Johnie N Jenkins

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114
papers1,985
citations26
h-index38
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ext. papers2,452
ext. citations3.1
avg, IF4.65
L-index

#	Paper	IF	Citations
114	Development of a 63K SNP Array for Cotton and High-Density Mapping of Intraspecific and Interspecific Populations of Gossypium spp. <i>G3: Genes, Genomes, Genetics</i> , 2015 , 5, 1187-209	3.2	117
113	Quantitative analysis and QTL mapping for agronomic and fiber traits in an RI population of upland cotton. <i>Euphytica</i> , 2009 , 165, 231-245	2.1	72
112	A MAGIC population-based genome-wide association study reveals functional association of GhRBB1_A07 gene with superior fiber quality in cotton. <i>BMC Genomics</i> , 2016 , 17, 903	4.5	71
111	Effects of chromosome-specific introgression in upland cotton on fiber and agronomic traits. <i>Genetics</i> , 2006 , 172, 1927-38	4	70
110	Quantitative trait loci analysis of fiber quality traits using a random-mated recombinant inbred population in Upland cotton (Gossypium hirsutum L.). <i>BMC Genomics</i> , 2014 , 15, 397	4.5	69
109	Association of Root-Knot Nematode Resistance Genes with Simple Sequence Repeat Markers on Two Chromosomes in Cotton. <i>Crop Science</i> , 2006 , 46, 2670-2674	2.4	51
108	SSR markers closely associated with genes for resistance to root-knot nematode on chromosomes 11 and 14 of Upland cotton. <i>Theoretical and Applied Genetics</i> , 2010 , 121, 1323-37	6	49
107	Genetic Effects of Thirteen Gossypium barbadense L. Chromosome Substitution Lines in Topcrosses with Upland Cotton Cultivars: II. Fiber Quality Traits. <i>Crop Science</i> , 2007 , 47, 561-570	2.4	49
106	Genetic Effects of Thirteen Gossypium barbadense L. Chromosome Substitution Lines in Topcrosses with Upland Cotton Cultivars: I. Yield and Yield Components. <i>Crop Science</i> , 2006 , 46, 1169-1	1 78	48
105	Transcriptome profiling, sequence characterization, and SNP-based chromosomal assignment of the EXPANSIN genes in cotton. <i>Molecular Genetics and Genomics</i> , 2007 , 278, 539-53	3.1	45
104	Primitive Accession Derived Germplasm by Cultivar Crosses as Sources for Cotton Improvement. <i>Crop Science</i> , 2004 , 44, 1226-1230	2.4	45
103	Trend Analysis and Forecast of Precipitation, Reference Evapotranspiration, and Rainfall Deficit in the Blackland Prairie of Eastern Mississippi. <i>Journal of Applied Meteorology and Climatology</i> , 2016 , 55, 1425-1439	2.7	43
102	QTLs for node of first fruiting branch in a cross of an upland cotton, Gossypium hirsutum L., cultivar with primitive accession Texas 701. <i>Euphytica</i> , 2008 , 163, 113-122	2.1	41
101	Phytochrome RNAi enhances major fibre quality and agronomic traits of the cotton Gossypium hirsutum L. <i>Nature Communications</i> , 2014 , 5, 3062	17.4	40
100	Use of fiber and fuzz mutants to detect QTL for yield components, seed, and fiber traits of upland cotton. <i>Euphytica</i> , 2010 , 172, 21-34	2.1	40
99	Identification of QTL regions and SSR markers associated with resistance to reniform nematode in Gossypium barbadense L. accession GB713. <i>Theoretical and Applied Genetics</i> , 2011 , 122, 271-80	6	39
98	F2 Hybrids of Host Plant Germplasm and Cotton Cultivars: I. Heterosis and Combining Ability for Lint Yield and Yield Components. <i>Crop Science</i> , 1993 , 33, 700-705	2.4	38

(2008-2008)

97	Cotton (Gossypium spp.) R2R3-MYB transcription factors SNP identification, phylogenomic characterization, chromosome localization, and linkage mapping. <i>Theoretical and Applied Genetics</i> , 2008 , 116, 1015-26	6	37	
96	Detection, Validation, and Application of Genotyping-by-Sequencing Based Single Nucleotide Polymorphisms in Upland Cotton. <i>Plant Genome</i> , 2015 , 8, eplantgenome2014.07.0034	4.4	35	
95	Genetic variance components and genetic effects among eleven diverse upland cotton lines and their F2 hybrids. <i>Euphytica</i> , 2009 , 167, 397-408	2.1	34	
94	F2 Hybrids of Host Plant Germplasm and Cotton Cultivars: II. Heterosis and Combining Ability for Fiber Properties. <i>Crop Science</i> , 1993 , 33, 706-710	2.4	31	
93	Primitive Accession Derived Germplasm by Cultivar Crosses as Sources for Cotton Improvement. <i>Crop Science</i> , 2004 , 44, 1231-1235	2.4	30	
92	Genetic dissection of chromosome substitution lines of cotton to discover novel Gossypium barbadense L. alleles for improvement of agronomic traits. <i>Theoretical and Applied Genetics</i> , 2010 , 120, 1193-205	6	27	
91	Genetic and transcriptomic dissection of the fiber length trait from a cotton (Gossypium hirsutum L.) MAGIC population. <i>BMC Genomics</i> , 2019 , 20, 112	4.5	26	
90	An additive-dominance model to determine chromosomal effects in chromosome substitution lines and other gemplasms. <i>Theoretical and Applied Genetics</i> , 2006 , 112, 391-9	6	26	
89	Whole genome sequencing of a MAGIC population identified genomic loci and candidate genes for major fiber quality traits in upland cotton (Gossypium hirsutum L.). <i>Theoretical and Applied Genetics</i> , 2019 , 132, 989-999	6	26	
88	Genetic association of yield with its component traits in a recombinant inbred line population of cotton. <i>Euphytica</i> , 2004 , 140, 171-179	2.1	24	
87	Molecular and cytological characterization of a cytoplasmic-specific mutant in pima cotton (Gossypium barbadense L.). <i>Euphytica</i> , 2004 , 139, 187-197	2.1	23	
86	Use of Primitive Derived Cotton Accessions for Agronomic and Fiber Traits Improvement: Variance Components and Genetic Effects. <i>Crop Science</i> , 2007 , 47, 100-110	2.4	22	
85	SSR markers for marker assisted selection of root-knot nematode (Meloidogyne incognita) resistant plants in cotton (Gossypium hirsutum L). <i>Euphytica</i> , 2012 , 183, 49-54	2.1	21	
84	Molecular cloning, differential expression, and functional characterization of a family of class I ubiquitin-conjugating enzyme (E2) genes in cotton (Gossypium). <i>Biochimica Et Biophysica Acta Gene Regulatory Mechanisms</i> , 2003 , 1625, 269-79		21	
83	Combining Ability Analysis of Root-Knot Nematode Resistance in Cotton. <i>Crop Science</i> , 1995 , 35, 373	2.4	21	
82	Seed trait evaluation of Gossypium barbadense L. chromosomes/arms in a G. hirsutum L. background. <i>Euphytica</i> , 2009 , 167, 371-380	2.1	20	
81	Cotton chromosome substitution lines crossed with cultivars: genetic model evaluation and seed trait analyses. <i>Theoretical and Applied Genetics</i> , 2010 , 120, 1473-83	6	20	
80	Defining the experimental unit for the design and analysis of site-specific experiments in commercial cotton fields. <i>Agricultural Systems</i> , 2008 , 96, 237-249	6.1	19	

79	BAC-derived SSR markers chromosome locations in cotton. <i>Euphytica</i> , 2008 , 161, 361-370	2.1	19
78	Evaluation of Reference Evapotranspiration Methods in Arid, Semiarid, and Humid Regions. <i>Journal of the American Water Resources Association</i> , 2017 , 53, 791-808	2.1	18
77	Small RNA sequencing and degradome analysis of developing fibers of short fiber mutants Ligon-lintles-1 (Li 1) and -2 (Li 2) revealed a role for miRNAs and their targets in cotton fiber elongation. <i>BMC Genomics</i> , 2016 , 17, 360	4.5	18
76	Assessment of Surface Water Resources in the Big Sunflower River Watershed Using Coupled SWATMODFLOW Model. <i>Water (Switzerland)</i> , 2019 , 11, 528	3	17
75	Genetic association of cotton yield with its component traits in derived primitive accessions crossed by elite upland cultivars using the conditional ADAA genetic model. <i>Euphytica</i> , 2008 , 161, 337-3	352 ¹	17
74	Genetic effects of nine Gossypium barbadense L. chromosome substitution lines in top crosses with five elite Upland cotton G. hirsutum L. cultivars. <i>Euphytica</i> , 2012 , 187, 161-173	2.1	16
73	Evaluation of genomic selection methods for predicting fiber quality traits in Upland cotton. <i>Molecular Genetics and Genomics</i> , 2020 , 295, 67-79	3.1	16
72	Molecular characterization and temporal expression analyses indicate that the MIC (Meloidogyne Induced Cotton) gene family represents a novel group of root-specific defense-related genes in upland cotton (Gossypium hirsutum L.). <i>Planta</i> , 2008 , 228, 111-23	4.7	15
71	Genetic diversity for agronomic and fiber traits in day-neutral accessions derived from primitive cotton germplasm. <i>Euphytica</i> , 2006 , 148, 283-293	2.1	15
70	Registration of 79 Day-Neutral Primitive Cotton Germplasm Lines. <i>Crop Science</i> , 1993 , 33, 351	2.4	15
69	Simulating soybean productivity under rainfed conditions for major soil types using APEX model in East Central Mississippi. <i>Agricultural Water Management</i> , 2016 , 177, 379-391	5.9	15
68	Genetic association of lint yield with its components in cotton chromosome substitution lines. <i>Euphytica</i> , 2008 , 164, 199-207	2.1	14
67	A Model to Estimate Hydrological Processes and Water Budget in an Irrigation Farm Pond. <i>Water Resources Management</i> , 2017 , 31, 2225-2241	3.7	13
66	Detecting epistatic effects associated with cotton traits by a modified MDR approach. <i>Euphytica</i> , 2012 , 187, 289-301	2.1	13
65	A GENERALIZED APPROACH AND COMPUTER TOOL FOR QUANTITATIVE GENETICS STUDY. Conference on Applied Statistics in Agriculture,		13
64	Simulating the Fate of Fall- and Spring-Applied Poultry Litter Nitrogen in Corn Production. <i>Soil Science Society of America Journal</i> , 2015 , 79, 1804-1814	2.5	12
63	Characterization and promoter analysis of a cotton RING-type ubiquitin ligase (E3) gene. <i>Molecular Biotechnology</i> , 2010 , 46, 140-8	3	12
62	Molecular and SNP characterization of two genome specific transcription factor genes GhMyb8 and GhMyb10 in cotton species. <i>Euphytica</i> , 2007 , 159, 259-273	2.1	12

(2016-2006)

Effects of chromosome 5sh from Gossypium barbadense L. on flower production in G. hirsutum L <i>Euphytica</i> , 2006 , 152, 99-107	2.1	12	
Poultry Litter and Cover Crop Integration into No-till Cotton on Upland Soil. <i>Agronomy Journal</i> , 2019 , 111, 2097-2107	2.2	12	
Molecular confirmation of Gossypium hirsutum chromosome substitution lines. <i>Euphytica</i> , 2015 , 205, 459-473	2.1	11	
Integrative placement and orientation of non-redundant SSR loci in cotton linkage groups by deficiency analysis. <i>Molecular Breeding</i> , 2009 , 23, 693-707	3.4	11	
A novel root-specific gene, MIC-3, with increased expression in nematode-resistant cotton (Gossypium hirsutum L.) after root-knot nematode infection. <i>Biochimica Et Biophysica Acta Gene Regulatory Mechanisms</i> , 2002 , 1576, 214-8		11	
Rainwater Deficit and Irrigation Demand for Row Crops in Mississippi Blackland Prairie. <i>Soil Science Society of America Journal</i> , 2018 , 82, 423-435	2.5	11	
The P450 gene CYP749A16 is required for tolerance to the sulfonylurea herbicide trifloxysulfuron sodium in cotton (Gossypium hirsutum L.). <i>BMC Plant Biology</i> , 2018 , 18, 186	5.3	11	
Subsurface Band Placement of Pelletized Poultry Litter in Cotton. <i>Agronomy Journal</i> , 2016 , 108, 1356-	1366	10	
Broiler Litter Type and Placement Effects on Corn Growth, Nitrogen Utilization, and Residual Soil Nitrate-Nitrogen in a No-Till Field. <i>Agronomy Journal</i> , 2012 , 104, 43-48	2.2	10	
TESTING VARIANCE COMPONENTS BY TWO JACKKNIFE METHODS. Conference on Applied Statistics in Agriculture,		10	
Rain Water Deficit and Irrigation Demand of Major Row Crops in the Mississippi Delta. <i>Transactions of the ASABE</i> , 2018 , 61, 927-935	0.9	10	
Molecular characterization of the reniform nematode C-type lectin gene family reveals a likely role in mitigating environmental stresses during plant parasitism. <i>Gene</i> , 2014 , 537, 269-78	3.8	9	
Registration of 16 Day Length-Neutral Flowering Primitive Cotton Germplasm Lines. <i>Crop Science</i> , 2002 , 42, 1755-1756	2.4	9	
Estimating the ratio of pond size to irrigated soybean land in Mississippi: a case study. <i>Water Science and Technology: Water Supply</i> , 2016 , 16, 1639-1647	1.4	9	
Overexpression of MIC-3 indicates a direct role for the MIC gene family in mediating Upland cotton (Gossypium hirsutum) resistance to root-knot nematode (Meloidogyne incognita). <i>Theoretical and Applied Genetics</i> , 2015 , 128, 199-209	6	8	
Genetic analysis without replications: model evaluation and application in spring wheat. <i>Euphytica</i> , 2013 , 190, 447-458	2.1	8	
Clustering, haplotype diversity and locations of MIC-3: a unique root-specific defense-related gene family in Upland cotton (Gossypium hirsutum L.). <i>Theoretical and Applied Genetics</i> , 2010 , 120, 587-606	6	8	
Optimum Poultry Litter Rates for Maximum Profit versus Yield in Cotton Production. <i>Crop Science</i> , 2016 , 56, 3307-3317	2.4	8	
	Euphytica, 2006, 152, 99-107 Poultry Litter and Cover Crop Integration into No-till Cotton on Upland Soil. Agranomy Journal, 2019, 111, 2097-2107 Molecular confirmation of Gossypium hirsutum chromosome substitution lines. Euphytica, 2015, 205, 459-473 Integrative placement and orientation of non-redundant SSR loci in cotton linkage groups by deficiency analysis. Molecular Breeding, 2009, 23, 693-707 A novel root-specific gene, MIC-3, with increased expression in nematode-resistant cotton (Gossypium hirsutum L.) after root-knot nematode infection. Biochimica Et Biophysica Acta Gene Regulatory Mechanisms, 2002, 1576, 214-8 Rainwater Deficit and Irrigation Demand for Row Crops in Mississippi Blackland Prairie. Soil Science Society of America Journal, 2018, 82, 423-435 The P450 gene CYP749A16 is required for tolerance to the sulfonylurea herbicide trifloxysulfuron sodium in cotton (Gossypium hirsutum L.). BMC Plant Biology, 2018, 18, 186 Subsurface Band Placement of Pelletized Poultry Litter in Cotton. Agronomy Journal, 2016, 108, 1356-Broiler Litter Type and Placement Effects on Corn Growth, Nitrogen Utilization, and Residual Soil Nitrate-Nitrogen in a No-Till Field. Agranomy Journal, 2012, 104, 43-48 TESTING VARIANCE COMPONENTS BY TWO JACKKNIFE METHODS. Conference on Applied Statistics in Agriculture, Rain Water Deficit and Irrigation Demand of Major Row Crops in the Mississippi Delta. Transactions of the ASABE, 2018, 61, 927-935 Molecular characterization of the reniform nematode C-type lectin gene family reveals a likely role in militating environmental stresses during plant parasitism. Gene, 2014, 537, 269-78 Registration of 16 Day Length-Neutral Flowering Primitive Cotton Germplasm Lines. Crop Science, 2002, 42, 1755-1756 Estimating the ratio of pond size to irrigated soybean land in Mississippi: a case study. Water Science and Technology: Water Supply, 2016, 16, 1639-1647 Overexpression of MIC-3 indicates a direct role for the MIC gene family in mediating Upland cotton (Gossypium hirsutum L.). T	Euphytica, 2006, 152, 99-107 Poultry Litter and Cover Crop Integration into No-till Cotton on Upland Soil. Agronomy Journal, 2019, 111, 2097-2107 Molecular confirmation of Gossypium hirsutum chromosome substitution lines. Euphytica, 2015, 205, 459-473 Integrative placement and orientation of non-redundant SSR loci in cotton linkage groups by deficiency analysis. Malecular Breeding, 2009, 23, 693-707 A novel root-specific gene, MIC-3, with increased expression in nematode-resistant cotton (Gossypium hirsutum L.) after rook-knot nematode infection. Biochimica Et Biophysica Acta Gene Regulatory Mechanisms, 2002, 1576, 214-8 Rainwater Deficit and Irrigation Demand for Row Crops in Mississippi Blackland Prairie. 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Gene, 2014, 537, 269-78 Registration of 16 Day Length-Neutral Flowering Primitive Cotton Germplasm Lines. Crop Science, 2002, 42, 1755-1756 Estimating the ratio of pond size to irrigated soybean land in Mississippi: a case study. Water Science and Technology: Water Supply, 2016, 16, 1639-1647 Overexpression of MIC-3 indicates a direct role for the MIC gene family in mediating Upland cotton (Gossypium hirsutum)	Poultry Litter and Cover Crop Integration into No-till Cotton on Upland Soil. Agronomy Journal, 2019, 111, 2097-2107 Molecular confirmation of Gossyplum hirsutum chromosome substitution lines. Euphytica, 2015, 205, 459-473 Molecular confirmation of Gossyplum hirsutum chromosome substitution lines. Euphytica, 2015, 205, 459-473 An ovel root-specific gene, MIC-3, with increased expression in nematode-resistant cotton (Gossyplum hirsutum L) after root-knot nematode infection. Biochimica Et Biophysica Acta Gene Regulatory Mechanisms, 2002, 1576, 214-8 Rainwater Deficit and Irrigation Demand for Row Crops in Mississippi Blackland Prairie. Soil Science Society of America Journal, 2018, 82, 423-435 The PASO gene CYPTA9A16 is required for tolerance to the sulfonylurea herbicide trifloxysulfuron sodium in cotton (Gossypium hirsutum L). BMC Plant Biology, 2018, 18, 186 Subsurface Band Placement of Pelletized Poultry Litter in Cotton. Agronomy Journal, 2016, 108, 1356-1366 In Broiler Litter Type and Placement Effects on Corn Growth, Nitrogen Utilization, and Residual Soil Nitrate-Nitrogen in a No-Till Field. Agronomy Journal, 2012, 104, 43-48 TESTING VARIANCE COMPONENTS BY TWO JACKKNIFE METHODS. Conference on Applied Statistics in Agraculture, Rain Water Deficit and Irrigation Demand of Major Row Crops in the Mississippi Delta. 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43	A novel variant of Gh_D02G0276 is required for root-knot nematode resistance on chromosome 14 (D02) in Upland cotton. <i>Theoretical and Applied Genetics</i> , 2019 , 132, 1425-1434	6	8
42	Individual and combined contributions of the Ren barb1, Ren barb2, and Ren barb3 quantitative trait loci to reniform nematode (Rotylenchulus reniformis Linford & Oliveira) resistance in upland cotton (Gossypium hirsutum L.). <i>Euphytica</i> , 2017 , 213, 1	2.1	7
41	Effects of inter-species chromosome substitution on cottonseed mineral and protein nutrition profiles. <i>Agronomy Journal</i> , 2020 , 112, 3963-3974	2.2	7
40	Genetic mapping of non-target-site resistance to a sulfonylurea herbicide (Envoke□) in Upland cotton (Gossypium hirsutum L.). <i>Molecular Breeding</i> , 2014 , 33, 341-348	3.4	7
39	A recursive approach to detect multivariable conditional variance components and conditional random effects. <i>Computational Statistics and Data Analysis</i> , 2006 , 50, 285-300	1.6	7
38	Consequences of pelletized poultry litter applications on soil physical and hydraulic properties in reduced tillage, continuous cotton system. <i>Soil and Tillage Research</i> , 2019 , 194, 104309	6.5	6
37	Tri-Species Shuffling of Chromosomes to Study the Effects on Fiber Traits Using Chromosome Substitution Lines. <i>Crop Science</i> , 2017 , 57, 1211-1226	2.4	6
36	A categorical, improper probability method for combining NDVI and LiDAR elevation information for potential cotton precision agricultural applications. <i>Computers and Electronics in Agriculture</i> , 2012 , 82, 15-22	6.5	6
35	Designing experiments to evaluate the effectiveness of precision agricultural practices on research fields: part 1 concepts for their formulation. <i>Operational Research</i> , 2010 , 10, 329-348	1.6	6
34	Management Strategies on an Upland Soil for Improving Soil Properties. <i>Communications in Soil Science and Plant Analysis</i> , 2020 , 51, 413-429	1.5	6
33	Harvest Management Effects on Tifton 44 Bermudagrass Phosphorus Removal and Nutritive Value. <i>Agronomy Journal</i> , 2018 , 110, 879-889	2.2	6
32	Morph-physiological responses of cotton interspecific chromosome substitution lines to low temperature and drought stresses. <i>Euphytica</i> , 2018 , 214, 1	2.1	6
31	Genotypic comparisons of chromosomes 01, 04, and 18 from three tetraploid species of Gossypium in topcrosses with five elite cultivars of G. hirsutum L <i>Euphytica</i> , 2017 , 213, 1	2.1	5
30	A cotton-fiber-associated cyclin-dependent kinase a gene: characterization and chromosomal location. <i>International Journal of Plant Genomics</i> , 2012 , 2012, 613812		5
29	QTL mapping for flowering-time and photoperiod insensitivity of cotton Gossypium darwinii Watt. <i>PLoS ONE</i> , 2017 , 12, e0186240	3.7	5
28	Genetic Effects of Chromosomes 1, 4, and 18 from Three Tetraploid Gossypium Species in Topcrosses with Five Elite Cultivars. <i>Crop Science</i> , 2017 , 57, 1338-1346	2.4	4
27	Molecular evolution of the clustered MIC-3 multigene family of Gossypium species. <i>Theoretical and Applied Genetics</i> , 2011 , 123, 1359-73	6	4
26	Genetic effects of individual chromosomes in cotton cultivars detected by using chromosome substitution lines as genetic probes. <i>Genetica</i> , 2010 , 138, 1171-9	1.5	4

(1999-2020)

25	High-Temperature and Drought-Resilience Traits among Interspecific Chromosome Substitution Lines for Genetic Improvement of Upland Cotton. <i>Plants</i> , 2020 , 9,	4.5	4
24	Application of the Cottonscope for determining fiber maturity and fineness of an upland cotton MAGIC population. <i>Crop Science</i> , 2020 , 60, 2266-2279	2.4	3
23	Introgression of Gossypium barbadense L. into Upland cotton germplasm RMBUP-C4S1. <i>Euphytica</i> , 2018 , 214, 1	2.1	3
22	Effects of Interspecific Chromosome Substitution in Upland Cotton on Cottonseed Micronutrients. <i>Plants</i> , 2020 , 9,	4.5	3
21	Comparative assessment of genetic diversity in cytoplasmic and nuclear genome of upland cotton. <i>Genetica</i> , 2016 , 144, 289-306	1.5	3
20	Genetic diversity of day-neutral converted landrace Gossypium hirsutum L. accessions. <i>Euphytica</i> , 2018 , 214, 1	2.1	3
19	Field experimental design comparisons to detect field effects associated with agronomic traits in upland cotton. <i>Euphytica</i> , 2015 , 206, 747-757	2.1	2
18	Analysis of Cotton Chromosome 11 and 14 Root-Knot Nematode Resistance Quantitative Trait Loci Effects on Root-Knot Nematode Postinfection Development, Egg Mass Formation, and Fecundity. <i>Phytopathology</i> , 2020 , 110, 927-932	3.8	2
17	Registration of four upland cotton germplasm lines with elevated levels of seed oil oleic acid. Journal of Plant Registrations, 2020 , 14, 64-71	0.7	2
16	Coupling of MIC-3 overexpression with the chromosomes 11 and 14 root-knot nematode (RKN) (Meloidogyne incognita) resistance QTLs provides insights into the regulation of the RKN resistance response in Upland cotton (Gossypium hirsutum). <i>Theoretical and Applied Genetics</i> , 2016 ,	6	2
15	Use of Bacillus thuringiensis Genes in Transgenic Cotton To Control Lepidopterous Insects. <i>ACS Symposium Series</i> , 1993 , 267-280	0.4	2
14	Effects on Selected Soil Properties of Subsurface Banding and Surface Broadcasting Pelletized Poultry Litter on Cotton. <i>Soil Science</i> , 2018 , 183, 112-120	0.9	2
13	Managing harvest of Russelland Tifton 44 Dermudagrass receiving broiler litter for nutritive value and phosphorus removal. <i>Crop, Forage and Turfgrass Management</i> , 2020 , 6, e20013	0.5	1
12	Effect of chromosome substitutions from Gossypium barbadense L., G. tomentosum Nutt. Ex Seem and G. mustelinum Watt into G. hirsutum L. on cottonseed protein and oil content. <i>Euphytica</i> , 2020 , 216, 1	2.1	1
11	The effect of cotton growth stage on response to a sublethal concentration of 2,4-D. <i>Weed Technology</i> , 2019 , 33, 321-328	1.4	1
10	Simulating Potential Weekly Stream and Pond Water Available for Irrigation in the Big Sunflower River Watershed of Mississippi Delta. <i>Water (Switzerland)</i> , 2019 , 11, 1271	3	1
9	Caloric Analyses of the Distribution of Energy in Ripened Cotton (Gossypium hirsutum L.). <i>Journal of Agricultural and Food Chemistry</i> , 1997 , 45, 3258-3261	5.7	1
8	Use of Cross-Species Simple Sequence Repeat (SSR) Primers for Developing Polymorphic DNA Markers. <i>Journal of New Seeds</i> , 1999 , 1, 25-37		1

7	Assessing the Weed-Suppressing Potential of Cotton Chromosome Substitution Lines Using the Stair-Step Assay. <i>Plants</i> , 2021 , 10,	4.5	1	
6	Alteration of root and shoot morphologies by interspecific replacement of individual Upland cotton chromosome or chromosome segment pairs. <i>Euphytica</i> , 2021 , 217, 1	2.1	1	
5	Registration of RMBHMTUP-C4, a Random-Mated Cotton Population Containing Alleles from Four Gossypium Species. <i>Journal of Plant Registrations</i> , 2019 , 13, 411-415	0.7	1	
4	Registration of Two CS-B17-derived Upland Cotton Recombinant Inbred Lines with Improved Fiber Micronaire. <i>Journal of Plant Registrations</i> , 2018 , 12, 97-100	0.7	1	
3	Genomic interrogation of a MAGIC population highlights genetic factors controlling fiber quality traits in cotton <i>Communications Biology</i> , 2022 , 5, 60	6.7	0	
2	Effect of physical damage to early- and late-maturing cotton cultivars. <i>Agronomy Journal</i> , 2020 , 112, 1140-1147	2.2		
1	Pelleted biosolids and cover crop effects on major Southern row crops. <i>Journal of Plant Nutrition</i> , 2021 , 44, 2677-2690	2.3		