Note

CHARACTERIZATION OF RUST, EARLY AND LATE LEAF SPOT RESISTANCE IN WILD AND CULTIVATED PEANUT GERMPLASM

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ABSTRACT: Groundnut (*Arachis hypogaea*) has an AB genome and is one of the most important oil crops in the world. The main constraints of crop management in Brazil are fungal diseases. Several species of the genus *Arachis* are resistant to pests and diseases. The objective of our experiments was to identify wild species belonging to the taxonomic section *Arachis* with either A or B (or "non-A") genomes that are resistant to early leaf spot (*Cercospora arachidicola*), late leaf spot (*Cercosporidium personatum*) and rust (*Puccinia arachidis*). For the identification of genotypes resistant to fungal diseases, bioassays with detached leaves were done in laboratory conditions, with artificial inoculation, a controlled temperature of 25°C and a photoperiod of 10 h light/14 h dark, for 20–42 days, depending on the fungi species. Most of the accessions of wild species were more resistant than accessions of *A. hypogaea* for one, two or all three fungi species studied. *Arachis monticola*, considered to be a possible tetraploid ancestor or a derivative of *A. hypogaea*, was also more susceptible to *Cercosporidium personatum* and *Puccinia arachidis*, as compared to most of the wild species. Therefore, wild germplasm accessions of both genome types are available to be used for the introgression of resistance genes against three fungal diseases of peanut.

Key words: Puccinia arachidis, Cercospora arachidicola, Cercosporidium personatum, groundnut, Arachis spp.

CARACTERIZAÇÃO DA RESISTÊNCIA À FERRUGEM, MANCHA PRETA E MANCHA CASTANHA EM GERMOPLASMA SILVESTRE E CULTIVADO DE AMENDOIM

RESUMO: O amendoim (*Arachis hypogaea*) possui genoma AB e é uma das mais importantes culturas oleaginosas em todo o mundo. Os principais problemas da cultura no Brasil são as doenças fúngicas. Várias espécies do gênero *Arachis* são resistentes a pragas e doenças. Este trabalho visou a identificar espécies silvestres pertencentes à seção *Arachis* associadas aos genomas A ou B (ou "não-A") do amendoim que são resistentes à mancha castanha (*Cercospora arachidicola*), mancha preta (*Cercosporidium personatum*) e ferrugem (*Puccinia arachidis*). Para a identificação de genótipos resistentes a doenças fúngicas, bioensaios utilizando folhas destacadas foram realizados em condições de laboratório, com inoculação artificial, temperatura controlada de 25°C e fotoperíodo de 10h luz/14h escuro, por 20–42 dias, de acordo com a espécie fúngica. A maioria dos acessos das espécies silvestres foram mais resistentes que os acessos de *A. hypogaea* para uma, duas ou todas as espécies fúngicas estudadas. *Arachis monticola*, considerada como o possível ancestral tetraplóide ou como um derivativo de *A. hypogaea*, também mostrou-se mais suscetível a *Cercosporidium personatum* e *Puccinia arachidis*, quando comparado à maioria das espécies silvestres. Portanto, acessos de germoplasma silvestre com genoma A ou B estão disponíveis para serem utilizados na introgressão de genes de resistência a doenças fúngicas no amendoim.

Palavras-chave: Puccinia arachidis, Cercospora arachidicola, Cercosporidium personatum, Arachis spp.

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INTRODUCTION

Peanut (*Arachis hypogaea* L.) is the fourth most important oleaginous plant in the world. It is used mainly for oil and candy production, or for consumption *in natura*. World production is thought to be more than 30 million tons per year (CONAB, 2003). Brazil has about 90,000 ha planted with peanut, with a production of about 220,000 tons in 2006. The main constraints of the peanut production in Brazil, and indeed in the world, are fungal diseases, such as web blotch (*Phoma arachidicola* Marasas, Pauer & Boerema), early leaf spot (*Cercospora arachidicola* Hori), late leaf spot (*Cercosporidium personatum* (Berk & Curt.) Deighton), rust (*Puccinia arachidis* Speg.) and scab (*Sphaceloma arachidis* Bitancourt & Jenkins) (Godoy et al., 1999).

Species in the genus *Arachis* have potential for peanut improvement (Fávero et al., 2006). Several species have higher resistance levels to diseases when compared to *A. hypogaea* germplasm accessions (Pande & Rao, 2001; Stalker & Moss, 1987). The genus has nine taxonomic sections, and *A. hypogaea* is in the section *Arachis*, along with 30 wild species (Krapovickas & Gregory, 1994; Valls & Simpson, 2005).

The objective of the present study was to test diploid and tetraploid wild species within the section Arachis, with A and/or B or "non A" genomes, as well as several A. hypogaea varieties, for resistance against three fungal diseases - early and late leaf spot and rust, with the aim of future introgression of disease resistance genes into a breeding program of the cultivated peanut. Different accessions of a particular species may have different rates of resistance to fungal diseases. Although several previous publications have demonstrated the resistance of assorted wild Arachis germplasm against fungal diseases, old and newly collected accessions of many species were put together in the present work, and were tested with fungi isolates from São Paulo State, where 80% of the total Brazilian peanut acreage is concentrated. Isolates from Brazil may be different from those from other countries, so, this work was necessary as a basic step to implement a Peanut Pre-breeding project in Brazil.

MATERIAL AND METHODS

Accessions used in this study are shown in Table 2 and were obtained from the Arachis germplasm bank of Embrapa Recursos Genéticos e Biotecnologia, Brasília, Brazil. Only accessions of the section Arachis were selected because crossability with the cultivated peanut has not been accomplished so far with species of other taxonomic sections. The three diploid species of section Arachis with 2n = 18 chromosomes Arachis praecox, A. decora and A. palustris (Peñaloza & Valls, 1997; Lavia, 1998) were not included. Diploid species with 2n = 20 were assigned the A or B (non-A) genome according to the available documentation of the presence or absence of the small "A" chromosome pair in at least one accession of each in the literature (Fernández & Krapovickas, 1994; Lavia, 1998; Lavia, 1999; Peñaloza & Valls, 2005). The tetraploid wild species Arachis monticola Krapov. & Rigoni (2n = 40) is considered to share the same AB genome of A. hypogaea. Seeds from 102 accessions were treated with Carboxim and Thiram (0.5 g L^{-1}) fungicide and germinated at 25°C in germitest paper immersed in Ethrel solution (6 mL L⁻¹) to break dormancy. Plants were maintained under screenhouse conditions, with four replications per each accession.

Cercospora arachidicola and *Cercosporidium personatum* spores were collected from infected plants at the Experimental Station of Ribeirão Preto (21°10' S, 47°48' W), from the Agronomic Institute of São Paulo State, Brazil (IAC), and *Puccinia arachidis* spores were collected from infected plants at the Experimental Station of Pindorama, (21°11' S, 48°54' W), from the Agronomic Institute of São Paulo State, Brazil (IAC).

For *Cercosporidium personatum* and *Cercopora arachidicola*, fungi cultures were grown in oat-agar medium (Moraes & Salgado, 1979). In addition, some fungi from the IAC bank of fungal isolates (1436–1 and 1595–0) were used (Table 1). New isolates were given the numbers 11576–0 and 11576–1, respectively. The *P. arachidis* spores were collected and stored in jelly capsules in refrigerator (about 5°C), for about one week. It was periodically necessary to inoculate susceptible peanut leaves and re-isolate the spores.

Table 1 - Accession code at Agronomic Institute of São Paulo State, Brazil (IAC) fungi collection, fungi species names and municipality of São Paulo State where they were collected.

Code	Name	Region	Coordinates
1436-1	Cercospora arachidicola	Pompéia	22°06' S; 50°10' W
11576-1	C. arachidicola	Ribeirão Preto	21°10' S; 47°48' W
1595-0	Cercosporidium personatum	Jaú	22°17' S; 48°33' W
11576-0	C. personatum	Ribeirão Preto	21°10' S; 47°48' W

Species	Accession	Cercospora arachidicola	Cercosporidium personatum	Puccinia arachidis		Σ
		0-f (0-6)	0-d	Group 1 (0-d)	Group 2 (0-b)	
A. batizocoi	K 9484	sl *(0)	sl (0)	-	-	0*
A. cardenasii	GKP 10017	sl (0)	sl (0)	0.00228a (1)	-	1
A. aff. diogoi	VSPmSv 13774	0.00035a (1)	sl (0)	sl (0)	-	1
A. helodes	CoSzSv 6862	sl (0)	sl (0)	0.00337a (1)	-	1
A. helodes	Pa s/n	0.00055a (1)	sl (0)	sl (0)	-	1
A. helodes	VPoJSv 10470	sl (0)	sl (0)	0.000813a (1)	-	1
A. helodes	VK 12083	0.00142a (1)	sl (0)	sl (0)	-	1
A. hoehnei	VMPzW 13985	-	sl (0)	0.00276a (1)	-	1
A. linearifolia	VPoBi 9401	sl (0)	sl (0)	0.00631a (1)	-	1
A. simpsonii	VSPmSv 13710	0.00049a (1)	sl (0)	sl (0)	-	1
A. stenosperma	VSMGeSv 7379	0.00134a (1)	sl (0)	-	-	1*
A. stenosperma	VGaRoSv 12488	sl (0)	sl (0)	0.00474a (1)	-	1
A. diogoi	GK 10602	0.00054a (1)	sl (0)	0.00397a (1)	-	2
A. duranensis	VNvEv 14167	-	0.00064a (1)	0.00282a (1)	-	2
A. hoehnei	KG 30006	sl (0)	sl (0)	0.00974b (2)	-	2
A. hoehnei	VPoBi 9094	sl (0)	-	0.02177b (2)	-	2*
A. kempff-mercadoi	V 13250	sl (0)	0.00046a (1)	0.00491a (1)	-	2
A. kuhlmannii	VRGeSv 7639	sl (0)	-	0.00875b (2)	-	2*
A. kuhlmannii	VPoBi 9235	0.00035a (1)	-	0.00709a (1)	-	2*
A. kuhlmannii	VPoJSv 10506	0.00020a (1)	0.00029a (1)	sl (0)	-	2
A. microsperma	VRGeSv 13545	-	sl (0)	0.01195b (2)	-	2*
A. microsperma	VMPzW 14042	0.00011a (1)	sl (0)	0.00322a (1)	-	2
A. stenosperma	HLK 408	-	sl (0)	0.02340b (2)	-	2*
A. stenosperma	Lm 5	sl (0)	sl (0)	0.01380b (2)	-	2
A. stenosperma	VSStGdW 7762	sl (0)	sl (0)	0.02588b (2)	-	2
A. gregoryi	VSGr 6389	0.00275b (2)	0.00289a (1)	sl (0)	-	3
A. helodes	VSGr 6325	0.00020a (1)	0.00009a (1)	0.00067a (1)	-	3
A. kuhlmannii	VSGr 6344	0.00091a (1)	0.00745a (1)	0.00040a (1)	-	3
A. kuhlmannii	VSGr 6352	0.00122a (1)	sl (0)	0.02217b (2)	-	3
A. kuhlmannii	VSGr 6380	0.00112a (1)	sl (0)	0.01785b (2)	-	3
A. kuhlmannii	VKSSv 8916a	0.00012a (1)	sl (0)	0.01876b (2)	-	3
A. kuhlmannii	VPoBi 9470	0.00006a (1)	sl (0)	0.01698b (2)	-	3
A. kuhlmannii	VPoBi 9479	0.00026a (1)	sl (0)	0.01977b (2)	-	3
A. kuhlmannii	VSW 9912	0.00004a (1)	sl (0)	0.02308b (2)	-	3
A. kuhlmannii	VSPmSv 13721	0.00014a (1)	0.0000818a (1)	0.00039a (1)	-	3
A. magna	KGSSc 30097	0.00377b (2)	sl (0)	0.00125a (1)	-	3
A. stenosperma	Jt 2	sl (0)	sl (0)	0.03807c (3)	-	3
A. stenosperma	Lm 3	0.00080a (1)	sl (0)	0.01186b (2)	-	3
A. stenosperma	SvW 3755	sl (0)	sl (0)	0.06715c (3)	-	3
A. stenosperma	VKSSv 9010	0.00028a (1)	sl (0)	0.01302b (2)	-	3
A. stenosperma	VMiSv 10229	sl (0)	sl (0)	0.04220c (3)	-	3
A. stenosperma	VSPmSv 13832	sl (0)	0.00372a (1)	0.02715b (2)	-	3

 Table 2 - Species names, collector's number, the screening for resistance to Cercospora arachidicola, Cercosporidium personatum and Puccinia arachidis and sum of points related to resistance to all three diseases.

Continue...

A. stenosperma	WPz 422	0.00033a (1)	sl (0)	0.02057b (2)	-	3
A. villosa	VGoMrOv 12812	0.00011a (1)	0.00215a (1)	0.00072a (1)	-	3
A. batizocoi	K 9484 mut	0.00862c (3)	-	0.00134a (1)	-	4*
A. cruziana	WiSVg 1302	0.00940c (3)	sl (0)	0.00471a (1)	-	4
A. duranensis	K 7988	0.00091a (1)	-	0.05935c (3)	-	4*
A. hoehnei	VPoBi 9146	0.00071a (1)	sl (0)	0.03226c (3)	-	4
A. kuhlmannii	VSGr 6351	0.00016a (1)	0.00048a (1)	0.02434b (2)	-	4
A. kuhlmannii	VKSSv 8979	sl (0)	0.00415a (1)	0.04841c (3)	-	4
A. kuhlmannii	VPoBi 9230	0.00767c (3)	0.00091a (1)	sl (0)	-	4
A. kuhlmannii	VPoBi 9394	0.00203b (2)	0.00017a (1)	0.00314a (1)	-	4
A. magna	VSPmSv 13751	0.00071a (1)	0.00016a (1)	0.01040b (2)	-	4
A. magna	VSPmSv 13765	sl (0)	sl (0)	0.10415d (4)	-	4
A. simpsonii	VSPmSv 13716	0.00016a (1)	sl (0)	0.05585c (3)	-	4
A. simpsonii	VSPmSv 13728	0.00301b (2)	sl (0)	0.01221b (2)	-	4
A. simpsonii	VSPmSv 13745	0.00188b (2)	0.00012a (1)	0.00054a (1)	-	4
A. stenosperma	Lm 1	0.00065a (1)	sl (0)	0.04328c (3)	-	4
A. stenosperma	VSSv 7382	0.00009a (1)	sl (0)	0.04287c (3)	-	4
A. stenosperma	VSv 10309	0.00018a (1)	sl (0)	0.04461c (3)	-	4
A. stenosperma	VSPmSv 13670	sl (0)	sl (0)	0.16580d (4)	-	4
A. stenosperma	VKSSv 9017	0.01119c (3)	0.00024a (1)	0.00119a (1)	-	5
A. stenosperma	WPz 421	0.00133a (1)	0.00179a (1)	0.05112c (3)	-	5
A. kuhlmannii	VSGr 6413	0.00359b (2)	0.00397a (1)	0.01262b (2)	-	5
A. kuhlmannii	VPoBi 9375	0.00252b (2)	0.00168a (1)	0.00837b (2)	-	5
A. magna	VSPmSv 13748	0.00275b (2)	sl (0)	0.10855c (3)	-	5
A. schininii	VSW 9923	0.00686c (3)	0.00067a (1)	0.00034a (1)	-	5
A. stenosperma	SvPzSz 3042	0.00005a (1)	sl (0)	0.11090d (4)	-	5
A. stenosperma	VGaSv 12646	0.00032a (1)	sl (0)	0.16580d (4)	-	5
A. stenosperma	VSPmWiSv 13262	0.00048a (1)	0.00103a (1)	0.06378c (3)	-	5
A. stenosperma	VSPmSv 13672	sl (0)	sl (0)	-	0.03711a (5)	5
A. stenosperma	VSPmSv 13693	0.00085a (1)	sl (0)	0.19504d (4)	-	5
A. stenosperma	VSPmW 13828	sl (0)	sl (0)	-	0.03166a (5)	5
A. stenosperma	VSPmW 13844	0.00134b (2)	sl (0)	0.05125c (3)	-	5
A. valida	VPoBi 9153	-	sl (0)	-	0.01718a (5)	5*
A. valida	VPzRcSgSv 13514	0.00532b (2)	sl (0)	0.06793c (3)	-	5
A. hypogaea	Mf 1538	0.00073a (1)	-	-	0.01112a (5)	6*
A. kuhlmannii	VPoBi 9214	0.01858d (4)	0.00041a (1)	0.00221a (1)	-	6
A. magna	VSPmSv 13761	sl (0)	0.00657a (1)	-	0.03247a (5)	6
A. stenosperma	VGaRoSv 12575	0.00022a (1)	sl (0)	-	0.04907a (5)	6
A. stenosperma	VSSv 13258	0.00034a (1)	0.01924b (2)	0.05798c (3)	-	6
A. stenosperma	VSPmW 13824	0.00007a (1)	-	-	0.03898a (5)	6*
A. williamsii	WiDc 1118	0.00112a (1)	sl (0)	-	0.07156a (5)	6
A. ipaënsis	KGBPScS 30076	0.00287b (2)	-	-	0.06128a (5)	7*
A. kuhlmannii	VPoBi 9243	0.14491d (4)	0.00004a (1)	0.01002b (2)	-	7
A. stenosperma	SvSz 2411	0.00035a (1)	sl (0)	_	0.20452b (6)	7
A. stenosperma	SvW 3712	0.00030a (1)	0.00029a (1)	-	0.06331a (5)	7

Table 2 - Continuation.

Continue...

VSPmW 13796	0.00338b (2)	sl (0)	-	0.05784a (5)	7
VPoBi 9157	0.00580b (2)	-	-	0.07219a (5)	7*
VPzRcSgSv 13516	0.00042a (1)	-	-	0.20794b (6)	7*
US 224	0.00251b (2)	0.00073a (1)	-	0.02885a (5)	8
VOa 14165	0.00776c (3)	-	-	0.03144a (5)	8*
VGaRoSv 12549	0.00802c (3)	0.01083b (2)	-	0.04417a (5)	10
cv. Tatu	0.01446d (4)	0.00946b (2)	-	0.03613a (5)	11
Mf 1560	0.01706d (4)	0.02179b (2)	-	0.00968a (5)	11
VGaRoSv 12548	0.01423d (4)	0.01859b (2)	-	0.02358a (5)	11
cv. Tatuí	0.02693e (5)	0.01587b (2)	-	0.03717a (5)	12
cv. Caiapó	0.02663e (5)	0.03075c (3)	-	0.03125a (5)	13
Mf 1678	0.05183f (6)	0.01747b (2)	-	0.03713a (5)	13
Runner 886	0.02812e (5)	0.03901c (3)	-	0.03840a (5)	13
F 1334	0.02391e (5)	0.05477d (4)	-	0.07898a (5)	14
cv. BR1	0.05034f (6)	0.05972d (4)	-	0.07430a (5)	15
	VSPmW 13796 VPoBi 9157 VPzRcSgSv 13516 US 224 VOa 14165 VGaRoSv 12549 cv. Tatu Mf 1560 VGaRoSv 12548 cv. Tatuí cv. Caiapó Mf 1678 Runner 886 F 1334 cv. BR1	VSPmW 137960.00338b (2)VPoBi 91570.00580b (2)VPzRcSgSv 135160.00042a (1)US 2240.00251b (2)VOa 141650.00776c (3)VGaRoSv 125490.00802c (3)cv. Tatu0.01446d (4)Mf 15600.01706d (4)VGaRoSv 125480.01423d (4)cv. Tatuí0.02693e (5)cv. Caiapó0.02663e (5)Mf 16780.05183f (6)Runner 8860.02391e (5)cv. BR10.05034f (6)	VSPmW 137960.00338b (2)sl (0)VPoBi 91570.00580b (2)-VPzRcSgSv 135160.00042a (1)-US 2240.00251b (2)0.00073a (1)VOa 141650.00776c (3)-VGaRoSv 125490.00802c (3)0.01083b (2)cv. Tatu0.01446d (4)0.00946b (2)Mf 15600.01706d (4)0.02179b (2)VGaRoSv 125480.01423d (4)0.01859b (2)cv. Tatuí0.02693e (5)0.01587b (2)cv. Tatuí0.02663e (5)0.03075c (3)Mf 16780.05183f (6)0.01747b (2)Runner 8860.02812e (5)0.03901c (3)F 13340.02391e (5)0.05477d (4)cv. BR10.05034f (6)0.05972d (4)	VSPmW 137960.00338b (2)sl (0)-VPoBi 91570.00580b (2)VPzRcSgSv 135160.00042a (1)US 2240.00251b (2)0.00073a (1)-VOa 141650.00776c (3)VGaRoSv 125490.00802c (3)0.01083b (2)-cv. Tatu0.01446d (4)0.00946b (2)-Mf 15600.01706d (4)0.02179b (2)-VGaRoSv 125480.01423d (4)0.01859b (2)-cv. Tatuí0.02693e (5)0.01587b (2)-cv. Tatuí0.02663e (5)0.03075c (3)-mf 16780.05183f (6)0.01747b (2)-Runner 8860.02812e (5)0.03901c (3)-F 13340.02391e (5)0.05477d (4)-cv. BR10.05034f (6)0.05972d (4)-	VSPmW 137960.00338b (2)sl (0)-0.05784a (5)VPoBi 91570.00580b (2)0.07219a (5)VPzRcSgSv 135160.00042a (1)0.20794b (6)US 2240.00251b (2)0.00073a (1)-0.02885a (5)VOa 141650.00776c (3)0.03144a (5)VGaRoSv 125490.00802c (3)0.01083b (2)-0.04417a (5)cv. Tatu0.01446d (4)0.00946b (2)-0.03613a (5)Mf 15600.01706d (4)0.02179b (2)-0.00968a (5)VGaRoSv 125480.01423d (4)0.01859b (2)-0.02358a (5)cv. Tatuí0.02693e (5)0.01587b (2)-0.03717a (5)cv. Caiapó0.02663e (5)0.03075c (3)-0.03125a (5)Mf 16780.05183f (6)0.01747b (2)-0.03713a (5)Runner 8860.02812e (5)0.03901c (3)-0.03840a (5)F 13340.02391e (5)0.05477d (4)-0.07430a (5)

Table 2 - Continuation.

sl means no lesion. Number inside parenthesis means the punctuation of the genotypes to do the ranking. They are related to letters obtained in the average test (Scott & Knott test). The signal * means that there is data missing so with this data the ranking may change.

Five isolates were prepared: two of *Cercospora* arachidicola, two of *Cercosporidium personatum* and one of *P. arachidis*. For *Cercospora arachidicola* and *Cercosporidium personatum* isolates were replicated by the use of oat-agar medium into Erlenmeyers that were shaked for about 20 minutes. The cultures were replicated at each 14 days, during approximately three months.

A detached leaf technique was used for the establishment of bioassays (Moraes & Salgado, 1982) mostly using leaves of the main stem of the plants. Cotton and germitest paper layers and a slide were used to keep the leaves well conditioned in Petri dishes. Destilated water was used to maintain leaves alive and turgid for several weeks. For the inoculation of Cercospora arachidicola and Cercosporidium personatum, the first expanded leaves were placed in Petri dishes, with the adaxial side upward. For Puccinia arachidis, the abaxial side was set upward. Inoculation was done with a mixture of the isolates of each leaf spot by spraying Tween 20 at 0.5% in a concentration of 50.000 spores mL⁻¹. Plates were maintained at 23-25°C, photoperiod of 10 h light and 14 h darkness, in four random blocks. During the first 48 h, the Petri dishes were sealed in a plastic bag.

Puccinia arachidis bioassays were analysed at 20 and 27 days, *Cercospora arachidicola* at 27 days and *Cercosporidium personatum* at 42 days. For *P. arachidis* evaluation, the number of lesions per leaflet area (mm²) was recorded and the lesions scored for presence or absence of spores. For *Cercospora arachidicola* and *Cercosporidium personatum*, the ratio of lesion area to leaflet area was evaluated (Foster et al., 1981).

The SAS program was used for the analysis of mathematic model according to the random blocks experiments. Different transformations were used for each group of data, according to ANOVA conditions. For Cercospora arachidicola, the transformation was $\arcsin (x + 0.5)$. For *Cercosporidium personatum* and *P. arachidis* a $\sqrt{\log(x+1)}$ transformation was used. The ANOVA presuppositions were verified: independence, homogeneity, and normal distribution of residues. Only data from accessions with lesions were used in the analyses. The Scott & Knott method (Scott & Knott, 1974) was used at 5% probability of Type I Error for the multiple comparisons among averages of different accessions. A sum of points, adapted to an index of selection (rank sum) of Mulamba & Mock (1978), was done to determine the most resistant and most susceptible genotypes. The t test was used to observe significant differences between the three groups, the species that have the A genome, species with non-A genome, and species that possess the AB genome. For Cercospora arachidicola and Cercosporidium personatum, this test was done using the average data of the species. For P. arachidis, the t test was used in the punctuation data to have the same screening for accessions that had lesions without pustule and lesions with pustule.

RESULTS AND DISCUSSION

Results for all diseases studied are summarized in Table 2.

Cercospora arachidicola

From 97 accessions investigated, 22 were shown to be highly resistant (Table 2 - third column)

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without fungal lesions. From these accessions, seven are perennial and with A genome (A. kuhlmannii Krapov. & W.C. Greg. (2 accessions), A. cardenasii Krapov. & W.C. Greg. (1), A. helodes Mart. ex Krapov. & Rigoni (2), A. kempff-mercadoi Krapov., W.C. Greg. & C.E. Simpson (1), A. linearifolia Valls, Krapov. & Simpson (1)), ten are annual and with A genome (A. stenosperma Krapov. & W.C. Greg.) and five are annual and with "non-A" genome (A. magna Krapov., W.C. Greg. & C.E. Simpson (2), A. hoehnei Krapov. & W.C. Greg. (2) and A. batizocoi Krapov. & W.C. Greg. (1)). Therefore, there are highly resistant wild species associated to both genomes of A. hypogaea.

From 75 accessions considered in the statistical analysis, 43 were in the "a" group, with a smaller number of lesions. In this group, there were A genome species (A. stenosperma (17 accessions), A. kuhlmannii (11), A. helodes (3), A. diogoi Hoehne (1), A. aff. diogoi (1), A. simpsonii Krapov. & W.C. Greg. (2), A. duranensis Krapov. & W.C. Greg. (1), A. microsperma Krapov., W.C. Greg. & Valls (1), A. villosa Benth. (1)), and "non-A" genome species (A. magna (1), A. hoehnei (1), A. valida Krapov. & W.C. Greg. (1) and A. williamsii Krapov. & W.C. Greg. (1) and one accession of A. hypogaea var. hirsuta Köhler (Mf 1538). The "b" group included 14 accessions distributed among the A genome species A. kuhlmannii (3), A. simpsonii (2), A. stenosperma (2), and the "non-A" genome species A. magna (2), A. valida (2), A. gregoryi C.E. Simpson, Krapov. & Valls (1) and A. ipaënsis Krapov. & W.C. Greg. (1), and one accession of A. hypogaea var hypogaea (US 224). In the "c" group, seven accessions were observed, three with A genome (A. kuhlmannii (1), A. stenosperma (1)) and A. schininii Krapov., Valls & C.E. Simpson (1), two with "non-A" genome, (A. batizocoi and A. cruziana Krapov., W.C. Greg. & C.E. Simpson) and two were alotetraploids (A. monticola (1) and A. hypogaea (1)). The "d" group presented accessions of A. kuhlmannii (2) and A. hypogaea (3). The "e" and "f" groups showed accessions with more lesions than the other groups, so these were the most susceptible. In these two final groups, only accessions of A. hypogaea (6) were observed. These results were already expected, confirming the greater resistance of many wild species when compared to A. hypogaea and their potential use in the improvement of the cultivated peanut, like A and "non-A" peanut genome substitutes. Resistance to C. arachidicola was also found in accessions of A. stenosperma, A. diogoi, A. correntina (Burkart) Krapov. & W.C. Greg. and A. duranensis by Foster et al. (1981).

The resistances are very heterogeneous among accessions of the same species, as in *A. kuhlmannii* and *A. stenosperma*. So, a precise analysis of each accession is necessary, as the data do not support the species as always resistant or susceptible. A high coefficient of variation of 49.68%, was obtained. However, it is commom to find values as great as this in disease evaluation data.

Cercosporidium personatum

From 91 accessions submitted to the C. personatum resistance test, 54 appeared highly resistant. No lesion was observed (Table 2 -fourth column) in these 54 accessions of different species of Arachis, from A genome species (A. stenosperma (24 accessions), A. kuhlmannii (6), A. helodes (4), A. simpsonii (3), A. diogoi (1), A. aff. diogoi (1), A. microsperma (2), A. linearifolia (1), A. cardenasii (1)) and "non-A" genome species (A. cruziana (1), A. hoehnei (3), A. magna (3), A. valida (2), A. batizocoi (1), A. williamsii (1)). As for resistance to Cercospora arachidicola, it was observed that wild germplasm accessions associated to both A. hypogaea genomes are available to be used in the introgression of *Cercosporidium personatum* resistance genes into the cultivated peanut. The results may suggest that the above accessions are immune to the pathogen. However, it seems too hasty to come to this conclusion based just on a laboratory test. It would be necessary to repeat the bioassay or to test the material in the field.

In the "a" group, which includes the most resistant of those accessions statistically analyzed, 22 accessions associated to the A genome species were observed, A. kuhlmannii (11 accessions), A. stenosperma (5), A. duranensis (1), A. helodes (1), A. kempffmercadoi (1), A. schininii (1), A. simpsonii (1), A. villosa (1) and three of "non-A" genome, A. magna (2) and A. gregoryi (1). An accession of A. hypogaea (US 224) showed higher resistance than the others, also being located in the "a" group. It is interesting to mention that this peanut accession, from Rondonia State, in Brazil, which also presented some resistance to C. arachidicola, is the source of resistance to Tomato Spotted Wilt Virus/TSWV incorporated in the Tamrun 96 cultivar (Smith et al., 1998). The "b" group only includes one accession of A. stenosperma and six of A. hypogaea. The "c" and "d" groups only include accessions of A. hypogaea (4), confirming that the cultivated species is significantly more susceptible than the wild species used in the experiment.

The coefficient of variation was 59.06%. As in the tests for *Cercospora arachidicola* resistance, it is normal to find high values like this in disease evaluation data.

Puccinia arachidis

In the rust resistance tests, seven out of 100 accessions appeared highly resistant, without evidence of any lesion (Table 2 - fifth column), 67 accessions showed different lesion levels, however, without pustules but with hypersensitivity reactions (Table 2 - fifth column), and 26 accessions were shown to be more susceptible, presenting pustules (Table 2 - sixth column).

Six of the highly resistant accessions belong to A genome species, *A. helodes* (2 accessions), *A. kuhlmannii* (2), *A. aff. diogoi* (1), *A. simpsonii* (1), and a single one to the "non-A" genome species, *A. gregoryi* (1).

In Table 2, from 67 accessions that had just hypersensibility reactions, 23 were in the "a" group, being the most resistant among those encompassed by the statistical analysis. The accessions found in the "a" group were in 19 species with "A" genome - A. kuhlmannii (5 accessions), A. helodes (3), A. stenosperma (2), A. duranensis (1), A. cardenasii (1), A. diogoi (1), A. kempff-mercadoi (1), A. linearifolia (1), A. microsperma (1), A. schininii (1), A. simpsonii (1), A. villosa (1) - and four with "non-A" genome species - A. batizocoi (1), A. cruziana (1), A. hoehnei (1), A. magna (1). In the "b" group, other 23 accessions were observed: 20 with A genome - A. kuhlmannii (11), A. stenosperma (7), A. microsperma (1), A. simpsonii (1) - and three with "non-A" genome - A. magna (1), A. hoehnei (2). The "c" group involved 16 accessions, 13 with A genome - A. stenosperma (10), A. duranensis (1), A. kuhlmannii (1), A. simpsonii (1) and three with "non-A" genome - A. hoehnei (1), A. magna (1) and A. valida (1). The "d" group presented four accessions of A. stenosperma (A genome) and one of A. magna ("non-A" genome). Such results are reasonably in line with those of Subrahmanyam (1983), who found immunity against P. arachidis in A. batizocoi (accession K 9484), A. cardenasii (GKP 10017), A. diogoi (GK 10602) and high resistance in A. stenosperma (HLK 408). He also found susceptibility in A. monticola.

In the same way as for resistance to *Cercospora arachidicola* and *Cercosporidium personatum*, wild germplasm accessions relating to both genomes of *A. hypogaea* were observed, and are available to be used in the introgression of *P. arachidis* resistance genes in the cultivated peanut.

All the accessions in the sixth column showed sporulation of *P. arachidis*. In the "a" group, there were 12 accessions of *A. hypogaea*, 6 of *A. stenosperma*, 2 of *A. valida*, 1 of *A. monticola*, 1 of *A. magna*, 1 of *A. ipaënsis* and 1 of *A. williamsii* (Table 2). In the "b" group, only 2 accessions were observed, 1 of *A. stenosperma* and 1 of *A. valida*. All accessions of *A. hypogaea* showed pustules, so the cultivated peanut is more susceptible than most of the wild species accessions in this study. *Arachis monticola*, an allotetraploid wild species, considered by some authors to be the ancestor or, alternatively, a derivative of *A. hypogaea*, also showed susceptibility to *P. arachidis. Arachis ipaënsis*, one of the original diploid ancestors of *A. hypogaea* (Fávero et al., 2006), also showed susceptibility to *P. arachidis*. The variation coefficient was 71.43%.

A sum of points was done to determine the most resistant and most susceptible genotypes and the added values for each accession are shown in Table 2 (Seventh column). Some data were missed, so with these data, the ranking may change in some cases. For Cercospora arachidicola, the letters from Scott & Knott test varied from "a" to "f", and some genotypes were not included in the variance analysis as they presented no lesion, therefore earning a zero score. To rank all the lesions and all the genotypes, the categorization by letters was converted to numbers: where it varied from zero to "f", the ranking varied from zero to six. For *Cercosporidium personatum* the values were from zero to "d", so they became zero to four. For P. arachidis, group one included the categories zero to "d", changing to 0 to 4; for group two, "a" and "b" groups were replaced by, respectively, scores 5 and 6. So, all test values obtained for each genotype were added, according to the conversions above. Most of the accessions of wild species were more resistant than the accessions of A. hypogaea. The allotetraploid A. monticola, also has been more susceptible to Cercosporidium personatum and P. arachidis when compared with most of the wild species. Arachis monticola was not tested for C. personatum because the leaves were not in appropriate condition for the analysis, nor were other accessions without group letters. Arachis *ipaënsis*, one of the diploid ancestors of A. hypogaea, was also shown to be more susceptible to Cercospora arachidicola and P. arachidis when compared to many of the wild species.

For *Cercospora arachidicola*, *Cercosporidium personatum* and *P. arachidis*, the results presented by Stalker & Moss (1987) in a table of species and respective resistance results for several diseases and pests show a marked similarity to those found in the present study.

The resistance to late leaf spot and rust were studied by Pande & Rao (2001) in 74 accessions of wild species of *Arachis* under greenhouse conditions. The accession KG 30006 of *A. hoehnei* did not show symptoms of either of the two diseases. Twenty-six accessions were classified as resistant to late leaf spot.

Table 3 - Comparison among A and "non-A" genome wild species and *A. hypogaea* and between A and "non-A" genome species for *Cercospora arachidicola, Cercospora arachidicola* and *Puccinia arachidis* based on P-values obtained by the use of the t test. Values lower than 0.05 indicate difference between genomes.

Genomes	Cercospora arachidicola	Cercospora arachidicola	Puccinia arachidis
$A \times AB$ (A. hypogaea)	0.0012	0.0015	0.0000
"Non-A" \times AB (A. hypogaea)	0.0008	0.0014	0.0001
$A \times$ "Non-A"	0.7461	0.9049	0.0913

Sixty-eight accessions were considered rust resistant. Although most of the accessions appraised by Pande & Rao (2001) are not the same as those used in the present work, results can be extrapolated in some cases, as there are several common sites for the collections. Our accession of *A. monticola* (V 14165), although collected more recently, but from the same site of Pande & Rao's accession, and the coincident accession of *A. ipaënsis* (K 30076) were also susceptible to rust, corroborating the results of Pande & Rao (2001).

There is resistance to *Cercosporidium personatum*, *Cercospora arachidicola* and *Puccinia arachidis* in many accessions of wild species and these resistances may be different among accessions of the same species.

Species with A genome are more resistant than *A. hypogaea* under the conditions of the present *Cercospora arachidicola, Cercosporidium personatum* and *Puccinia arachidis* bioassays (Table 3). The same was observed for species that have "non-A" genome. On the other hand, species with A genome were not significantly different from species with "non-A" genome, showing that resistance genes for the three fungal diseases are at both genomes, and it is possible to introgress them from the both genomes, doing the gene piramidization.

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