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Biodiversity assessment based on cpDNA and crossability analysis in selected species of Allium subgenus Rhizirideum

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Abstract The chloroplast DNA diversity of 33 accessions belonging to 16 species of five sections in Allium subgenus Rhizirideum was studied by analysing the sequence of three fragments: the trnL-F intergenic spacer, the rps 16 intron and rbcL (rubisco large subunit). The three sections Cepa, Schoenoprasum and Rhizirideum, representing the majority of the included species, each possess a separate clade after phylogenetic analysis. Exceptions to this general rule are the placement of Allium pskemense (section Cepa) connected to Allium senescens (section Rhizirideum) and Alium roylei, taking an intermediate position between sections Cepa and Schoenoprasum. Both species were located in their own section after nuclear DNA analysis. A range of crossing experiments has been carried out. The different position of A. roylei when comparing cpDNA and nDNA diversity was not confirmed with the production of hybrid seeds after crossing A. roylei with species other than those of section Cepa. The different position of A. pskemense in the cpDNA and the nDNA tree can not be compared to its crossability, since only a few crossing experiments are reported for this species. The hypothesis that a shorter distance between two species in a cpDNA tree compared to their distance in a nDNA tree will indicate interfertility at a certain level, is neither confirmed nor rejected by the currently available results.

Keywords Allium subgenus Rhizirideum · Allium cepa · Allium roylei · Sequence analysis \cdot trnL \cdot rps16 \cdot rbcL \cdot Onion · Phylogeny reconstruction · cpDNA

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Introduction

The genus *Allium* comprises approximately 700 species, arranged in four large subgenera with 100–300 species each (Allium, Rhizirideum, Amerallium and Melanocrommyum) and two much smaller subgenera (*Bromatorrhiza* and Caloscordon; Hanelt et al. 1992). The main cultivated species are found in the subgenera Allium (garlic, leek) and Rhizirideum (onion, shallot, bunching onion, chives, chinese chives), whereas a range of ornamentals originates from subgenus Melanocrommyum. Most research has been focused on the improvement of these cultivated species. Especially onion (Allium cepa), as the most important cultigen in the genus, received much attention with respect to the relationship with wild species (El-Gadi and Elkington 1977; Hanelt 1990; van Raamsdonk and de Vries 1992a) and to crossability (Saini and Davis 1967; McCollum 1971, 1974; Gonzalez and Ford-Lloyd 1987; van der Meer and de Vries 1990; van Raamsdonk et al. 1992; Keller et al. 1996). Diversity studies based on DNA variation have been carried out to study the position of A. cepa (Havey 1992; Wilkie et al. 1993; Bradeen and Havey 1995; Linne von Berg et al. 1996; Dubouzet et al. 1997; Mes et al. 1997, 1999; van Raamsdonk et al. 1997, 2000; Fritsch et al. 2001; Lilly and Havey 2001). Although several studies of selected sets of species of Allium subgenus Rhizirideum were based on chloroplast (cp) DNA, detailed sequence analyses of specified cpDNA fragments, or a comparison of cpDNA and nDNA based phylogenies of exactly the same set of species were rarely published (Friesen et al. 1999, 2000; Mes et al. 1999; Fritsch et al. 2001).

Chloroplast DNA is in general maternally inherited, as is the case in Allium (Havey 1995), and its lack of recombination makes it suitable for the study of hybridisation and introgression (Rieseberg and Soltis 1991; Rieseberg 1995). The importance of cpDNA for studying the aforementioned phenomena has recently been emphasized by Hollingworth et al. (1999). In Allium subgenus *Rhizirideum* the inheritance of nuclear (n) DNA variation in hybrids has been studied by means of

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the Hybrid Distance approach (van Raamsdonk et al. 2000). A cpDNA study of the same set of species will give much more insight about events of hybrid speciation, will yield information on the crossability relationship to be expected and provides the opportunity to compare phylogenies based on biparentally and maternally inherited DNA.

In the present study, sequences of four cpDNA fragments will be used for studying the diversity in cpDNA subjected to cladistic analysis. The study includes equal amounts of accessions of the sections Cepa, Rhizirideum and Schoenoprasum in order to reach a balanced share in the resulting phylogeny. The use of more than one accession for most species of sections Rhizirideum and Schoenoprasum allows us to compare the infraspecific variation with the interspecific variation. Some species of section Oreiprason (three species), of section *Petroprason* (one) and of section *Reticulato*bulbosa (one) were added for comparison. Allium tuberosum will be used as an outgroup. The aim of the present study is: (1) to analyse the cpDNA diversity and unravel the maternally based relationships in Allium subgenus Rhizirideum, (2) to study the occurrence of introgression events and of hybrid speciation, and (3) to analyse whether a shorter distance between two species in a cpDNA tree compared to their distance in a nDNA tree will indicate interfertility.

Materials and methods

Plant material

The species and accessions included in the analysis are listed in the Appendix.

Vouchers are deposited in either the herbaria of Wageningen University (WAG), the Royal Botanic Gardens in Kew (K), the Institute für Pflanzengenetik und Kulturpflanzenforschung in Gatersleben (GAT), and the Plant Research international (formerly the Institute for Horticultural Plant Breeding, Wageningen, WAHO). All accessions labeled as "BG" originate from an originally wild population.

DNA-isolation

Genomic DNA was isolated from leaf material with a midi DNA preparation procedure (Beek et al. 1992) with some minor modifications: after hooking the DNA out of the isopropanol mixture, the DNA was washed overnight in 70% EtOH and 100 mM NH₄Ac, and resuspended in 200 μ l or more-sterile TE (10 mM Tris-HCl pH = 8.0 and 1 mM EDTA). DNA concentrations were measured using a fluorometer. DNA could also be isolated with the mini prep DNA-isolation method: approximately 0.25 g of fresh leaf material was collected in Eppendorf tubes, frozen and ground in liquid nitrogen, and stored at -50 °C; 750 μ l of DNAisolation buffer (IB) with $Na₂S₂O₅$ (3.8 g/l) was added to the leaf material. This mixture was incubated for 60 min at 65 \degree C with occasional inverting of the tubes [IB = lysis buffer:extraction buffer:sarkosyl $(5\% \text{ w/v}) = 2.5:2.5:1$; lysis buffer = 0.2 M Tris-HCl $pH = 7.5$, 0.05 M EDTA, 2 M NaCl, 2% w/v CTAB; extraction buffer = 0.35 M Sorbitol, 0.1 M Tris HCl pH = 7.5 , 5 mM EDTA]. The DNA was further purified by adding $750 \mu l$ of chloroform/ isoamylalcohol (24:1), inverting the tubes (10–20 times) and centrifuging for 5 min at 14,000 rpm. After transfer of the

supernatant to a new tube the DNA was precipitated by the addition of 400- μ l isopropanol (–20 °C). The DNA could either be hooked out or it had to be pelleted for 5 min at 14,000 rpm. The DNA samples were washed once with 70% ethanol and re-suspended in 100 μ l of TE.

cpDNA sequencing

Four sequences of the cpDNA genome were amplified by specific primers: the trnL-F intergenic spacer and the trnL exon (Taberlet et al. 1991), the rps 16 intron (Oxelman et al. 1997) and the rbcL (rubisco large subunit). Four rbcL primers were designed from three Allium sequences available in the EMBL databases. The best results were obtained with primers 2 and 4, amplifying a sequence of approximately 1,100 nucleotides. Primers are listed in Table 1.

The PCR reaction mixture (total volume 25μ) contained 25 ng of template DNA, 25 ng of the primers (Table 1), 10 mM of each dNTP, 0.1 μ l (0.5 units) of superTaq polymerase, 50 mM of KCl, 10 mM of Tris-HCl, 1.5 mM of MgCl₂ and 0.01% gelatine (pH = 8.3). The mixture was covered by two drops of mineral oil. PCR was performed with a thermal cycler (Perkin Elmer) as follows: 35 cycles at 94 \degree C for 1 min (denaturation), annealing for 2 min, from 35 °C to 72 °C with a ramp of 1 °C/5 s, 72 °C for 2 min (extension); and one final cycle at 72 °C for 10 min. Optimal annealing temperatures were 60 °C for the trnL-F intergenic spacer, 55 °C for the trnL exon, 50 °C for the rps 16 intron and 60 °C for the rbcL sequence. Electrophoresis was performed through 1.5% 1 \times TBE agarose gels, for 6 to 8 h at 4 V/cm, stained with Ethidium bromide. Small fragments of the gel containing the DNA fragment were cut out and the DNA was extracted from the gel using the Qiaex protocol (QIAGEN corporation). The DNA amount was measured and 150–400 ng of DNA, depending on the size of the fragment, was used for the sequencing reaction. The fragment was amplified with a sequence mix, containing fluorescence-labelled nucleotides, and one of the primers in a PCR reaction of 25 cycles. The amplified fragment was sequenced in an ABI sequencer.

Scoring of data

All sequences were aligned manually. The resulting raw-data matrix consisted of 369 nucleotide positions for the trnL-F intergenic spacer, 339 for the trnL exon, 861 for the rps 16 intron (between primers R2 and 1A) and 1,128 for the rbcL (between primers 2 and 4). All differences were scored as 0, 1, 2 etc. depending on the number of states per mutation site. A mutation could either consist of a single nucleotide (single-site mutation) or a deletion or duplication of a stretch of nucleotides (indel). The direction of all mutations was estimated on the basis of the situation found in the outgroup (A. tuberosum).

Data analysis

All analyses are based on a matrix with digits (0, 1, 2, etc.) indicating independent states of each character per accession. All Table 2 Sequences for 33 accessions of Allium subgenus Rhizirideum of the trnL-F intergenic spacer. Mutations are indicated by capitals (nucleotide changes) or by lower-case letters (duplications or deletions) in the bottom rows of the table. Letters in the

sequences are given according to the IUB codes: A: adenosine, C: cytidine, G: guanosine, T: thymidine, R: A or G, Y: C or T, K: G or T, M: A or C, and n : any base

characters were extracted from the original data matrix with nucleotide positions. The phylogenetic reconstruction was carried out using the PAUP software package 3.1 (Swofford 1993). Data were treated as unordered and equally weighted. Consensus trees were based on all most-parsimonious trees of equal length after a heuristic search of trees using the TBR and MULPARS options. Bootstrap analysis has been carried out with 500 replicates.

Hybridisation experiments

Flowers of the female parents were emasculated by hand prior to anthesis. Mature pollen was transferred from anthers of the pollen donor to the stigma of the female parent by means of a paint brush. All pollination was carried out in insect-free glasshouses. Ripe F1 seeds were harvested during summer and sown at late autumn. Seedlings were raised during winter in frost-free glasshouses. All presumed hybrids were checked on their hybrid background by using RAPDs.

Results

cpDNA diversity

Single nucleotide mutations or duplications/deletions were found in three of the four sequences investigated.

The trnL-F intergenic spacer showed 15 single nucleotide mutations, and 12 duplications and deletions of stretches longer than one nucleotide (Table 2). Only mutations that appeared to be unambiguous after checking the original sequence data are indicated and used for phylogenetic analysis. This spacer between the trnL intron and the $3'$ end of the trnF gene includes 369 nucleotide positions in Allium subgenus Rhizirideum. All species except A. tuberosum share a duplication of nine nucleotides at position 87 in Table 2 (CAAAnAAAA). The representatives of sections Cepa and Schoenoprasum except for Allium pskemense share the absence of any further duplication in that region, whereas e.g. the species of section Rhizirideum share a further duplication of five nucleotides (TGGTT) in front of the first-mentioned duplication. It is unlikely that these two duplications arose independently, since independent duplications would imply that the five nucleotides TGGTT were shifted nine positions to the right after the duplication of the nine nucleotides. It is more likely to assume a duplication of 14 nucleotides in section Rhizirideum and at least one other duplication of nine nucleotides in the sections *Cepa* and Schoenoprasum. In this explanation it is not possible to

Table 2 (continued)

 110

 120

 130

 140

 D

EFG

i

 150

 160

 170

 180

 190

prove whether the duplication in section Cepa is the same as found in section Schoenoprasum or not. There is at least one difference between these sections at the fifth nucleotide of the nine-nucleotide duplication (position 91 in Table 2: G or T, respectively). Duplications at position 75 and at 162–164 show synapomorphisms between species of section *Oreiprason* (Allium *globosum* and Allium saxatile) and a species of section Petroprason (Allium obliquum). A duplication of 25 nucleotides is found in the sequences of A. cepa and of Allium vavilovii at position 249–273 (Table 2).

The intron of the rps16 gene sequenced with the R2 primer showed good results, but the reverse amplification with the F primer stopped at a region with nine G nucleotides. A new primer (20-mer) was designed covering the G region, which successfully amplified the remaining part of the rps16 intron sequence. This part, 175 nucleotides shorter than the part of the intron enclosed by R2 and F, showed seven single-site mutations and 15 duplications or deletions (sequences not shown). A duplication of 43 nucleotides is found in the sequences of A. cepa and of A. vavilovii, which is supported by the duplication of 25 nucleotides in the sequences of the trnL-F intergenic spacer of the same species. Agarose-gel electrophoresis showed that the intron of the rps16 gene is present in a large and a small copy, whereas Allium

 H

Fig. 1 Results of amplification of the rps16 intron for a range of species. Left: amplification at 55 °C, right: amplification at 60 °C. Note the shorter fragment in A. *fistulosum* and A. *altaicum*

fistulosum and Allium altaicum show two copies of shorter size (Fig. 1). Sequence analysis showed that the main but smaller copy in these two species is 160 nucleotides shorter than the main copy in the other species. The short copy of Allium schoenoprasum accession 96152 possesses the same deletion of 160 nucleo-

200

 \mathbf{T}

Table 2 (continued)

1052

 $\mathbf 1$

 $\begin{array}{cc} K\mathbf{L} & \mathbf{M} \\ \mathbf{I} & \mathbf{k} \end{array}$

tides. A duplication of seven nucleotides is erratically found in a range of different species. This set of seven nucleotides is included at one end of the already mentioned duplication of 43 nucleotides in A. cepa and A. vavilovii. The seven-nucleotide duplication is found at five locations in the tree of Fig. 2 (character 31), which indicates that it apparently does not influence the structure of the tree. It either might have arisen simultaneously in several events or it can be the remnant of a basic, larger duplication, that vanished in most species for its larger part during evolution. The large subunit of the rubisco gene (rbcL) is very conservative, and at the level of sections and species hardly any mutations were found. Four single-site mutations were recovered. The trnL exon with a size of 339 base pairs did not show any polymorphisms between the accessions included in the dataset.

Phylogeny

The phylogenetic analysis of the dataset including 53 mutations for 33 accessions resulted in 52 most-parsimonious trees of 78 steps with a consistency index (Cl) = 0.805 (CI_{uninformative} = 0.754) and a retention index (RI) = 0.915. One of the most-parsimonious trees is shown in Fig. 2. Section Cepa is basically divided in three clades. One consists of Allium galanthum, a second one of A. cepa and A. vavilovii, the last clade combines A. fistulosum/A. altaicum with Allium oschaninii. The species Allium pskemense of section Cepa is connected to the branch of one of the accessions Allium senescens (section Rhizirideum), but it shares some characters with certain branches of section Cepa, e.g. character 31 and 32b (Fig. 2). A. obliquum shares two duplications of three and seven nucleotides respectively in the trnL-F intergenic spacer with A. globosum and A. saxatile. A duplication of seven nucleotides in the rps16 intron is shared by A. saxatile and Allium hymenorrhizum. The 80% majority rule consensus tree based on the 52 most-parsimonious trees is shown in Fig. 3. The polytomies are caused by the synapomorphisms as indicated by the tree in Fig. 2. Sections Cepa and Schoenoprasum are closely connected, together with Allium roylei. The species of section Rhizirideum do not share a synapomorphism unambiguously, although some synapomorphisms are indicated in the individual tree of Fig. 2. Sections Petroprason and Oreiprason are not clearly separated. Bootstrap analysis

Table 2 (continued)

 $\,$ N

showed weak support to some clades as indicated in Fig. 3. Although the separate position of A. hymenorrhizum, the clade with Allium pskemense/A. senescens and the position of A. galanthum at the root of the Cepa clade are weakly supported, these positions are all supported by more than 80% of the most-parsimonious trees. The final, conclusive tree is exclusively based on branches supported after bootstrap analysis (Fig. 4b).

Crossability analysis

Twenty one crosses are carried out in 16 combinations between eight species. The results are listed in Table 3. No hybrid seeds have been obtained in the current crossability program, which might be caused by the fact that almost all crosses involve combinations of species belonging to different sections.

Table 3 Results of crossing experiments between species of Allium subgenus Rhizirideum

| Parents | Flowers | Seeds |
|---|----------------|----------------|
| Crossing Experiments in 1997 | | |
| roylei × senescens All 1122 | 10 | 0 |
| roylei × altyncolicum Tax 42 | 13 | 0 |
| roylei × karelinii Tax 536 | 7 | 0 |
| roylei × tuberosum 86001 | 18 | 0 |
| roylei × obliquum Tax 94 | 16 | 0 |
| roylei × schoenoprasum Tax 1605 | 16 | 0 |
| tuberosum 86001 \times karelinii Tax 536 | 18 | $\overline{0}$ |
| tuberosum 86001 × roylei | 15 | 0 |
| <i>tuberosum</i> 86001 self poll. | 40 | 43 |
| Crossing experiments in 1998 | | |
| cepa 96211 \times senescens All 1122 | 120 | 0 |
| cepa 96211 \times schoenoprasum Tax 1605 | 140 | 0 |
| cepa 96211 \times tuberosum 86001 | 250 | 0 |
| roylei 95001 × altyncolicum Tax 42 | 53 | 0 |
| roylei 95001 × schoenoprasum Tax 1605 | 46 | θ |
| roylei 95001 \times tuberosum 86001 | 51 | $\overline{0}$ |
| roylei 95001 self poll. | 120 | 99 |
| senescens All $1122 \times cepa$ 96030 | 60 | θ |
| senescens All 1122 × roylei 95001 | 73 | 0 |
| senescens All $1122 \times$ tuberosum 86001 | 78 | 0 |
| altyncolicum Tax $42 \times$ senescens All 1122 | 37 | 0 |
| tuberosum 86001 \times senescens All 1122 | 41 | 0 |
| tuberosum 86001 \times schoenoprasum Tax 1605 | 35 | 0 |
| tuberosum 86001 \times altyncolicum Tax 42 | 67 | 0 |

Most parsimonious cladogram

Fig. 2 One of the 52 most-parsimonious trees of 78 steps-long of representatives of Allium subgenus Rhizirideum (CI = 0.808 ; CI uninformative = 0.756 ; RI = 0.911). The *indices* of the bars indicate the position of the mutation: 1–12: deletions/duplications in the trnL-F intergenic spacer; 13–27: single-site mutations in the trnL-F intergenic spacer; 28–42: deletions/duplications in the rps16 intron; 43–49: single-site mutations in the rps16 intron; 50–53: single-site mutations in the rubisco large subunit

Discussion

Phylogeny

The division of the included species in clades is generally in agreement with the accepted taxonomic division based on morphology, cytogenetics or chemical compounds (El-Gadi and Elkington 1977; Hanelt 1990; Hanelt et al.

Fig. 3 Eighty percent majority rule consensus tree of 52 mostparsimonious trees showing the evolutionary position of 33 accessions of Allium subgenus Rhizirideum. Bootstraps values are indicated at the branches. Section names are abbreviated: Cepa: section Cepa; Rhiz: section Rhizirideum; Schoe: section Schoenoprasum; Orei: section Oreiprason; Petro: section Petroprason; R-b: section Reticulato-bulbosa; Buto: section Butomissa

1992; van Raamsdonk and de Vries 1992a; Hanelt and Fritsch 1994). The combination of species of sections Oreiprason and Petroprason in one cluster is also found by Dubouzet et al. (1997). This position is supported by the close resemblance of the species of these two sections in morphology and geographical distribution (Hanelt et al. 1992).

All species included with more than one accession (i.e. belonging to sections Schoenoprasum and Rhizirideum) appear to be monophyletic; or else due to a lack of discriminating characters monophyly was neither confirmed nor rejected (A. cepalA. vavilovii and A. schoenoprasum/Allium altyncolicum). The same conclusion has been reached for all wild species of section Cepa, but cultigens were intermixed with their immediate ancestor (van Raamsdonk et al. 1997). Monophyly of species was also found after nDNA analysis except for one accession of A. senescens that was placed in one clade with the only accession of Allium angulosum (van Raamsdonk et al. 2000). These results agree with the results of Dubouzet et

Fig. 4 Comparison of the nDNA phylogeny (van Raamsdonk et al. 2000) with the cpDNA phylogeny (current results) of a set of species of Allium subgenus Rhizirideum

al. (1997) where all accessions per species were placed in the same cluster.

The three alliances in section *Cepa* as proposed by Hanelt (1990) cannot completely be recognised in the current results (Figs. 2 and 3). The three groups, i.e. A. fistulosum and A. altaicum, A. galanthum and A. pskemense, and A. cepa, A. vavilovii and A. oschaninii, do arise after morphological analysis (van Raamsdonk and de Vries 1992a), and after analysis of nDNA variation (Bradeen and Havey 1995; van Raamsdonk et al. 2000). The species with a deviating position are A. oschaninii and A. pskemense. The position of A. oschaninii close to A. fistulosum and A. altaicum is in concordance with chromosome C-banding patterns (de Vries and Jongerius 1988), but disagrees with other data (van Raamsdonk 1992b). A distant position of A. oschaninii was also found after cpDNA analysis (Havey 1992; Lilly and Havey 2001). The current results support the main division between the clade of A. cepa/A. vavilovii and A. fistulosum/A. altaicum ("subsection Phyllodolon"). The distinction of the A. galanthum/A. pskemense alliance is not supported by the consensus tree since A. pskemense possess an A. senescens-like cpDNA type with some modifications and one notable reversal that is shared with other members of section Cepa (Fig. 2).

Each of the four main subgenera of the genus Allium posses a separate cluster after phenetic analysis of cpDNA diversity in the study of Linne von Berg et al. (1996). Species of sections Cepa and Schoenoprasum are located in the same cluster in their results, whereas the two species included in their study belonging to section Rhizirideum were placed at a larger distance. This seems to be in concordance with the present results, but different data sets can not be reliably compared by using different approaches, i.e. phylogenetic and phenetic analyses.

A phylogenetic analysis of section Cepa, including A. roylei, was carried out by Havey (1992). A. oschaninii and A. pskemense were placed at the root of the clade of section *Cepa*. Nothing can be said about the relationship between A. pskemense and any representative of section

Rhizirideum, since none of the latter were included in his analysis.

It is most important for a proper comparison of differently based phylogenies that the same set of material is used in the datasets to be compared (e.g. Sanderson and Donoghue 1989). Such a comparison was carried out using the cpDNA dataset of Havey (1992) and the RAPDnDNA dataset of van Raamsdonk et al. (1997) for the species exclusively belonging to section *Cepa*. There appeared to be a striking correlation between the cpDNA tree (Havey 1992) and the crossability dendrogram (van Raamsdonk et al. 1992), but differences in the level of crossability should be correlated with differences in species position between nDNA and cpDNA phylogenies, because of the different modes of inheritance of these types of DNA (van Raamsdonk et al. 1997). In order to extend a view on the crossability of A. cepa and A. roylei beyond the border of section Cepa, it is important to have different phylogenies available based on the same set of material.

Comparison of nDNA and cpDNA phylogenies and the analysis of introgression events

The set of species used in the current cpDNA study is identical to the set of species as used in the nDNA study of van Raamsdonk et al. (2000) except for A. oschaninii, due to the fact that most probably the pre-amplification of the nDNA of A. oschaninii did not work out well. As in the nDNA consensus tree (van Raamsdonk et al. 2000), sections Cepa, Schoenoprasum and Rhizirideum each possess a separate position in the tree (Figs. 3 and 4). The clade with A. globosum, A. saxatile and A. obliquum, combining species of sections Oreiprason and Petroprason, is also recognisable in both trees (Fig. 4). The relationships at the level of sections is not congruent between the nDNA and cpDNA-based phylogenies. For instance, the relatively close relationship between the sections Cepa and Schoenoprasum after cpDNA analysis (current results: Fig. 3) was not found after nDNA analysis (van Raamsdonk et al. 2000). The relative distances between the different sections in the nDNA tree were almost equal to each other, as was shown in the dendrogram after phenetic analysis (van Raamsdonk et al. 2000: Fig. 4).

The main differences between the cpDNA and the nDNA-based phylogenies (Fig. 4) are the remote positions of A. roylei, A. pskemense, A. lineare and A. hymenorrhizum. It is clear that A. *pskemense* possesses the A. senescens-like cpDNA, notwithstanding the fact that A. pskemense shares two mutations with accessions of the section Cepa. Considering the situation that the cpDNA genome is not recombined during gametogenesis, some sort of chloroplast-capture occurred during the evolution of A. pskemense, followed by a re-establishment of a nDNA composition that fits with those of the species of section Cepa. The same conclusion might be applied to the situation of A. roylei at the root of section Schoenoprasum, since no sharing of mutations is found with representatives of the clade of section Cepa (Fig. 2). The differences between section Cepa and section Schoenoprasum, together with A. roylei, however, are too weak to be supported by bootstrap analysis (Fig. 3). The same situation might explain the position of A. lineare and A. hymenorrhizum, but their position in the final cpDNA tree is not supported after bootstrap analysis (Fig. 4b). The relatively separated position of A. hymenorrhizum found after cpDNA as well as nDNA analysis, is comparable to the results of Dubouzet et al. (1997).

Several other analyses are based on cpDNA as well as nDNA. These studies exclude most species of section Cepa (Mes et al. 1999; Friesen et al. 2000) or are exclusively based on species of section Cepa without an outgroup from outside the section (Friesen et al. 1999). A correlation between haplotypes and geographic distribution has been found in several species of Allium, indicating either parallel adaptation or gene flow. Incongruence between datasets could not be tested for the structure of the ITS dataset (Mes et al. 1999). The nDNA and the cpDNA phylogenetic trees in the study of Friesen et al. (2000) differ only for the position of Allium kingdonii. Hybridisation as well as sample errors are presumed as a cause (Friesen et al. 2000).

Crossability and the prediction of interfertility

The currently presented crossability study is the first one that covers a range of species belonging to different sections of the subgenus Rhizirideum without using artificial pollination methods. Earlier studies focused primarily on the combination of A. cepa with other species. Hybrid plants with A. galanthum, A. fistulosum, A. vavilovii and A. roylei were obtained, whereas crosses of A. cepa with A. pskemense, A. oschaninii, A. angulosum, A. senescens and A. schoenoprasum failed (Saini and Davis 1967; McCollum 1971, 1974; Gonzales and Ford-Lloyd 1987; van Raamsdonk et al. 1992). Hybrids between A. *cepa* and a range of species are being

Fig. 5 Overview of the currently known crossability results between species of sections Cepa, Rhizirideum, Schoenoprasum, Oreiprason, Petroprason, Reticulato-bulbosa and Butomissa

produced by means of embryo-rescue techniques (Keller et al. 1996). All the currently known results are summarised in Fig. 5.

The hypothesis is proposed that a shorter distance between two species in a cpDNA tree compared to their distance in a nDNA tree will predict inter-fertility at a certain level. On the other hand, no conclusion can be drawn from a larger distance in the cpDNA tree (van Raamsdonk et al. 1997). This hypothesis is based on the situation that chloroplast capture, i.e. a short distance in a cpDNA tree, is the result of a certain level of effective inter-fertility. Introgression between species as basic mechanism can result in replacement of the cytoplasmic DNA of the recipient parent, whereas only a small portion of donor nDNA will finally be included in the recipient gene pool. The differences between the effect of introgression of nuclear and cytoplasmic-types of DNA are due to the absence of recombination in the latter (van Raamsdonk et al. 1997; compare Rieseberg and Brunsfeld 1992). The present comparison of nDNA- and cpDNAbased phylogenies with the same set of species and accessions allows a detailed analysis of the differences between bi-parentally and maternally inherited genomes, and can be used for the prediction of inter-fertility. Validation of the hypothesis depends on the availability of crossing results. The position of A. roylei (nDNA: section Cepa) close to section Schoenoprasum in the cpDNA phylogeny should indicate a certain level of crossability with representatives of this section, but the currently

available crossing results neither support not reject this assumption (Fig. 5). Band sharing of the nDNA profiles of A. roylei with those of other species revealed a larger percentage of bands shared with representatives of section Rhizirideum than of section Schoenoprasum (van Raamsdonk et al. 2000). Most probably a considerable secondary evolution after the putative hybridisation and introgression event took place in A. roylei. Therefore, the effect of parallel evolution can not completely be ruled out. A. roylei is close to an A. cepa accession with the S-type cytoplasm (Lilly and Havey 2001). This CMS onion might have been originated after introduction of an alien cytoplasm from a A. roylei-like plant relating to some representative of section Schoenoprasum. The position of A. pskemense close to A. senescens in the cpDNA tree might also predict some interfertility between

Appendix: list of plant material used

this species and A. senescens. However, this prediction can not be approved by any crossability result of A. pskemense with a representative of section Rhizirideum (Fig. 5). Further crossability results are necessary in order to find supportive evidence for the hypothesis that a shorter distance between two species in a cpDNA tree compared to their distance in a nDNA tree might indicate a certain level of inter-fertility.

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