substantially free of microorganisms;
   c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition of step b);
   d) selecting one or more composition from step c) if it is observed to impart one or more beneficial property to the one or more plants.

9. A method for the selection of a composition which is capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:
   a) culturing one or more microorganisms selected by a method of claim 1 and/or claim 3 in one or more media to provide one or more culture;
b) inactivating the one or more culture of step a) to provide one or more composition containing one or more inactivated microorganisms;

c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition of step b);

d) selecting one or more composition from step c) if it is observed to impart one or more beneficial property to the one or more plants.

10. A method for the selection of one or more microorganisms which are capable of producing a composition which is capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:

a) culturing one or more microorganism selected by a method of claim 1 and/or claim 3 in one or more media to provide one or more culture;

b) separating the one or more microorganism from the one or more media in the one or more culture from step a) after a period of time to provide one or more composition substantially free of microorganisms;

c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition from step b);

d) selecting the one or more microorganisms associated with one or more composition observed to impart one or more beneficial property to the one or more plants.

11. A method for the selection of one or more microorganisms which are capable of producing a composition which is capable of imparting one or more beneficial property to a plant, the method comprising at least the steps of:

a) culturing one or more microorganisms selected by a method of claim 1 and/or claim 3 in one or more media to provide one or more culture;

b) inactivating the one or more culture of step a) to provide one or more composition containing one or more inactivated microorganisms;

c) subjecting one or more plant (including for example seeds, seedlings, cuttings, and/or propagules thereof) to the one or more composition of step b);

d) selecting the one or more microorganisms associated with one or more composition observed to impart one or more beneficial property to the one or more plants.
12. A method for assisting in the improvement of one or more plants, comprising
arranging for the evaluation of said plant(s) in the presence of one or more microorganisms
and/or compositions, wherein the method comprises at least the steps of a method of claim
1 or claim 3.

13. A system for implementing the method as claimed in claim 7.

14. One or more microorganism acquired, selected or isolated by a method according
to any one of claims 1, 3, 10 or 11.

15. A composition comprising one or more microorganism according to claim 14.

16. A composition made by a method of claim 7 or selected by a method of claim 8 or
15 9.

17. The use of one or more microorganism as claimed in claim 14 for imparting one or
more beneficial property to one or more plant.

18. The use of a method according to claim 1 or 3 in a plant breeding programme.

19. A plant breeding programme comprising conducting a method of claim 1 or 3.

20. A method for identifying one or more plant marker that is associated with one or
more beneficial trait the method comprising at least the step of conducting a method claim
1 or claim 3 and identifying the one or more plant marker.

21. A method for identifying one or more microorganism marker that is associated with
one or more beneficial plant trait the method comprising at least the step of conducting a
method of claim 1 or claim 3 and identifying the one or more microorganism marker.

22. A method for identifying a combination of one or more plant marker and one or
more microorganism marker comprising at least the step of conducting a method of claim
1 or claim 3 and identifying the combination of one or more plant marker and one or more microorganism marker.

23. The use of a microorganism selected by a method of claim 10 or claim 11 as the first set of microorganisms in a method of claim 1 or 3.
FIGURE 1

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FIGURE 2

21 Identify plant

22 Send Request

23 Process Request

24 Parameter Selection

25 Transfer to Grower

26 Selection Process

27 Response to Request
INTERNATIONAL SEARCH REPORT

International application No.
PCT/Z2014/000044

A. CLASSIFICATION OF SUBJECT MATTER

C12Q 1/24 (2006.01) A01H 3/00 (2006.01) A01H 17/00 (2006.01)

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

WPI, EPDOC, MEDLINE, AGRICOLA, HCAPPLUS, BIOSIS & keywords: beneficial microorganism, endophyte, select, plant, marker and like terms

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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× Further documents are listed in the continuation of Box C

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Date of the actual completion of the international search 21 May 2014
Date of mailing of the international search report 21 May 2014

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