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Pait et al.

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(54) **LOBLOLLY PINE TREE NAMED ‘CF LP1-7696’**

(50) Latin Name: *Pinus taeda*
Varietal Denomination: **CF LP1-7696**

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(51) **Int. Cl.**
A01H 5/00 (2006.01)

(52) **U.S. Cl.** **Plt./213**

(58) **Field of Classification Search** **Plt./213**
See application file for complete search history.

(56) **References Cited**

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(57) **ABSTRACT**

A new and distinctive variety of a loblolly pine tree which has been denominated varietally as ‘CF LP1-7696’ which is distinguished by high growth rate, good resistance to fusiform rust, excellent stem straightness, medium crown width, medium number of whorls, medium branch angle and medium branch diameter.

2 Drawing Sheets

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Latin name: *Pinus taeda*.

Variety denomination: ‘CF LP1-7696’.

BACKGROUND

A new variety of loblolly pine tree (*Pinus taeda*), has been discovered. This selection has been designated as ‘CF LP1-7696.’

This new variety is a progeny of two second generation selections. The female parent is a progeny of two first generation selections made in Cherokee County, Tex. and Tyler County, Tex. The male parent is a progeny of an open pollinated first generation selection made in Montgomery County, Tex.

Cross pollination occurred in early 2000 followed by induction and cryopreservation of embryogenic tissue in 2001. First somatic seedlings were produced in 2002 and planted in early 2003 in three field experiments. A total of 30 ramets were planted at 10 ramets per field experiment. The field experiments are located in Texas and Louisiana.

BRIEF SUMMARY

A new and distinct cultivar of loblolly pine (*Pinus taeda*) is distinctly characterized by high growth rate, good resistance to fusiform rust, excellent stem straightness, medium crown width, medium number of whorls, medium branch angle, medium branch diameter and which is mature for commercial

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harvesting sooner than conventionally grown trees under the ecological conditions prevailing in the Gulf Coastal Plains of the United States.

The *Pinus taeda* plants of this variety were asexually propagated using an advanced form of micropropagation called somatic embryogenesis carried out at a production facility in Victoria, Canada. Somatic embryogenesis uses a complex process which relies on the splitting of one embryo into many identical embryos. Somatic embryos can then be grown into plants which are all identical genetically. The asexual propagation occurs at an earlier stage in the plant’s life cycle than most other micropropagated plants. The detailed methods for somatic embryogenesis used for asexually propagating conifers in general are described in U.S. Pat. No. 6,372,496 and for loblolly pine in particular in U.S. patent application Publication No. 2004/0203150.

BRIEF DESCRIPTION OF THE DRAWINGS

The drawings are color photographs showing the new variety of loblolly pine.

FIG. 1 is a photograph showing ‘CF LP1-7696’ ramet # 3 planted in Beulah, Tex. The picture was taken after five field growing seasons. The picture shows superiority of growth and medium crown width.

FIG. 2 is a photograph showing ‘CF LP1-7696’ ramet # 5 planted in Beulah, Tex. The picture was taken after five field growing seasons. The picture shows excellent stem straight-

ness, medium number of whorls per unit stem length, medium angle between the stem and the branches, and medium branch diameter (relative to the size of the stem).

DETAILED BOTANICAL DESCRIPTION

The botanical details of this new and distinct variety of loblolly pine tree follow. All color descriptions are made in reference to The Royal Horticultural Society (R.H.S.) Colour Chart (2005).

Parentage:

Female parent.—(Unknown) progeny of two first generation selections made in Cherokee County, Tex. and Tyler County, Tex.

Male parent.—(Unknown) progeny of an open pollinated first generation selection made in Montgomery County, Tex.

Leaf: Evergreen needles, 6 to 9 inches long, with (usually) three yellow-green needles per fascicle. The color of the foliage was taken from field established trees aged 6 years, but was not found to vary significantly with age. The color of the foliage was RHS 137A (60%) and 136A (40%).

Flower: Monoecious; males long cylindrical, red to yellow, in clusters at branch tips; females yellow to purple.

Fruit: Ovoid to cylindrical, 3 to 6 inch red-brown cones; umbo is armed with a short spine, maturing in early fall. Cones are sporadic in 5-7 year old plants.

Branch: Orange-brown in color, fine to moderately stout; buds are narrowly ovoid, light reddish brown.

Bark: Initially red- to gray-brown and scaly; older trees are ridged and furrowed, with somewhat rounded scaly plates; very old trees have red-brown, flat scaly plates.

Shape: A medium to large tree can reach well over 100 feet tall, self-prunes well and develops a fairly straight trunk and an oval, somewhat open crown.

Compared to unimproved loblolly pine trees, 'CF LP1-7696' is characterized by high growth rate, good resistance to fusiform rust (caused by *Cronartium quercuum* (Berk.) Miyabe ex Shirai f. sp. *fusiforme* (Cumm.) Burds. et Snow), excellent stem straightness, medium crown width, medium number of whorls, medium branch angle and medium branch diameter.

Average height: 20 ft after 5 field growing seasons

Maximum height: 24 ft after 5 field growing seasons

Average trunk diameter at breast height (4.5 feet above the soil level): 3.8 inches after 5 field growing seasons and 4.7 inches to 6.0 inches after 6 field growing seasons.

Sampling of Branch Characteristics: In order to sample branches from a consistent position from one tree to the next the following methodology was utilized. From a point nine feet from the base of the tree, the first complete whorl of limbs below was labeled "Whorl One" and the first complete whorl of limbs above labeled "Whorl Two". This sampling point was chosen because it is the midpoint of the basal sixteen foot log of each tree. A complete whorl was defined as one with at least three branches. All measurements were taken commencing on the South side of the tree and progressing anticlockwise around the stem. When more than three branches were available for measurement on the whorl the largest three branches, by basal diameter, were used for sampling. The following branch characteristics were measured after six field growing seasons.

Branch diameter: Diameter of each measured branch was taken at its base. Using a caliper the diameter of the branch, over bark, was measured to the closest 16th of an inch. 'CF

LP1-7696' has an average branch diameter of 0.63 inches at the base of the branch. Zygotic seedlings of the same genetic origin have an average branch diameter of 0.98 inches at the base of the branch.

5 Branch angle: Utilizing a large protractor, the angle of each branch was measured as its deviation from horizontal. Branch angles were recorded for the portion of the branch emerging from the stem of the tree with data rounded to the closest 10 degrees. 'CF LP1-7696' has an average branch angle of 29.50 degrees from horizontal. Zygotic seedlings of the same genetic origin have an average branch angle of 29.44 degrees from horizontal.

Branch length: The length of each sampled branch was measured directly with a graduated measurement pole. Branch lengths were recorded to the closest 0.5 feet. 'CF LP1-7696' has an average branch length of 6.48 feet. Zygotic seedlings of the same genetic origin have an average branch length of 8.8 feet.

20 Crown diameter: The width of the crown, at the point where branch measurements were taken, was directly measured with the use of a graduated measurement pole. A radial measurement was taken on the East and West side of each tree. Crown radius was measured to the closest 0.5 feet. Crown width data is presented as diameter of the crown. 'CF LP1-7696' has an average crown diameter of 8.65 feet. Zygotic seedlings of the same genetic origin have an average crown diameter of 11.37 feet.

Internode length: In proximity to the area of the stem utilized for branch measurements the mean internode length was determined for each tree. Internode distances for the calculation of the mean were directly measured from the stem of the tree using a graduated measurement pole. 'CF LP1-7696' has an average internode length of 1.22 feet. Zygotic seedlings of the same genetic origin have an average internode length of 1.25 feet.

Maximum trunk diameter at breast height (4.5 feet above the soil level): 4.4 inches after 5 field growing seasons

Percent stem fusiform rust infection at age 5:0

40 Percent branch fusiform rust infection at age 5:5

Percent branch and stem fusiform rust infection at age 5:0

Percent dead ramets due to fusiform rust infection at age 5:0

Percent stem fusiform rust infection in the USDA Resistance Screening Center (Asheville, N.C.) tests after artificial inoculation with rust spores: 31% (compared to 76% infection in unimproved seedlings)

Propagation: propagated by somatic embryogenesis

Seeds: none produced at age 5 years of age, plants are not yet mature. Expected seed production by 12-15 years of age.

50 Use: high yield industrial plantations

Although the new variety of loblolly pine tree possesses the detailed characteristics noted above as a result of the growing conditions prevailing in the test locations, it is to be understood that the variations of the usual magnitude and characteristics incident to changes in growing conditions, irrigation, fertilization, pruning, pest control, climatic variations and the like are to be expected. An example of 'CF LP1-7696' can be found at The Campbell Group Beulah year 2003 line trial, Angelina county, Tex.

COMPARISON WITH PARENTS BY MICROSATELLITE ANALYSIS

65 Microsatellite markers (SSR's) were used to generate a unique DNA fingerprint for the variety. Young foliage samples from 6 ramets of LP1-7696 variety and from the

parental trees used to make the LP1 cross were collected for DNA fingerprinting. The DNA extraction protocol of Doyle and Doyle (1987) was used after slight modifications. DNA fingerprinting of parents and the LP1-7696 variety was conducted using a set of nine microsatellite markers (Elsik et al., 2000; Auckland et al., 2002; Echt et al., 2008). Table 1 shows the sequences and conditions for each primer.

TABLE 1

ID's, sequences and conditions of SSR primers used in loblolly pine LP1-7696 variety.				
Primer full ID	UniSTS #	GenBank accession	SEQUENCE (5'-3') (SEQ ID NO:)	
PtTX3011	508455	BV728852	F: AATTTGGGTGTATTTTCT TAGA (SEQ ID NO: 1) R: AAAAGTTGAAGGAGTTG GTGATC (SEQ ID NO: 2)	
PtTX3025	508459	BV728855	F: CACGCTGTATAATAACAA TCTA (SEQ ID NO: 3) R: GGATAACAATTTACACA GGTTCTATATTCGCTTTAG TTTC (SEQ ID NO: 4)	
PtTX3034	508463	BV728857	F: CACGACGTTGTAAAACGA CTCAAATGCAAAAGACG (SEQ ID NO: 5) R: ATTAGGACTGGGGATGA T (SEQ ID NO: 6)	
PtTX3049	508467	BV728826	F: GAAGTGATAATGGCATA GCAAAAT (SEQ ID NO: 7) R: GCAGACCCGTGAAAGTA ATAAACAT (SEQ ID NO: 8)	
PtTX3105	508475	BV728847	F: TGTCGGTGGAGTTGGCAG TAGACT (SEQ ID NO: 9) R: GCCCAGCGTTTCCTG (SEQ ID NO: 10)	
PtTX3116	508479	BV728848	F: CACGACGTTGTAAAACGA CCTCCCAAAGCCTAAAGAA T (SEQ ID NO: 11) R: CATACAAGGCCTTATCTT ACAGAA (SEQ ID NO: 12)	
PtTX3127	508483	BV728849	F: ACCCTTACTTTCAGAGA GGATA (SEQ ID NO: 13) R: GGATAACAATTTACACA GGAATTGGGGTTCACTATT CTATTA (SEQ ID NO: 14)	
PtSIFG_0566	516281	BV728755	F: CACGACGTTGTAAAACGA CACTTAGTGGGAAAGGGGG AA (SEQ ID NO: 15) R: GTTCTTTTCCTCAGCCA AAAGCTCTC (SEQ ID NO: 16)	
PtSIFG_4233	516353	BV728685	F: CACGACGTTGTAAAACGA CAGGGAAACCGCGGATTAT AG (SEQ ID NO: 17) R: GTTCTTCCGGAATGAAG ATTGCAGTT (SEQ ID NO: 18)	
Primer full ID	LABEL TAIL (F/R); E (end labeled)		MgCl ₂ (mM)	Ta (° C)
PtTX3011	E		2.5	55
PtTX3025	R		2.5	61
PtTX3034	F		2.5	55
PtTX3049	E		2.5	59 → 49
PtTX3105	E		2.5	59 → 49
PtTX3116	F		2.5	55 → 45
PtTX3127	R		2.5	61
PtSIFG_0566	F		2.5	65 → 55
PtSIFG_4233	F		2.5	65 → 55

Ta = primer annealing temperature.

Microsatellite products were detected by M13 tailed primer (Oetting et al., 1995) or infrared dye(IRD)-labeled primer. The amplification products were electrophoresed on

5.5% Long Ranger polyacrylamide gels using a LiCor 4200 automated sequencer (LiCor Inc., Lincoln, Neb.).

The observed parental genotypes and their expected offspring's genotypes at nine studied SSR loci are presented in Table 2. LP1-7696 fingerprint based on nine loci is presented in Table 3.

TABLE 2

Parental genotypes and their expected offspring's genotypes at nine SSR loci.					
Primer	Genotype		Expected offspring genotypes		
	Female	Male			
PtTX3011	157/193	157/193	157/157	157/193	193/193
PtTX3025	277/289	274/277	277/274	227/227	289/274 289/277
PtTX3127	207/210	204/207	207/204	207/207	210/204 210/207
PtTX3034	228/228	216/220	228/216	228/220	
PtTX3049	311/313	323/325	311/323	311/325	313/323 313/325
PtTX3116	147/150	159/180	147/159	147/180	150/159 150/180
PtTX3105	169/190	169/184	169/169	169/184	190/169 190/184
SIFG0566	133/139	145/145	133/145	139/145	
SIFG4233	127/127	129/137	127/129	127/137	

TABLE 3

LP1-7696 genotypes at nine SSR loci. Allelic sizes include LiCor primer tails for M13 tailed primers.					
PtTX3011		PtTX3025		PtTX3127	
Allele1	Allele2	Allele1	Allele2	Allele1	Allele2
157	193	274	289	204	207
PtTX3034		PtTX3049		PtTX3116	
Allele1	Allele2	Allele1	Allele2	Allele1	Allele2
216	228	311	325	150	180
PtTX3105		SIFG0566		SIFG4233	
Allele1	Allele2	Allele1	Allele2	Allele1	Allele2
169	190	139	145	127	137

References:
Auckland, L., T. Bui, Y. Zhou, M. Shepherd and C. Williams. 2002. Conifer Microsatellite Handbook Corporate Press, Raleigh, N.C., USA.
Doyle, J.J. and J.L. Doyle. 1987. A rapid DNA isolation procedure for small quantities of fresh tissue. Phytochemical bulletin 19:11-15.
Echt, C.S., Saha, S. and Nelson, C.D. 2008. Southern Institute of Forest Genetics, USDA Forest Service Southern Research Station, 23332 Mississippi 67, Saucier, MS 39574, USA. On-line genetic database: <http://www.ncbi.nlm.nih.gov/unists>
Elsik, C.G., Minihan, V.T., Hall, S.E., Scarpa, A.M. and Williams, C.G. 2000. Low-copy microsatellite markers for *Pinus taeda* L. Genome 43(3):550-555.
Oetting, W.S., H.K. Lee, D.J. Flanders, G.L. Wiesner, T.A. Sellers and R.A. King. 1995. Linkage analysis with multiplexed short tandem repeat polymorphisms using infrared fluorescence and M13 tailed primers. Genomics 30:450-458.

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We claim:

1. A new and distinct variety of loblolly pine tree named ‘CF LP1-7696’ substantially as described and illustrated.

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FIG. 1



FIG. 2