NEW EUROPEAN PATENT SPECIFICATION

Selective restriction fragment amplification: a general method for DNA fingerprinting
Selektive Restriktionsfragmentenamplifikation: generelles Verfahren für DNS-Fingerprinting
Amplification sélective des fragments de restriction: procédé général pour le "fingerprinting" d'ADN

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References cited:
WO-A-90/08821
WO-A-90/11372
WO-A-91/05861
WO-A-91/14003

- ROSENTHAL A ET AL: "GENOMIC WALKING AND SEQUENCING BY OLIGO-CASSETTE MEDIATED POLYMERASE CHAIN REACTION"
1. FIELD OF THE INVENTION

[0001] This invention relates to applications of DNA fingerprinting and the use of DNA markers in a number of different fields including, but not limited to, plant and animal breeding, variety or cultivar identification, diagnostic medicine, disease diagnosis in animals and plants, identification of genetically inherited diseases in humans, family relationship analysis, forensic analysis, and microbial typing.

[0002] More specifically, this invention relates to methods for DNA fingerprinting and for detecting specific DNA markers in genomes ranging from microorganisms to higher plants, animals and humans. The invention also relates to synthetic DNA molecules and products based thereon which are used in the methods of the invention in the different fields of application.

2. BACKGROUND OF THE INVENTION

2.1. DNA fingerprinting

[0003] DNA fingerprinting or DNA typing, as well as other methods of genotyping, profiling and DNA identification analysis, refer to the characterization of either similarities or one or more distinctive features in the genetic make up or genome of an individual, a variety or race, or a species. The general rule is that the closer the genetic relationship is, the greater the identity or more appropriate the similarity of genomes, and consequently distinctive features in the genome will be rarer. These similar or distinctive features can be revealed by analyzing the DNA of an organism after clearing the DNA with a restriction endonuclease. Restriction endonucleases are enzymes which recognize short nucleotide sequences, usually 4 to 8 bases in length and cleave the two DNA strands, thereby producing fragments of DNA of discrete length. Because of their high degree of sequence specificity, restriction endonucleases will cleave DNA molecules in a very specific fashion. The result is that a reproducible set of DNA fragments will be produced. DNA fragments can be fractionated according to their length on porous matrices, or gels, yielding typical banding patterns, which constitutes a DNA fingerprint of the organism's genetic makeup.

2.2. DNA polymorphisms

[0004] When the fingerprints of very closely related species, varieties or races are compared, the DNA fingerprints can be identical or very similar. When differences are observed within otherwise identical DNA fingerprints, such differences are referred to as DNA polymorphisms: these are new DNA fragments which appear in a fingerprint. The DNA is said to be polymorphic at that position and the novel DNA fragment can be used as a DNA marker. DNA polymorphisms detected in DNA fingerprints obtained by restriction enzyme cleavage can result from any of the following alterations in the DNA sequence: mutations abolishing the restriction endonuclease target site, mutations creating new target sites, insertions, deletions or inversions between the two restriction sites.

[0005] Such DNA polymorphisms are generally referred to as RFLP, Restriction Fragment Length Polymorphisms. Such mutational changes will behave as bona fide genetic markers when they are inherited in a mendelian fashion. Consequently, DNA polymorphisms can be used as genetic markers in much the same way as other genetic markers: in parentage analysis, in genetic studies on the inheritance of traits, in the identification of individuals.

2.3. DNA fingerprinting techniques

[0006] For almost all living organisms, except viruses, restriction digests of the total genomic DNA of the organisms yield so many bands that it is not possible to score individual bands. Therefore, all methods for DNA fingerprinting are based on the principle that only a small fraction of the DNA fragments are visualized so as to yield a simple banding pattern which constitutes the DNA fingerprint.

[0007] The most widely utilized method involves digesting the DNA of the organism with restriction endonucleases, fractionating the restriction fragments by gel electrophoresis, transferring and binding the fractionated DNA fragments onto membranes and hybridizing the membrane with a specific DNA fragment ("probe"). The DNA fragment will form double-stranded DNA molecules with the DNA fragment (or fragments) on the membrane which has (have) complementary nucleotide sequences. When the probe is tagged with a visualizable marker, the DNA fragment to which the probe is attached can be visualized. This procedure is generally referred to as "Southern hybridization". When differences are observed in the sizes of the corresponding restriction fragments to which the probe attaches in closely related genomic DNA molecules, these differences are referred to as DNA polymorphisms, more specifically restriction fragment length polymorphisms. The restriction fragment length differences correspond to the different allelic forms of the
genetic locus recognized by the DNA probe. Although the Southern hybridization method for DNA fingerprinting has been widely used, the method is laborious and time consuming. [0008] Furthermore, the method has a low resolution and can thus only be used to score single loci or a few loci at most in a single reaction.

2.4. Polymerase chain reaction

[0009] The Polymerase Chain Reaction (PCR) technique is a method for synthesizing specific DNA fragments in vitro. The method relies on the use of specific oligonucleotides which will attach to unique sequences on a DNA molecule and a thermostable DNA polymerase. The oligonucleotides are designed in such a way that they can anneal to the opposite strands of the DNA and serve as primers in a DNA synthesis reaction in such a way that each will direct the synthesis of new DNA strands. Hence, in one round of synthesis a complete copy of the DNA molecule between the primers will be made, so that the DNA between the primers is duplicated. Each round of DNA synthesis results in the doubling of the amount of DNA, hence leading to the amplification of the DNA comprised between the two primers. Consequently, the PCR technique allows one to synthesize a precise DNA segment using a small amount of "substrate DNA". WO 90/11372 discloses detecting a test nucleotide in a known DNA sequence. The detection involves using primers defined starting from the DNA sequence adjacent to the test nucleotide. WO 90/08821 relates to microdissection of chromosomal DNA and further digestion of the obtained DNA. The digests are then at both ends ligated to oligonucleotide duplexes which are used to anchor primers corresponding to one of their strands.

3. SUMMARY OF THE INVENTION

[0010] In the present invention we have devised a new method to amplify, with the PCR method, restriction fragments obtained after cleaving the DNA of an organism with at least one restriction enzyme. In this novel application of the PCR method the oligonucleotides used are not directed against a known DNA sequence but are designed such that they recognize the ends of the restriction fragments. To this end it is necessary to modify the ends of the restriction fragments by adding oligonucleotide linkers (or adaptors) to the ends. The reason for this is that the ends of restriction enzymes have only usually few nucleotides in common, i.e. 2 to 8 nucleotides, too short to be used to design primers for PCR amplification.

[0011] The invention is based on the use of a novel application of polymerase chain reaction technique (PCR) for amplifying one or more restriction fragments from complex mixtures of DNA fragments obtained by digesting genomic DNA molecules with restriction endonucleases. One particular advantage of the invention is to enable the amplification of DNA restriction fragments in situations where the nucleotidic sequence of the 5' and 3' ends of the restriction fragments are not determined. In such cases the usual sequence specific primers hybridizing to each strand of a restriction fragment to be amplified can not be defined and therefore one cannot use the methods known in the art for amplification purposes.

[0012] The method of the invention can be used for instance in two different ways, leading to two different types of applications:

(1) Methods for DNA fingerprinting of genomes by randomly selecting subsets of one or more restriction fragments to be amplified by the PCR technique. The invention also covers synthetic oligonucleotides for use in said methods and some applications of said methods can be forensic typing, microbial identification, varietal identification, pedigree analysis and screening of DNA markers linked to genetic traits;

(2) Methods for identifying one or more preselected DNA fragments which can be polymorphic, by PCR amplification. The invention also covers specific synthetic oligonucleotides for use in said methods and some applications of said methods can be the screening of genetically inherited diseases in humans, monitoring the inheritance of agronomic traits in plant and animal breeding and the detection of infections agents in diseases.

4. BRIEF DESCRIPTION OF THE DRAWINGS

[0013] Figure 1 provides a graphic outline for the general method of PCR amplification of tagged restriction fragments obtained by digesting genomic DNA molecules with a restriction enzyme. Figure 2 depicts the ligation of adaptors to different ends of restriction fragments: flush ends and staggered ends. Figure 3 depicts the PCR amplification of tagged restriction fragments. The boxed areas depict the adaptors which are ligated to the restriction fragment, and the primers which are used in the PCR amplification. The arrows indicate
5. DETAILED DESCRIPTION OF THE INVENTION

5.1. Definitions

In the description and examples that follow, a number of terms are used herein. In order to provide a clear and consistent understanding of the specification and claims, including the scope to be given such terms, the following definitions are provided.

- **Restriction Endonuclease**: a restriction endonuclease or restriction enzyme is an enzyme that recognizes a specific base sequence (target site) in a double-stranded DNA molecule, and will cleave both strands of the DNA molecule at every target site.

- **Restriction Fragments**: the DNA molecules produced by digestion with a restriction endonuclease are referred to as restriction fragments. Any given genome will be digested by a particular restriction endonuclease into a discrete set of restriction fragments. The DNA fragments that result from restriction endonuclease cleavage are separated and detected by gel electrophoresis.

- **Restriction Fragment Length Polymorphism (RFLP)**: the genomic DNA of two closely related organisms, for example, will exhibit differences in their nucleotide sequence composition at many sites. When these differences occur in the target site for a restriction endonuclease, the modified target site will not be cleaved at that point. Likewise, a nucleotide sequence variation may introduce a novel target site where non exists in the other organism, causing the DNA to be cut by the restriction enzyme at that point. Alternatively, insertions or deletions of nucleotides occurring in one organism between two target sites for a restriction endonuclease will modify the distance between those target sites. Because of this, digestion of the two organism's DNA will produce restriction fragments having different lengths. A polymorphism in the length of restriction fragments produced by digestion of the DNA of the

Figure 4 provides a graphic outline for the PCR amplification of tagged restriction fragments.

Figure 5 shows the general design of the selective primers used in the PCR amplification of tagged restriction fragments. The boxes denote the inverted repeat sequences at the ends of the restriction fragments. The selectivity of the primers is illustrated in two examples where there is respectively a perfect match and a total mismatch between the selective base sequence and that of the restriction fragment template DNA.

Figure 6 shows the principle of selective PCR amplification using a PCR primer which selects template DNA molecules having a trinucleotide sequence adjacent to the adaptor sequence.

Figure 7 depicts the selective PCR amplification of tagged restriction fragments.

Figure 8 shows the principle of fragment specific amplification using a combination of two PCR primers each comprising 6 selective bases. Each primer forms a double-stranded structure in the different strand of the restriction fragment and thereby forms a primer/template complex from which DNA synthesis can be initiated (represented by the arrows).

Figure 9 depicts the general sequence elements which are recognized with the method of selective restriction fragment amplification, including the two nucleotide sequences which are recognized and the distance separating the two sequences.

Figure 10 depicts the types of nucleotide sequence variations which are detected in the method of identifying amplified fragment length polymorphisms.

Figure 11 shows a 1.0% agarose gel with the analysis of the results of the amplification of Tomato DNA restricted with PstI, using primers of increasing selectivity.

Figure 12 shows a 1.0% agarose gel with the analysis of the results of specific amplification of 3 different PstI fragments of Tomato DNA using fragment specific primers.

Figure 13 shows a 2.5% polyacrylamide/1% agarose gel with DNA fingerprints obtained by Selective Restriction Fragment amplification of two Tomato lines.

Figure 14 shows part of a 4.5% denaturing polyacrylamide gel with DNA fingerprints of 4 Tomato lines using SRFA with the enzyme combination PstI/MseI.

Figure 15 shows part of a 4.5% denaturing polyacrylamide gel with DNA fingerprints of 10 Lactuca lines using SRFA with the enzyme combination PstI/MseI.

Figure 16 shows part of a 4.5% denaturing polyacrylamide gel with DNA fingerprints of 2 Corn lines using SRFA with the enzyme combinations PstI/TaqI and EcoRI/TaqI.

Figure 17 shows part of a 4.5% denaturing polyacrylamide gel with DNA fingerprints of 26 Xanthomonas campestris strains using SRFA with the enzyme combination ApaI/TaqI.

Figure 18 shows part of a 4.5% denaturing polyacrylamide gel with DNA fingerprints of different individuals of 4 domestic animals Chicken, Pig, Cow and Horse using SRFA with the enzyme combination SseI/MseI.
two organisms will result.

- Gel Electrophoresis: To detect restriction fragments, an analytical method for fractioning double-stranded DNA molecules on the basis of size is required. The most commonly used technique for achieving such fractionation is gel electrophoresis. The rate at which DNA fragments move in such gels depends on their size; thus, the distances travelled decrease as the fragment lengths increase. The DNA fragments fractionated by gel electrophoresis can be visualized directly by a staining procedure if the number of fragments included in the pattern is small.

- Synthetic oligonucleotides: the single-stranded DNA molecules having preferably from almost 10 to almost 50 bases, which can be synthesized chemically are referred to as synthetic oligonucleotides. In general, these synthetic DNA molecules are designed to have a unique nucleotide sequence, although it is possible to synthesize families of molecules having related sequences and which have different nucleotide compositions at specific positions within the nucleotide sequence. The term synthetic oligonucleotide will be used to refer to DNA molecules having a unique nucleotide sequence. The term mixed synthetic oligonucleotides will be used to refer to families of related synthetic oligonucleotides.

- Ligation: the enzymatic reaction catalyzed by the enzyme ligase in which two double-stranded DNA molecules are covalently joined together is referred to as ligation. In general, both DNA strands are covalently joined together, but it is also possible to prevent the ligation of one of the two strands, through chemical or enzymatic modification of one of the ends. In that case the covalent joining will occur in only one of the two DNA strands.

- Adaptors: short double-stranded DNA molecules, with a limited number of base pairs, e.g. 10 to 30 base pairs long, which are designed in such a way that they can be ligated to the ends of restriction fragments. Adaptors are composed of two synthetic oligonucleotides which have nucleotide sequences which are in part complementary to each other. When mixing the two synthetic oligonucleotides, they will form a double-stranded structure in solution under appropriate conditions. One of the ends of the adaptor molecule is designed so that it can be ligated to the end of a restriction fragment, the other end is designed so that it cannot be ligated.

- Polymerase Chain Reaction (PCR): the enzymatic reaction in which DNA fragments are synthesized from a substrate DNA in vitro is referred to as PCR. The reaction involves the use of two synthetic oligonucleotides, which are complementary to nucleotide sequences in DNA molecules which are separated by a short distance of a few hundred to a few thousand base pairs, and the use of a thermostable DNA polymerase. The chain reaction consists for example of a series of 10 to 30 cycles. In each cycle the substrate DNA is first denatured at high temperature. After cooling down the synthetic oligonucleotides which are present in vast excess will form double-stranded structures with the substrate DNA molecules in solution at specific sites on the substrate DNA molecule that have complementary nucleotide sequences. The oligonucleotide-substrate DNA complexes will then serve as initiation sites for the DNA synthesis reaction catalyzed by the DNA polymerase, resulting in the synthesis of a new DNA strand complementary to the substrate DNA strand.

- DNA amplification: the term DNA amplification will be used to denote the synthesis of double-stranded DNA molecules in vitro using Polymerase Chain Reaction (PCR). The products of the PCR reaction will be referred to as amplified DNA fragments.

- Primers: in general, the term primer refers to a DNA strand which can prime the synthesis of DNA. DNA polymerase cannot synthesize DNA de novo without primers: they can only extend an existing DNA strand in a reaction in which the complementary strand is used as a template to direct the order of nucleotides to be assembled. We will refer to the synthetic oligonucleotide molecules which are used in the PCR reaction as primers.

- Southern Hybridization Procedure: the purpose of the Southern hybridization procedure, also referred to as Southern blotting, is to transfer physically DNA fractionated by agarose gel electrophoresis onto a support such as nylon membrane or nitrocellulose filter paper while retaining the relative positions of DNA fragments resulting from the fractionation procedure. The methodology used to accomplish the transfer from agarose gel to the support is to draw the DNA from the gel into the support by capillary action.

- Nucleic Acid Hybridization: Nucleic acid hybridization is used to detect related DNA sequences by hybridization of single-stranded DNA on supports such as nylon membrane or nitrocellulose filter papers. Nucleic acid molecules that have complementary base sequences will reform the double-stranded structure if mixed in solution under the proper conditions. The double-stranded structure will be formed between two complementary single-stranded nucleic acids even if one is immobilized on a support. In the Southern hybridization procedure, the latter situation occurs.

- Hybridization Probe: to detect a particular DNA sequence in the Southern hybridization procedure, a labelled DNA molecule or hybridization probe is reacted to the fractionated DNA bound to a support such as nylon membrane or nitrocellulose filter paper. The areas on the filter that carry DNA sequences complementary to the labelled DNA probe become labelled themselves as a consequence of the reannealing reaction. The areas of the filter that exhibit such labelling can then be detected according to the type of label used. The hybridization probe is generally produced by molecular cloning of a specific DNA sequence from the maize genome.
The invention thus relates to a Process for the amplification by Polymerase Chain Reaction of at least one restriction fragment obtained from a starting DNA which contains a plurality of restriction sites for at least one specific determined Restriction endonuclease, which process comprises:

(a) digesting said starting DNA with at least one restriction endonuclease, to fragment it into restriction fragments;
(b) ligating the restriction fragments obtained with at least one double-stranded synthetic oligonucleotide adaptor having one end which is compatible to be ligated to one or both of the ends of the restriction fragments to thereby produce tagged restriction fragments;
(c) contacting said tagged restriction fragments under hybridizing conditions with at least one oligonucleotide primer;

wherein said primer or primers include a nucleotide sequence having the same nucleotide sequence as the terminal parts of the strands at the ends of said tagged restriction fragments, including the nucleotides involved in the formation of the target sequence for said restriction endonuclease and including at least part of the nucleotides present in the ligated adaptors, wherein at least one of said primers includes at its 3' end, a selected sequence comprising at least one nucleotide located immediately adjacent to the nucleotides involved in the formation of the target sequence for said restriction endonuclease
(d) amplifying said tagged restriction fragments hybridized with said primer or primers in the presence of the required nucleotides and DNA polymerase or elongating the hybridized primers along those tagged restriction fragments, and
(e) identifying or recovering amplified or elongated DNA fragments as produced in step (d).

5.2. Description of the preferred embodiments

This invention relates more particularly to a process and means which enable the polymerase chain reaction (PCR) to be applicable to the detection of restriction fragment polymorphisms (RFPs) including length polymorphisms. This invention comprises methods for detecting RFPs, synthetic oligonucleotides for use in the methods of the invention, kits comprising means for detecting RFPs, and applications of the methods and procedures of the invention for plant and animal breeding, diagnostics of genetically inherited diseases, identification of organisms, and forensic typing, etc...

Specifically, this invention provides means for the identification of either individual genomic restriction fragments or of sets of genomic restriction fragments from any organism, microorganism, plant, animal or human, which are either individually genetically linked to one or more particular traits or that collectively provide a fingerprint of the genome that can be used to identify an organism, a variety or an individual.

The general method of the invention for production and for identification of restriction fragments involves the use of restriction endonucleases, ligation of synthetic oligonucleotides to the restriction fragments, and PCR amplification of restriction fragments (figure 1). Restriction endonucleases cleave genomic DNA molecules at specific sites, target sites, thereby generating restriction fragments.

PCR amplification of restriction fragments no matter whether one knows the nucleotidic sequence of the ends of the restriction fragments or not, can be achieved according to the invention, by first ligating synthetic oligonucleotides (adaptors) to the ends of restriction fragments, thus providing each restriction fragment with two common tags which will serve as an anchor base for the primers used in PCR amplification.

Typically, restriction enzymes produce either flush ends, in which the terminal nucleotides of both strands are base paired, or staggered ends in which one of the two strands protrudes to give a short single strand extension (figure 2). In the case of restriction fragments with flush ends, adaptors are used with one flush end. In the case of restriction fragments with staggered ends adaptors are used which have a single stranded extension which is complementary to the single stranded extension of the restriction fragment. Consequently, for each type of restriction fragment specific adaptors are used, which differ only in one of the ends so as to allow the adaptor to be ligated to the restriction fragment. Typically, the adaptors are composed of two synthetic oligonucleotides which are in part complementary to each other, and which are usually approximately 10 to 30 nucleotides long, preferably 12 to 22 nucleotides long and which form double-stranded structures when mixed together in solution. Using the enzyme ligase the adaptors are ligated to the mixture of restriction fragments. When using a large molar excess of adaptors over restriction fragments one ensures that all restriction fragments will end up carrying adaptors at both ends. Restriction fragments prepared with this method will be referred to as tagged restriction fragments and the method will be further referred to as restriction fragment tagging.

The adaptors can now serve as templates for the primers having the characteristics hereabove defined, used in the subsequent PCR amplification reaction. In a preferred embodiment of the invention, the restriction fragment carries the same adaptor at both of its ends and a single PCR primer can be used to amplify the restriction fragment as illustrated in figure 3. Since in such a case all restriction fragments are tagged in the same way, it is obvious that PCR amplification of a mixture of tagged restriction fragments will amplify all restriction fragments in a synchronous...
the fragment. When the primer matches with only one end, the amplification will be linear rather than exponential, and
be amplified when the selective bases of the primers used recognize both complementary sequences at the ends of
selection of these bases renders the primer selective. It must be realized that a tagged restriction fragment will only
will also use the term selective bases to denote the nucleotides in the variable sequence part thus showing that the
the selectivity of the primers increases with the number of nucleotides in the variable (selected) sequence part. We
illustrated in figure 5.
the variable DNA sequence can perfectly base pair with the template strand of the tagged restriction fragment, as
under appropriate conditions the primers will only initiate DNA synthesis on those tagged restriction fragments in which
preselection of tagged restriction fragments which will be amplified in the PCR step: when the PCR reaction is performed
of selected bases which are different among the primers thus formed.
distinguished primers. In such a case, primers can have the same constant sequence and variable sequences made
a particular embodiment of the invention, several sequences of selected bases can be used, in order to define several
ments should coamplify synchronously in the PCR reaction. In a preferred embodiment of the invention applicable to
genomic DNAs of large sizes, we have used a general principle to limit the number of restriction fragments which are
to be amplified. This is done by preselecting a subset of tagged restriction fragments so that only a relatively small
number of tagged restriction fragments will be amplified during the PCR amplification reaction.
[0025] The selective principle defined in this embodiment of the invention resides in the design of the oligonucleotides
which are used as primers for the PCR amplification, as is illustrated in figure 5.
[0026] Tagged restriction fragments have the following general structure: a variable DNA sequence (corresponding
to the restriction fragment before tagging), flanked on both sides by an inverted DNA sequence (constant sequence).
The inverted DNA sequence (constant DNA sequence) is composed of part of the target sequence of the restriction
endonuclease and of the sequence of the adaptor attached to both ends of the restriction fragment. The variable
sequences of the restriction fragments comprised between the constant DNA sequences are usually unknown, and
will thus have a random sequence composition. Consequently, the nucleotide sequences flanking the constant DNA
sequence will be totally random in a large mixture of restriction fragments.
[0027] The present invention therefore also provides specific PCR primers which comprise a constant nucleotide
sequence part and in the embodiment of the invention relying to the amplification of a restricted subset of the restriction
fragments obtained, a variable sequence part. In the constant sequence part the nucleotide sequence is designed so
that the primer will perfectly base pair with the constant DNA sequence of one of the DNA strands at the end of the
restriction fragment. The variable sequence part comprises a randomly chosen nucleotide sequence ranging from 1
to 10 bases chosen.
[0028] The expression “variable sequence” more exactly designates a sequence consisting of selected nucleotides
forming a sequence which will then remain constant for the purpose of amplifying a subset of restriction fragments. In
a particular embodiment of the invention, several sequences of selected bases can be used, in order to define several
distinguished primers. In such a case, primers can have the same constant sequence and variable sequences made
of selected bases which are different among the primers thus formed.
[0029] It is the addition of these variable (selected) sequences to the 3' end of the primers which will direct the
preselection of tagged restriction fragments which will be amplified in the PCR step: when the PCR reaction is performed
under appropriate conditions the primers will only initiate DNA synthesis on those tagged restriction fragments in which
the variable DNA sequence can perfectly base pair with the template strand of the tagged restriction fragment, as
illustrated in figure 5.
[0030] The selection is determined by the number of nucleotides residing in the variable sequence part of the primer:
the selectivity of the primers increases with the number of nucleotides in the variable (selected) sequence part. We
will also use the term selective bases to denote the nucleotides in the variable sequence part thus showing that the
selection of these bases renders the primer selective. It must be realized that a tagged restriction fragment will only
be amplified when the selective bases of the primers used recognize both complementary sequences at the ends of
the fragment. When the primer matches with only one end, the amplification will be linear rather than exponential, and
the product will remain undetected.

[0031] It is possible to estimate beforehand the degree of selectivity obtained with variable sequences with different numbers of selective bases, using the general formula $4^{2n}$, where $n$ equals the number of selective bases: using 1 selective base, 1 out of 16 tagged fragments will be amplified, using 2 selective bases, 1 out of 256, using 3 selective bases, 1 out of 4,096, using 4 selective bases, 1 out of 65,536, and so on, will be amplified. One preferred embodiment of the present invention thus allows one to selectively amplify a random subset of tagged restriction fragments from any genomic DNA digest regardless of the number of fragments produced by the restriction enzyme used.

[0032] In a preferred embodiment, the number of selective nucleotides is chosen so that the number of restriction fragments which will be amplified is limited to 5 to 200. Although this number can be calculated by dividing the number of fragments by $4^{2n}$, a precise prediction is not possible because not all restriction fragments can be amplified with equal efficiency. Hence, in practice, one finds less fragments of the amplification than theoretically expected. It should also be pointed out that mixtures of two (or more) primers can be used. This will allow the amplification of the fragments recognized by each primer and in addition, the fragments recognized by the two primers. Finally, it should be pointed out that the selection based on the base pairing between the selective nucleotides of the primer and the complementary template is strongly influenced by the temperature chosen for the annealing step in the PCR reaction when this temperature is below or too close to the melting temperature of the primer/template complex, primers will anneal the imperfectly matching template sequences allowing a mismatch to occur in the complex. This should be avoided because it will lead to the amplification of many more fragments than predicted, producing more variable results.

[0033] The PCR products obtained in accordance with the invention can be identified using standard fractionation methods for separating DNA molecules according to size followed by staining of the DNA molecules with appropriate agents. Alternatively, the primers used for the PCR amplification can be tagged with a suitable radio-active label or fluorescent chromophore thus allowing the identification of the reaction products after size fractionation. In a preferred embodiment of the invention the PCR products are fractionated by gel electrophoresis using standard gel matrices such as, but not limited to, agarose, polyacrylamide or mixed agarose/polyacrylamide. The PCR products obtained according to the invention will be denoted further by the term Amplified Restriction Fragments (ARF).

[0034] The means and method of the present invention can be used to generate sets of ARF from restriction digests of any complex genome. The invention permits the number of restriction fragments obtained to be tuned in accordance with the resolution of the gel fractionation system used to separate the ARFs. In one particular embodiment the selective primers are designed to produce 5 to 10 ARFs which are then separated by agarose gel electrophoresis. Another particular embodiment involves the use of selective primers which are designed to produce 20 to 50 ARFs which are then separated on a high resolution gel electrophoresis system such as, but not limited to, polyacrylamide gels or mixed polyacrylamide-agarose gels.

[0035] In one preferred embodiment the restriction enzyme or enzymes are chosen to yield restriction fragments in the size range of 20 to 1000 base pairs, because as is generally known for PCR amplification, this fragment size range is amplified most effectively. Although much fragments can be fractionated on various standard gel matrices, best results are obtained by fractionation on denaturating polyacrylamide gel systems as are currently used for DNA sequencing.

[0036] In accordance with the invention, different sets of ARFs are obtained with each different selective primer in the PCR amplification reaction. The patterns of ARFs identified after separation constitute unique and perfectly reproducible fingerprints of the genomic DNA. Such fingerprints can have several applications such as, but not limited to, forensic typing, the diagnostic identification of organisms, and the identification of species, races, varieties or individuals. The level of identification will be determined by the degree of similarity (the degree of variability) exhibited by different members of a specific group. The variability or similarity is determined by the degree of variation in the nucleotide composition of the related genomes. The underlying principle of the invention is that in each Amplified Restriction fragment two nucleotide sequences are detected which are separated from each other by a given distance, as is illustrated in figure 9. Each of the two nucleotide sequences is composed of two parts: (a) the target site for the restriction endonuclease and (b) the nucleotide sequence adjacent to the target site which is included in the selective primer. In related organisms, species, varieties, races or individuals these sequence elements and their relative distances will be conserved to a greater or lesser degree. Hence, the fingerprints constitute a basis for determining the degree of sequence relationships between genomes. On the other hand, differences in the ARF patterns can be used to distinguish genomes from each other. The particular advantages of the present invention over other methods for fingerprinting genomes is the high resolution that can be obtained with the method: several tens or even hundreds of ARFs can be compared simultaneously.

[0037] Another particular application of the present invention involves the screening and identification of restriction fragment polymorphisms (RFP). Changes in the nucleotide composition of genomic DNA often result in polymorphisms of restriction fragments: insertions or deletions affect the size of the restriction fragments containing them (figure 10), nucleotide changes can result in the elimination of restriction endonuclease target sites or the creation of new restriction endonuclease target sites. The most commonly used techniques for identifying such changes are Southern blotting
experiments using cloned DNA probes, a technique usually referred to as restriction fragment length polymorphism (RFLP) detection. This technique involves the extensive screening of randomly cloned DNA fragments in Southern blotting experiments for associated RFLPs among different genomes. In accordance with the method of the present invention, RFPs can be identified directly by comparing the ARFs obtained from different genomes. In principle, the method of the present invention is more sensitive for detecting RFPs because not only differences in the target sites of the restriction endonuclease are detected, but also differences in the adjacent nucleotide sequences comprised in the selective endonuclease fragment mixture. Consequently, the method of the present invention constitutes a far superior method for detecting RFLPs.

RFLPs are now currently used for several applications including forensic typing, monitoring of genetically inherited diseases in humans and monitoring the inheritance of agronomic traits in plant and animal breeding. The underlying principle is that certain DNA polymorphisms which are closely linked with specific genetic traits can be used to monitor the presence or absence of specific genetic traits.

According to the method of the present invention, the analysis of ARF patterns can be used to define the genetic linkage of polymorphic ARFs with specific genetic traits. Such polymorphic ARFs will be further referred to as Amplified Fragment Length Polymorphisms (AFLP®) to distinguish them from RFLP type DNA polymorphisms detected in Southern blotting experiments using cloned DNA probes.

One particular application of the present invention involves the detection of AFLPs linked to specific genetic traits. The application involves the analysis of ARF patterns obtained with different selective primers in restriction digests of genomic DNA of closely related individuals exhibiting differences in the specific genetic trait and the use of analysis techniques that can find correlations between the inheritance of one or more AFLP® and the phenotype exhibited by the specific genetic traits.

A second preferred embodiment of the present invention involves the use of the method of the invention to identify one or more specific restriction fragments. One specific restriction fragment can be amplified from a complex mixture of tagged restriction fragments by first determining the nucleotide sequence of the first 8-12 bases at each end of the restriction fragment. Based on these sequences one can design two primers with each 5 to 10 selective nucleotides exhibiting a sequence complementary to that of the sequence flanking the restriction site of the complementary strand of the restriction fragment. Using such sets of primers one can obtain, after PCR amplification, a single amplified fragment. The restriction fragment used in this method can be either a cloned restriction fragment or an amplified restriction fragment. Since not many restriction fragments cannot be amplified very efficiently, the preferred method of the invention for identifying polymorphic DNA markers involves first amplifying randomly chosen set of fragments and identifying AFLPs which yield strong bands after PCR amplification. These AFLP® can be characterized by sequencing to develop restriction fragment specific primers. Typically, the AFLP® will be isolated by cutting out the corresponding DNA band from the gel, and determining the nucleotide sequences at both ends to establish the sequence of the first 5 to 10 nucleotides adjacent to the restriction endonuclease target sites. Once these nucleotide sequences are known, restriction fragment specific primers can be designed which will only amplify a single restriction fragment from a genomic DNA digest. In this particular embodiment of the invention, one set of two different selective primers can be used for detecting a specific restriction fragment. In each of the two selective primers of one set the selective bases are chosen such that they are complementary to the nucleotide sequence adjacent to the restriction endonuclease target site, as is illustrated in figure 8. The number of selective bases to be included in each primer depends upon the complexity of the restriction endonuclease fragment mixture.

The PCR technique has developed tremendously over the past few years and is rapidly becoming one of the most widely used diagnostic methods in human health care. Its application includes amongst others detection of infectious diseases and detection of genetically inherited diseases. Each diagnostic test is based on the use of two specific synthetic oligonucleotides which are used as primers in the PCR reaction to obtain one or more DNA fragments of specific lengths. In disease detection the test will detect the presence of as little as one DNA molecule per sample, giving the characteristic DNA fragment. In the case of genetically inherited diseases the primers are designed such that their products can discriminate between normal and disease alleles. The distinction either relies on sequence differences in the DNA segment in the genome which is complementary to the primer or, on distance differences between the two primers.

Because the primers exhibit an extremely high degree of specificity, it is possible to monitor different diseases simultaneously, a method often referred to as multiplex PCR. The multiplex PCR method, however, suffers from the limitation that generally only few, 5 to 8, different traits can be monitored simultaneously. The scientific basis for this limitation is that the optimal conditions for PCR amplification (annealing temperature, Mg²⁺ concentration, primer concentration) vary considerably depending on the pair of primers used. In multiplex PCR compromise conditions have to be established under which all primer pairs yield detectable products. In addition, superimposed upon this phenomenon there is the phenomenon of strong differences in the efficiency of amplification of different fragments. Consequently, one often has encountered the problem that products of certain primer pairs are not detectable in multiplex PCR reactions.
The methods of the present invention in essence overcome these limitations of multiplex PCR, because all the primers used in the present invention have a substantial part of their nucleotide sequence in common. Furthermore, by selecting AFLP®, we select DNA markers that are amplified with equal efficiency. Hence, the optima of the PCR amplification conditions for the different selective primers exhibit much less variation than is observed with commonly used sequence specific primers. In essence, ideal compromise between the number of bases in the synthetic oligonucleotide which are necessary to obtain the required specificity of detecting a single DNA fragment of a given size in a complex genome, which is calculated above, and the length and composition of the oligonucleotide which is optimal for efficient PCR amplification. The method of the invention thus provides a far superior method for multiplex PCR.

The present invention provides a general method for isolating DNA markers from any genome and for using such DNA markers in all possible applications of DNA fingerprinting.

The following examples and figures provide an illustration of the invention which is nevertheless not limited to these examples.

EXAMPLES

EXAMPLE 1: SELECTIVE RESTRICTION FRAGMENT AMPLIFICATION OF TOMATO DNA USING PSTI

A) Isolation and modification of the DNA

Total Tomato DNA (Lycopersicon esculentum c.v. Moneymaker) was isolated from young leaves as described by Bernatzki and Tanksley (Theor. Appl. Genet. 72, 314-321). The typical yield was 50 - 100 µg DNA per gram of fresh leaf material. The DNA was restricted with PstI (Pharmacia) and double-stranded (ds) PstI-adapters were ligated to the restriction fragments following the procedure described below. These adapters had the following structure:

5'- CTCGTAAGCTGCCTAGCATGCA -3'
3'- CATCTGACGCATGT -5'

The 3’TGCA-overhang in these adapters anneals to the staggered ends created by PstI. The PstI recognition sequence CTGCAG is not restored upon ligation of this adapter, because the 5’ C-residue is replaced by A. The ligation reaction was designed in such a way that the end result is almost exclusively DNA fragment- to-adapter molecules. This was achieved by: 1. using nonphosphorylated adapters, which excludes adapter- to-adapter ligation, 2. Performing the ligation and restriction reaction at the same time. The latter procedure results in restriction of any fragment-to-fragment ligation product, thereby eliminating these products almost completely. Adapter-to-fragment ligation products cannot be restricted by the restriction enzyme, because the PstI recognition sequence is not restored in these products. The reaction conditions used for the adapter ligation were:

2 µg Tomato DNA
0.2 µg adapters
20 units PstI
1 unit T4 DNA-ligase
10 mM Tris.HAc pH 7.5, 10 mM MgAc, 50 mM KAc,
2 mM dithiotreitol, 0.5 mM ATP

The ligation reaction was performed in a reaction volume of 20 µl for 3 hours at 37°C. After the adapter ligation, non-ligated adapters were removed by selective precipitation. For this purpose the reaction mixture was increased to 100 µl and NH4Ac was added to a final concentration of 2.5 M. 100 µl ethanol of -20°C was added and the mixture was incubated for 5 minutes at room temperature. The DNA was collected by centrifugation for 10 minutes at 14000 rpm in a cooled eppendorf centrifuge at 4°C. The DNA pellet was washed once with 0.5 ml of 70% ethanol at room temperature, and dissolved in 40 µl of TO.1E (10 mM Tris.HCl pH 8.0, 0.1 mM EDTA). The DNA was stored at -20°C.

B) The amplification reaction

The DNA prepared above was used as template for amplification of the PstI-fragments. The reaction mixture for the PCR contained:

[0044] [0045] [0046] [0047] [0048] [0049] [0050]
The reaction mixture was covered with 20 µl of light mineral oil to prevent evaporation during the amplification reaction. The PCR was performed on a Perkin Elmer DNA Thermal Cycler using the following cycle profile: 1 minute at 94°C, 1 minute at 60°C, a temperature increase from 60°C to 72°C at a rate of 1 °C/5 seconds, and 2½ minutes at 72°C. A total of 33 cycles were performed. After the reaction 20 µl chloroform was added, and 10 µl of loading dye, in this case 50% sucrose with 0.1% w/v of the dye Orange G (Merck). This was then mixed well with the reaction mixture and briefly centrifuged to separate the organic fase (mineral oil and chloroform) from the reaction mixture supplemented with the loading dye. 20 µl of this reaction mixture was analysed on a 1.0% agarose gel.

C) Amplification of Tomato DNA with primers of increasing selectivity

Tomato DNA restricted with PstI and tagged with the PstI-adapter was amplified using the conditions specified above. Four different primers were selected with the sequences:

1. 5-CTCGTAGACTGCGTACA-3
2. 5-GACTGCGTACATgcaAA-3
3. 5-GACTGCGTACATgcAC-3
4. 5-GACTGCGTACATgcAC-3

Primer 1 is part of the the top strand of the adapter used to modify the DNA, and therefore should amplify all PstI-fragments. Primer 2 contains part of the adapter sequence, the PstI-recognition sequence (lower case letters) and one selective nucleotide (bold) and should amplify theoretically about 1/16 part of all PstI-fragments. Primers 3 and 4 are similar to primer 2, but contain 2 and 3 selective nucleotides respectively, and therefore are expected to amplify about 1/256 and 1/4096 of the PstI-fragments. Part of the reaction mixtures were analysed on a 1.0 % agarose gel, which is shown in Figure 11. Lanes 1 and 6 of this figure contain DNA markers, of which the sizes are indicated at the left. Lanes 2, 3, 4 and 5 contain the PCR's obtained with primers 1, 2, 3 and 4 respectively. The results indicate that only in case of the primer with 3 selective nucleotides, the number of amplified fragments was such that a clear band pattern was obtained. The other 3 primers gave band patterns, which could not be resolved on agarose gels, because to many PCR products were generated. Within these many PCR products always some fragments predominate, and are seen as bands on a background smear of the other PCR products. Probably these stronger products are present in higher copy numbers on the Tomato genome, or amplify more efficient than the other products. It was anticipated that primers with 3 selective nucleotides had to be used to generate a clear band pattern on agarose gels, because of the total number of PstI-fragments of Tomato genomic DNA (20.000 to 100.000).

D) Analysis of amplified fragments on Southern blots

The amplified fragments were tested on Southern blots to verify that these fragments corresponded to bona-fide restriction fragments of the same size. For this purpose four individual fragments obtained with primer 4, were cut out of the agarose gel. The DNA was purified from these gel slices by means of absorption to glass beads (Gene Clean, manufacturer Bio 101), and part of the purified DNA was reamplified to obtain about 1 µg of each of the four DNA fragments. The reamplification reactions were subsequently electrophoresed on a 1.0% preparative agarose gel, and the desired DNA fragments were purified. 200 ng of each fragment was labeled with (α-32P)dATP using a random hexamer labelling kit according to procedures advised by the manufacturer (Boehringer Mannheim). Total Tomato DNA was restricted with PstI, and electrophoresed on a 1.0% agarose gel. Four clearly separated lanes each containing
about 3 µg of restricted DNA were used. Next, the agarose gel was blotted to a Genescreen+ hybridisation membrane as indicated by the manufacturer (New England Nuclear). After blotting the gel was cut in four slices, each containing one lane of the Tomato DNA restricted with PstI. These four slices were each hybridised to one of the four DNA probes following the procedure described by Klein-Lankhorst et al. (Theor. Apll. Genet. 81, 661-667). The hybridised blots were autoradiographed for 40 hours using Kodak XAR5 films. The results obtained showed that all genomic DNA fragments recognised by the four DNA probes, had the same length as these probes. This demonstrated that the amplified fragments, used as probes, originated from the fragments detected on the blots.

E) Selective amplification of a single restriction fragment

[0055] Three sets of primers were designed for 3 corresponding random PstI-fragments from Tomato genomic DNA, of which the sequence next to the PstI-recognition sequence was known. Sets of primers with 5 selective nucleotides were made as shown below:

Primer set 1:

\[
\text{Sequence 1:} \quad 5'-\text{ctgca}G\text{CAT}C\text{GC}A\text{CAGC}-----\text{CGCGCAC}T\text{Gctgcag}-3' \\
5'-\text{TGC}G\text{TAAC}T\text{gcagCAGTA}-3' \quad 3'-\text{TGGACgacgtACATGCGT}-5' \\
\text{Primer 1.1} \quad \text{Primer 1.2}
\]

Primer set 2:

\[
\text{Sequence 2:} \quad 5'-\text{ctgca}G\text{CGAATCCTCT}-----\text{AGTGAGT}T\text{AGctgcag}-3' \\
5'-\text{TGC}GT\text{ACAtgcagCGGA}-3' \quad 3'-\text{CAATCgacgtACATGCGT}-5' \\
\text{Primer 2.1} \quad \text{Primer 2.2}
\]

Primer set 3:

\[
\text{Sequence 1:} \quad 5'-\text{ctgca}A\text{ATACCAAGA}-----\text{GCAAGC}C\text{AGCtgcag}-3' \\
5'-\text{TGC}G\text{TAACAtgcagCTATG}-3' \quad 3'-\text{GTGTCgacgtACATGCGT}-5' \\
\text{Primer 3.1} \quad \text{Primer 3.2}
\]

[0056] Tomato DNA was digested with PstI and adapters were ligated to the ends of the restriction fragments as described above. This DNA was used as template in PCR's with Primer sets 1 or 2 or 3, using the conditions as described in one of the previous sections. The reaction products of each PCR were analysed on a 1.0% agarose gel. This gel is shown in Figure 12. Figure 12 shows 13 lanes, of which lanes 1, 2, 12 and 13 are DNA markers. The sizes in kilobases of these markers are indicated at both sides of the gel. Lanes 3, 6 and 9 show plasmid DNA with each of the three PstI-fragments restricted with PstI, which yields the vector fragment, pUC18 (Yanisch-Perron et al., Gene 33, 103-119), and the inserted PstI-fragment. Lanes 4 and 5 show amplification with primer set 1 of 5 fg of the corresponding plasmid DNA and 1 ng of total genomic DNA respectively. Lanes 7 and 8 show amplification with primer set 2 of plasmid DNA and total genomic DNA, and lanes 10 and 11 show amplification with primer set 3. These results demonstrate that it is possible to amplify a single PstI-fragment out of a mixture of at least 20,000 fragments using the selective restriction fragment amplification technique with primers having 5 selective nucleotides.
F) Identification of DNA polymorphisms using SRFA

[0057] In the previous sections it was clearly demonstrated that with the selective restriction fragment amplification technique it is possible to amplify restriction fragments, either at random, or specific fragments, when sequence information is available. Hence, it should be possible to search for restriction site polymorphisms between two individuals of the same species. This is described below for two Tomato lines, which are very related but differ in the presence of the root knot nematode resistance gene, Mi, in one of the lines. This Mi-gene originates from Lycopersicon peruvianum, a species distantly related to the edible Tomato L.esculentum. It has been introduced into the L.esculentum line by crossing, and subsequent back crossing 12 times to the L.esculentum parent, and selecting the offspring for presence of the Mi-gene. Therefore, the two Tomato lines differ only in a small portion of their genetic material, i.e. the Mi-gene and surrounding region. The Mi-region was calculated to constitute < 1% of the genome of this line, using classical genetic methods.

[0058] DNA was isolated from the two Tomato lines (line 83M-71392, Mi-sensitive, and line 83M-71398, Mi-resistant, obtained from De Ruiter Seeds, Bleiswijk, The Netherlands) and subsequently restricted with PstI and provided with adapters as described above. A large number of amplification reactions were performed using primers, which differed in their extension of selective nucleotides. Three selective nucleotides were used, and apart from single primers also combinations of two different primers were used. The reactions were analysed on mixed polyacrylamide/agarose gels: 2.5% polyacrylamide and 1.0% agarose was used, with a ratio acrylamide to bisacrylamide of 20:1. Gels were run on a Protean II gel unit (Biorad), using spacers of 1.5 mm. A total of 16 different primers was used giving 16 reactions with a single primer, and 120 reactions with all possible combinations of two primers. A typical example of a gel with six of these combinations is shown in Figure 13. Lanes 1 and 14 of this gel contain DNA markers, of which the sizes in kilobases are indicated at the right side of the gel. Lanes 2 and 3, 4 and 5, 6 and 7 etc contain amplifications with a specific primer or primer pair of the two Tomato lines. The screening for restriction site polymorphisms yielded a number of fragments, three of which were very prominent and which are depicted in lanes 9, 11 and 12 of figure 13 (indicated by a small circle). The polymorphic bands in lanes 9 and 11 were expected to be the same, because the same primer was present in both reactions (the difference is the presence of a second primer in lane 11). The two polymorphic fragments of lanes 11 and 12 were cut out of the gel, the gel slices were crushed by forcing them through a 18 gauge needle and the DNA was eluted from the gel slices by elution through diffusion in 200 µl of 100 mM Tris.HCl pH 8.0, 10 mM EDTA. 2 µl was used for reamplification of these fragments as described above. 200 ng of each fragment was made blunt end using T4 DNA polymerase and subsequently ligated to 100 ng of plasmid vector pUC18 (Yanisch-Perron et al., Gene 33, 103-119) restricted with Smal. The ligation mixture was transformed to E.coli and for each fragment one recombinant E.coli clone was selected for sequence analysis. All these manipulations were performed using standard procedures as described by Sambrook, Fritsch and Maniatis in: Molecular Cloning, A Laboratory Manual (Cold Spring Harbor Laboratory Press, New York).

[0059] Two sets of primers with 6 selective nucleotides were synthesised based on the sequences of the two fragments as described above. We were able to amplify each fragment specifically using these primer sets. Fragments were only amplified from the Tomato line, from which they originated. Hence, these primer sets exhibited the same polymorphism, initially found with the primers with 3 selective nucleotides used to find this polymorphism.

EXAMPLE 2: SELECTIVE RESTRICTION FRAGMENT AMPLIFICATION OF TOMATO DNA WITH TWO RESTRICTION ENZYMES

[0060] In example 1 the principle of selective restriction fragment amplification (SRDA) is exemplified using Tomato DNA and the restriction enzyme PstI. In this example SRFA using two different restriction enzymes, PstI and MseI, will be illustrated.

Isolation and modification of the DNA

[0061] Total Tomato DNA was isolated from young leaves as described in example 1. Two pairs of so-called isogenic lines were used as source of the DNA, named GemR and GemS, and GCR26 and GCR151 respectively (These lines are described in the following references: Denby and Williams, [1962], Can. J. Plant Sci. 42, 681-685, Smith and Ritchie, [1983], Plant Mol. Biol. Rep. 1, 41-45). The two individuals of each pair of isogenic lines are genetically very similar, but differ in the presence of a trait conferring resistance to the fungal pathogen Verticillium albo-atratum.

[0062] The first step of the modification of the DNAs comprised the restriction of the DNAs with the two enzymes PstI and MseI. The restriction of the DNA, and also the subsequent ligation of the adapters to the DNA-fragments was carried out in the same buffer, which was named RL-buffer (restriction-igation buffer), and which contained: 10 mM Tris.HAc/10 mM MgAc/50 mM KAc/5 mM DTT, pH 7.5.
Restriction of the DNAs with PstI and Msel

[0063]

2.5 µg DNA  
12.5 units PstI (Pharmacia, 10 units/µl)  
12.5 units Msel (N.E. Biolabs, 4 units/µl)  
5 µl 10 x RL-buffer  
H2O to 50 µl

[0064] Incubation was carried out at 37°C for 1 hr.  
[0065] The next step in the modification of the DNAs was the ligation of adapter molecules to the ends of the DNA fragments. First appropriate double-stranded adapter molecules had to be prepared.

Preparation of adapters

[0066]

Msel-adapter: 5-GACGTGAGTCCCTGAG-3  
3-TACCTCAGGACTCAT-5

[0067] For preparation of a solution of 50 pMoles/µl of this adapter 8 µg (1430 pMoles) of the 16-mer 5-GACGTGAGTCCCTGAG-3 was mixed with 7 µg (1430 pMoles) of the 14-mer 5-TACCTCAGGACTCAT-3 in a total volume of 28.6 µl of H2O.

PstI-adapter: 5-bio-CTCGTAGACTGCGTACATGCA-3  
3-CATCTGACGCTGATGT-5

[0068] For preparation of a solution of 5 pMoles/µl of this adapter 5.25 µg (715 pMoles) of the biotinylated 21-mer 5-bio-CTCGTAGACTGCGTACATGCA-3 was mixed with 3.5 µg (715 pMoles) of the 14-mer 5-TGTACGCAGTCTAC-3 in a total volume of 143 µl of H2O.

Ligation of the adapter molecules

[0069] To the restricted DNA a mix of 10 µl was added containing:

1 µl PstI bio-adapter (= 5 pMol)  
1 µl Msel adapter (= 50 pMol)  
1.2 µl 10 mM ATP  
1 µl 10 x RL-buffer  
1 µunit T4 DNA ligase (Pharmacia, 5 units/µl)  
H2O to 10 µl

[0070] The resulting reaction mix of 60 µl was incubated for 3 hours at 37°C.  
[0071] The adapters were designed in such a way that the restriction sites were not restored after ligation. In this way fragment-to-fragment ligation was prevented, since fragment concatamers are restricted, because the restriction enzymes were still active during the ligation reaction. Adapter-to-adapter ligation was not possible because the adapters were not phosphorylated (see also example 1).

Selection of biotinylated DNA-fragments

[0072] Preparation of the template-DNAs for SRFA using two restriction enzymes generally involved an extra step
not used when using SRFA with a single enzyme. In this step the DNA-fragments to which a biotinylated adapter was ligated were separated from all other fragments.

[0073] Biotinylated fragments were separated from non-biotinylated fragments (MseI-MseI-fragments) in this step, by binding to paramagnetic streptavidine beads (Dynal). 10 µl beads were washed once in 100 µl STEX (100 mM NaCl/10 mM Tris.HCl/1 mM EDTA/0.1 % Triton X-100 pH 8.0), and resuspended in 140 µl STEX. The beads were subsequently added to the ligation mixture, to give a final volume of 200 µl. This was incubated for 30 minutes with gentle agitation at room temperature, to ensure proper binding of the biotinylated DNA-fragments to the beads. The beads were collected by holding the tubes containing the beads close to a magnet. This prevented the beads from being pipetted when the supernatant was transferred to another tube. The beads were washed once, and subsequently transferred to a fresh tube. Then the beads were washed 3 times with 200 µl STEX. Finally the beads were resuspended in 200 µl T01.E (10 mM Tris/0.1 mM EDTA, pH 8.0), and transferred to a fresh tube. The DNA was kept a 4°C.

[0074] The DNAs restricted with the restriction enzymes, provided with adapters, attached to the paramagnetic streptavidine beads and purified from the MseI-MseI fragments prepared as described above will be referred to as template-DNAs in the following steps.

Amplification of PstI-MseI fragments

[0075] The template-DNAs prepared as described above should contain all PstI-MseI fragments from the mentioned Tomato lines, and in addition a small amount of PstI-PstI-fragments with no internal MseI-fragments. In this experiment a number of these PstI-MseI fragments were visualised by amplification, essentially as described in example 1. Gel analyses of the amplification products was performed on denaturing acrylamide gels (Maxam and Gilbert, Proc. Natl. Acad. Sci. U.S.A. 74, 560-564), because the kind of fragments obtained by the procedure described in this example were much smaller than the ones described in example 1. In addition these types of gels allowed the separation of up to 100 bands per lane, which was about ten times more than the agarose gels described in example 1. The fragments were visualised by labeling one of the PCR-primers at the 5’end with (°32P)ATP and polynucleotide kinase.

Labeling of the PCR-primer

[0076] The primer selected for labeling was the 19-mer 5-GATGAGTCCTGAGTAAgaa-3 which was named MseI-primer-1, and in which the selective nucleotides are indicated with lower case letters. The labeling was performed in the following way:

- 3.0 µl 18-mer (from solution of 50 ng/µl = 150 ng)
- 5.0 µl (°32P)-ATP (from solution of 10 µCi/µl = 50 µCi)
- 3.0 µl 250 mM Tris.HCl/100 mM MgCl₂/50 mM DTT, pH 7.5
- 0.5 µl T4-kinase (Pharmacia 10 units/µl)
- 18.5 µl H2O

This gave a total volume of 30 µl, which was incubated at 37°C for 30 minutes. For each PCR 1 µl of this 5’labeled primer was added.

[0077] A total of 28 PCRs were performed, in which each of the 4 template-DNAs were amplified with 7 primer combinations. Each primer combination had the same MseI-primer (MseI-primer-1, described above), but varied in the choice of the PstI-primer. A total of 7 different primers were chosen (As with the MseI-primer the selective nucleotides are indicated with lower case letters):

- **PstI-primer-1**: 5-GACTGCATCACAGGaa-3
- **PstI-primer-2**: 5-GACTGCATCACAGGgt-3
- **PstI-primer-3**: 5-GACTGCATCACAGGg-3
- **PstI-primer-4**: 5-GACTGCATCACAGGag-3
All PCR-primers were dissolved in H₂O at a concentration of 50 ng/µl.

The amplification reaction

The PCR-mixture consisted of:

2.0 µl of template-DNA
1.0 µl of 5’labeled MseI-primer (5 ng)
0.5 µl unlabeled MseI-primer (25 ng)
0.6 µl PstI-primer (30 ng)
2.0 µl of 100 mM Tris.HCl/15 mM MgCl₂/500 mM KCl, pH 8.5
0.8 µl of 5 mM dNTPs
0.1 µl of Taq polymerase (Cetus Perkin Elmer, 5 units/µl)
13.0 µl of H₂O

All components of the reaction were added and mixed well, an essential component of the PCR, generally the enzyme, was added last. Subsequently the reaction was started as soon as possible.

The amplifications were performed on a Perkin Elmer 9600 thermal cycler. The cycle profile was as follows:

<table>
<thead>
<tr>
<th>Cycle</th>
<th>Denaturation</th>
<th>Annealing</th>
<th>Extension</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 cycle</td>
<td>30 sec at 94°C</td>
<td>30 sec at 65°C</td>
<td>60 sec at 72°C</td>
</tr>
<tr>
<td>11 cycles</td>
<td>Denaturation : 30 sec at 94°C</td>
<td>Annealing :</td>
<td>Extension :</td>
</tr>
<tr>
<td></td>
<td>lower annealing temperature 0.7°C each cycle,</td>
<td>60 sec at 72°C</td>
<td>60 sec at 72°C</td>
</tr>
<tr>
<td></td>
<td>64.3°C, 63.6°C, 62.9°C, 62.2°C, 61.5°C, 60.8°C, 60.1°C, 59.4°C, 58.7°C, 58.0°C, 57.3°C. Incubate for 30 seconds at each temperature</td>
<td></td>
<td></td>
</tr>
<tr>
<td>23 cycles</td>
<td>Denaturation : 30 sec at 94°C</td>
<td>Annealing : 30 sec at 56°C</td>
<td>Extension : 60 sec at 72°C</td>
</tr>
</tbody>
</table>

Gel analysis of amplified fragments

The reaction products were analyzed on 4.5% denaturing polyacrylamide gels. 50 x 38 cm gels were used, of which the gel cassettes to prepare these gels were purchased from Biorad. 100 ml of gel solution was used containing 4.5% w/v acrylamide/0.225% w/v bisacrylamide/7.5 M Urea/50 mM Tris/50 mM Boric acid/1 mM EDTA, pH 8.3. 100 ml gel solution was mixed with 500 µl 10% Ammonium persulfate and 100 µl TEMED immediately before casting the gel. A Tris/Boric acid/EDTA-buffer was used as electrophoresis buffer and contained: 100 mM Tris/100 mM Boric acid/2 mM EDTA, pH 8.3. The reaction mixtures were mixed with an equal volume (20 µl) of 98% formamide/10 mM EDTA/0.01% w/v bromo phenol blue/0.01% w/v xylene cyanol. The resulting mixtures were heated for 3 minutes at 95°C,
and then quickly cooled on ice. 2 µl of each sample was loaded on the gel. Gels were run at constant power of 110 Watts to give a constant heat development during electrophoresis. Under these conditions the field strength of the gels corresponded to 40 to 50 Volt/cm.

The results of the SRFA reactions are shown in figure 14. The lanes are numbered from 1 to 28, and contain each time the four Tomato lines with one of the 7 primer combinations. The order of the Tomato lines on the gel is: 1. GCR26, 2. GCR151, 3. GemR, 4. GemS. Lanes 1 to 4 contain these DNAs amplified with Msel-primer-1 and PstI-primer-1, lanes 5 to 8 contain these DNAs amplified with Msel-primer-1 and PstI-primer-2, lanes 9 to 12 contain these DNAs amplified with Msel-primer-1 and PstI-primer-3, lanes 13 to 16 contain these DNAs amplified with Msel-primer-1 and PstI-primer-4, lanes 17 to 20 contain these DNAs amplified with Msel-primer-1 and PstI-primer-5, lanes 21 to 24 contain these DNAs amplified with Msel-primer-1 and PstI-primer-6, and lanes 25 to 28 contain these DNAs amplified with Msel-primer-1 and PstI-primer-7. The gel contains no size markers but the DNA fragments visualised correspond with ±200 nucleotides at the bottom of the Figure to ±500 nucleotides at the top.

EXAMPLE 3: SELECTIVE RESTRICTION FRAGMENT AMPLIFICATION OF DNA OF VARIOUS LACTUCA SPECIES WITH TWO RESTRICTION ENZYMES

In example 2 the principle of selective restriction fragment (SRFA) amplification using two restriction enzymes is exemplified for Tomato DNA. In this example we will illustrate that similar results are obtained using DNAs of various Lactuca species using the same two restriction enzymes PstI and Msel.

Isolation and modification of the DNA

DNAs were isolated as described in example 1 using young leaf material of various Lactuca species. As indicated below these plants include a commercial lettuce (L.sativa) variety, and several individuals of two wild Lactuca species, L.saligna and L.virosa. The plants were arbitrarily designated the following names:

1. L.saligna, nr. 21, plant 1
2. L.saligna, nr. 21, plant 2
3. L.saligna, nr. 22, plant 1
4. L.saligna, nr. 22, plant 2
5. L.virosa, nr. 01, plant 1
6. L.virosa, nr. 01, plant 2
7. L.virosa, nr. 02
8. L.virosa, nr. 03, plant 1
9. L.virosa, nr. 03, plant 2
10. L.sativa, a commercial butterhead variety

The genetic material analysed thus represented 6 different plant types, including two different individuals of 4 of these plants.

Modification of the Lactuca DNAs to generate the templates for the SRFA was performed identical to the procedure described in example 2.

Amplification of PstI-Msel fragments

The DNAs prepared as described above were used as templates for SRFA reactions. Two primer combinations were used employing a single Msel-primer and two different PstI-primers. These primers (selective nucleotides depicted in lower case letters) were:

Msel-primer: 5-GATGAGTCCTGAGTAAac-3
PstI-primer-1: 5-GACTGCCTACATGCAGa-3
PstI-primer-2: 5-GACTGCGTACATGCAca-3

Amplification of PstI-Msel fragments using the primers depicted above was carried out exactly as described in example 2, and the generated fragments were visualised on denaturing polyacrylamide gels as described in example
2. The band patterns obtained are shown in Figure 15. Lanes 1 to 10 show DNAs 1 to 10 amplified with the MseI-primer in combination with PstI-primer-I, lanes 11 to 20 show DNAs 1 to 10 amplified with the MseI-primer in combination with the PstI-primer 2. Size markers (not visible in this Figure) in nucleotides are indicated to the right of the gel. The differences in band patterns reflects the differences in relatedness of the various plants.

EXAMPLE 4: SELECTIVE RESTRICTION FRAGMENT AMPLIFICATION OF CORN INBRED LINES WITH A VARIETY OF RESTRICTION ENZYME COMBINATIONS

[0090] In example 2 and 3 the principle of selective restriction fragment (SRFA) amplification using two restriction enzymes is exemplified using Tomato DNA and Lettuce (Lactuca species) DNAs respectively. In this example it will be illustrated that similar results are obtained with Corn (Zea mais) lines. In addition it will be illustrated that a variety of restriction enzyme combinations can be used to obtain DNA fingerprints of in this case Corn lines.

Isolation and modification of the DNA

[0091] Two corn inbred lines were used, named 1 and 2. The source of these lines is irrelevant, because in our experience any selected line gave good DNA fingerprints using SRFA. DNA of these lines was isolated from young leaf material as described by Saghai-Mahoot et al. (1984), Proc. Natl. Acad. Sci. U.S.A. 81, 8014-8018. The following restriction enzyme combinations (EKs) were used to make the template-DNAs: PstI/TaqI, EcoRl/TaqI, AseI/TaqI, Sse8387-I/TaqI. All enzymes were purchased from Pharmacia, except AseI which was purchased from New England Biolabs, and Sse8387-I which was purchased from Amersham. Template DNAs were prepared essentially as described in examples 2 and 3, with the following exceptions:

[0092] Restriction of the DNA was performed by first incubating with TaqI at 65°C for one hour, and subsequently incubating with the second enzyme, PstI, AseI, EcoRl or Sse8387-I, for an additional hour at 37°C. Ligation of adapters was as described in example 2 using the following adapters:

**TagI-adapter:**

5-GACGATGAGTCTCTGAC-3  
3-TACTCAGGACTGGC-5

**PstI & Sse8387-I-adapter:**

5-bio-CTCGTAGACTGAGGTAC-3  
3-CATCTGACGCATGT-5

**AseI-adapter:**

5-bio-CTCGTAGACTGAGTACC-3  
3-CTGACGCATGGAT-5

**EcoRI-adapter:**

5-bio-CTCGTAGACTGACGTACC-3  
3-CTGACGCATGGTTAA-5

Amplification of restriction fragments

[0093] Amplification of restriction fragments was performed as described in example 2. The primers selected for labeling of the amplification products were the following TagI-primers having 3 selective nucleotides (indicated by lower case letters):
These 4 primers were used for detection of amplification products with all four enzyme combinations. For each enzyme combination 4 primers for the other enzyme were selected to give a total of 16 combinations for each enzyme. These primers are indicated below (selective nucleotides shown in lower case letters). For EcoRI and AseI primers with 3 selective nucleotides were selected, for PstI primers with 2 selective nucleotides were chosen, and for SseI primers with a single selected nucleotide were chosen. For enzymes cutting less frequently in the Corn genomic DNA, primers were selected containing extensions with fewer selective nucleotides.

EcoRI-primers:  
1. 5-CTGCCTAACATATCaa-3  
2. 5-CTGCCTAACATATCaca-3  
3. 5-CTGCCTAACATATCaac-3  
4. 5-CTGCCTAACATATCcag-3

AseI-primers:  
1. 5-GACTGCGTACATGAc-3  
2. 5-GACTGCGTACATGAcg-3  
3. 5-GACTGCGTACATGAcac-3  
4. 5-GACTGCGTACATGAcacc-3

PstI-primers:  
1. 5-GACTGCGTACATGAcac-3  
2. 5-GACTGCGTACATGAcgac-3  
3. 5-GACTGCGTACATGAcgcc-3  
4. 5-GACTGCGTACATGAcggt-3

Sse8387-I-primers:  
1. 5-GACTGCGTACATGAcgca-3  
2. 5-GACTGCGTACATGAcgcc-3  
3. 5-GACTGCGTACATGAcggc-3  
4. 5-GACTGCGTACATGAcggt-3

A total of 128 PCRs were performed (2 DNAs x 4 enzyme combinations x 16 primer combinations), following the protocol described in example 2. The reaction products of these PCRs were analysed on 3 gels (containing 48 lanes/gel) as described in example 2. All primer combination gave DNA fingerprints of 50 to 100 bands per lane, except for the combination SseI/TagI, which gave only 10 to 15 bands per lane. An example of one of the gels is shown in Figure 16. This Figure shows part of the gel with the analysis of DNA fingerprints obtained with the enzyme combinations PstI/TagI and EcoRI/TagI. Lanes 1 to 8 show DNA-fingerprints of the two Corn DNAs obtained by SRFA with TagI.
EXAMPLE 5: SELECTIVE RESTRICTION FRAGMENT OF BACTERIAL DNAs

[0096] In example 2, 3 and 4 the principle of selective restriction fragment (SRFA) amplification using two restriction enzymes is exemplified for Tomato, Lettuce (Lactuca species) and Corn DNAs respectively. In this example it will be illustrated that this technique can also be used to characterise bacterial DNAs. A number of Xanthomonas campestris strains were obtained from the Laboratory of Microbiology in Gent, Belgium, to illustrate the usability of the technique in bacteria.

Isolation and modification of the DNA

[0097] All DNAs were prepared from Xanthomonas campestris strains isolated from a variety of origins, mostly from infected plants. These strains, numbered 1 to 26 are listed below, and may be obtained from the Laboratory of Microbiology in Ghent, Belgium.

<table>
<thead>
<tr>
<th>DNA subspecies</th>
<th>pathovar</th>
<th>isolate</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. albilineans</td>
<td></td>
<td>494</td>
</tr>
<tr>
<td>2. fragariae</td>
<td></td>
<td>708</td>
</tr>
<tr>
<td>3. oryzae</td>
<td>oryzae</td>
<td>5047</td>
</tr>
<tr>
<td>4. oryzae</td>
<td>populi</td>
<td>5743</td>
</tr>
<tr>
<td>5. maltophilia</td>
<td></td>
<td>958</td>
</tr>
<tr>
<td>6. campestris</td>
<td>campestris</td>
<td>568</td>
</tr>
<tr>
<td>7. campestris</td>
<td>alfalfa</td>
<td>497</td>
</tr>
<tr>
<td>8. campestris</td>
<td>coracanae</td>
<td>686</td>
</tr>
<tr>
<td>9. campestris</td>
<td>citri</td>
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</tr>
<tr>
<td>10. campestris</td>
<td>citri</td>
<td>9658</td>
</tr>
<tr>
<td>11. campestris</td>
<td>citri</td>
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</tr>
<tr>
<td>12. campestris</td>
<td>citri</td>
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<tr>
<td>13. campestris</td>
<td>citri</td>
<td>8654</td>
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<tr>
<td>14. campestris</td>
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<tr>
<td>15. campestris</td>
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</tr>
<tr>
<td>26. campestris</td>
<td>citri</td>
<td>9160</td>
</tr>
</tbody>
</table>

[0098] DNA of these bacterial strains was isolated as described by Marmur (J. Mol. Biol. 3, 208-218). The DNAs were restricted essentially as described in example 4, with the exception that TaqI and ApaI were chosen as restriction enzymes. Ligation of adapters was as described in example 4 using the following adapters:
Amplification of restriction fragments

Amplification of restriction fragments was performed as described in example 2. The primers selected for SRFA were the TaqI-primer 5'-CGATGAGTCCTGACGGAg-3' (having one selective nucleotide indicated in lower case letter), and the Apal-primer 5'-GACTGCGTACAGGCCCg-3' (having one selective nucleotide indicated in lower case letter). The Apal-primer was labeled at the 5'end for detection of the amplified fragments as described in example 2.

Each of the 26 DNAs was amplified using the primer set described above. Amplification conditions were as described in example 2, except that the last 9 cycles of the PCR were omitted, because of the lower complexity of the DNAs compared to the plant DNA in examples 2, 3 and 4.

The DNA fingerprints obtained with the bacterial DNAs as described in this example are shown in Figure 17. Lanes 1 to 26 represent bacterial DNAs 1 to 26. The sizes of marker DNAs (not visible on the gel) in nucleotides are indicated to the right of the gel. This figures shows clearly that the relatedness of the bacterial strains is reflected by the similarity of the band patterns.

EXAMPLE 6: SELECTIVE RESTRICTION FRAGMENT AMPLIFICATIONS OF DNA OF VARIOUS ANIMALS WITH TWO RESTRICTION ENZYMES

In the previous examples selective restriction fragment amplification (SRFA) was exemplified for plant DNA of various sources. Here we illustrate the efficacy of the procedure using random samples of DNA obtained from different domestic animals. The animal species tested are: Gallus domesticus (chicken); Sus scrofa domestica L. (pig); Bos taurus (cow); Equus caballus (horse). Restriction enzymes used are Sse8387I and Msel.

Isolation and modification of the DNA

DNAs were isolated from blood samples following procedures described by Maniatis et al. (1982). DNA samples 1 to 3 (chicken), 4 to 7 (pig), 8 to 11 (cow) and 12 to 15 (horse) were digested by restriction enzymes Sse8387I and Msel. The DNA fragments were ligated to adapters as described in example 2. Since the restriction enzymes Sse8387I and PstI generate compatible 3' overhangs we could use the PstI- and Msel-adapter described in example 2.

Amplification of restriction fragments

Template DNAs named above and prepared as described in example 2 served as templates in SRFA reactions. The primer combinations used consisted of a single Msel-primer and different SseI-primers:

**Msel-primer:** 5'-GATGAGTCCTGAGTAAAtac-3

**Sse8387I-primer-1:** 5'-GACTGCATGACAGGaa-3

**Sse8387I-primer-2:** 5'-GACTGCATGACAGGag-3

Amplification of Sse8387I-Msel fragments using primer pairs described above was carried out using the protocol described in example 2. Reaction products were run on denaturing polyacrylamide gels also described in
example 2. An autoradiograph showing fingerprints of the above samples is shown in Figure 18. Lanes 1 through 15 show fingerprints of DNAs 1 to 15 amplified with the Msel-primer paired with Sse83871-primer-1, lanes 16 through 30 show similar patterns obtained with the Msel-primer combined with Sse83871-primer-2. Differences in fingerprints between animals of one species reflect heterogeneity in animal populations; overall-patterns are characteristic for a specific species.

The invention relates to a Process for the amplification by Polymerase Chain Reaction of at least one restriction fragment obtained from a starting DNA which contains a plurality of restriction sites for at least one specific determined Restriction endonuclease, which process comprises:

(a) digesting said starting DNA with at least one restriction endonuclease, to fragment it into restriction fragments;
(b) ligating the restriction fragments obtained with at least one double-stranded synthetic oligonucleotide adaptor having one end which is compatible to be ligated to one or both of the ends of the restriction fragments to thereby produce tagged restriction fragments;
(c) contacting said tagged restriction fragments under hybridizing conditions with at least one oligonucleotide primer;
(d) amplifying said tagged restriction fragments hybridized with said primer or primers in the presence of the required nucleotides and DNA polymerase or elongating the hybridized primers along those tagged restriction fragments, and
(e) identifying or recovering amplified or elongated DNA fragments as produced in step (d).

In a particular embodiment of this process, the terminal nucleotide of at least one of said primers in the direction of the elongation sought corresponds to the last of the nucleotides involved in the restriction site for said specific endonuclease, and which process comprises identifying or recovering the restriction fragments of said starting DNA which have been amplified.

In another particular embodiment of this process, at least one of said primers includes a selected sequence comprising a determined number (one or several nucleotides) extending beyond the last of the nucleotides involved in the restriction site for said specific endonuclease in the direction of its own elongation within the corresponding restriction fragments during the amplification step.

In a specific embodiment of the above-described process, double-stranded DNA-linker contains several sites for different specific endonucleases which are all distinct from one another, which processes comprise repeating, on a same starting DNA the steps of the process defined above with one of these restriction endonucleases yet with another of said distinct specific endonucleases and upon using primers whose nucleotide sequences are selected as defined in the above description, yet with respect to said other specific endonuclease.

The process described above or of the oligonucleotide of the invention, is appropriate, for the identification of polymorphisms in determined DNAs originating from the same live species, e.g. genomic DNAs of a microbial, plant or animal, including humans, or of fragments thereof, either among themselves or relative to a corresponding determined DNA standard, which use comprises subjecting the DNAs under study to the process or to the contact of the oligonucleotide in conditions allowing an amplification or elongation reaction, comparing the restriction patterns obtained starting from each of said DNAs and, optionally, of said standard DNA and relating the existence and, where appropriate, the localization of that DNA polymorphism to the differences observed between the sizes of the restriction fragments of the different DNAs.

The invention also relates to a fragmented DNA whose different fragments have sequences which all correspond to initial digests of the unfragmented starting DNA from which they are produced with a same determined specific endonuclease, characterized in that all of said fragments were tagged at their 5’ and 3’ ends respectively by determined 3’ and 5’ adaptors corresponding to the cleaved part of a same starting DNA linker which initially included a single restriction site for said specific endonuclease, and optionally prolonged with determined constant sequences. The fragmented DNA can be in the form of a pattern of migration bands on a suitable support, e.g. gel support, in which its fragments had initially been caused to migrate under the influence of an electric field.

The fragmented DNA can also comprise end portions including oligonucleotide characterized by the following composition, starting from the 5’ end:

(i) a nucleotide sequence (constant sequence) of at least 10 bases, but not longer than 30 bases, complementary
The invention further relates to a kit for the fragmentation of determined DNAs by at least one specific restriction endonuclease into fragments and analysis of these fragments which comprises:

- the specific restriction endonuclease;
- a double-stranded DNA oligonucleotide linker including itself a single site within its own nucleotidic sequence for said specific endonuclease to thereby cleave said linker in corresponding 5' and 3' adaptors respectively, wherein said double-stranded DNA linker had a sufficient size to provide 5' and 3' parts which may subsequently provide templates for the PCR primers of this kit;
- PCR primers which respectively comprise, on the one hand the same sequences as the strands of the 5' and 3' adaptors complementary to the strands subsequently acting as templates for said primers wherein said primers further include the nucleotides complementary to those which are involved in the formation of the site for said determined specific restriction endonuclease in the template strands;
- if appropriate, oligonucleotide segments of determined (constant) sequences for generating sites of sufficient length for hybridization with said primers, for the elongation of the 5' ends of said 5' adaptors or the 3' ends of said 3' adaptors or both, prior to digestion of said linker by said specific restriction endonuclease to produce said 5' and 3' adaptors respectively, or alternatively for the elongation of the tagged fragments obtained subsequent to the ligation of said 5' and 3' adaptors to the extremities of the fragments of the starting DNA;
- optionally a fragmented DNA standard corresponding to the determined DNA subject to a fragmentation study, whereby the fragments of said DNA standard were obtained by digesting it with said specific endonuclease.

A particular embodiment of this kit is such that said oligonucleotide segments for the elongation of both said 5' and 3' adaptors or 5' and 3' ends of the tagged DNA fragments, have identical nucleotide sequences.

In another embodiment, the linker of the kit contains several respective unique sites for specific endonucleases all different from one another, said kit further including primers corresponding to each of the 3' and 5' adaptors formed by cleavage of said linker with said different specific endonucleases respectively, wherein said primers are respectively as defined in claim 8, in respect of the 3' and 5' adaptors which are produced in said linker by cleavage thereof by each of said specific endonucleases.

Also in a particular embodiment the kit can comprise fragmented DNA standards as defined above in respect of the corresponding specific restriction endonucleases, wherein each of said fragmented DNA standards is in respect of each of the determined specific restriction enzymes.

Claims

1. Process for the amplification by Polymerase Chain Reaction of at least one restriction fragment obtained from a starting DNA, which contains a plurality of restriction sites for at least one specific determined restriction endonuclease which process comprises:

(a) digesting said starting DNA with at least one restriction endonuclease, to fragment it into restriction fragments;
(b) ligating the restriction fragments obtained with at least one double-stranded synthetic oligonucleotide adaptor having one end which is compatible to be ligated to one or both of the ends of the restriction fragments to thereby produce tagged restriction fragments;
(c) contacting said tagged restriction fragments under hybridizing conditions with at least one oligonucleotide primer;
wherein said primer or primers include a nucleotide sequence having the same nucleotide sequence as the terminal parts of the strands at the ends of said tagged restriction fragments, including the nucleotides involved in the formation of the target sequence for said restriction endonuclease and including at least part of the nucleotides present in the ligated adaptors, wherein at least one of said primers includes at its 3' end, a selected sequence comprising at least one nucleotide located immediately adjacent to the nucleotides involved in the formation of the target sequence for said restriction endonuclease
(d) amplifying said tagged restriction fragments hybridized with said primer or primers in the presence of the
required nucleotides and DNA polymerase or elongating the hybridized primers along those tagged restriction fragments, and
(e) identifying or recovering amplified or elongated DNA fragments as produced in step (d).

2. Process according to claim 1 wherein in step (c), the primer or primers perfectly base pair with a constant DNA sequence composed of part of the target sequence of the restriction endonuclease and of the sequence of the adaptor.

3. A process of claim 1, wherein at least one primer comprises a selected sequence of 1 to 10 nucleotide(s).

4. A process according to anyone of claims 1 to 3, wherein the primers comprise 10 to 50 nucleotides, and wherein the target sequence of the restriction endonuclease comprises 4 to 8 nucleotides.

5. A process according to anyone of claims 1 to 4, wherein the nucleotide sequence of the primers base pairing with the adaptor comprises from 10 to 30 nucleotides.

6. A process according to anyone of claims 1 to 5, wherein the adaptors are designed in such a way that the corresponding target sequence of the endonuclease is not restored when the adaptor hybridizes with the tagged restriction fragment(s) and, wherein the primer or primers are oligonucleotides comprising a constant sequence perfectly base pairing to at least one end of the sequence of at least one tagged restriction fragment from a mixture, which constant sequence comprises nucleotides base pairing to part of the target sequence for said restriction endonuclease, and at least one of the said primers includes adjacent to the 3' end of the nucleotides involved in the formation of said target sequence, 1 to 10 additional selected nucleotides.

7. A process according to anyone of claims 1 to 6 wherein the nucleic acid sequence of the starting DNA is at least in part unknown.

8. A process according to anyone of claims 1 to 7, wherein two different restriction endonucleases are used.

9. A process according to anyone of claims 1 to 8, wherein all the primers have the same constant nucleotide sequence.

10. A process according to anyone of claims 1 to 8, wherein a mixture of different primers is used.

11. A process according to anyone of claims 1 to 8, wherein at least one primer contains 1, 2, or 3, selected nucleotides at their 3' end.

12. A process according to anyone of claims 1 to 10, wherein 2 different restriction endonucleases are used in step (a) and wherein at least one double-stranded synthetic oligonucleotide adaptor for one of the endonucleases is biotinylated.

13. A process according to anyone of claims 1 to 12 wherein at least one primer is labelled.

14. A process according to anyone of claims 1 to 13 wherein the number of selective nucleotides is chosen so that the number of restriction fragments which will be amplified is limited to 5 to 200.

15. A process for the preparation of a set of amplified restriction fragments comprising:
   a) amplifying a subset of restriction fragments from a mixture of generated restriction fragments obtained from a starting DNA using a process according to anyone of claims 1 to 14,
   b) separating the amplified tagged restriction fragments.
   c) identifying the amplified restriction fragments.

16. A process according to anyone of claims 1 to 14 wherein said starting DNA to be amplified is genomic DNA from a human or animal biological sample, especially a tissue or a blood sample.

17. A process according to anyone of claims 1 to 14 wherein said starting DNA to be amplified is genomic DNA from plant, or plant tissue.
18. A process according to anyone of claims 1 to 14 wherein said starting DNA to be amplified is DNA of a microorganism.

19. Process according to anyone of claims 1 to 15 for the identification of polymorphisms between different starting DNAs originating from the same live species, which comprises identifying differences observed between the amplified restriction fragments of the different starting DNAs.

20. Process of anyone of claims 1 to 14 for the identification of polymorphisms in genomic DNAs of a microorganism, plant or animal, including humans, or of fragments thereof, either among themselves or relative to a corresponding determined DNA standard.

21. Process as claimed in anyone of claims 1 to 14, for identifying DNA polymorphisms associated with genetically inherited traits in humans, DNA polymorphisms associated with genetically inherited traits in animals, or for identifying DNA polymorphisms associated with genetically inherited traits in plants, and for identifying DNA markers based on such DNA polymorphisms.

22. Process according to anyone of claims 1 to 14, further comprising comparing two or more amplified or elongated DNA fragments identified or recovered in step (e) of said process for detecting similarities between plant or animal varieties, species, cultivars, microorganisms, or for evaluating genetic distances and characterizing such plant or animal varieties, species, cultivars, microorganisms.

23. Process according to anyone of claims 1 to 14, wherein the amplified or elongated DNA fragments as produced in step (e) are identified or recovered as DNA fingerprints.

24. A process for identification of DNA markers linked to a genetic trait, said process comprising identifying polymorphisms according to claim 22, between starting DNAs originating from the same live species which exhibit differences in said genetic trait and correlating said polymorphisms with the phenotype exhibited by said genetic trait.

25. The process according to claims 1 to 14 further comprising the steps of:

(f) isolating at least one DNA fragment identified or recovered in step (e);
(g) determining the nucleotide sequence of the first 8-10 nucleotide residues internally adjacent to the target sequence of the restriction endonuclease at both ends of said DNA fragments;
(h) designing oligonucleotide primers having a nucleotide sequence according to the primers of step (c) of claim 1 wherein the selected nucleotide sequence comprises 5 to 10 nucleotide residues which correspond to the first 8-12 nucleotide residues internally adjacent to the restriction sites at both ends of said DNA fragment.

Patentansprüche

1. Verfahren zur Amplifizierung mit Polymerase Kettereaktion von mindestens einem Restriktionsfragment, erhalten von einer Ausgangs-DNA, daß eine Pluralität Restriktion Stätten umfäßt mit einem spezifisch Restriktionendonuklease bestimmt, wobei das Verfahren umfaßt:

a) die Ausgangs-DNA mit mindestens einer Restriktionsendonuklease zu verdauen, um sie in Restriktionsfragmente zu fragmentieren,
b) die erhaltenen Restriktionsfragmente mit mindestens einem doppelsträngigen synthetischen Oligonukleotid-Adaptor, der ein Ende aufweist, das kompatibel ist, mit einem oder beiden der Enden der Restriktionsfragmente ligiert zu werden, zu ligieren, um dadurch mit einem "Tag" versehene Restriktionsfragmente zu erzeugen,
c) die mit einem "Tag" versehenen Restriktionsfragmente unter Hybridisierungsbedingungen mit mindestens einem Oligonukleotidprimer in Berührung zu bringen, wobei der oder die Primer eine Nukleotidsequenz umfassen, mit der gleichen Nukleotidsequenz wie die terminalen Teile der Stränge an den Enden der mit einem "Tag" versehenen Restriktionsfragmente einschließlich der Nukleotide, die an der Bildung der Zielsequenz für die Restriktionsendonuklease beteiligt sind, und einschließlich mindestens eines Teils der in den ligierten Adapten vorliegenden Nukleotiden umfassen, wobei mindestens einer der Primer an seinem 3'-Ende eine ausgewählte, mindestens ein Nukleotid umfassende Sequenz, die sich unmittelbar benachbart zu den an der Bildung der Zielsequenz für die Restriktionsendonuklease beteiligten Nukleotiden befindet, umfasst
d) die mit dem Primer oder den Primern hybridisierten, mit einem "Tag" versehenen Restriktionsfragmente in Gegenwart der erforderlichen Nukleotide und von DNA-Polymerase zu amplifizieren oder die hybridisierten Primer entlang jenen mit einem "Tag" versehenen Restriktionsfragmenten zu verlängern und
e) amplifizierte oder verlängerte DNA-Fragmente, die in Schritt (d) erzeugt worden sind, zu identifizieren oder zu gewinnen.

2. Verfahren nach Anspruch 1, wobei in Schritt (c) der oder die Primer mit einer konstanten DNA-Sequenz, gebildet aus einem Teil der Zielsequenz der Restriktionsendonuklease und der Sequenz des Adaptors, eine perfekte Basenpaarung ausbilden.

3. Verfahren nach Anspruch 1, wobei mindestens ein Primer eine ausgewählte Sequenz von 1 bis 10 Nukleotid(en) umfaßt.

4. Verfahren nach einem der Ansprüche 1 bis 3, wobei die Primer 10 bis 50 Nukleotide umfassen und wobei die Zielsequenz der Restriktionsendonuklease 4 bis 8 Nukleotide umfaßt.

5. Verfahren nach einem der Ansprüche 1 bis 4, wobei die Nukleotidsequenz der Primer, die eine Basenpaarung mit dem Adaptor ausbildet, 10 bis 30 Nukleotide umfaßt.

6. Verfahren nach einem der Ansprüche 1 bis 5, wobei die Adaptoren so gestaltet sind, daß die entsprechende Zielsequenz der Endonuklease nicht wiederrhergestellt wird, wenn der Adaptor mit dem oder den mit einem "Tag" versehenen Restriktionsfragmenten hybridisiert und wobei der oder die Primer Oligonukleotide sind, die eine konstante Sequenz umfassen, die eine perfekte Basenpaarung mit mindestens einem Ende der Sequenz von mindestens einem mit einem "Tag" versehenen Restriktionsfragment aus einer Mischung ausbildet, welche konstante Sequenz Nukleotide umfaßt, die eine Basenpaarung zu einem Teil der Zielsequenz für die Restriktionsendonuklease ausbilden, und mindestens einer der Primer angrenzend an das 3'-Ende der an der Bildung der Zielsequenz beteiligten Nukleotide 1 bis 10 zusätzliche ausgewählte Nukleotide umfaßt.

7. Verfahren nach einem der Ansprüche 1 bis 6, wobei die Nukleinsäuresequenz der Ausgangs-DNA mindestens teilweise unbekannt ist.

8. Verfahren nach einem der Ansprüche 1 bis 7, wobei zwei unterschiedliche Restriktionsendonukleasen verwendet werden.

9. Verfahren nach einem der Ansprüche 1 bis 8, wobei alle Primer die gleiche konstante Nukleotidsequenz aufweisen.

10. Verfahren nach einem der Ansprüche 1 bis 8, wobei eine Mischung unterschiedlicher Primer verwendet wird.

11. Verfahren nach einem der Ansprüche 1 bis 8, wobei mindestens ein Primer 1, 2 oder 3 ausgewählte Nukleotide an ihrem 3'-Ende enthält

12. Verfahren nach einem der Ansprüche 1 bis 10, wobei zwei unterschiedliche Restriktionsendonukleasen in Schritt (a) verwendet werden und wobei mindestens ein doppelsträngiger synthetischer Oligonukleotidadaptor für eine der Endonukleasen biotinyliert ist.

13. Verfahren nach einem der Ansprüche 1 bis 12, wobei mindestens ein Primer markiert ist.

14. Verfahren nach einem der Ansprüche 1 bis 13, wobei die Anzahl von selektiven Nukleotiden so ausgewählt wird, daß die Anzahl von Restriktionsfragmenten, die amplifiziert werden, auf 5 bis 200 begrenzt ist.

15. Verfahren zur Herstellung eines Satzes von amplifizierten Restriktionsfragmenten, umfassend:

   a) eine Untergruppe von Restriktionsfragmenten aus einer Mischung von erzeugten Restriktionsfragmenten, erhalten von einer Ausgangs-DNA unter Verwendung eines Verfahrens nach einem der Ansprüche 1 bis 14, zu amplifizieren,
   b) die amplifizierten, mit einem "Tag" versehenen Restriktionsfragmente aufzutrennen,
   c) die amplifizierten Restriktionsfragmente zu identifizieren.
16. Verfahren nach einem der Ansprüche 1 bis 14, wobei die zu amplifizierende Ausgangs-DNA genomische DNA aus einer menschlichen oder tierischen biologischen Probe, speziell einem Gewebe oder einer Blutprobe, ist.

17. Verfahren nach einem der Ansprüche 1 bis 14, wobei die zu amplifizierende Ausgangs-DNA genomische DNA aus einer Pflanze oder einem Pflanzengewebe ist.

18. Verfahren nach einem der Ansprüche 1 bis 14, wobei die zu amplifizierende Ausgangs-DNA DNA eines Mikroorganismus ist.

19. Verfahren nach einem der Ansprüche 1 bis 15 zur Identifizierung von Polymorphismen zwischen verschiedenen Ausgangs-DNAs, die von der gleichen lebenden Spezies stammen, welches umfaßt, Unterschiede, die zwischen den amplifizierten Restriktionsfragmenten der verschiedenen Ausgangs-DNAs beobachtet werden, zu identifizieren.

20. Verfahren nach einem der Ansprüche 1 bis 14 zur Identifizierung von Polymorphismen in genomischen DNAs eines Mikroorganismus, einer Pflanze oder eines Tiers, einschließlich eines Menschen, oder von Fragmenten davon entwederuntiereinander oder in Bezug auf einen entsprechend bestimmten DNA-Standard.


22. Verfahren nach einem der Ansprüche 1 bis 14, welches ferner umfaßt, zwei oder mehr amplifizierte oder verlängerte DNA-Fragmente, die in Schritt (e) des Verfahren identifiziert oder gewonnen wurden, zum Nachweisen von Ähnlichkeiten zwischen Pflanzensorten oder Tierarten, -Spezies, -Kultivaren, Mikroorganismen oder zum Auswerten von genetischen Abständen und Charakterisieren solcher Pflanzensorten oder Tierarten, -Spezies, -Kultivare, Mikroorganismen.

23. Verfahren nach einem der Ansprüche 1 bis 14, wobei die amplifizierten oder verlängerten DNA-Fragmente wie erzeugt in Schritt (e) als DNA-Fingerprints identifiziert oder gewonnen werden.

24. Verfahren zur Identifizierung von DNA-Markern, die mit einem genetischen Merkmal assoziiert sind, wobei das Verfahren umfaßt, Polymorphismen zwischen Ausgangs-DNAs, die von der gleichen lebenden Spezies stammen und Unterschiede in diesem genetischen Merkmal aufweisen, gemäß Anspruch 22 zu identifizieren und diese Polymorphismen mit dem von diesem genetischen Merkmal bewirkten Phänotyp zu korrelieren.

25. Verfahren nach den Ansprüchen 1 bis 14, umfassend ferner die Schritte:

   (f) mindestens ein in Schritt (e) identifiziertes oder gewonnenes DNA-Fragment zu isolieren,
   (g) die Nukleotidsequenz der ersten 8 bis 10 Nukleotidreste, die im Inneren an die Zielsequenz der Restriktionsendonuklease an beiden Enden der DNA-Fragmente angrenzen, zu bestimmen,
   (h) Oligonukleotidprimer, die eine Nukleotidsequenz gemäß den Primern von Schritt (c) von Anspruch 1 aufweisen, zu gestalten, wobei die ausgewählte Nukleotidsequenz 5 bis 10 Nukleotidreste umfaßt, die den ersten 8 bis 12 Nukleotidresten, die im Inneren and die Restriktionsstellen an beiden Enden des DNA-Fragments angrenzen, entsprechen.

Revendications

1. Procédé pour l’amplification par une réaction de polymérisation en chaîne, d’au moins un fragment de restriction obtenu à partir d’un ADN de départ, qui contient une pluralité de sites de restriction pour au moins une endonucléase de restriction spécifique déterminée, le procédé comprenant :

   (a) la digestion dudit ADN de départ avec au moins une endonucléase de restriction, pour le fragmenter en des fragments de restriction ;
   (b) la ligation des fragments de restriction obtenus avec au moins un oligonucléotide adaptateur synthétique double brin ayant une extrémité qui est compatible pour être ligaturée à une ou deux des extrémités des
fragments de restriction produisant par ce moyen des fragments de restriction repérés ;
(c) la mise en contact desdits fragments de restriction repérés dans des conditions d'hybridation avec au moins une amorce oligonucléotidique ; dans lequel la ou lesdites amorces incluent une séquence de nucléotides ayant la même séquence de nucléotides que les parties terminales des brins aux extrémités desdits fragments de restriction repérés, incluant les nucléotides impliqués dans la formation de la séquence cible pour ladite endonucléase de restriction et incluant au moins une partie des nucléotides présents dans les adaptateurs ligaturés, dans lequel au moins une desdites amorces inclut à son extrémité 3’, une séquence sélectionnée comprenant au moins un nucléotide situé immédiatement adjacent aux nucléotides impliqués dans la formation de la séquence cible pour ladite endonucléase de restriction.
(d) l'amplification desdits fragments de restriction repérés hybridés avec la ou lesdites amorces en présence des nucléotides et de l'ADN polymérase requis ou l'allongement des amorces hybridées le long de ces fragments de restriction repérés, et
(e) l'identification ou la récupération des fragments amplifiés ou allongés d'ADN comme produits dans l'étape (d).

2. Procédé selon la revendication 1, dans lequel dans l'étape (c), la ou les amorces effectuent des appariements de bases parfaits avec une séquence d'ADN constante composée d'une partie de la séquence cible de l'endonucléase de restriction et de la séquence de l'adaptateur.

3. Procédé selon la revendication 1, dans lequel au moins une amorce comprend une séquence sélectionnée de 1 à 10 nucléotides.

4. Procédé selon l'une quelconque des revendications 1 à 3, dans lequel les amorces comprennent de 10 à 50 nucléotides, et dans lequel la séquence cible de l'endonucléase de restriction comprend 4 à 8 nucléotides.

5. Procédé selon l'une quelconque des revendications 1 à 4, dans lequel la séquence de nucléotides des amorces réalisant un appariement de bases avec l'adaptateur, comprend de 10 à 30 nucléotides.

6. Procédé selon l'une quelconque des revendications 1 à 5, dans lequel les adaptateurs sont conçus de manière telle que la séquence cible correspondante de l'endonucléase ne soit pas restaurée lorsque l'adaptateur s'hybride avec le ou les fragments de restriction repérés et, dans lequel la ou les amorces sont des oligonucléotides comprenant une séquence constante effectuant parfaitement un appariement de bases avec au moins une extrémité de la séquence d'au moins un fragment de restriction repéré du mélange, laquelle séquence constante comprend des nucléotides effectuant un appariement de bases à une partie de la séquence cible pour ladite endonucléase de restriction, et au moins une desdites amorces comprend adjacents à l'extrémité 3’ des nucléotides impliqués dans la formation de ladite séquence cible, 1 à 10 nucléotides additionnels sélectionnés.

7. Procédé selon l'une quelconque des revendications 1 à 6, dans lequel la séquence d'acides nucléiques de l'ADN de départ est au moins en partie inconnue.

8. Procédé selon l'une quelconque des revendications 1 à 7, dans lequel deux endonucléases de restriction différentes sont utilisées.

9. Procédé selon l'une quelconque des revendications 1 à 8, dans lequel toutes les amorces possèdent la même séquence de nucléotides constante.

10. Procédé selon l'une quelconque des revendications 1 à 9, dans lequel un mélange de différentes amorces est utilisé.

11. Procédé selon l'une quelconque des revendications 1 à 8, dans lequel au moins une amorce contient 1, 2, ou 3, nucléotides sélectionnés, à leur extrémité 3’.

12. Procédé selon l'une quelconque des revendications 1 à 10, dans lequel 2 endonucléases de restriction différentes sont utilisées dans l'étape (a) et dans lequel au moins un oligonucléotide adaptateur synthétique double brin pour une des endonucléases est biotinylé.

13. Procédé selon l'une quelconque des revendications 1 à 12, dans lequel au moins une des amorces est marquée.
14. Procédé selon l'une quelconque des revendications 1 à 13, dans lequel le nombre de nucléotides sélectifs est choisi de sorte que le nombre de fragments de restriction qui seront amplifiés soit limité de 5 à 200.

15. Procédé pour la préparation d'un ensemble de fragments de restriction amplifiés comprenant :

a) l'amplification d'un sous-ensemble de fragments de restriction à partir d'un mélange de fragments de restriction engendrés, obtenu à partir d'un ADN de départ utilisant un procédé selon l'une quelconque des revendications 1 à 14,

b) la séparation des fragments de restriction repérés amplifiés ;

c) l'identification des fragments de restriction amplifiés.

16. Procédé selon l'une quelconque des revendications 1 à 14, dans lequel le dit ADN de départ à amplifier est l'ADN génomique en provenance d'un échantillon biologique humain ou animal, en particulier un tissu ou un échantillon de sang.

17. Procédé selon l'une quelconque des revendications 1 à 14, dans lequel le dit ADN de départ à amplifier est un ADN génomique végétal ou de tissu végétal.

18. Procédé selon l'une quelconque des revendications 1 à 14, dans lequel le dit ADN de départ à amplifier est un ADN de micro-organisme.

19. Procédé selon l'une quelconque des revendications 1 à 15, pour l'identification de polymorphismes entre différents ADNs de départ provenant des mêmes espèces vivantes qui comprend l'identification des différences observées entre les fragments de restriction amplifiés des différents ADNs de départ.

20. Procédé selon l'une quelconque des revendications 1 à 14, pour l'identification de polymorphismes dans les ADN génomiques de micro-organisme, de végétal ou d'animal, incluant les humains, ou des fragments de ceux-ci, soit parmi eux-mêmes, soit apparentés à un ADN étalon déterminé correspondant.

21. Procédé selon l'une quelconque des revendications 1 à 14, pour identifier des polymorphismes d'ADN associés à des caractéristiques génétiques héréditaires chez les humains, des polymorphismes d'ADN associés à des caractéristiques génétiques héréditaires chez les animaux, ou pour identifier des polymorphismes d'ADN associés à des caractéristiques génétiques héréditaires chez les végétaux, et pour identifier des marqueurs d'ADN basés sur de tels polymorphismes d'ADN.

22. Procédé selon l'une quelconque des revendications 1 à 14, comprenant en outre la comparaison de deux ou plusieurs fragments d'ADN amplifiés ou allongés identifiés ou récupérés dans l'étape (e) dit procédé pour déte

23. Procédé selon l'une quelconque des revendications 1 à 14, dans lequel les fragments d'ADN amplifiés ou allongés sont identifiés dans l'étape (e) tels que produit dans l'étape (e) sont identifiés ou récupérés comme empreintes d'ADN.

24. Procédé pour l'identification de marqueurs d'ADN liés à une caractéristique génétique, le dit procédé comprenant l'identification de polymorphismes selon la revendication 22, entre les ADN de départ provenant des mêmes espèces vivantes qui présentent des différences dans ladite caractéristique génétique et la mise en corrélation desdits polymorphismes avec le phénotype présenté par ladite caractéristique génétique.

25. Procédé selon l'une quelconque des revendications 1 à 14 comprenant de plus les étapes consistant :

(f) à isoler au moins un fragment d'ADN identifié et récupéré dans l'étape (e) ;

(g) à déterminer la séquence de nucléotides des 8 à 10 premiers résidus de nucléotides adjacents, de manière interne, à la séquence cible de l'endonucléase de restriction aux deux extrémités desdits fragments d'ADN ;

(h) à concevoir des amorces oligonucléotidiques ayant une séquence de nucléotides selon les amorces de l'étape (c) de la revendication 1 dans laquelle la séquence de nucléotides sélectionnée comprend 5 à 10 résidus de nucléotides qui correspondent aux 8 à 12 premiers résidus de nucléotides adjacents de manière interne aux sites de restriction au niveau des deux extrémités dudit fragment d'ADN.
Fig. 1 tagging of genomic restriction fragments

genomic DNA

restriction enzyme digest

adaptor ligation
Fig. 2 adaptor ligation

**flush ends**

- ex. PvuII
  - CTGN
  - GACN
  - NCAG
  - NGTC

- +adaptor

- NNNN
  - NNNN

**staggered ends**

- 5' extension
  - ex. BglII
    - GATC
    - CTAG

- +adaptor

- GATC

- 3' extension
  - ex. PstI
    - ACGT
    - TGCA

- +adaptor

- ACGT

- TGCA
Fig. 3 PCR amplification of tagged restr. fragments

primer
Fig. 4 Multiplex restriction fragment amplification
Fig. 5 Selective PCR primers

selective primers:

constant sequence selective bases

variable sequence

constant sequence

template selection:

matching template

NNNNNNNNNNNNNNNGATCC GAG
NNNNNNNNNNNNNNNCCTAGG CTCNNNNNNNNNNNNNNNNNNNNNNNNNNN

mismatch template

NNNNNNNNNNNNNNNGATCC G
NNNNNNNNNNNNNNTCTAGG AAGNNNNNNNNNNNNNNNNNNNNNNNNNNN
Fig. 6 Selective PCR amplification of restr. fr.

Adaptor

Restriction fragment
Fig. 7 Selective restriction fragment amplification
Fig. 8 Fragment specific selective PCR primers

**selective primer 1**

constant sequence  selective b.  constant sequence

**selective primer 2**

constant sequence  selective b.  constant sequence

template selection:
FIGURE 14: SRFA OF FOUR TOMATO LINES USING PStI AND HaeIII AS RESTRICTION ENZYMES.
FIGURE 13: SRFA OF LACTUCA DNA USING Pst I AND Xba I AS RESTRICTION ENZYMES.

-340
-250
-200
-150
FIGURE 16: SIEPA OF CORN DEA5 USING PSEU/TAC1 AND ECOE1/TAC1
FIGURE 17: EXPERIMENTAL RESULTS USING ALFA AND TAQ AS RESTRICTION ENZYMES.