

Simple sequence repeat (SSR)-based diversity analysis of groundnut (*Arachis hypogaea* L.) germplasm resistant to bacterial wilt

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Abstract

Groundnut is one of the most important oilseed crops in the world. Bacterial wilt, caused by *Ralstonia solanacearum* E. F. Smith, is one of the major biotic constraints to groundnut production particularly in South-East Asia and East Africa. Several sources of resistance to bacterial wilt have been identified through field screening of groundnut germplasm. The aim of the present study was to quantify the genetic diversity among selected bacterial wilt-resistant lines, in comparison with the levels of variation observable within the cultivated *A. hypogaea* gene pool. Thirty-two SSR markers were used to assess the degree of molecular polymorphism between 46 selected genotypes revealing 107 alleles, of which 101 (99.4%) were polymorphic with gene diversity scores ranging from 0.103 to 0.669, averaging 0.386. Cluster and multidimensional scaling analysis revealed two distinct groups within the germplasm broadly corresponding to the two subspecies (*hypogaea* and *fastigiata*) of *A. hypogaea*. However, accessions of varieties *peruviana* and *aequatoriana* grouped together with the varieties from subsp. *hypogaea*, rather than grouping with the other varieties of subsp. *fastigiata*. Analysis of molecular variance (AMOVA) revealed that 15% of the total observed variation was accounted for by disease response groups. This analysis will be useful in the selection of parental genotypes for mapping populations and breeding programmes attempting to broaden the genetic base of future groundnut cultivars. In particular, this opens up significant opportunities for the development of intraspecific mapping populations that will be highly relevant to modern groundnut breeding programmes.

Keywords: bacterial wilt; genetic diversity; groundnut; SSRs

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Introduction

Groundnut (*Arachis hypogaea* L.), also known as peanut, is one of the most important oilseed crops in the world. It is grown extensively in the Americas, Africa and Asia with a total annual global area of nearly 24 million hectares yielding 33.5 million tonnes. Around 53% of the global

production is crushed for edible oil, 32% for confectionery consumption and the remaining 15% is used for feed and seed production. Cultivated groundnut consists of two subspecies, *hypogaea* and *fastigiata*, which are further classified into six botanical varieties largely based on growth habit, flowering and branching patterns, presence of hairs on leaf surfaces and numbers of seeds per pod (Krapovickas and Gregory, 1994) and with lesser support by protein and genomic analysis (Smartt and Simmonds, 1995), with a number of recent genomic studies refuting this classification (He and Prakash, 2001; Raina *et al.*, 2001; Ferguson *et al.*, 2004a; He *et al.*, 2005; Tallury *et al.*, 2005). Botanical varieties *hypogaea* (Virginia) and *hirsuta* (Peruvian) are currently classified under subsp. *hypogaea* while varieties *fastigiata* (Valencia), *peruviana*, *aequatoriana* and *vulgaris* (Spanish) are assigned within subsp. *fastigiata*.

Bacterial wilt (BW), caused by *Ralstonia solanacearum* E. F. Smith, is a major biotic factor affecting groundnut production particularly in South-East Asia and East Africa (Hayward, 1990). It also infects many other crop plants including potato (*Solanum tuberosum* L.), tomato (*Lycopersicon esculentum* Mill), tobacco (*Nicotiana* spp.), pepper (*Capsicum* spp.), eggplant (*Solanum melongina* L.) and ginger (*Zingiber officinale* Rosc.). The bacterial species have been isolated and classified into five races based on host range (Buddenhagen and Kelman, 1964; He *et al.*, 1983) and five biovars based on biochemical characteristics (Hayward, 1964; He *et al.*, 1983). Race 1 causes wilt in groundnut, in addition to many other leguminous and solanaceous plants. Biovar 1 causes wilt in groundnut and predominantly occurs in America, whereas Biovars 3 and 4 cause wilt of groundnut in Asia and Africa (Hayward, 1991).

Extensive screening of groundnut germplasm, largely based on field evaluations at disease 'hot-spots' in Indonesia and China, has resulted in the identification of many BW-resistant lines. Most of these sources of BW resistance originate from China or Indonesia. Up to a quarter of germplasm accessions have shown some level of resistance to BW (Singh *et al.*, 1997; Pande *et al.*, 1998); resistance has been identified across different botanical types and also in some wild species. Despite the broad range of BW-resistant donor genotypes, only a few of the resistant landraces have been successfully used in breeding in China and Indonesia (Liao *et al.*, 1998). The majority of BW-resistant cultivars released in China were derived from just three sources (Xiekangqing, Taishan Sanlirou or Taishan Zhenzhu) (Liang, 1998; Liao *et al.*, 1998), while a single resistance donor source (Schwartz 21) has been the basis of the majority of cultivars released in Indonesia. Thus, there is a progressive narrowing of genetic diversity in BW-resistant breeding programmes that creates a major risk

for groundnut production should the pathogen overcome these sources of resistance. In addition, the currently available BW-resistant cultivars are low yielding and have poor tolerance to drought. Identification and utilization of a broad spectrum of genetically diverse sources of BW resistance is, therefore, critical for the development of a new generation of broad-based high-yielding BW-resistant groundnut cultivars. Limited knowledge about the genetic diversity of the BW-resistant germplasm and deleterious linkage drag have impeded the utilization of a wide spectrum of BW resistance donors.

Diversity studies in groundnut have generally revealed extensive phenotypic variation amongst varieties (Upadhyaya *et al.*, 2001, 2003) yet limited variation at the molecular level (Halward *et al.*, 1991, 1992; Kochert *et al.*, 1991, Paik-Rao *et al.*, 1992; He and Prakash, 1997; Subramanian *et al.*, 2000; Moretzsohn *et al.*, 2004). It is hypothesized that this may be due to the selective neutrality of the molecular markers utilized, while phenotypic traits have been subjected to intense selection (He and Prakash, 1997). It has also been suggested that the lack of molecular polymorphism revealed to date within the cultivated groundnut gene pool could be due to the inadequacy of the material studied and the range of techniques used (Singh *et al.*, 1998).

Microsatellite markers, also known as simple sequence repeat (SSR) markers, have been reported to detect high levels of polymorphism even amongst closely related cultivated germplasm (Gianfranceschi *et al.*, 1998). For this reason, considerable efforts have been recently made to develop a large number of SSR markers in groundnut (Hopkins *et al.*, 1999; He *et al.*, 2003, 2005; Ferguson *et al.*, 2004b; Moretzsohn *et al.*, 2004, 2005). It has been demonstrated in previous studies that SSR markers are more variable within genomes than other marker types (e.g. Belaj *et al.*, 2003). Additionally, SSRs have the advantage of being co-dominant, only requiring very small amounts of DNA and hence have been widely applied in many plant genetics studies, e.g. for evaluating genetic diversity (Zhebentyayaeva *et al.*, 2003; Fahima *et al.*, 1998), genome mapping and gene tagging, e.g. in rice (Chen *et al.*, 1997), wheat (Röder *et al.*, 1998), barley (e.g. Künzel and Waugh, 2002) and tomato (Broun and Tanksley, 1996). The recent development of groundnut-specific SSRs (Hopkins *et al.*, 1999; He *et al.*, 2003, 2005; Ferguson *et al.*, 2004b; Moretzsohn *et al.*, 2004, 2005) now offers new and exciting opportunities for groundnut genomics.

In the present study we have used groundnut-specific SSRs to analyse a diverse range of cultivated groundnut accessions encompassing all six botanical varieties. The purpose of this study was to investigate the level of molecular polymorphism amongst BW-resistant accessions and to compare this with the genetic diversity across

the cultivated *A. hypogaea* gene pool. This analysis is important for the selection of genetically diverse parental genotypes for mapping populations and BW resistance breeding programmes aimed at the development of broad-based cultivars with durable disease resistance.

Materials and methods

Plant material and DNA extraction

Thirty-one groundnut genotypes from the Oil Crops Research Institute (OCRI) of the Chinese Academy of Agricultural Sciences (CAAS) and 15 genotypes from the International Crops Research Institute for the Semi-Arid Tropics (ICRISAT), consisting of landraces, released cultivars and intraspecific derivatives representing all six varieties of *A. hypogaea*, were selected for the present study (Table 1).

Total genomic DNA was extracted using a CTAB-based procedure reported previously, with 3% (v/v) β -mercaptoethanol in a 3% (w/v) CTAB buffer (Mace *et al.*, 2003). The quantity and quality of DNA were determined electrophoretically through comparison with known concentrations of uncut λ DNA standards and spectrophotometric analysis at 260/280 nm, and subsequently diluted to 5 ng/ μ l.

SSR amplification

Thirty-two SSR markers were assayed for their ability to detect polymorphism among the 46 cultivated groundnut accessions selected (Table 2). The 32 SSRs were selected on the basis of prescreening approximately 200 groundnut SSRs, based on the level of polymorphism revealed between BW-resistant and -susceptible genotypes and the reliability and quality of amplicon detection.

PCR reactions were conducted in 20 μ l volumes using a GeneAmp PCR System 9700 (Applied Biosystems). The PCR reaction mixtures contained between 5 and 15 ng of genomic DNA, 10–30 pmol of each primer, 100–125 μ M of dNTP, 0.6–1.2 U/ μ l of *Taq* DNA polymerase (Amersham), 1 \times PCR buffer (10 mM Tris-HCl pH 8.3, 50 mM KCl) and 0.5–2.5 mM MgCl₂.

The fixed-temperature PCR programmes consisted of an initial denaturation step for 2 min at 94°C, followed by 35 cycles of denaturation for 45 s (94°C), annealing for 1 min (57–64°C; see Table 2) and extension for 1 min 30 s (72°C). The PCR products were then incubated at 72°C for a further 10 min to ensure complete extension. A second PCR programme using the touchdown approach was also used for selected SSRs (see Table 2) with the following conditions: initial denaturation for 2 min at 94°C, followed by 10 cycles: 94°C for 45 s, 65°C

(–1°C/cycle) for 1 min and 72°C for 1 min 30 s. This was then followed by 20 cycles of 94°C for 45 s, 55°C for 1 min and 72°C for 1 min 30 s, followed by a final extension step of 10 min (72°C).

Electrophoresis and data collection

PCR amplification products were separated on 6% non-denaturing polyacrylamide gels and revealed using a silver staining procedure based on ammoniacal solutions of silver, modified from Kolodny (1984). The size of the allele scored was determined through comparison with the 100 bp DNA ladder (Amersham) included on all gels. Estimates of similarity were based on two different measurements: (1) Nei and Li's (1979) definition of similarity: $S_{ij} = 2a/(2a + b + c)$, where S_{ij} is the similarity between two individuals, i and j , a is the number of bands present both in i and j , b is the number of bands present in i and absent in j , and c is the number of bands absent in i and present in j ; (2) Jaccard's coefficient (Jaccard, 1908): $S_{ij} = a/(a + b + c)$. The similarity matrices were then analysed using the clustering method UPGMA (unweighted pair group method; Sokal and Michener, 1958) using the NTSYS 2.1 software (Rohlf, 2001). The dendrograms were created with the tree program of NTSYS, and the goodness of fit of the clustering to the data was calculated using the COPH and MXCOMP program. Multidimensional scaling (MDS) (Kruskal and Wish, 1978) was also performed to confirm whether the observed molecular variation indicated evidence of clustering among accessions, as compared to the clustering by UPGMA.

An analysis of molecular variance (AMOVA) was undertaken to partition genetic variability using Arlequin software version 2.0 (Schneider *et al.*, 2000), and significance values assigned to variance components based on the random permutation (10,000 times) of individuals assuming no genetic structure. Additionally, the gene diversity (GD) of each SSR was determined as described by Weir (1990). $GD = 1 - \sum P_i^2$, where P_i is the frequency of the i th allele in the examined genotypes.

Results

All 32 SSRs successfully generated at least one allele in the region of the expected size in all 46 cultivated groundnut genotypes (Fig. 1; Table 2). A total of 107 alleles were observed following amplification of 29 polymorphic and three monomorphic (pPGPseq-13B06, pPGPseq-3D09 and A1-275) SSR loci, of which 101 (99.4%) were polymorphic. The total number of alleles revealed per polymorphic SSR locus ranged from two (pPGPseq-1B09) to 10 (pPGPseq-7H6) with an average of 3.34 alleles per locus. The observed allele sizes ranged from 131 bp

Table 1. Groundnut genotypes included in the current study with different levels of resistance (R) and susceptibility (S) to bacterial wilt (BW) (botanical variety and geographic origin also indicated)

Accession	Botanical variety	R/S to BW	Origin	Code
93-76 (Zhonghua No. 6)	<i>vulgaris</i>	R	China	1
Gouliaozhong	<i>hypogaea</i>	R	China	2
Qingmiaodou	<i>hirsuta</i>	R	China	3
Zao18	<i>vulgaris</i>	S	China	4
Dayebentianzi	<i>hirsuta</i>	S	China	5
9102	<i>vulgaris</i>	R	China	6
Yueyou200	<i>vulgaris</i>	R	China	7
Xiekangqing	<i>vulgaris</i>	R	China	8
Luao Wanhuasheng	<i>hirsuta</i>	R	China	9
Wuchang Laohuasheng	<i>hirsuta</i>	R	China	10
Zhonghua No. 2	<i>vulgaris</i>	R	China	11
Changsha Tuzihuasheng	<i>hirsuta</i>	R	China	12
Feilongxiang	<i>hirsuta</i>	R	China	13
Shitang Dahuasheng	<i>hirsuta</i>	R	China	14
Zhonghua 212	<i>vulgaris</i>	R	China	15
Taishan Zhenzhu	<i>vulgaris</i>	R	China	16
Jiangtianzhong	<i>hypogaea</i>	R	China	17
Qidong Dahuasheng	<i>hirsuta</i>	R	China	18
Nanning Sanjindou	<i>hirsuta</i>	R	China	19
Lingui Make	<i>hirsuta</i>	R	China	20
Chico (China)	<i>vulgaris</i>	S	China	21
Bobai Dahuasheng	<i>hirsuta</i>	R	China	22
Mashanguling	<i>hirsuta</i>	R	China	23
QiongxianXiaohongmao	<i>hirsuta</i>	R	China	24
Ehua No. 5	<i>vulgaris</i>	R	China	25
Zhongxingchi	<i>hirsuta</i>	R	China	26
Bobai Shiyadou	<i>hirsuta</i>	R	China	27
Taishan Sanlirou	<i>fastigiata</i>	R	China	28
91-074	<i>vulgaris</i>	R	China	29
Jiankang (89-15 048)	<i>vulgaris</i>	R	China	30
ICG 1704	<i>peruviana</i>	R	Peru	31
ICG 7894	<i>peruviana</i>	R	Peru	32
ICG 5276	<i>vulgaris</i>	R	Russia	33
ICG 14 159	<i>vulgaris</i>	Unknown	Vietnam	34
J 11	<i>vulgaris</i>	S	India	35
Gangapuri	<i>fastigiata</i>	S	India	36
ICG15222-1	<i>hypogaea</i>	R	China	37
ICG15222-2	<i>hypogaea</i>	R	China	38
Chico	<i>vulgaris</i>	S	USA	39
ICG 15 208	<i>hirsuta</i>	Unknown	Mexico	40
ICG 15 213	<i>hirsuta</i>	Unknown	Mexico	41
ICG 12 625	<i>aequatoriana</i>	Unknown	Ecuador	42
ICG 12 722	<i>aequatoriana</i>	Unknown	Ecuador	43
ICG 2381	<i>hypogaea</i>	Unknown	Brazil	44
ICG 3027	<i>hypogaea</i>	Unknown	India	45
Zhonghua No.5	<i>vulgaris</i>	S	China	46

(pPGPseq-8D9) to 531 bp (A1-193). The GD scores of the 29 polymorphic SSR loci ranged from 0.103 (pPGPseq-13E9) to 0.669 (pPGPseq-3A08) (Table 2).

Genetic diversity analysis of cultivated groundnut germplasm

Groundnut is a complex polyploid with two distinct genomes that largely segregate in isolation and is, therefore,

termed an amphidiploid (Burow *et al.*, 2001). This has significant implications in the appropriate choice of biometric analysis. In particular, SSRs may not always retain their codominant nature as defining allelic relationships becomes difficult when three or more alleles are detected in a single individual, however, this does not occur in any of the polymorphic loci scored in this study, with the exception of A1-275. For this reason cluster analysis was undertaken based on two different similarity measures: Jaccard (for dominant datasets) and

Table 2. Details of SSRs used in the groundnut diversity analysis (SSRs were developed by Ferguson *et al.*, 2004a unless indicated otherwise)

Primer name	Repeat motif	T _m (°C)	Expected product size (bp)	Observed product size(s) (bp)	Total no. of alleles observed	Gene diversity (GD)
pPGPseq-1B09	GA	64	282	268; 269	2	0.306
pPGPseq-2B10	TAA	58	259	268; 269	2	0.297
pPGPseq-2D12B	TAA	60	265	289; 300; 323; 333	4	0.57
pPGPseq-2E06	GA	60	250	269; 289; 306; 323	4	0.504
pPGPseq-2G03	TAA	64	215	254; 269; 281	3	0.459
pPGPseq-2G04	TAA	60	289	269; 289; 300; 333	4	0.47
pPGPseq-3A01	TAA	64	238	257; 269; 277; 289	4	0.304
pPGPseq-3A08	TAA	64	152	173; 178; 191; 197	4	0.669
pPGPseq-3B06	GA	61	244	157	1	0
pPGPseq-3B08	TAA	56	266	289; 300; 314; 323	4	0.555
pPGPseq-3D09	GA,GT	63	292	281	1	0
pPGPseq-4A06	AT	63	126	167; 173; 177	3	0.402
pPGPseq-7G2	TATC	65	225	223; 239; 246; 250; 257; 262	6	0.61
pPGPseq-7H6	CTT	60	300	308; 310	2	0.297
pPGPseq-8D9	CTT	61	132	131; 135; 146	3	0.468
pPGPseq-8E12	TTG,TAA	59	198	204; 207; 210; 214	4	0.485
pPGPseq-10H1A	CTT	58	139	193; 200	2	0.296
pPGPseq-12F7	TAA	57	290	305; 310	2	0.375
pPGPseq-13E9	TAA	59	299	323; 333	2	0.103
pPGPseq-13A7	TAA	58	265	289; 291; 293	3	0.44
pPGPseq-14A7	CTT,CTG	60	173	167; 173; 177	3	0.402
pPGPseq-14F4	TAA	60	163	173; 178; 184	3	0.402
pPGPseq-14H6	GT	59	285	269; 271; 284; 297; 306; 310; 312; 319; 333; 348	10	0.618
pPGPseq-15C10	TAA	64	203	212; 220	2	0.427
pPGPseq-16G8	TAA	60	194	214; 217; 227; 229	4	0.32
pPGPseq-18A5A	AT,TAA	60	268	300; 328	2	0.468
Lec-1 ^a	AT	65–55	120,125	243; 250; 261; 281; 300	5	0.473
Ah4-26 ^a	CT	65–55	160	173; 178; 184	3	0.616
A1-041 ^b	Unknown	65–55	230,350	269; 281; 293	3	0.351
A1-193 ^b	Unknown	65–55	460	510; 520; 531	3	0.403
A1-275 ^b	Unknown	65–55	190,300	181; 195; 305; 330	4	0
A1-745 ^b	Unknown	65–55	150,250	224; 226; 236	3	0.57

^a Microsatellites developed by Hopkins *et al.* (1999).

^b Microsatellites developed by Moretzsohn *et al.* (2004).

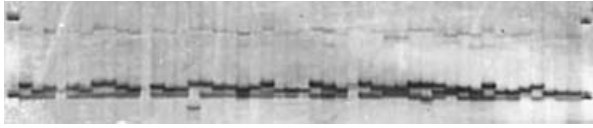


Fig. 1. Polymorphism detected by SSR pPGPseq-8E12 across 46 cultivated groundnut genotypes.

Dice/Nei and Li (for codominant datasets). The correlation coefficients were broadly similar from both Jaccard's similarity coefficient ($r = 0.89$) and Nei and Li's similarity coefficient ($r = 0.87$). Figure 2 shows the dendrogram produced by Jaccard's similarity coefficient using the UPGMA clustering method, with clear evidence of two separate clusters (A and B) at a level of approximately 36% similarity. The presence of two primary clusters within the data set was also confirmed through MDS (Fig. 3), where clusters A and B appear clearly separated on two axes. The 46 genotypes were all uniquely identified based on the 32 SSR loci, with the exception of Feilongxiang and Shitang Dahuashe, both botanical variety *birsuta*, resistant to bacterial wilt. Of the 46 genotypes, seven exhibit susceptibility to bacterial wilt, 32 show resistance and the response of the remaining seven genotypes to bacterial wilt is unknown. The two subspecies of *A. hypogaea* are equally represented with 23 genotypes belonging to subsp. *hypogaea* (17 to variety *birsuta* and six to variety *hypogaea*) and 23 genotypes belonging to subsp. *fastigiata* (17 to variety *vulgaris* and two representatives each of varieties *fastigiata*, *peruwiana* and *aequatoriana*). The 46 genotypes group into two separate clusters; cluster A containing 19 genotypes, 18 of which belong to subsp. *fastigiata*, and cluster B containing 27 genotypes, 22 of which belong to subsp. *hypogaea*. The 19 genotypes in cluster A include 15 accessions of variety *vulgaris*, two of variety *fastigiata* and two germplasm lines (Chico and ICG15222-2). Three subclusters could be further identified within cluster A at a level of 45% similarity. The first subcluster, A(I), contained genotypes belonging to only variety *vulgaris* and the line ICG15222-2, which were all resistant to BW. The second subcluster, A(II), consisted predominantly of variety *vulgaris* genotypes, with the exception of one accession belonging to variety *fastigiata* (Taishan Sanlirou) and one variety *aequatoriana* accession (ICG12722); of these, six were resistant and four susceptible to BW. The third subcluster, A(III), consisted of two accessions susceptible to BW; an accession belonging to variety *fastigiata* (Gangapuri) and a germplasm line.

Cluster B contained 27 accessions representing five botanical varieties, with botanical variety *fastigiata* unrepresented. Three subclusters could be identified within cluster B, at the level of 42% similarity. The first subcluster, B(I), consisted of 21 genotypes in total, the majority (15) belonging to variety *birsuta*, and all resistant to

bacterial wilt with the exception of Zao 18 (variety *vulgaris*). The second subcluster, B(II), consisted of five landraces originating from South and Central America, including variety *birsuta*, but surprisingly also including variety *peruwiana* and variety *aequatoriana*. The third subcluster, B(III), consisted of only one genotype, Zhonghua No. 5, belonging to variety *vulgaris* which is susceptible to BW.

Overall there was a clear distinction between the BW-resistant lines belonging to subsp. *fastigiata* (cluster A) and the BW-resistant lines of subsp. *hypogaea* (cluster B). However, not all the accessions susceptible to BW followed this pattern; the two BW-susceptible variety *vulgaris* accessions belonging to *A. hypogaea* subsp. *fastigiata* clustered instead with the *A. hypogaea* subsp. *hypogaea* genotypes in cluster B.

AMOVA was performed on the data set in order to partition the total genetic variation within and between three parameters: (i) within and between resistant and susceptible types, (ii) within and between botanical variety, and (iii) within and between country of origin. The AMOVA revealed that only 15% of the total variation observed was accounted for between BW resistance and susceptible types, whereas the majority of variation (85%) was observed within each disease response group (Table 3A). In contrast, upon partitioning the total genetic variation between and within the botanical varieties (Table 3B), 50% of the total variation was accounted for between the different botanical varieties, indicating a clear differentiation based on botanical variety compared to BW response groups. Finally, 86% of the variation was accounted for within the 11 different countries of origin of the genotypes included in this study, rather than between the countries of origin.

Putative association of SSR loci with BW resistance

A locus-by-locus AMOVA was performed to calculate the contribution of each locus to the differentiation of resistant and susceptible groups within the germplasm tested. Six of the 107 alleles (Table 4) were found to contribute significantly to the differentiation between the BW-resistant and -susceptible genotypes. Furthermore, markers pPGPseq-16G8 (size: 229 bp) and pPGPseq-12F7 (size: 305 bp) contributed 74.91% of the total genetic difference between the two disease-response groups.

Discussion

SSRs have proven to be powerful tools for the detection of molecular genetic diversity amongst the cultivated groundnut germplasm included in this study, representing all

six botanical varieties within the *A. hypogaea* gene pool. A clear distinction was observed between the two subspecies, subsp. *hypogaea* and subsp. *fastigiata*. Accessions of var. *hypogaea* and *hirsuta* (subsp. *hypogaea*) grouped together in cluster B (Fig. 2), whereas var. *vulgaris* and *fastigiata* (subsp. *fastigiata*) grouped together in cluster A. However, accessions of var. *peruviana* and var. *aequatoriana* grouped together with the varieties from subsp. *hypogaea* in cluster B, rather than grouping with the other varieties of subsp. *fastigiata* in cluster A. These results support recent conclusions from amplified fragment length polymorphism (AFLP) analysis (He and Prakash, 2001) that var. *aequatoriana* and *peruviana* are much closer to subsp. *hypogaea* than to subsp. *fastigiata*.

Most previous reports of diversity analysis in cultivated groundnut have readily detected morphological variation but consistently failed to detect a parallel level of molecular genetic variability, however recent studies (Ferguson *et al.*, 2004a; Moretzsohn *et al.*, 2004, 2005) using more recently developed SSRs have found higher levels of genetic diversity in the six botanical varieties of *A. hypogaea* than reported previously. Significantly, the SSR screening reported in this study also detected a substantial level of molecular genetic variation between genotypes of all botanical varieties. However, AMOVA showed an equal level of diversity within and between botanical varieties. This may suggest that botanical variety designations (largely based on morphological traits) are not truly reflective of gross genetic diversity.

There was some degree of clustering of accessions from similar geographic origins within subspecies, e.g. four landraces and one breeding line from South and Central America grouped together at a level of approximately 50% similarity in cluster B and were clearly differentiated from accessions derived from other geographic regions. Additionally, the BW-resistant accessions Bobai Shiyadou and Bobai Dahuasheng both originate from Bobai County in China and were observed to group together at a level of approximately 82% similarity in cluster B. However, AMOVA estimated 86% of the SSR variation is accounted for within countries. So although there may be some influence of geographic isolation on genetic polymorphism, as has been reported previously (e.g. He and Prakash, 2001), it is likely that breeding selection pressure has had greater impact. However, it should be noted that the groundnut accessions studied here were selected primarily based on their response to BW and thus may not be truly reflective of groundnut variability in each geographical region.

Regarding the genetic variation observed within and between the BW-resistant and -susceptible accessions, it was observed that the BW-resistant lines belonging to subsp. *fastigiata* and subsp. *hypogaea* were clearly differentiated, and clustered within subspecies. In contrast, two BW-susceptible accessions (Zao18 and Zhonghua No. 5) did not fall in the expected cluster (subsp. *fastigiata* var. *vulgaris*), but were grouped in cluster B (predominately subsp. *hypogaea*). This anomaly

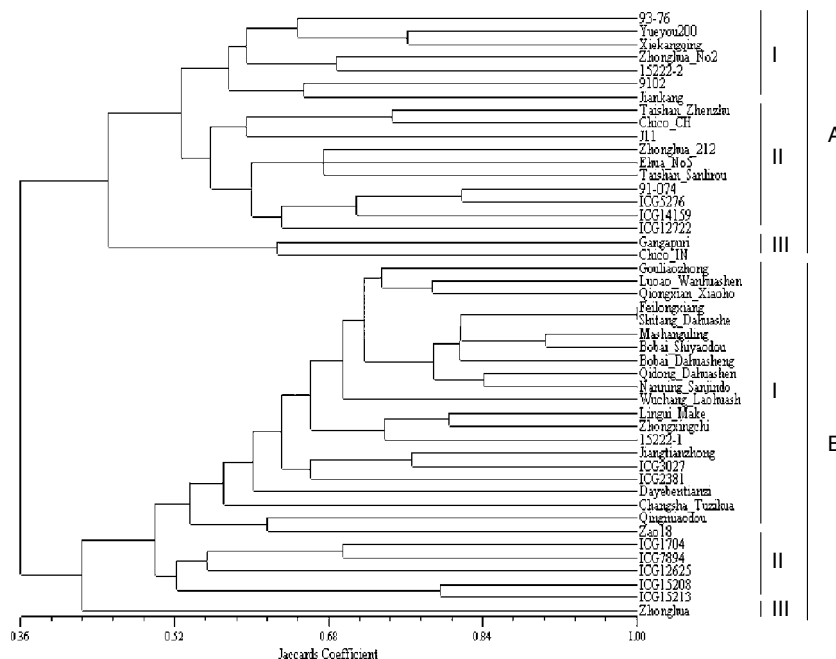


Fig. 2. Dendrogram constructed using Jaccard's similarity coefficient and UPGMA clustering, for the 46 groundnut genotypes. Two main clusters (A and B) and subclusters are identified.

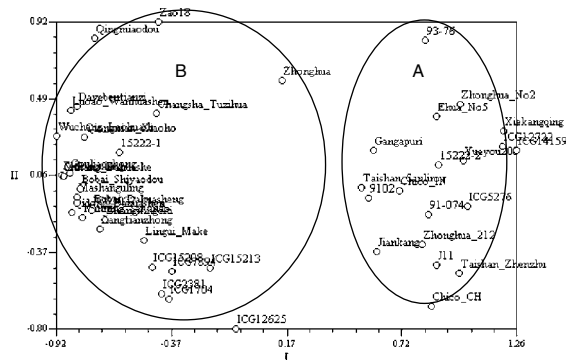


Fig. 3. Multidimensional scaling (MDS) analysis of SSR data across 46 cultivated groundnut genotypes with varying levels of resistance to bacterial wilt with two clusters of genotypes indicated as 'A' and 'B', corresponding to clusters identified in Fig. 2.

may be largely explained by the pedigree of these two breeding lines which included accessions from subsp. *hypogaea*. It should be noted that the BW-susceptible var. *vulgaris* accessions within cluster B (subclusters B(II) and B(III)) grouped only at a level of approximately 60% similarity with the resistant accessions belonging to subsp. *hypogaea*. These results could indicate that the accessions susceptible to BW are more genetically diverse than their resistant counterparts, suggesting that the selection for adaptation to the bacterial wilt pathogen may have contributed to reduced genetic variation in the germplasm which may be due to a combination of pathogen pressure and breeder's selection pressure.

Pairwise dissimilarities of up to 64% were observed between cultivated groundnut genotypes screened in this study. This compares with 41% genetic differentiation revealed between cultivated groundnut genotypes as revealed by randomly amplified polymorphic DNA (RAPDs; Dwivedi *et al.*, 2001) and 52% as revealed by AFLPs (He and Prakash, 2001). Thus, SSRs are clearly

the most powerful tools for revealing genetic variation within the cultivated groundnut gene pool.

Several SSR alleles were also found to be significantly associated with BW resistance and these represent candidates for marker-assisted selection (MAS) following validation in traditional, segregating mapping populations. Molecular breeding offers the potential for improving the speed, precision and cost of groundnut disease resistance breeding programmes (Dwivedi *et al.*, 2003). In addition, MAS offers the potential for pyramiding different sources of resistance together with resistance to multiple diseases which is difficult or impossible to achieve through conventional approaches (Mohan *et al.*, 1997). The paucity of molecular polymorphism previously revealed in cultivated groundnut posed a considerable obstacle to genetic mapping and MAS. The first and only currently available genetic linkage map of the tetraploid groundnut genome (Burow *et al.*, 2001) was only made possible through the use of a synthetic amphidiploid, TxAG-6, capturing a high level of genetic diversity from divergent diploid species (Simpson, 1991; Simpson *et al.*, 1993). Although SSRs now offer a substantially higher level of detectable variation, it is still essential to base the selection of parental genotypes for mapping populations on detailed diversity analysis. It is hoped that the result of this study will help molecular breeders in selecting the most appropriate parental genotypes for mapping BW resistance.

It is now important to distinguish which genotypes possess different mechanisms of resistance and to search for additional novel sources of resistance to BW. Landraces and varieties from Indonesia, China and Vietnam (where BW disease pressures are highest) would appear to be the most likely targets for identifying new sources of resistance. On this basis, marker-assisted gene-pyramiding programmes can aim to develop high-yielding varieties with more durable resistance to this devastating disease.

Table 3. AMOVA (Excoffier *et al.*, 1992) for (A) 39 genotypes of two disease response types, bacterial wilt-resistant and -susceptible, employing 107 SSR alleles; and for (B) 46 genotypes in six botanical varieties employing 107 microsatellite alleles (nested analysis was carried out on populations grouped as above)

Source of variation	df	SSD ^a	Variance components	Variation (%)	Fixation index FST
(A)					
Among populations	1	40.857	2.58 815 Va	15.08***	0.15 078
Within populations	37	539.348	14.57 699 Vb	84.92 ***	
Total	38	580.205	17.16 513	100.00	
(B)					
Among populations	5	336.005	9.46 528 Va	46.19***	0.46 193
Within populations	7	407.949	11.02 564 Vb	53.81***	
Total	42	743.953	20.49 092	100.00	

^a Sum of squared deviations.

Table 4. List of alleles putatively linked with bacterial wilt resistance as determined through locus-by-locus AMOVA

Primer name	Allele size (bp)	Contribution value (%) to differentiation between resistant and susceptible groups
pPGPseq-16G8	229	74.91
pPGPseq-12F7	305	74.91
pPGPseq-14H6	310	62.31
pPGPseq-14H6	306	60.85
pPGPseq-7G2	246	59.07
A1-745	226	59.07

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