**SCREENING METHOD FOR SELECTING PLANTS THAT SHOW A REDUCED WOUND-INDUCED SURFACE DISCOLOURATION AND PLANT AND PLANT PARTS THUS OBTAINED**

SCREENING-VERFAHREN ZUR AUSWAHL VON PFLANZEN, DIE EINE VERMINDERTE VERWUNDUNGSTIMULIERTE OBERFLÄCHENVERFÄRBUNG AUFWEISEN UND SO ERHALTENE PFLANZEN UND PFLANZENSTEILE

PROCÉDÉ DE CRIBLAGE PERMETTANT DE SÉLECTIONNER DES PLANTES PRÉSENTANT UNE DÉCOLORATION DE SURFACE INDUITE PAR BLESSURE RÉDUITE ET PLANTE ET PARTIES VÉGÉTALES AINSI OBTENUES

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- DEMEKE T ET AL: "Effect of germination, seed abrasion and seed size on polyphenol oxidase assay activity in wheat" PLANT BREEDING, vol. 120, no. 5, October 2001 (2001-10), pages 369-373, XP002428728 ISSN: 0179-9541

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Description

Field of the invention

[0001] The present invention relates to plants and parts derived therefrom showing an absent or reduced wound-induced surface discolouration and to parts and progeny thereof.

Background of the invention

[0002] Due to increasing demand, processing of fresh produce, in particular lettuce, has expanded significantly over recent years. The harvesting and processing of lettuce involves expensive cutting of the leaves, which induces a strong wound response. This wound response leads to a rapid deterioration of the processed product. This deterioration is manifested by discolouration due to enzymatic browning or pinking at and around the wound surface, respiration and desiccation due to transpiration. Especially the enzymatic browning or pinking is considered of significant importance, determining directly or indirectly the overall quality of the fresh cut, packaged lettuce.

[0003] Moreover, as a consequence of the deterioration, micro-organisms can significantly increase in number, which may compromise food safety. The highly perishable nature of processed lettuce leads to a strong off-colour, off-odour and off-texture perception by the consumer which is hampering a faster than current growth of the so called convenience market.

[0004] In order to inhibit the deterioration process, many chemical or physical post-harvest treatments have been developed which can be applied to decelerate the deterioration of the processed lettuce.

[0005] Amongst these are the packaging of fresh cut lettuce under a modified atmosphere, application of edible coatings, heat shock treatment and addition of chemicals, which inhibit the enzymatic browning. When fresh cut lettuce is packaged under an atmosphere of reduced oxygen at low temperatures, the enzymatic browning can substantially be reduced. However such modified, low oxygen environment leads to anaerobic respiration, which creates an off-flavour and off-odour of the produce which is perceived as very unattractive.

[0006] Edible coatings are thin layers of materials, which act as physical insulation barrier and which effectively protect the produce from different forms of deterioration such as evaporation and browning. These coatings can for example be made of resins, polysaccharides or protein.

[0007] It has further been demonstrated that browning of fresh cut lettuce can be prevented by applying a brief heat shock of 90 seconds at 45°C, immediately after processing. Possibly, the heat shock diverts protein biosynthesis from the enzymes involved in discoloration towards heat shock proteins thereby reducing the enzymatic browning capacity. Alternatively, the effect of heat shock treatment on browning may be explained by thermosensitivity of enzymes involved in the discoloration pathway.

[0008] Chemicals, which can be applied can for example be reducing agents like vitamin C, chelating agents like EDTA, complexing agents like cyclodextrin and enzymatic inhibitors like L-cysteine. Application of chemicals in fresh food obviously involves food safety issues and requires regulatory approval. Combinations of the post-harvest technologies described above can be thought of and ultimately the applied procedure is a trade-off between technological efficacy, cost and food safety.

[0009] Irrespective of the technology applied, improvement of post-harvest quality of processed lettuce will come at a cost and therefore a clear need in the art exists to provide alternatives, which eliminate or reduce the need to apply physical or chemical post-harvest technologies.

Summary of the invention

[0010] It is the object of the present invention to provide plants, that show a reduced wound-induced discolouration response and to provide plants and progeny derived therefrom that are resistant to post-harvest processing disorders such as enzymatic browning or pinking. It is a further object of the invention to provide plants that show a significantly reduced pinking or browning discoloration upon wounding. Discolouration upon wounding can also be visible in parts of the plants, such as stems, seeds, fruits, leaves, flowers, tubers, shoots.

[0011] The invention thus provides a lettuce plant having a reduced or absent wound-induced surface discoloration obtainable by crossing a plant with the NCIMB accession numbers 41454 or 41441 with another plant of the same species and testing plants resulting from the cross by subjecting the plants or plant parts thereof to a screening method which comprises:

a) creating a wound surface on the plants or plant parts to be screened and on the control plants or plant parts;
b) incubating the wound surfaces to allow for discoloration to occur therein or thereon;
c) observing the wound surface discoloration in or on the plants or plant parts;
d) comparing the observed wound surface discoloration in or on the plants or plant parts to be screened with the discoloration that is observed on or in the control plant or plant part to identify plants or plant parts that show no discoloration or a discoloration that is reduced as compared to the control plant or plant part.

[0012] The screening method as described herein is intended for identifying plants that have a reduced wound-induced surface discolouration in one or more of their parts or tissues. For the screening, it is therefore very practical to use the part or tissue that is prone to discolouration. In lettuce, this may be the leaf or a part thereof, such as a punch.

[0013] The method is in particular useful for selecting plants belonging to the family Asteraceae, in particular plants of the genus Lactuca and more in particular to the species Lactuca sativa that show an absence or reduction of wound-induced surface discolouration.

[0014] The method is suitably performed with plant parts having a wound surface. Very useful test samples are discs that are punched from a leaf, the so-called leaf discs. Alternatively, the midrib tissue of veined leafy vegetables can be used. Suitably, discs are cut from such ribs.

[0015] Incubation takes suitably place in an aqueous environment. The method can be very well practised with leaf discs that are incubated on or between wetted filter paper. The discolouration response is then very well visible around the edges of the wound on the paper. In the case of plants of the genera Lactuca the discolouration is the pinking response.

[0016] Alternatively, the aqueous environment comprises water or a solution. In a specific embodiment that will be further illustrated below the solution contains L-3,4-dihydroxyphenylalanine. This compound is converted to the production of the black pigment melanin by the enzyme polyphenol oxidase. Alternative compounds which can be used in this respect include but are not limited to chlorogenic acid, isochlorogenic acid, L-tyrosine and catechol.

[0017] The invention relates to a plant showing a reduced wound-induced surface discoloration, which plant is obtainable by subjecting a population of plants to the screening method of the invention and selecting plants from the population that show no surface discolouration in the screening or show a surface discolouration in the screening that is reduced as compared to a control plant.

[0018] The plant is a leafy vegetable plant, more in particular a plant, which belongs to the genus Lactuca and in particular to the species Lactuca sativa.

[0019] Preferably, the plant of the invention is identified in the screen and subsequently selected as having a reduced or absent wound-induced surface discolouration but is subsequently tested to determine whether it has a normal habit. More in particular, the plant should preferably not show negative pleiotropic effects.

[0020] According to the invention plants were identified and selected that show no or a significantly reduced wound-induced surface discolouration. Progeny of seeds of these plants were deposited with the NCIMB and have been given the accession number as listed in Table 1. Details about seed descendence of the deposits are given in Example 4 and in Example 6. These deposits are made because they have the single specific characteristic of no or significantly reduced wound-induced discolouration. They were not tested for

[0021] DUS-criteria for variety registration, i.e. distinguishability, uniformity, stability on all registration characteristics, and are not expected to meet these criteria in any way.

Table 1

<table>
<thead>
<tr>
<th>plant no.</th>
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<th>NCIMB accession number</th>
<th>deposit date</th>
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<td>06D.210202</td>
<td>06D.863B2</td>
<td>41454</td>
<td>3 Jan. 2007</td>
</tr>
<tr>
<td>05D.202539</td>
<td>07G.9979</td>
<td>41441</td>
<td>10 Oct. 2006</td>
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[0022] The invention further relates to plants having a reduced or absent wound-induced surface discolouration and that are obtainable by crossing a plant of the invention with another plant of the same species. The feature "reduced or absent wound-induced surface discolouration" can thus be brought into other plants that originally do not have the feature. Whether the plants resulting from such a cross are indeed plants of the invention can be tested by subjecting these plants to the screening method of the invention. Preferably, the plants that are selected as plants of the invention are then also tested for having a normal habit.

[0023] The invention further relates to progeny of a parent plant of the invention that retains the absence or reduction of wound-induced leaf discoloration as found in the parent plant. Such progeny may be many generations removed from the parent. As long as the feature "reduced or absent wound-induced surface discoloration" is retained, the plant is a plant of the invention.

[0024] The invention further relates to parts of the plants of the invention. The plants parts like lettuce or endive heads or leaves are usually the parts that have a cut surface that may be subject to discoloration.

[0025] Plant parts of the invention can be used in tissue culture to regenerate plants that retain the absence or reduction of wound-induced leaf discoloration as found in the plant from which the tissue for the tissue culture is derived. Such
regenerated plants are also part of this invention.

[0026] The invention further relates to seed of a plant of the invention. From the seed plants can be grown that also have the feature "reduced or absent wound-induced surface discoloration". Whether or not the seeds and thus the plants grown therefrom have retained that feature can be tested in the screening method of the invention. The invention also relates to further generation seeds that retain the absence or reduction of wound-induced leaf discoloration as found in the original seeds.

[0027] The invention is commercially very interesting for the processed vegetable market. As explained above, discoloration of produce, in particular fresh fruits and vegetables, is considered undesirable since the discoloured product is rejected by the consumer. The feature "reduced or absent wound-induced surface discoloration" of the invention is thus suitable found in processed vegetable products, like cut lettuce, endive or witloof or combinations thereof. When screened using the method of the invention processed vegetables of the invention show no or a limited wound-induced leaf discoloration. All processed vegetables, in particular processed lettuce, endive and witloof that meet the screen are part of the invention since they are demonstrated to have a modification leading to a reduced PAL and/or PPO dependent metabolic flux.

**Detailed description of the invention**

[0028] When lettuce is harvested and processed by cutting, many leaf wound surfaces are generated which leads to a significant response of the plant or plant parts, manifested by a brown or pink discoloration at or adjacent to the wound surface. Pinking can also be observed at sites distant from the wound surface at the midrib of the leaf as well as the butt. Sometimes pinking can also be observed at stages just prior to harvest which is considered to be due to abiotic stress or over-maturity of the crop.

[0029] The different forms of discoloration are effected by enzymatic activity, which is strongly enhanced as a consequence of wounding and which generates several forms of polyphenols and reaction products derived therefrom.

[0030] An important enzymatic activity involved in the browning reaction is PPO. PPO activity in relation to enzymatic browning is not restricted to lettuce but has been described for many other plant species to be involved in post-harvest deterioration like in apple, banana and potato. In fact, PPO is widely recognised to be one of the most important enzymes involved in post-harvest deterioration of many processed fresh fruits and vegetables.

[0031] For this reason PPO has been the target of many technologies, which aim at the reduction or prevention of its activity in order to increase post-harvest quality of food products. PPO catalyses a reaction in which polyphenols residing in the plant tissue are oxidised to give rise to the formation of o-quinones. Subsequently, enzymatic and non-enzymatic reactions lead to the formation of brown or black pigments.

[0032] In many plant species PPO is encoded by a small gene family of which the individual members may have different temporal and spatial expression patterns indicative of functional divergence. It has for example been shown that lettuce contains different PPO isoforms in the photosynthetic and vascular tissue of the leaf.

[0033] The natural substrate of PPO can differ between the different species. In lettuce, caffeic acid derivatives like chlorogenic and isochlorogenic acid mainly act as PPO substrate.

[0034] The level of PPO enzyme is not specifically induced upon wounding of plant tissues but it resides inactively in the chloroplast. Upon wounding PPO is activated which is manifested due to the fact that the phenolic substrate residing in the vacuoles is brought into contact with PPO due to tissue disruption.

[0035] In lettuce, the production of polyphenols, which are the substrate of PPO, is induced upon wounding. Therefore, the browning potential of lettuce tissue seems not to be limited by the amount of PPO in the leaf tissue but rather by the rate of polyphenol biosynthesis upon wounding.

[0036] In this respect the situation may differ between crops. For example, in apple the amount of polyphenols is sufficient to generate a browning response of the fruits within one hour after wounding whereas in lettuce the browning reaction may take a few days due to the fact that in lettuce the polyphenol pool largely needs to be synthesised de novo upon wounding.


[0038] In lettuce, wounding of leaves leads to a strong induction of PAL gene expression and PAL activity. The formation of polyphenols is correlated with this enzymatic activity, which suggests that PAL activity induced by wounding of lettuce is an important factor responsible for browning (Campos, R. et al. (2004) Physiologia Plantarum 121, 429-438 and references therein). However, it is currently unclear which other factors determine the final outcome of the wound-induced discoloration reaction. For example, the activity of peroxidases (POD) has been suggested to be important as well in establishing the final level of discoloration (Fukumoto, L.R. et al. (2002) J. Agric. Food Chem. 540, 4503-4511; Martin-Diana A. et al (2005) Biosci. Biotechnol. Biochem. 69, 1677-1685).
As the enzyme activity depends on the availability of internal hydrogen peroxide, the contribution of POD to discoloration may be limited.

It is further evident that wounding is somehow perceived by the plant and subsequently a signal is generated through a cascade, which is currently poorly defined for lettuce. It seems obvious that these activities will primarily be targeted towards wound healing and defence against pathogens. Therefore, it is likely that many genetic factors are involved in mounting the discoloration response of wounded lettuce tissue and each of these are potential targets for genetic modification to reduce or eliminate the wound-induced discoloration.

Most of these genetic factors are currently unknown and for those known to be involved it is unclear to what extent these factors play a specific role in the discoloration reaction or perhaps have a more general function in relation to the wound physiology of the plant.

For example, although wound-induced PAL activity is considered to be determining the browning level of lettuce, products of the phenyl propanoid pathway are known to be involved in inter alia cell wall biosynthesis or defence response as well. Therefore reducing the wound-induced PAL activity in order to reduce browning potential may compromise other functions besides wound-induced browning which may be less desirable in relation to other aspects of lettuce cultivation.

Likewise, PPO activity has been implied to be involved in defence response and therefore reducing the browning potential by reducing PPO levels may increase the susceptibility to pathogens (Thipyapong, P. et al (2004) Planta 220, 105-117). Therefore, it was reasoned by the inventors that a more unbiased approach may be more successful in this respect. Such approach comprises the following steps:

1. Generation of a variant population of plants, in particular a mutant population. Such mutant population can be generated by treatment of seeds or plant tissues with mutagenic agents like ethyl methane sulfonate (EMS) or X-rays.
2. The set-up of an efficient phenotypic screen in which selection is based on a wound response-induced discoloration of the plant, in particular a leafy vegetable, more in particular lettuce, endive or witloof, which is channelled through PAL and/or PPO.
3. Characterisation of the mutants modified in their wound-response with respect to post-harvest discoloration potential and absence of pleiotropic effects of the modification, which compromise growing and processing of the plant, in particular a leafy vegetable, more in particular lettuce, endive or witloof, according to common practice.

Disclosed herein is an unbiased screening method for identifying, selecting and obtaining a plant showing a reduced wound-induced discoloration and post-harvest processing disorders such as enzymatic browning or pinking.

Also disclosed is a more biased method that uses a substrate that is converted into a pigment by PPO. In this assay, the screen is specifically to PPO mutants. A suitable substrate is L-DOPA which is converted into the black pigment melanin.

The method is illustrated herein referring to leafy vegetable, like lettuce, but can also be practised in the same way with other plants as indicated above.

The invention relates to plants or plant parts with altered genotypes, which plants or plant parts show a reduced susceptibility towards physiological post-harvest processing disorders, such as enzymatic browning or pinking. The invention relates to plants or plant parts, which have in their genome genetic information which is responsible for the reduced susceptibility towards post-harvest processing disorders, such as enzymatic browning or pinking, and is found in the genome of a lettuce plant.

Progeny of the plants as claimed are also part of this invention. "Progeny" as used herein is intended to encompass all plants having the same or a similar reduced susceptibility towards post-harvest processing disorders, in particular enzymatic browning or pinking, as the original plants described herein and being derived therefrom in any way, such as by sexual reproduction, like self-fertilisation or cross-fertilisation with another plant of the same genus, or vegetative reproduction such as cutting, tissue culture, haploid culture, protoplast culture, protoplast fusion, or other techniques. Such progeny is not only the first generation of plants derived by one or more of these techniques, but also every further generation of plants derived by one or more of these techniques, provided that the derived plants have the reduced susceptibility.

In order to carry out the phenotypic screen of the invention, a wound surface must be generated as the enzymatic discoloration reaction is induced upon wounding. Wounding is the irreversible disturbance of the natural plant, tissue and/or cell structure by methods like cutting, punching, slicing, abrasion, squashing, breaking, peeling, crushing, pressing, slashing, grinding, fluid injection, osmotic shock, detaching, mowing, shredding, rubbing and tearing.

Subsequently, a phenotypic characteristic must become manifest which is diagnostic for the pathway leading to tissue discoloration and which can be used very efficiently in a screen of the mutant population.

It was surprisingly found that such phenotypic characteristics can be obtained by taking leaf parts of lettuce plants and incubating them under very specific conditions which favour different forms of wound surface discoloration to occur. Subsequently, such assays can be applied to large numbers of mutant plants in order to select those plants which show a reduction of the wound-induced discoloration response.
One embodiment of the method used for selecting plants of this invention is based on the surprising finding that when discs from leaves, in particular lettuce leaves, are taken and incubated between wetted filter papers at 5°C, after approximately 4 days the formation of a pink dye at the edges of the leaf discs becomes apparent. Suitable filter paper is filter paper type 1450 CV, Ref.no. 10 313 281 from Schleier & Schuell, Microscience GmbH, Dassel, Germany. Upon further incubation, the signal intensifies and after approximately one week the maximum intensity has been reached. The formation of the pink dye occurs specifically at wound surfaces.

The discolouration can be measured by scoring on a visual scale from 0, which means no browning or pinking, to 10, which means browning and pinking like a standard lettuce variety (L. sativa). In the present example the L. sativa variety ‘Troubadour’ is used as a standard for 10. If desired, pictures can be used for comparison to score the intermediate classes between 0 and 10. In addition, digital pictures can be made of the filter paper with the pink dye, followed by counting per leaf disc position the number of pixels with an intense pink colour. Using one of these measurements, simple statistical analyses like a t-test, well-known by persons skilled in the art, can be performed to establish whether a plant or group of plants is significantly less pinking than the standard, like cv. ‘Troubadour’. The applied significance level of a one-sided test is 0.001.

For mutants, the statistical comparison can be made between the pinking scores of the original variety, which is the best available standard, and the pinking scores of the individual mutants and/or their offspring.

Further, it was shown that the wound-induced discolouration response can be obtained using many different types of tissue of leaves of different developmental stages. For example, midrib tissue can also be induced to give this response upon wounding. When applied to different types of lettuces, such as butterhead, iceberg, cos, batavia or oakleaf, no individual accessions were found that showed significantly less pinking than the rest of the investigated population.

It was further demonstrated that a specific inhibitor of PPO, L-cysteine, when applied during the reaction, strongly suppressed the formation of the pink dye. In addition, it was found that the formation of the pink dye was inhibited by cinnamaldehyde, which is an inhibitor of PAL activity and browning of fresh cut lettuce (Fujita, N. et al (2006) Biosci. Biotechnol. Biochem. 70, 672-676). These findings show that the pink discolouration response of lettuce is PAL and PPO dependent.

Enzymatic browning of fresh cut lettuce is known to be very effectively prevented by the application of a brief heat shock. The observed effect can be explained by assuming re-routing of protein biosynthesis from the phenyl propanoid pathway towards heat shock proteins thereby reducing the metabolic flux towards the formation of polyphenols.

Alternatively, the effect may be explained by assuming that the enzymes involved in polyphenol oxidation, such as PPO and POD, are inactivated by the heat shock treatment. When the heat shock is applied to lettuce, which is subsequently assayed for the pinking response, it was shown that this response, like the enzymatic browning, was effectively inhibited. This demonstrates that the pinking response of lettuce, which is part of this invention is physiologically very similar to the well known enzymatic browning response.

This finding was further substantiated by applying L-cysteine as a reducing agent. L-cysteine, besides being an inhibitor of PPO, is also known to react with coloured o-quinones and convert them back into colourless diphenols in a chemical reduction reaction. When the pink dye formed by lettuce leaf discs is treated with L-cysteine, it was demonstrated that the pink compound was converted into a colourless compound. It seems therefore likely that the pink dye is an o-quinone formed by PPO.

This was corroborated by the finding that reducing agents like ascorbic acid or glutathion also convert the pink dye into a colourless compound.

In addition, when plants, which are taken from the field, which show pinking, are treated with L-cysteine, the pink discolouration is also eliminated. This demonstrates that the leaf disc pinking response is representing the natural occurring pinking phenomenon, which can sometimes be seen on plants growing under field conditions.

The invention is illustrated by the following experiment. Parts of a lettuce leaf of a head are produced by cutting and incubated at 16°C in air. As a response the wound surface turns brown after approximately 4 days. Especially at the wound surface of the main vein the browning can clearly be observed. Furthermore, the browning reaction can also be observed at the whole plant level upon damaging leaves by cutting or abrasion.

All of these browning reactions can completely be inhibited by L-cysteine, an inhibitor of PPO, which demonstrates that these phenotypes are manifested through PPO activity and therefore can be considered diagnostic for post-harvest browning as observed during processing and packaging of lettuce.

These wound-induced browning reactions can be generated in an efficient manner which can be exploited in a phenotypic screening procedure to identify mutant plants which are reduced in wound-induced browning potential.

A further method disclosed herein is based on the discolouration at wound surfaces of lettuce tissues induced by applying substrates which can be converted by the phenol oxidising enzymes into coloured compounds.

For example, when lettuce leaf discs are incubated with the PPO substrate L-3,4-dihydroxyphenylalanine (L-DOPA), a dark brown to black discolouration is observed at the wound surface which is the manifestation of the formation of melanine through PPO. When L-cysteine was applied simultaneously, the black discolouration was com-
Although L-DOPA is not considered to be a natural substrate for lettuce PPO it can be useful in assays aimed at the identification of mutants, which show reduced wound-induced discoloration. In a similar manner as described for L-DOPA other substrates can be applied in order to raise a discoloration response. These include but are not limited to chlorogenic acid, isochlorogenic acid, L-tyrosine, and catechol. Taken together, the formation of the different dyes at wound surfaces generated in plants monitors modifications in a pathway starting by the induction of a wound signal, channelled through PAL and PPO and leading to discoloration. As described, these wound-induced discoloration reactions can readily be assessed by visual inspection which allows a very efficient mutant screening procedure. The rationale underlying the method described by this invention is illustrated in Figure 1. According to this invention it was thus found that the wound-induced discoloration pathway of leaf discs in vitro largely overlaps with the wound-induced discoloration of lettuce processed at industrial scale and can therefore be considered diagnostic for this process. This is corroborated by the notion that inhibitors of PAL or PPO inhibit the enzymatic browning of processed and packaged lettuce under practical, industrial conditions. Importantly, as the procedure comprises the inducing step i.e. wounding and one of the final metabolic conversions mediated by PPO, the procedure allows to capture all genetic factors directly or indirectly involved in this physiological process. Moreover, as this response can be generated using a whole range of leaf tissues of leaves of different developmental stages, mutant screens can be targeted towards these different stages or tissues when considered relevant. Mutant plants, which have been identified as being modified with respect to the physiological process leading from wounding to a PAL- and PPO-dependent discoloration based on one or more of the phenotypic assays described above can be further characterised. Such characterisation can be done at different levels e.g. at the molecular, biochemical, physiological and phenotypic level. It is obvious to those skilled in the art that variable levels of discoloration may be observed which may reflect either the presence of different mutant loci or different allelic forms of identical loci affecting the discoloration trait in the original population. In case recessive mutations these two possibilities can easily be distinguished by carrying out allelism tests, which comprise the crossing of the two mutant plants and determining the phenotype of the hybrid. In case of allelism of the mutations, the reduced discoloration trait will be apparent in the F1 whereas in case the phenotype in the mutants is determined by different recessive loci this will not be the case. As random mutagenesis was applied to generate the starting population, mutations in the genetic background may also contribute to the variation of the phenotype under the experimental conditions. In order to discriminate between single mutations of different strengths and a combined effect of mutations in the genetic background, backcrosses should be performed to create uniform genetic backgrounds for the different reduced discoloration events. Such procedure is further relevant in order to determine whether mutations at specific loci involved in wound-induced discoloration display pleiotropic effects. The M2 plants thus selected on the basis of a reduced discoloration response are used to grow M3 seeds. Subsequently, the inbred lines descending from the reduced discoloration events are re-evaluated for their reduced response wounding. In addition, the reduced browning or pinking can be assessed in different genetic backgrounds and under different conditions of crop cultivation and processing. Biochemical studies can be performed to address questions related to the pathways affected by the genetic modification. Molecular studies can be performed to determine if candidate genes putatively involved in the enzymatic browning or pinking response like genes encoding PAL, PPO or peroxidases have been modified. Genetic analysis will subsequently be carried out to demonstrate if the modification found in a candidate gene is causative with respect to the altered phenotype. Although induced mutagenesis is the preferred method to be used in this invention, it is known to the person skilled in the art that technology exists which allows to modify gene targets residing in the genome of a plant in a specific manner. For example, chimeric oligonucleotides have been demonstrated to be effective mutagens with a specific mode of action. Another approach is to modify gene targets through homologous recombination or gene targeting. Using such approach, a fragment of a gene is exchanged by an introduced DNA fragment containing a desired modification. Transgenic approaches are also feasible in which modified target genes are introduced which compete with the endogenous product. This may lead to dominant negative effects. Moreover specific downregulation of the expression of genes is feasible through RNA interference. In case mutagenic oligonucleotides, gene targeting or transgenic approaches are used to modify a genetic factor involved in wound-induced discoloration response, obviously, the primary structure of the relevant genes should be known. This invention relates to lettuce mutants and progeny derived therefrom which were identified on the basis of the wound-induced pink discoloration of lettuce leaf discs. Applying this pinking assay to plants of an M2 population
which contain random, ems-induced mutations resulted in the identification of a number of mutants, which showed a significant reduction of the pinking response as compared to the control plants, which do not contain the ems-induced mutations.

[0082] Most of such mutants showed a dwarfed and often chlorotic phenotype. It was however surprisingly found that some mutants with a reduced pinking response showed a normal growth habitus, i.e. a size, shape, growth and colour very much similar to the control plants.

[0083] When progeny plants of this particular mutant grown from seeds obtained through self-fertilisation are assayed for pinking, a similar reduction as found for the originally identified mutant is observed. This demonstrates that a reduced pink discoloration response can be heritable and caused by a modification of the genome.

[0084] A further surprising finding was the fact that when the progeny plants are grown to maturity and tested for enzymatic browning of wounded midrib tissue, this response is also strongly inhibited.

[0085] This shows that the leaf disc pinking assay is causally related to enzymatic browning in lettuce and that the pinking assay can be used to predict the level of enzymatic browning of a mature lettuce plant.

[0086] Therefore, the leaf disc pinking assay can be used as a selection tool to identify lettuce plants with a reduced enzymatic browning potential. Such tool can be used to identify lettuce plants with reduced enzymatic browning potential from any kind of plant population irrespective of the cause of the genetic variation, which resides in such population. For example, in addition to ems populations, one can use natural accessions or breeding populations.

[0087] One or more of the screening methods disclosed herein can be applied to any leafy vegetable species for which post-harvest processing quality needs improvement.

[0088] The relates phenotypic feature of the present invention can be detected in a plant by performing one of the screening methods that are disclosed herein. Plants of the invention are those plants which, compared to a control plant, show the absence of or a reduction in wound-induced surface discoloration. The presence of the feature is determined by means of one or more of three discoloration tests, namely the occurrence of pinking or browning or the ability to convert the substrate L-DOPA to melanin. A plant of the invention is a plant, which in at least one of these test shows a discolouration that is at least reduced as compared to a control.

[0089] The "control" as used herein is any plant of which it is known that it shows one or more of the discoloration reactions pinking, browning and conversion of L-DOPA to melanin, which reactions can be inhibited by L-cystein or cinnamaldehyde. "Suitably a plant is used of which a leaf disc when incubated between wetted filter paper at 5°C for 7 days shows pink discoloration around the edges of the disc.

[0090] The present invention will be further illustrated in the Examples that follow and that are not intended to limit the invention in any way. In the Examples reference is made to the following figures.

[0091] Figure 1: Schematic outline of the rationale behind the design of the mutant screening procedure of lettuce populations for reduced post-harvest enzymatic discoloration. The input signal of the screen is wounding of leaf tissue, which is sensed by the plant and which generates a divergent signalling response leading to a number of physiological processes including senescence, respiration and tissue discoloration. This input signal can be combined with the application of phenolic compounds as PPO substrates.

[0092] The output signal of the screen is a brown or pink discoloration, depending on the conditions applied, of the wound surface diagnostic for post-harvest browning and pinking. This is inferred from the fact that the output signal is completely inhibited by cinnamaldehyde and L-cysteine, which are specific inhibitors of PAL and PPO, respectively.
presence of a pink dye. The lower right panel shows a disc taken from the leaf showing pinking symptoms after treatment with 1 mM L-cysteine for 30 minutes at room temperature. The lower left panel shows a similar leaf disc after treatment with water for 30 minutes at room temperature.

**Figure 9:** Panel A: Phenotypic analysis of individual lettuce M2 plants (grouped in pools) for leaf disc discoulouration according to the method described by this invention. A total of 138 samples out of 12000 is shown in this panel of which the one indicated by an arrow showed a strongly reduced pinking discoulouration. Panel B: Re-testing of the selected individual indicated in panel A confirmed the near absence of the formation of the pink discoulouration (sample in the middle position) as compared to control samples which show a clear discoulouration response.

**Figure 10:** Phenotypes of M2 lettuce plants. The plants labelled 1, 2, 4, 5, 7, 10 and 12 show reduced pink leaf disc discoulouration using the assay according to this invention. Plants 3, 6, 8, 9, and 11 are plants, which showed a level of pink leaf disc discoulouration comparable to the wild type control. Plant 1 is the only example of a mutant which shows a strong reduction in pink discoulouration and a normal growth habitus. Plants 2, 4, 5, 7, 10 and 12 show reduced pink discoulouration and a dwarfed, bleached phenotype.

**Figure 11:** Progeny testing of a mutant of lettuce showing a reduced discoulouration. On the left, 25 control samples are shown which show a normal wound-induced discoulouration response. On the right, a group of samples is shown which is taken from a series of 35 progeny plants derived from a single mutant, which is severely reduced in its wound-induced discoulouration response.

**Figure 12:** Representative image of the output phenotype of the screen based on brown discoulouration of leaf midrib parts taken from mature lettuce plants. The picture shows lettuce outer leaf midrib tissue discs after incubation for 3 days at 16°C. The typical brown discoulouration can clearly be observed at the wound surface. Each dish contains 3 discs taken at different positions of the midrib (green, light green and white). The number above the dish indicates the mM concentration of L-cysteine which was added to the filter.

**Figure 13:** Conversion of L-DOPA at the lettuce leaf surface into melanin. Panel A shows the assay in a 1.5 mM L-DOPA solution. The upper tube is the negative control, the other 3 tubes are identical. Panel B shows the result of the incubation of leaf discs between wetted filter papers which contain 1.5 mM L-DOPA.

**Figure 14:** Progeny testing of a mutant of lettuce showing a reduced wound-induced pink discoulouration on midrib browning. Panel A shows the midrib discs of 8 progeny plants, numbered 1 to 8 (3 discs per plant) of the reduced pinking mutant. Panel B shows the midrib discs of 8 control plants numbered 9 to 16 (3 discs per plant) which show a normal browning response.

**Figure 15:** Assessment of a mutant of lettuce showing a reduced wound-induced pink discoulouration or browning response after cutting and packaging under ambient atmosphere. Leaf pieces of the head of a control plant are shown on the left and leaf pieces of the reduced discoulouration mutant is shown on the right. The fresh cut leaf material was stored for 6 days at 4°C. The brown discoulouration can be clearly observed in the control samples whereas the mutant samples remain unchanged.

**EXAMPLES**

**EXAMPLE 1**

**Genetic modification of lettuce using ems**

[0093] Approximately 2000 seeds of the lettuce varieties Troubadour, Apache, Yorvik and Roderick were incubated in an aerated solution of either 0.05% (w/v) or 0.07% (w/v) ems during 24 hours at room temperature. After the ems treatment the M1 seeds were rinsed water and planted in a greenhouse at 20°C at 16 hours light, 8 hours dark regime to grow the mature plants and to induced bolting and flowering in order to produce M2 seeds. After maturation, M2 seeds were harvested, bulked and stored until further use. The mutation frequency was estimated on the basis of the relative number of individual plants with a bleached phenotype which are disturbed in the chlorophyll biosynthesis.

**EXAMPLE 2**

**Development of a phenotypic screen diagnostic for the wound-induced discoulouration of lettuce based on pink pigment formation**

[0094] A phenotypic assay was developed in which leaf discoulouration of lettuce induced by wounding can readily be assessed. This approach allows young plant screening for discoulouration. Leaf discs of 5 mm diameter were taken from young or mature plants and placed between wetted filter papers in a tray. The system was incubated at 5°C for 7 days. During the incubation a pink dye developed at the wound site of the leaf disc which became clearly visible as a printed circle on the filter paper (Figure 2).

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In order to demonstrate that the production of the pink dye requires an active phenylpropanoid pathway, the effect of inhibitors of PAL (cinnamaldehyde, Figure 3) and PPO (L-cysteine, Figure 4) were tested in this assay. When cinnamaldehyde was applied during the assay, the pink discolouration was completely inhibited at a concentration of 0.01% or higher.

A similar result was obtained using L-cysteine at a concentration of 0.001% and higher, while other amino acids like L-leucine or L-alanine did not show any effect. This demonstrates that L-cysteine can inhibit the pinking response of the lettuce leaf discs and that the inhibitory effect of L-cysteine is specific.

In order to demonstrate that L-cysteine is indeed acting as an inhibitor of PPO activity in this system, the lettuce leaf discs were incubated with the PPO substrate L-3,4-dihydroxyphenylalanine (L-DOPA). Although L-DOPA is not considered to be a natural substrate for lettuce PPO, a dark brown to black discolouration was observed at the wound surface which is the manifestation of the formation of melanin through PPO. When 1 mM or a higher concentration of L-cysteine was applied simultaneously, the discoloration was completely inhibited as shown in Figure 5.

The lettuce leaf disc pinking response was further characterised by applying a heat shock before inducing the wound response. Detached leaves were incubated during 90 seconds at 21, 40, 50 and 60°C. After this treatment leaf discs were taken and assayed for pinking. The pinking response was completely inhibited when the heat shock was carried out at a temperature of 50°C or higher. This result is shown in Figure 5.

As L-cysteine is known to react with o-quinones, which are PPO products, by converting them back into colourless diphenols, the effects of L-cysteine on the pink dye coming from lettuce leaf discs was determined. In parallel the effect of L-cysteine was determined on the melanin formation upon incubation with L-DOPA.

Leaf discs were taken and incubated according to the procedures described above. After the wound response was completed, a concentration series of L-cysteine was added to the leaf disc and the change in colour was monitored. The result is shown in Figure 7. The experiment clearly demonstrated that L-cysteine was converting the pink dye back into a colourless compound whereas the black melanin formed in the L-DOPA assay was not affected by the L-cysteine. This demonstrates that the pink dye is very likely an o-quinone which is formed by the lettuce polyphenol oxidation system.

To demonstrate that the observed in vitro response reflects a response which is physiologically relevant the L-cysteine based discoloration was applied to field-grown plant material. This was carried out by harvesting a leaf from a field-grown lettuce plant which showed severe pinking symptoms along the veins. This is typically observed when plants have been stressed, for example by conditions of severe water logging. The leaf was used to prepare leaf discs which were incubated immediately by 1 mM L-cysteine. After approximately 30 minutes incubation at room temperature, the pink discolouration disappeared as shown in Figure 8.

Taken together, these experimental data show that lettuce leaf discs can be induced by wounding to produce pink discoloration which is PAL and PPO dependent. This phenotype allows an efficient and effective screening procedure for lettuce mutants which have a modified wound response induced discoloration channelled through PAL, PPO or both.

### EXAMPLE 3

**Screening for mutants with a reduced wound-induced discoloration**

In order to identify lettuce mutants with low wound-induced enzymatic browning or pinking potential, the leaf disc assay described in Example 2 was applied to plants of a lettuce mutant population. 12000 Plants were grown in a greenhouse (Location: De Lier, the Netherlands; sowing 28 March; planting 18 April; growing under regular lettuce grower’s conditions) and from each individual plant a leaf disc was taken (sampling from 15 May onwards) and incubated as pools of (on average) 25 samples between wetted filter papers at 5°C for 7 days. A visual score was given to each leaf disc depending on the intensity of the pink discoloration. On the basis of this assessment, plants were selected of which the leaf discs showed no or a relatively low degree of wound surface discoloration. The plant with hardly visible traces of discoloration was numbered 06D.210202.

Of the 12000 plants, 1 plant was finally selected which showed only traces of discoloration which were hardly visible and 11 which showed a relatively low level of discoloration. The result of one of these assays is shown in Figure 9.

The discolouration assay was repeated for the initially selected 12 individuals and for most individual cases the original result was confirmed. Only the confirmed individuals were selected for further analysis and seed production.
EXAMPLE 4

Screening for mutants with a reduced wound-induced discolouration

[0107] In order to identify lettuce mutants with low wound induced enzymatic browning potential the leaf disc assay described in Example 2 was applied to plants of a lettuce mutant population. 8500 plants were grown in a greenhouse until 3 weeks old (6-8 leaf stage) and from each individual plant a leaf disc was taken and incubated between wetted filter papers at 5°C for 7 days.

[0108] A visual score was given to each leaf disc depending on the intensity of the pink discolouration. On the basis of this assessment, plants were selected of which the leaf discs showed no or a relatively low degree of wound surface discolouration. Of the 8500 plants 8 plants were selected which did not show any visible discolouration and 10 showed a relatively low discolouration. The discolouration assay was repeated for the 18 individuals that were initially selected and for most individual cases the original result was confirmed. Twelve individuals are shown in Figure 10. Only the confirmed individuals were selected for further analysis and seed production. One mutant plant without pleiotropic side-effects (e.g. bleaching, dwarfing) was given number 05D.202539. The seed produced by selfing of this plant was numbered 05D.810596. The seed produced by selfing of three plants grown from seeds of 05D.810596 was numbered 07G.9979 and deposited at NCIMB. The NCIMB-number is 41441 (deposited on 10 October 2006).

EXAMPLE 5

Phenotypic analysis of the selected mutants showing reduced, wound-induced discolouration

[0109] Of the 12 mutants selected from the screen presented in Example 3, 6 showed a strong reduced growth phenotype and bleaching. Other mutants developed normally i.e. according to the type of the starting population of the mutagenesis experiment.

[0110] The dwarfed and bleached mutants are probably disturbed in chloroplast function. As PPO resides in these cellular organelles this may explain the relatively low response in the leaf disc assays. As such pleiotropic mutations are undesirable, these mutants were considered to be less relevant.

[0111] The mutant plant 06D.210202 which showed the strongest reduction of leaf disc discolouration showed a normal phenotype and the mutation is therefore considered specific for the discolouration without strong pleiotropic effects.

EXAMPLE 6

Confirmation of the near absence of discolouration phenotype in offspring

[0112] To demonstrate that the reduced discolouration of lettuce mutants like plant 06D.210202 from Examples 3 and 5 is caused by a genetic effect generated by the mutagenesis treatment described herein, seed were produced by selfing. The seed produced by selfing of plant 06D.210202 was numbered 06D.819784. The seeds were germinated in soil and the plants were tested for discolouration using the leaf disc pinking assay.

[0113] This experiment clearly showed that the altered phenotype had a genetic basis as all progeny plants showed a similar phenotype i.e. a strong reduction in pink discolouration, like the mutant which was used to produce the seeds. This result is illustrated in Figure 11.

[0114] The seed produced by selfing of three plants grown from seeds of 06D.819784 was numbered 06D.863B2 and deposited at NCIMB. The NCIMB-number is 41454 (deposited on 3 January 2007).

EXAMPLE 7

Development of a phenotypic screen diagnostic for the wound-induced discolouration of lettuce based on brown pigment formation

[0115] Lettuce plants were grown to maturity and parts of outer leaves were taken by cutting discs from the rib tissue. The discs were incubated on wetted filter paper at 16°C. After approximately 72 hours, the wound surface had turned brown. In the presence of 10 mM L-cysteine the browning response was inhibited indicating that the observed discolouration is PPO mediated. A representative outcome of such experiment is shown in Figure 12.

[0116] As the response as displayed in Figure 12 is a PPO mediated browning response, the screening procedure as described in this example can considered to be effective and unbiased in order to screen for mutants showing reduced brown discoloration which occurs during the processing of lettuce.
EXAMPLE 8

Development of a phenotypic screen diagnostic for the wound-induced discolouration of lettuce based on the conversion of L-3,4-dihydroxyphenylalanine (L-DOPA) into a black pigment called melanin

[0117] In addition to assays which address wound-induced discolouration in a broad sense, the method according to this invention also allows to screen in a more specific manner for mutants which have a reduced PPO activity. A phenotypic assay which is indicative for PPO activity was developed by using leaf discs which were incubated in the presence of 1.5 mM L-DOPA. As a black discolouration became apparent it can be concluded that L-DOPA can readily be converted by the polyphenol oxidising system at the wound surface of lettuce leaves into a black pigment called melanine. L-DOPA is converted by PPO into the reactive L-DOPA-quinone which is converted non-enzymatically via dopachrome and indol quinone into the black melanin.

[0118] Furthermore, it was shown that the reaction can be inhibited by adding 1 mM L-cysteine during the reaction (Figure 5). Therefore, this assay enable the identification of mutants which are modified in the ability to mount a PPO activity at a wound surface of a leaf. The response of lettuce leaf discs to the presence of L-DOPA can be observed both in solution as well as between wetted filter papers as shown in Figure 13.

EXAMPLE 9

Assessing the offspring of a mutant showing a near absence of discolouration for reduced browning response using the ribdisc browning assay

[0119] To demonstrate that the reduced discolouration of lettuce mutants like plant number 06D.210202 from Examples 3, 5 and 6 which is significantly reduced in pink discolouration of wound surfaces of leaf discs is also effectively reduced in the wound-induced browning response, a number progeny plants were grown to maturity.

[0120] At this stage of development, 3 midrib discs are taken from the outer leaves of a number of progeny plants. These rib discs are incubated according to the procedure described in Example 7. It is shown that the progeny plants of the mutant which were earlier shown to be strongly reduced in wound-induced pink discolouration are also strongly reduced in wound-induced midrib browning. The result of this experiment is shown in Figure 14.

EXAMPLE 10

Assessing the offspring of a mutant showing near absence of discolouration for reduced browning response using fresh cut lettuce heads packaged in plastic bags

[0121] Mature heads of the lettuce plants grown from seed number 06D.819784 from Example 6, which is significantly reduced in pink discolouration of wound surfaces of leaf discs, were cut into pieces using a knife and packaged in a plastic bag containing an ambient atmosphere. Control plants which show a normal leaf disc pink discoloration response were treated in an identical manner. The bags were stored at 4°C during 6 days after which the leaf material was assessed for its browning response.

[0122] It is shown by this experiment that the progeny plants of the mutant which were earlier shown to be strongly reduced in wound-induced pink discoloration are also strongly reduced in wound-induced midrib browning when processed and stored in plastic bags using an ambient atmosphere. The result of this experiment is shown in Figure 15.

Claims

1. Lettuce plant having a reduced or absent wound-induced surface discolouration, obtainable by crossing a plant with the NCIMB accession number 41454 or 41441 with another plant of the same species and testing plants resulting from the cross by subjecting the plants or parts thereof to a screening method, which comprises

   a) creating a wound surface on the plants or plant parts to be screened and on the control plants or plant parts;
   b) incubating the wound surfaces to allow for discolouration to occur therein or thereon;
   c) observing the wound surface discolouration in or on the plants or plant parts;
   d) comparing the observed wound surface discoloration in or on the plants or plant parts to be screened with the discoloration that is observed on or in the control plant or plant part to identify the plants or plant parts that show no discoloration or a discoloration that is reduced as compared to the control plant or plant part.
2. Lettuce plant as claimed in claim 1, wherein the plant parts to be screened are selected from leafs, heads, shoots, roots, tubers, stems, flowers, seeds or pieces thereof and cells.

3. Lettuce plant as claimed in claim 1 or 2, wherein the plant parts to be screened are leaf discs.

4. Lettuce plant as claimed in claim 1 or 2, wherein the plant parts to be screened are discs from midrib tissue.

5. Lettuce plant as claimed in any one of the claims 1-4, wherein the incubation takes place in an aqueous environment.

6. Lettuce plant as claimed in claim 5, wherein the aqueous environment comprises wetted filter paper.

7. Lettuce plant as claimed in claim 5, wherein the aqueous environment comprises water or a solution.

8. Lettuce plant as claimed in claim 6 or 7, wherein the aqueous environment contains a compound selected from L-3,4-dihydroxyphenylalanine, chlorogenic acid, isochlorogenic acid, L-tyrosine and catechol.

9. Lettuce plant as claimed in any one of the claims 1-8, wherein the control plant is a plant of which a leaf disc when incubated between two sheets of wetted filter paper for 7 days at 5°C shows pink discolouration around the edges.

10. Lettuce plant as claimed in any one of the claims 1-9, wherein the plant shows a reduced wound-induced surface discolouration and no negative pleiotropic effects.

11. Progeny of a parent plant as claimed in any one of the claims 1-10 that shows the absence or reduction of wound-induced surface discolouration as found in the parent plant.

12. Part of a plant as claimed in any one of the claims 1-11, wherein the plant part shows a reduced wound-induced surface discolouration or can be grown into a plant that shows a reduced wound-induced surface discolouration.

13. Part of a plant as claimed in claim 12, which part is selected from leafs, heads, shoots, roots, tubers, stems, flowers, fruits, seeds or pieces thereof and cells.

14. Plant regenerated from a plant part as claimed in claim 13 that shows the absence or reduction of wound-induced surface discolouration as found in the parent.

15. Seed of a plant as claimed in any one of the claims 1-11, wherein plants that can be grown from the seed show a reduced wound-induced surface discolouration.

16. Progeny from a seed as claimed in claim 15 that shows the absence or reduction of wound-induced surface discolouration as found in the parent.

17. Vegetable product, comprising a lettuce plant as claimed in any one of the claims 1-11 or part thereof.

18. Vegetable product as claimed in claim 17, wherein the product is processed lettuce that shows no or a limited wound-induced surface discolouration.

Patentansprüche

1. Salatpflanze mit einer reduzierten oder fehlenden wundinduzierten Oberflächenentfärbung, erhältlich durch Kreuzen einer Pflanze mit der NCIMB Zugangs-Nummer 41454 oder 41441 mit einer anderen Pflanze der selben Art, und Testen der Pflanzen, die aus der Kreuzung resultieren, durch Unterwerfen der Pflanzen oder Teilen davon einem Screening-Verfahren, welches umfasst

   a) Schaffen einer Wundoberfläche auf Pflanzen oder Pflanzenteilen, die gescreent werden sollen, und auf die Kontrollpflanzen oder Pflanzenteilen;
   b) Inkubieren der Wundoberflächen, um zu ermöglichen, dass eine Entfärbung darin oder darauf entsteht;
   c) Beobachten der Wundoberflächenentfärbung in oder auf den Pflanzen oder Pflanzenteilen;
   d) Vergleichen der beobachteten
Wundoberflächenentfärbung in oder auf den Pflanzen oder Pflanzenteilen, die gescreent werden sollen, mit der Entfärbung, die an oder in der Kontrollpflanze oder Pflanzenteil beobachtet wird, um die Pflanzen oder Pflanzenteile zu identifizieren, die keine Entfärbung zeigen oder eine solche Entfärbung zeigen, die im Vergleich zu der Kontrollpflanze oder dem Pflanzenteil reduziert ist.

2. Salatpflanze wie in Anspruch 1 beansprucht, wobei die Pflanzenteile, die gescreent werden sollen, ausgewählt sind aus Blättern, Köpfen, Austrieben, Wurzeln, Knollen, Stämmen, Blüten, Samen, oder Teilen davon, und Zellen.

3. Salatpflanze wie in Anspruch 1 oder 2 beansprucht, wobei die Pflanzenteile, die gescreent werden sollen, Blatt scheiben sind.

4. Salatpflanze wie in Anspruch 1 oder 2 beansprucht, wobei die Pflanzenteile, die gescreent werden, Scheiben von Mittelrippe-Gewebe sind.

5. Salatpflanze wie in irgendeinem der Ansprüche 1 - 4 beansprucht, wobei die Inkubation in einer wässrigen Umgebung stattfindet.

6. Salatpflanze wie in Anspruch 5 beansprucht, wobei die wässrige Umgebung benetztes Filterpapier umfasst.

7. Salatpflanze wie in Anspruch 5 beansprucht, wobei die wässrige Umgebung Wasser oder eine Lösung umfasst.

8. Salatpflanze wie in Anspruch 6 oder 7 beansprucht, wobei die wässrige Umgebung eine Verbindung, ausgewählt aus L-3,4-Dihydroxyphenylalanin, Chlorogensäure, Isochlorogensäure, L-Tyrosin und Catechol umfasst.

9. Salatpflanze wie in irgendeinem der Ansprüche 1 - 8 beansprucht, wobei die Kontrollpflanze eine Pflanze ist, von welcher eine Blattscheibe, wenn sie zwischen zwei Blättern benetzten Filterpapiers für 7 Tage bei 5°C inkubiert wird, um die Ränder pinke Entfärbung zeigt.

10. Salatpflanze wie in irgendeinem der Ansprüche 1 - 9 beansprucht, wobei die Pflanze eine reduzierte wundinduzierte Oberflächenentfärbung zeigt, und keine negativen pleiotropen Effekte.

11. Nachkomme einer Elternpflanze wie in irgendeinem der Ansprüche 1 - 10 beansprucht, welcher die Abwesenheit oder Reduzierung einer wundinduzierten Oberflächenentfärbung zeigt, wie in der Elternpflanze gefunden.

12. Teil einer Pflanze wie in irgendeinem der Ansprüche 1 - 11 beansprucht, wobei der Pflanzenteil eine reduzierte wundinduzierte Oberflächenentfärbung zeigt, oder zu einer Pflanze gezogen werden kann, die eine reduzierte wundinduzierte Oberflächenentfärbung zeigt.

13. Teil einer Pflanze wie in Anspruch 12 beansprucht, welcher Teil ausgewählt ist aus Blättern, Köpfen, Austrieben, Wurzeln, Knollen, Stämmen, Blüten, Früchten, Samen, oder Teilen davon, und Zellen.


15. Samen einer Pflanze wie in irgendeinem der Ansprüche 1 - 11 beansprucht, wobei Pflanzen, die aus diesem Samen gezogen werden können, eine reduzierte wundinduzierte Oberflächenentfärbung zeigen.


17. Pflanzliches Erzeugnis, umfassend eine Salatpflanze wie in irgendeinem der Ansprüche 1 - 11 beansprucht, oder Teile davon.

18. Pflanzliches Produkt wie in Anspruch 17 beansprucht, wobei das Produkt verarbeiteter Salat ist, welcher keine oder eine limitierte wundinduzierte Oberflächenentfärbung zeigt.
Revendications

1. Plante de laitue ayant une décoloration de surface induite par une lésion réduite ou absente, pouvant être obtenue par croisement d'une plante avec le numéro d'accession NCIMB 41454 ou 41441 avec une autre plante de la même espèce et par le test des plantes résultant du croisement en soumettant les plantes ou des parties de celles-ci à une méthode de criblage qui comprend :
   a) la création d'une surface abîmée sur les plantes ou les parties de plante à cribler et sur des plantes ou des parties de plante de contrôle ;
   b) l'incubation des surfaces abîmées pour permettre à une décoloration de se produire sur ou dans la surface ;
   c) l'observation de la décoloration de la surface abîmée sur ou dans les plantes ou les parties de plante ;
   d) la comparaison de la décoloration des surfaces abîmées observées sur ou dans les plantes ou les parties de plante à cribler en fonction de la décoloration, qui est observée sur ou dans les plantes ou les parties de plante de contrôle afin d'identifier les plantes ou les parties de plante qui ne présentent pas de décoloration ou une décoloration qui est réduite par comparaison aux plantes ou parties de plante de contrôle.

2. Plante de laitue selon la revendication 1, dans lequel les parties de plante à cribler sont choisies parmi les feuilles, les inflorescences, les pousses, les tubercules, les tiges, les fleurs, les semences ou des parties de celles-ci et les cellules.

3. Plante de laitue selon la revendication 1 ou 2, dans lequel les parties de plante à cribler sont des disques de feuilles.

4. Plante de laitue selon la revendication 1 ou 2, dans lequel les parties de plante à cribler sont des disques provenant du tissu de la côte.

5. Plante de laitue selon l'une quelconque des revendications 1 à 4, dans lequel l'incubation a lieu dans un environnement aqueux.

6. Plante de laitue selon la revendication 5, dans lequel l'environnement aqueux comprend un papier filtre humidifié.

7. Plante de laitue selon la revendication 5, dans lequel l'environnement aqueux comprend de l'eau ou une solution.

8. Plante de laitue selon la revendication 6 ou 7, dans lequel l'environnement aqueux comprend un composé choisi parmi la L-3,4-dihydroxyphénylalanine, l'acide chlorogénique, l'acide isochlorogénique, la L-tyrosine et le catéchol.

9. Plante de laitue selon l'une quelconque des revendications 1 à 8, dans lequel la plante de contrôle est une plante dont un disque de feuille lorsqu'il est incubé entre deux feuilles de papier filtre humidifié pendant 7 jours à 5 °C présente une décoloration rose sur les pourtours.

10. Plante de laitue selon l'une quelconque des revendications 1 à 9, dans lequel la plante montre une décoloration de surface induite par une lésion réduite et pas d'effets pléiotropiques négatifs.

11. Descendance d'une plante mère selon l'une quelconque des revendications 1 à 10, qui montre l'absence ou la réduction de décoloration de surface induite par une lésion observée dans la plante mère.

12. Partie d'une plante selon l'une quelconque des revendications 1 à 11, dans laquelle la partie de plante montre une décoloration de surface induite par une lésion réduite ou peut devenir une plante qui montre une décoloration de surface induite par une lésion réduite.

13. Partie de plante selon la revendication 12, laquelle partie étant choisie parmi les feuilles, les inflorescences, les pousses, les tubercules, les tiges, les fleurs, les fruits, les semences ou des parties de celles-ci et les cellules.

14. Plante régénérée à partir d'une partie de plante selon la revendication 13 qui montre l'absence ou la réduction de décoloration de surface induite par une lésion observée dans la plante mère.

15. Semence d'une plante selon l'une quelconque des revendications 1 à 11, dans laquelle les plantes qui peuvent être cultivées à partir de la semence montrent une décoloration de surface induite par une lésion réduite.
16. Descendance à partir d'une semence selon la revendication 15 qui montre l'absence ou la réduction de décoloration de surface induite par une lésion observée dans la plante mère.

17. Produit végétal, comprenant un plant de laitue selon l'une quelconque des revendications 1 à 11, ou une partie de celui-ci.

18. Produit végétal selon la revendication 17, dans lequel le produit est une laitue traitée qui montre peu ou pas de décoloration de surface induite par une lésion.
FIG. 4

FIG. 5
FIG. 6
Control group

Progeny derived from the reduced discoloration mutant of lettuce

FIG. 11
FIG. 14

control plant  reduced discolouration mutant

FIG. 15
REFERENCES CITED IN THE DESCRIPTION

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Non-patent literature cited in the description