High artemisinin yielding plant genotype 'cim-aroga'

FIELD OF THE PRESENT INVENTION

[0001] The present invention was related to the development of a novel, distinct high herb and artemisinin yielding genotype of Artemisia annua obtained through systematic marker assisted breeding followed by selection of uniform population in a methodical way. This invention further relates to marker assisted breeding to reach the high artemisinin yielding genotype. The genotype is distinct, uniform and stably maintainable by continuous rouging of off types in the population using DNA marker at early seedling stage from nursery itself and suitable for commercial cultivation.

BACKGROUND AND PRIOR ART REFERENCES OF THE PRESENT INVENTION

[0002] Many Artemisia species are cited by early herbalists including Theophrastus in the third century B.C (Einarson and Link., 1976), Pliny (Bostock and Riley., 1855-1857) and Dioscorides (Gunther., 1959) in the first century BC. Wormwood (probably the species A. judaica) is mentioned in the Bible (Rev 8:10,11). In 340 AD, Ge Hong prescribed aerial part of Artemisia for the treatment of fever in the "Chinese hand book of prescriptions for emergency treatments" and in 1527, Li Shi Zhen, a Chinese herbalist/pharmacologists mentioned the use of huang hua hao (or yellow flower, later identified as A. annua) for treatment of children's fever and qinghao (A. apiacea) as a treatment for the disease now known as malaria.

[0003] The plant Artemisia annua (family: Asteraceae) produces a sesquiterpenoid lactone endoperoxide named artemisinin which is a promising antimalarial drug effective against Plasmodium falciparum, Plasmodium vivax at nanomolar concentration. Artemisinins are active against Schistosoma mansoni and S. japonicum in-vitro and in-vivo in experiments in animals. These schistosomes, like malarial parasites, degrade haemoglobin and produce hemozoin. These compounds are also active against Leishmania major, Toxoplasma gondii and Pneumocystis carinii in-vitro and against P. carinii in-vivo. Artemisinins have immunosuppressive activity and also potential anticancer activity. For these activities, the doses of artemisinin required are substantially higher than the dose for antimalarial activities. According to Meshnick et al., (1996) (Microbiological Reviews 6:301-315) the antimalarial endoperoxides including artemisinin, dihydroartemisinin and arteethers, are not likely to be useful for other therapeutic purposes except against malarial parasites.

[0004] Although artemisinin rapidly suppresses the activity of parasites like Plasmodium vivax and P. falciparum, problems with high rate of recrudescence (>10%
recrudescence infections), short half life persist. Hence, there is a need to develop new
 drugs against quinolone resistant pathogenic bacteria. It is a known fact that clinically
 used antibacterial broad spectrum compounds such as quinolones which exhibit DNA
 gyrase activity of Mycobacterium sp. (causing tuberculosis), Haemophilus sp. and
 Haemophilus influenzae are gradually becoming ineffective due to the occurrence of
 mutatious in gyrase genes and their natural selection under continuous use of such drug.
 The compound a arteether developed as antimalarial drugs by Central Drug Research
 Institute (CDRI), Lucknow, India and Central Institute of Medicinal & Aromatic Plants
 (CIMAP), Lucknow, India, after phase II clinical trial is a stable derivative of
 artemisinin. Earlier we have found a novel property of .alpha.-arteether as being
effective against the gyr A mutant strains of E. coli but ineffective against wild type
 strains (U.S. Pat. No. 6,127,405). Also we have developed a strategic and novel
 composition comprising .alpha. arteether and nalidixic acid or quinolone drugs which is
 useful as an advanced generation drug to counter the resistance development itself and
 having a potential to be used in treating infectious diseases and in inhibiting the
 resistance developed due to mutation in the gyr A gene of bacteria, particularly in those
 cases where drug resistant strains are known to appear very frequently (U.S. Pat. No.
 6,423,741). We have already reported a genotype `Jeevanraksha’ earlier yielding more
 than 1% artemisinin (Sushil Kumar, S Banedjee, S Dwivedi, M M Gupta, R K Verma,
 D C Jain, S P S Khanuja, A K Mathur, G D Bagchi, M Zehra, V K Mehta, A A Naqvi, S
 Paul, G Ram, M Ram, D Saikia, R S Sangwan, T R Santha Kumar, A K Shasany, M P
 Sci. 21: 47-48,) and the method to increase yield through harvesting management (U.S.
 Pat. No. 6,393,763).

[0005] It is always beneficial to have diversity in genotypes in different background
 than a single genotype for commercial cultivation. With this objective a novel genotype
 was developed through a novel method of DNA marker assisted breeding.

OBJECTS OF THE PRESENT INVENTION

[0006] The main object of the present invention is to develop a novel, distinct high herb
 and artemisinin yielding genotype of Artemisia annua.

SUMMARY OF THE PRESENT INVENTION

[0007] The present invention was related to the development of a novel, distinct high
 herb and artemisinin yielding genotype of Artemisia annua obtained through systematic
 marker assisted breeding followed by selection of uniform population in a methodical
 way. This invention further relates to marker assisted breeding to reach the high
 artemisinin yielding genotype. The genotype is distinct, uniform and stably
 maintainable by continuous rouging of off types in the population using DNA marker at
 early seedling stage from nursery itself and suitable for commercial cultivation.

DETAILED DESCRIPTION OF THE PRESENT INVENTION

[0008] Accordingly, the present invention relates to a new and distinct genotype of
 Artemisia annua `CIM-Arogya’, developed through marker assisted breeding,
 possessing the following combination of characters:
Genus.--Artemisia.
Species.--annua.
Family.--Asteraeaceae.
Common name.--Qinghao.
Plant height.--280-305 cm.
Plant canopy.--Oval.
Growth habit.--Erect.
Branching.--sympodial branching pattern.
Stem.--Single, round hard woody green (137D) Stem width 5-12 cm (app).
Number of branches:
Primary branches.--55-65.
Secondary branches per primary branch.--50-60.
Tertiary branches per secondary branch.--37-45.
Range of length of internodes:
Main stem.--4-6 cm at bottom, 10-12 cm at middle, 2-5 cm at the top.
Primary branch.--2-5 cm at bottom, 8-11 cm at middle, 2-4 cm at the top.
Secondary branch.--8-11 cm at the bottom, 5-7 cm at the middle, 1-3 cm at the top.
Tertiary branch.--3-5 cm at the bottom, 1-3 cm at the middle, 0.5-1.5 cm at the top.
Leaf.--Green (137 B).
Texture.--Thin and flexible.
Surface.--Smooth non pubescent.
Shape.--Pinnately compound leaf.
Margin.--Pinnetisected.
Tip.--Acute.
Petiol length.--2.5-3.50 cm.
Lamina length.--5-7 cm.

Lamina width.--4-5 cm.

Inflorescence.--Capitulum (head).

Flower.--arranged in whorls. Colour yellow group (7A).

Two types of flowers.--Disc florets and Ray florets. Disc florets are bisexual and ray florets are unisexual (female).

Colour.--Greenish yellow 2-3 mm in diameter.

Receptacles.--Glabrous.

Calyx.--Bracteates.

Corolla.--Sympelatous, tubular top split in to 5 lobes in Disc florets and 2-3 lobes in Ray florets (legulate).

Androecium.--5 stamens, Anther lobes are fused and filaments are free.

Colour.--Yellowish.

Gynoecium.--Unilocular, inferior bifid stigma.

Colour.--yellowish.

Time of flowering.--198 days.

Seed setting.--240 days (app).

Artemisinin content.--0.9 to 1.1%.

Artemesinic acid.--0.002-0.004%.

Oil content.--0.35-0.45%.

Alternate deeply dissected aromatic leaves ranging between 2.5 to 5.0 cm in length.

Tiny yellow nodding flowers (capitula) capitula in loose panicle containing numerous central bisexual florets and marginal pistillate florets

Receptacle is glabrous
[0057] florets and receptacle bear abundant 10-celled biseriate trichomes

[0058] globular canopy

[0059] dry leaf yield of about 50 Q per hectare

[0060] flowers (7A) arranged in a head or Capitulum,

BRIEF DESCRIPTION OF THE ACCOMPANYING DRAWINGS

[0061] FIG. 1 shows the plant along with its architecture "CIM-Arogya"

[0062] The research on genome analysis is being taken up as a necessity to understand the genomic constitution of individuals in terms of DNA content, nature and variations etc. The data from genome analysis are of direct relevance to molecular plant breeding in which morphological characters can be tagged to unique DNA sequences and then inheritance patterns of DNA markers can be utilized to confirm the presence of traits even before expression. Techniques are available to differentiate even similar looking individuals of a population on the basis of DNA sequence variation Some recent important discoveries from application point of view towards genetic analysis include Restriction endonucleases mapping and Polymerised Chain Reaction for amplifying DNA sequences from traces. These discoveries have led to the means and techniques used to study the differences or uniqueness in the DNA sequences otherwise known as Polymorphism in the DNA. The tools like RAPD, AFLP, RFLP, micro-satellite and many others were invented earlier and used in literature extensively for differentiating and marking the plants for different characters.

[0063] Development of Marker Correlating to High Content of Artemisinin in the Plant Artemisia annua.

[0064] Selection of Genotypes

[0065] The seeds of ten chemotypic accessions of the plant Artemisia annua were selected from Kashmir and further studies were carried out in Lucknow field station. Seeds of A. annua were sown in pots with mixture of soil and FYM (farmyard manure) in the ratio of 1:1 and germination in glass house conditions during the month of November of the years 1998-2001. The seedlings having 10 cm height were transplanted with spacing 50 cm between rows and 30 cm between plants. The soil of experimental field was sandy loam in texture and neutral in reaction (pH 7.6). The plots were fertilized with FYM (Farm yard manure) @20 kg.ha before transplanting for obtaining optimum performance. Plots were prepared 3 m.times.3 m size with irrigation channels. For Hybridization, six seed lots were selected out of 10 seedlots (obtained from Kashmir in the year 1998) were transplanted in alternative rows. From the next year (1999) on wards the progeny seedlings of the chemotypically selected plants were planted again in alternate rows. All the seedlings were checked for artemisinin content after extraction. About 0.1 g dry powdered plant material was extracted in 10 ml of hexane by heating at 60.degree. C. for 3 minutes and left for overnight at room temperature. Then extract was filtered and evaporated on water bath at 50.degree. C. After evaporation extract was dissolved in 1 ml hexane and used in TLC. Properly (20.times.20 cm E-MEREK) dissolved extract was spotted in TLC plates at 1 cm apart along with standard (1 mg.ml). Spotted TLC plate was dipped in
solvent (mobile phase) Hexane:Diethyl ether (1:1) Plate was dried in air and dipped in developing reagent Glacial acetic acid:conc. Sulphuric acid:Anisaldehyde (50:1:0.5 ml) and heated at 120 degree. C. for 10-15 minutes and then Stabilized and scanned (540 nm, visible) (Densitometer CAMAG :Switzerland). The TLC plates were scanned and the artemisinin content of individual progeny plants were quantified. From the analysis the plants producing trace (0.10% or less) artemisinin and the plants producing more than 0.4% artemisinin were selected and finally 10 plants from each category were taken for DNA analysis.

DNA Isolation and PCR Amplifications

DNA was isolated from the leaf tissue essentially according to the protocol described earlier (Khanuja S P S, Shasany A K, Darokar M P, Sushil Kumar (1999) Rapid Isolation of PCR Amplifiable DNA from the Dry and Fresh Samples of Plants Producing Large Amounts of Secondary Metabolites and Essential oils by Modified CTAB Procedure. Plant Molecular Biology Reporter, 17, 74.). Polymerase chain reactions (PCRs) were carried out in 25 μl volume. A reaction tube contained 25 ng of DNA, 0.2 unit of Taq DNA polymerase, 100 μM of each dNTPs, 1.5 mM MgCl₂ and 5 pmol of decanucleotide primers. The amplifications were carried out using the DNA Engine thermal cycler (MJ Research, USA) using 94 degree. C., 35 degree. C. and 72 degree. C. temperatures for 40 cycles (Khanuja S P S, Shasany A K, Srivastava A, Sushil Kumar (2000). Assessment of genetic relationships in Mentha species. Euphytica, 111, 121-125.). The amplified products were separated on 1.2% agarose gel containing 0.5 μg ml⁻¹ of ethidium bromide and photographed with Image master VDS (Pharmacia). The bands were analyzed using Image master 1D elite software and the graphic phenogram of the genetic relatedness among the accessions was produced by means of UPGMA (unweighted pair group method with arithmetic average) cluster analysis. Custom-made decanucleotide primers were synthesised in the laboratory on Applied Biosystems 392 DNA-RNA Synthesizer and were designated as MAP01 to MAP20.

The sequences of the primers MAP01 to MAP20 were AAATCGGAGC, GTCCCTACTCG, GTCCCTAGCG, TGGCGCAGTCG, AACGTACCGG, GCACCGCGGA, CACCCTCCGC, CTATCGCCGC, CGGGATCCGC, GCGAATTCCG, CCCTGCAGGC, CCAAGCTTGC, GATCACTGAG, AGGATACGTG, AAGATAGCGG, GGATCCTGAAC, TTGTCCTAGG, CATCCCCAAC, GGACTCCACG, AGCCTGACGC, respectively. The other sets of primers used included kit J, O and T, each consisting of 20 random decamer primers, procured from Operon Technologies Inc., USA.

All the RAPD profiles thus generated were analyzed for bands always appearing with all the high artemisinin containing genotypes (more than 0.4%) and absent in the genotypes containing trace or no artemisinin. We could detect a band at approximately 850 base pair region amplified with the primer 5’CCAAGCTTGC3’ (MAP 12, Sequence ID 1) which consistently showed its presence in the genotypes containing more than 0.4% artemisinin and absent in the genotypes with trace or no artemisinin. This finding was interesting considering the complex nature of the artemisinin biosynthetic pathway. For all other primers the amplified products showed variable positions in these genotypes and could not be correlated. The presence of the band in the segregating populations having high artemisinin could be ascertained as the samples of 10 analyzed plants having high artemisinin were drawn from different populations. Similarly, the
sample of 10 plants for trace or no artemisinin drawn from different populations could show always the absence of the band. As all the plants analyzed were from the same initial population the genes for artemisinin biosynthesis were assumed to be normal. So the presence and absence of the band could be correlated to the regulatory function associated with the expression of some of the genes associated with the biosynthetic pathway. But certainly the DNA band of about 850 base pair size could be correlated with the biosynthesis of more than 0.4% artemisinin in Artemisia annua. In the next steps the DNA fragment described earlier was eluted out from the agarose gel and (since the fragment was amplified with the primer containing Hind III restriction site) restricted with Hind III restriction enzyme (Recognition and restriction site 5'AAGCTT3'). Similarly, pBluescript II SK(+) procured from Stratagene Inc. was used to clone the fragment at the Hind III site using T4 DNA ligase enzyme available commercially. Escherichia coli strain DH5. alpha., procured from Stratagene Inc again was transformed with this constructed plasmid and transformed cells were isolated on agar plates containing nutrient broth and ampicillin. All the experiments were performed according to the protocol Sambrook et al (1988).

This fragment was sequenced completely with the help of M13 forward and T3 reverse primer (the sequence sites are present in the plasmid pBluescript II SK(+) and the nucleotide sequence is given below.

1 1 AAGCTTGCTG AACGCACTGG TGTATCTGCC GCAGCCCGTG AACTCAGCCT GTATGAAATCA 61 CAACCTTACA ACTGGCGCAG TAAACAGCAA ATACAGCAGC GCTTTCTGGA AGTGAAGCTG 121 GAGATGTCTA CCGAGATTGC ACGTCTCAA A GGCAGCTGG CAAAGGGGGA TGAAGAGCTG 181 GCTATCCTCC AAAAGGCGC GACATACTTC GCGAAGCGCC TGAATGAAG TATGCTTTA 241 TTGAAATCAG TCGCTGGAG TTTCGACATCA AGCAATGTG CCGCCTGCTC CCGGTGGGCCC 301 GCAGCGGCTG GTATAAGCTG GTGGTACGGG CGGACAAGGA TAAGCCCGCC TAAGCAGTTC 361 CGCAACACT GCAGCAGGGG TTTGCTTCGCC GGTTTTACCC CCGGTAAAC AAGCGTTACC 421 GGTGGGCCAC GTCTGACGGG ATGACCTGCG GTGCTCAAGG TTACCTTTTA ACGTAAAAA 481 CCCGTCGCGG CAAGCTTGGCC CGGTCGGGGA CTAGAAGGCAA AGGCCTCACC GGAGAGTCCG 541 CGGTCGAGAT ACCTGCACGC ACGGGCGGTCC CTGCTGCTAG AAATCTGTG GAGCAGATT 601 TTTCGCGCCA GTGGCCTCAGA CCCAGAATGG GCAGGAGACA TCACGTATCT ACGTACAGT 661 GAAAGCTTGGC TGCTTCTGGC AGTCGGTCTG ATGGTGTGCTCCATT 721 TCAATGGTGC CACGGCAGC GGGCGAATCTG GCTGGTGCTG CCGTGACAT GGGCTTGCTGG 781 CGCGTAAAGA GGGCCGGCAA CGTATCGTTC GAGAGGTGGCCAGAGTCCATGA GTGGCTGCTG 841 GCAGATTATC AGTCGGCAACT GAAAGCTTCGCTGCTC 901 TGCTGCTACGG TATAATGCCTG CACGTTGGGCA TT

Based on the sequence at the ends forward and reverse primers were synthesized with the sequence

2 Forward Primer 5'CCAAGCTTGCTGAACGCACTGG3' Reverse primer 5'CCAAGCTTGCCACGCAAGGATTATC3'
These sequences were used to amplify the genomic DNA of Artemisia annua (both high content of artemisinin and low content of artemisinin). The plant genomic DNA with high artemisinin content could generate a band of 936 bp whereas in plants containing low amount of artemisinin the absence of the band was conspicuous.

Use of the Marker to Generate a Population of Plants with High Artemisinin Content.

In the first year polycross nursery was designed with alternate male and female line chosen among the seed lots. These plants were randomly picked up from the nursery raised from the 6 selected seed lots. The plants, which were designated as female (270 plants), were analyzed for artemisinin content, which were selected for further experimentation. Seed sample were collected from these selected plants (13 in number) containing high amount of artemisinin (0.15 to 0.20%) and planted again in a polycross nursery in the second year. Next year 180 plants were analyzed for artemisinin content and 13 plants containing 0.45 to 0.50% artemisinin were selected for planting in the third year. At this point 10 plants with more than 0.4% artemisinin and 10 plants containing trace amount artemisinin were taken for DNA isolation to develop SCAR marker as described previously. The SCAR marker was used to select plants from the nursery raised from the seeds selected 13 seed lots, and 12 plants from each seed lots showing the presence of SCAR marker were selected for random crossing among the plants in the third year. Randomly plants were analyzed for artemisinin content and among 150 plants analyzed 20 plants having artemisinin 0.8 to 1.0% were selected for next year (fourth year) planting. The seeds from these plants were grown in the nursery and 12 SCAR positive plants from each seed lot were grown randomly to facilitate cross pollination. From these 200 plants were analysed for artemisinin content and 11 plants were selected having 1.0 to 1.16% artemisinin content. Simultaneously, increase in the mean artemisinin content of the plants analysed every year were calculated.

Genetic Advancement

The key metabolite synthesis (Artemisinin content) was studied for genetic advancement which showed an upward trend beginning with 17.33% increase in the mean artemisinin content for the first year, crossing 50% in second year, 60% in third year and remaining at 42.06% in fourth year. The advancement in artemisinin content was calculated as per Singh and Chaudhary (1977) (Singh R K and Chaudhary B D (1977). Biometrical methods in quantitative genetic analysis. Kalyani Publications, New Delhi.

The seeds obtained from these selected plants four single plant seed lots (CIMAP-G1, CIMAP-G2, CIMAP-G3, CIMAP-G4) were evaluated in the field at CIMAP Farm, Lucknow, Uttarpradesh, Lucknow, India.

Experimental (Agronomic) details Artemisia annua evaluation trials at CIMAP research farm.

| Component(s) | 2001-02 | 2002-03 | Design | RBD | RBD | Genotypes (with check) | Five | Five | Manuring (FYM) 10t/ha | 10t/ha | Fertilization (NPK) 80:40:40 | 80:40:40 | Row to row distance 50 cm | 50 cm | Plant to plant distance 30 cm | 30 cm | Plot size 12.5 sq m (Net) | 15.75 sq m (Net) | DOT 23/02/02 | 10/03/03 | DOH (I) 03/06/02 (100 DAP) | 02/06/03 | (84
Four selected genotypes from these lots were selected among each other and with the best check ’Jeevan Raksha’. Herb yield data with regard to advanced lines of Artemisia annua along with the check during the evaluation trials at CIMAP research farm, Lucknow has been provided below. The genotype CIMAP-G2 (CIM-Arogya) yielded maximum dry leaves compared to other genotypes in the trial.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>2001-02</th>
<th>2002-03</th>
<th>2001-02</th>
<th>2002-03</th>
</tr>
</thead>
<tbody>
<tr>
<td>CIMAP-G1</td>
<td>523.6</td>
<td>466.67</td>
<td>55.56</td>
<td>48.09</td>
</tr>
<tr>
<td>CIMAP-G2</td>
<td>553.0</td>
<td>478.45</td>
<td>58.40</td>
<td>48.89</td>
</tr>
<tr>
<td>CIMAP-G3</td>
<td>559.8</td>
<td>444.45</td>
<td>56.04</td>
<td>45.71</td>
</tr>
<tr>
<td>CIMAP-G4</td>
<td>422.0</td>
<td>439.69</td>
<td>42.00</td>
<td>46.27</td>
</tr>
<tr>
<td>Jeevan Raksha</td>
<td>438.4</td>
<td>426.19</td>
<td>43.60</td>
<td>42.30</td>
</tr>
</tbody>
</table>

F-value: 7.55** 6.2** 10.79** 6.31**

gm 499.36 451.11

51.12 46.25 sem 23.6 8.5 2.34 10.02 cv 9.45 3.77 9.16 4.41 cd (1%) 101.88 36.71 10.11 4.4 cd (5%) 72.7 26.19 7.21 3.14

The plant genotype of Artemisia annua of the invention was named as ‘CIM-Arogya’ and referred in the same name in the patent document. The genotype can be grown as a uniform population of high artemisinin yielding plants with rouging at nursery.

**Taxonomic Description of ‘CIM-Arogya’**

The plant is usually single stemmed reaching about 2 m in height with alternate branches and alternate deeply dissected aromatic leaves ranging from 2.5 to 5.0 cm in length. Tiny greenish yellow nodding flowers (capitula) only 2 or numerous imbricate bracts enclose 3 mm in diameter. Capitula is displayed in loose panicle containing numerous central bisexual florets and marginal pistillate florets, the latter extruding stigmas prior to the central flower. The receptacles is glabrous, not chaffy and triangular in shape. Both florets and receptacle bear abundant 10-celled biseriate glandular trichomes, which are the source of artemisinin and highly aromatic volatile oils (essential oil).

The colour codes are in accordance with the "RHS colour chart published by The Royal Horticultural Society, 80 Vincent Square, London SW1P 2PE, 1995.

The genotype ‘CIM-Arogya’ possessing the traits of increased herb yield than the other check varieties and genotypes. The genotype is having higher biomass leading to high artemisinin yield. Its genetic make up is distinct in terms of DNA profile. The genotype in the population has expressed a genetic enhancement of artemisinin content to a very high content of artemisinin through strategic marker aided selection indicating the distinctiveness from the parent genotype. The plant has a unique globular canopy.

Randomly Amplified Polymorphic DNA analysis: The RAPD analysis of the genotype ‘CIM-Arogya’ were unambiguously able to establish its distinct identity as completely different from the check genotypes. The 20 MAP primers (MAP 01 to MAP 20) synthesized in the laboratory using ABI 392 DNA synthesizer, with the sequence AAATCGGAGC, GTCTCTACCTG, GTTCTTACCGG, TGCGCGGATCG, AACGTACCGG, GCACGCCGGA, CACCCTTCGCC, CTATGCCTCG, CGGGATCCGC, GCGAAATCCCG, CCCTGACGCG, CCAAGCTTGC,
GTGCAATGAG, AGGATACGTG, AAGATAGCGG, GGATCTGAAC, TTGTCTCAGG, CATCCGAAC, GGAICTCCACG, AGCCTGACGC were used for the analysis to differentiate among the genotypes.

[0086] From RAPD analysis the profiles were studied and similarity indices were calculated which were put into a matrix. This matrix was used to produce a graphic phenogram by means of UPGMA (unweighted pair group method with arithmetic average) cluster analysis. As represented in the phenogram provided below (FIG. 1) the clone of the invention is quite different from the other varieties. Sequence CWU 1

23 1 10 DNA ARTIFICIAL MAP Primer 1 aatacggagc 10 2 10 DNA ARTIFICIAL MAP Primer 2 gtctacgcc 10 3 10 DNA ARTIFICIAL MAP Primer 3 gtcttaacgcg 10 4 10 DNA ARTIFICIAL MAP Primer 4 tgecgatgcg 10 5 10 DNA ARTIFICIAL MAP Primer 5 aagtacgcgg 10 6 10 DNA ARTIFICIAL MAP Primer 6 gacaacgctgga 10 7 10 DNA ARTIFICIAL MAP Primer 7 caccctgcggc 10 8 10 DNA ARTIFICIAL MAP Primer 8 gacaccggc 10 9 10 DNA ARTIFICIAL MAP Primer 9 cggagtacgcg 10 10 10 DNA ARTIFICIAL MAP Primer 10 gcaataacgcc 10 11 10 DNA ARTIFICIAL MAP Primer 11 cccgctgcgcg 10 12 10 DNA ARTIFICIAL MAP Primer 12 cggagtacgcg 10 13 10 DNA ARTIFICIAL MAP Primer 13 gtctactgcc 10 14 10 DNA ARTIFICIAL MAP Primer 14 agcctggagc 10 15 10 DNA ARTIFICIAL MAP Primer 15 aagcttgacgc 10 16 10 DNA ARTIFICIAL MAP Primer 16 gatactgcacgc 10 17 10 DNA ARTIFICIAL MAP Primer 17 tgtgcacgcg 10 18 10 DNA ARTIFICIAL MAP Primer 18 ccgctgcgcc 10 19 10 DNA ARTIFICIAL MAP Primer 19 gacttcacgc 10 20 10 DNA ARTIFICIAL MAP Primer 20 agcctggagc 10 21 10 DNA ARTIFICIAL MAP Primer 21 aagcttgacgc