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Genome composition of ‘Elatior’-begonias hybrids analyzed by genomic in situ hybridisation

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Abstract Interspecific hybridization of various tuberous *Begonia* species hybrids with *Begonia socotrana* results in so-called ‘Elatior’-begonias hybrids (*B.* × *hiemalis* Fotsch). In our study, genomic in situ hybridization (GISH) has been employed to assess the genome composition in eleven ‘Elatior’-begonias hybrids and their ancestor genotypes. Genomic DNA of tuberous *Begonia* was sonicated to 1–10-kb fragments, labelled by nick translation with digoxigenin-11-dUTP and used as a probe whereas *B. socotrana* DNA was autoclaved to 100 bp fragments and used as block. The genome of tuberous *Begonia* was clearly pronounced in ‘Elatior’-begonias when the probe concentration was ~ 3.75 ng/ μ l (150 ng/slide), with 30 times the excess of *B. socotrana* blocking DNA and stringency of post hybridization washings at 73% ($0.1 \times$ SSC at 42°C). In ‘Elatior’-begonias hybrids GISH distinguished two

groups comprising short (0.6–1.03 μ m in length) and relatively longer chromosomes (1.87–3.88 μ m) which represent *B. socotrana* and tuberous *Begonia* genomes, respectively. The number of chromosomes derived from tuberous *Begonia* ranged from 14 to 56 and for *B. socotrana* from 7 to 28 which suggest the presence of different ploidy levels in analyzed ‘Elatior’-begonia hybrids. Intergenomic recombination has not been detected through GISH in hybrids analyzed. Genomic in situ hybridization turned out to be useful to identify the genome constitution of ‘Elatior’-begonia hybrids and thus gain an insight into the origins of these cultivars. This knowledge on the ploidy level and genome composition is essential for further progress in breeding Begonias.

Keywords *Begonia hiemalis* Fotsch · Genome differentiation · Interspecific hybrids · Polyploidy · *Begonia*

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Introduction

The genus *Begonia*, a member of the family Begoniaceae, includes around 1500 species distributed mainly in tropical and subtropical regions (Doorenbos et al. 1998; Wagner 1999). *Begonia* species are widely hybridized for improving flowering and foliage of plants where interspecific hybridization is the most suitable method for the transmission into

new cultivars of important agricultural traits, such as winter flowering capacity, new flower shapes and colours and disease resistance. The current commercial assortment comprise mostly of ‘Elatior-hybrids’ (*B. × hiemalis* Fotsch) (over 100 varieties) representing around 88% of the total *Begonia* production (Kroon 1993). The early varieties of ‘Elatior’-begonias resulted from crosses between winter flowering *Begonia socotrana* Hook and ‘Viscountess Doneraile’, a very early hybrid tuberous Begonias (*B. × tuberhybrida* Voss) (Arends 1970; Doorenbos 1973). However, the majority of modern varieties of ‘Elatior’-begonias are the result of crossing various tuberous *Begonia* species hybrids (*B. × tuberhybrida* Voss) and *B. socotrana* Hook. f (Gleed 1961). Some varieties of ‘Elatior-hybrids’ have also been obtained from the following crosses: ‘Elatior’-begonia × *B. socotrana* and ‘Elatior’-begonia × tuberhybrid (Arends 1970).

Chromosome numbers have been reported for many species of *Begonia*, ranging from $2n = 18, 22, 26, 28, 30, 38, 41, 44, 46, 52$ to 82 (Okuno and Nagai 1954; Zeilinga 1962; Ye et al. 2004; Nakata et al. 2007) which suggests the high levels of polyploidy and aneuploidy in the genus. The progenitors of ‘Elatior’-begonia hybrids, the tuberous cultivars have originated from hybrids between a number of widely diverging wild species with $2n = 26$ and $2n = 28$ chromosomes (Legro and Doorenbos 1969). By breeding and selection *B. × tuberhybrida* has been brought to polyploid level (Kroon 1993). In the hybrid tuberous Begonias (*B. × tuberhybrida* Voss) several different chromosome numbers were found ($2n$ from 27 to 64) (Okuno and Nagai 1954; Legro and Haegeman 1971). *B. socotrana*, the second ancestral species of ‘Elatior’-begonia hybrids, comprises of $2n = 2 \times = 28$ chromosomes (Matsuura and Okuno 1936; Doorenbos and Legro 1968).

Chromosomes of begonia are relatively small ranging in length from $0.5 \mu\text{m}$ in *B. socotrana* (Arends 1970) to $5.43 \mu\text{m}$ in *B. coptidifolia* (Ye et al. 2004) and poorly differentiated (Legro and Haegeman 1971). Due to small size and similar morphology, the evaluation of chromosome number and the discrimination of genomes in hybrids can be liable to mistakes (Legro and Doorenbos 1969). For some *Begonia* genotypes different chromosome numbers have been established by different authors (Zeilinga 1962). For this reason verification of

Begonia hybrids based on chromosome number or karyomorphology may be problematic.

Genomic in situ hybridization (GISH), which uses total genomic DNA of one of the parents as the probe, is a very effective molecular cytogenetic method usually providing clear and unambiguous distinction between genomes. This technique is especially effective in plants with large chromosomes such as in *Lilium* (Barba-Gonzalez et al. 2006), *Alstroemeria* (Kamstra et al. 1999) or *Tulipa* (Marasek et al. 2006). However, it has been also successfully applied to differentiate genomes with small chromosomes such as in *Oryza* (Li et al. 2001), *Lycopersicon* (Haider Ali et al. 2002), *Brassica* (Hasterok et al. 2005), and *Arabidopsis* (Ali et al. 2004). As yet, genomic in situ hybridization has not been applied to analyze hybrids of the genus *Begonia*.

The aim of present study was to optimize GISH technique for *Begonia* chromosomes and to identify *B. socotrana* and Tuberous hybrids genomes in ‘Elatior’-begonia hybrids of different ploidy level thus gaining an insight into the origins of these cultivars as well as the basic chromosome number.

Materials and methods

Plant materials

Begonia socotrana, two tuberous *Begonia* hybrids and eleven different accessions of ‘Elatior’-begonias hybrids were used in this study for GISH analysis (Table 2). All genotypes were provided by Beekenkamp B.V., Maasdijk, the Netherlands. For analysis of chromosome morphology and adaptation of GISH technique for *Begonia* chromosomes, tetraploid tuberous *Begonia* characterized by erect plant, totally filled red flowers and tubers has been selected as a representative genotype of tuberous Begonias. The cuttings from all hybrids and species were rooted in greenhouse under standard growing condition applicable for begonia cultivation ($20\text{--}25^\circ\text{C}$) and being maintained in Wageningen UR, Plant Breeding, The Netherlands.

Chromosome preparation

Root tips were pre-treated with 0.05% colchicine for 2.5 h and then fixed in 3:1 ethanol:glacial acetic acid.

The roots were subjected to enzymatic digestion in a mixture comprising 0.2% (w/v) pectolyase Y23, 0.2% (w/v) cellulase RS at 37°C for about 2 h. Meristems were squashed in a drop of 45% acetic acid. After freezing in liquid nitrogen, cover slips were removed by using a razor blade and the preparations were dehydrated in absolute ethanol and air dried. The best slides were selected under a phase contrast microscope (Leica Dialux 20 EB) and stored at –20°C until use.

Preparation of probe and block DNA

Total genomic DNA of *B. socotrana* and tuberous Begonia was extracted from young leaves using modified method of Fulton et al. (1995). The isolation was preceded by the initial wash step of powdered leaf tissue with TE buffer (10 mM tris-HCL, 10 mM EDTH, pH 8) according to Kopperud and Einset (1995). Tuberous Begonia DNA were sonicated to 1–10-kb fragments and labeled by nick translation with digoxigenin-11-dUTP by a standard nick translation protocol (Roche Diagnostics GmbH, Mannheim, Germany). Block DNA of *B. socotrana* was obtained by autoclaving for 5 min to a fragments 100–300 bp.

Genomic in situ hybridization and detection

DNA denaturation and in situ hybridization steps were performed according to Hasterok et al. (2001) and Marasek et al. (2006). Slides were pre-treated with RNase A (100 µg/ml) for 1 h at 37°C, treated with 10 mM HCl at 37°C for 2 min followed by incubation in pepsin solution (5 µg/ml) for 10 min and post-fixed in 1% formaldehyde in PBS buffer for 10 min. The hybridization mixture consisted of 50% deionized formamide, 10% dextran sulphate, 2× SSC, 1% SDS, 150 ng of probe DNA per slide and block DNAs (*B. socotrana* DNA and herring sperm DNA) in 30- to 60-fold excess of labeled probe. The hybridization mix was denatured for 10 min at 75°C and placed on ice for 10 min. After the hybridization mix was add to the slides, a 4.5 min denaturation step at 70°C was carried out. Hybridization was done overnight in a humid chamber at 37°C. The post-hybridization washes were carried out for 15 min in 2× SSC at room temperature, followed by washes in 0.1× SSC at 42°C for 30 min (73% stringency) and

2× SSC for 15 min at room temperature. Digoxigenin-labelled DNA was detected with antidigoxigenin-FITC (sheep) (Boehringer, Mannheim, Germany) and amplified with anti-sheep-FITC (rabbit) (Vector Laboratories). The chromosomes were counterstained with 1 µg/ml 4,6-diamidino-2-phenylindole (DAPI, Sigma) in Vectashield (Vector Laboratories).

Image capturing and processing

Images of fluorescently stained chromosomes were acquired using a Canon digital camera attached to an Axiophot microscope with an appropriate filter and then processed using software (Axio Vision 4.2). For each genotype 8–15 metaphases have been analysed at different stages of chromosomes contraction. Chromosome length was determined using freeware application MicroMeasure available on the Internet at the <http://www.colostate.edu/Depts/Biology/MicroMeasure>.

Results

GISH technique in *Begonia*

In this study, we have attempted to isolate DNA from begonia leaves using several conventional procedures (Aljanabi and Martinez 1997; Murray and Thompson 1980; Fulton et al. 1995). None of these methods yielded above 5 µg DNA per gram fresh weight of begonia leaves both for *B. socotrana* and tuberous Begonia cultivar. The initial washing step of powdered leaf tissue in T10E10 buffer according to Kopperud and Einset (1995) increased the yield to 20–30 µg of DNA per gram fresh weight of leaves.

Initially, the total genomic DNA of both tuberous Begonia cultivar and *B. socotrana* labeled with digoxigenin-11-dUTP and biotin-11-dUTP respectively, were used as hybridization probes to differentiate the respective chromosomes in the cells of tuberous Begonia, *B. socotrana* and ‘Elatior’-begonias (Table 1). Since genomic probe of *B. socotrana* did not label chromosomes of *B. socotrana*, digoxigenin labeled genomic probe of tuberous Begonia were used exclusively for differentiation of the chromosomes in ‘Elatior’-begonias. The genome of tuberous Begonia was clearly pronounced in ‘Elatior’-begonias when the probe concentration was

Table 1 The genomic probes and blocking DNA concentration used in GISH experiment for differentiation chromosomes in ‘Elatior’-begonias hybrids

Slide	DNA probe of <i>B. socotrana</i> ^a	DNA probe of tuberous Begonia ^a	Block DNA of <i>B. socotrana</i> ^{bc}	Block DNA of tuberous Begonia ^{bc}	Remarks
<i>B. socotrana</i>	75 ng (biotin)	–	–	2.2 µg (1:30)	No labeling
<i>B. socotrana</i>	150 ng (biotin)	–	–	4.5 µg (1:30)	No labeling
<i>B. socotrana</i>	150 ng (biotin)	–	–	–	No labeling
Tuberous hybrid	–	150 ng (digoxigenin)	9 µg (1:60)	–	Equal labeling of chromosomes
‘Elatior’-begonia	150 ng (biotin)	–	–	4.5 µg (1:30)	No labeling
‘Elatior’-begonia	–	150 ng (digoxigenin)	4.5 µg (1:30)	–	Tuberous Begonia chromosomes fully discriminated

Post hybridization washings were carried out at the stringency 73% (0.1× SSC at 42°C) in all treatments

^a Amount of labeled probe per slide

^b Amount of blocking DNA per slide

^c In brackets the ratios of probe DNA to blocking DNA

~3.75 ng/µl (150 ng/slide), with 30 times the excess of *B. socotrana* blocking DNA and stringency of post hybridization washings at 73% (0.1× SSC at 42°C).

Chromosome morphology of the ancestral genotypes of ‘Elatior’-begonia hybrids

Figure 1 shows the somatic metaphase chromosomes of tuberous Begonia (2n = 4x = 56) and *B. socotrana* (2n = 2x = 28). Tuberous Begonia chromosomes

were labeled uniformly when using total genomic DNA of tuberous Begonia as a probe (FITC-green fluorescence) (Fig. 1a). Chromosomes were small ranging from 1.87 to 3.88 µm and little differentiated. Centromeres were distinct only for some chromosomes when they were less condensed at prometaphase stage. Figure 1b represents the chromosome complement of *B. socotrana* stained with DAPI (blue fluorescence). The chromosomes at metaphase were particularly small ranging from 0.6 to 1.03 µm in length. The

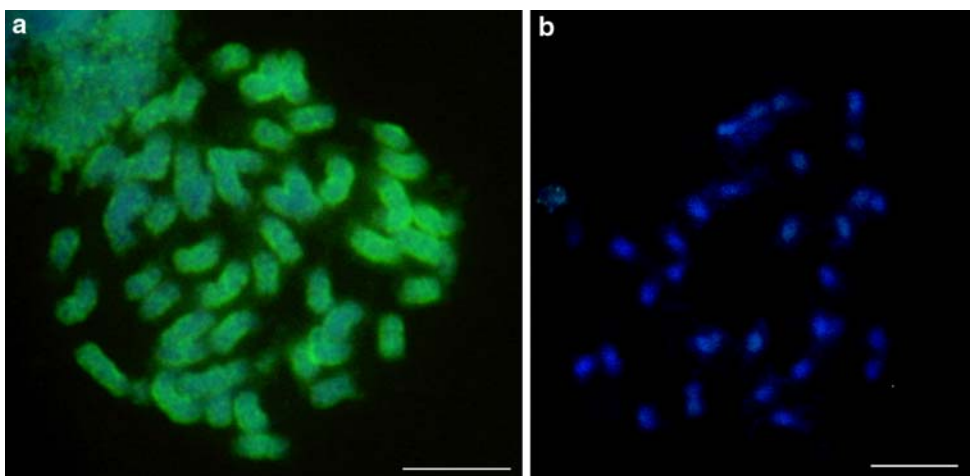


Fig. 1 a Tuberous Begonia chromosomes (2n = 4x = 56) probed with labeled nuclear DNA of tuberous Begonia with digoxigenin-11-dUTP and detected by anti-Dig FITC

(pale/green fluorescence). **b** *B. socotrana* chromosomes (2n = 2x = 28) detected by DAPI (dark/blue fluorescence). Bars = 5 µm (Color figure online)

position of the centromeres was not discernible. Both for chromosomes of tuberous Begonia and *B. socotrana* it seems impossible to identify homologous pairs due to the lack of differentiation of chromosomes.

Chromosomes differentiation in ‘Elatior’-begonia hybrids as revealed by GISH

The genome constitution of analyzed genotypes based on the results of genomic in situ hybridization and their origin are summarized in Table 2. GISH clearly distinguished two parental genomes in all ‘Elatior’-begonia hybrids analysed (Fig. 2a–j) except for genotype 04-122-04 ($2n = 44$) in which chromosomes derived from tuberous Begonia were found exclusively (Fig. 2k). Figure 2l represents tuberous Begonia hybrid 04-117-04 ($2n = 4x = 56$) resulted from crosses between tetraploid forms of tuberous Begonia. As genomic DNA of tuberous Begonia was used as a probe in this way green fluorescence (FITC) represented the chromosomes of tuberous Begonia whereas short *B. socotrana* chromosomes were stained by DAPI (blue fluorescence). In ‘Elatior’-begonia hybrids, the number of chromosomes derived from tuberous Begonia ranged from 14 to 56 and from *B. socotrana* from 7 to 28. In three out of 12 genotypes analyzed (genotypes HO-7256-3, HO-7256-4, H08-124-01) GISH revealed that 14 chromosomes were inherited from tuberous Begonia and 14 chromosomes from *B. socotrana* (Fig. 2a–c). Four

genotypes of ‘Elatior’-begonia (‘Netja Dark’, H07-102-13, H07-III-02, H08-143-03) had in total $2n = 42$ chromosomes, 28 long chromosomes belonging to tuberous Begonia (green fluorescence) and 14 short chromosomes of *B. socotrana* (blue fluorescence) (Fig. 2d–g). In hybrid 04-187-23 ($2n = 63$) it was possible to distinguish 56 chromosomes of tuberous Begonia and 7 chromosomes of *B. socotrana* (Fig. 2h). In genotype H08-132-01, GISH showed $2n = 56$ chromosomes of which 28 represented tuberous Begonia genome and 28 *B. socotrana* genome (Fig. 2i) whereas in H08-143-01 ($2n = 46$) 27 chromosomes were derived from tuberous Begonia genome and 18 from *B. socotrana* (Fig. 2j).

For some genotypes with less condensed metaphases tuberous Begonia chromosomes with satellites were observed (Fig. 2, indicated by arrows). Recombinant chromosomes were not detected via GISH in any ‘Elatior’-begonia hybrids analyzed.

Discussion

Karyomorphology of ‘Elatior’-begonia hybrids

Begonia is characterized by small and morphologically little differentiated chromosomes. In this study the position of the centromeres was not discernible for *B. socotrana* chromosomes whereas in chromosomes derived from tuberous Begonia centromeres

Table 2 The origin and genome composition of Begonia genotypes analyzed by GISH

Accession No.	Origin ^a	Chromosome number	Chromosomal constitution	Ploidy ^b
<i>B. socotrana</i>	SS	28	28S	2x (4x)
Tuberous begonia	TTTT	56	56T	4x (8x)
04-117-04	TTTT × TTTT	56	56T	4x (8x)
HO-7256-3	TT × SS	28	14T + 14S	2x (4x)
HO-7256-4	TT × SS	28	14T + 14S	2x (4x)
H08-124-01	TT × SS	28	14T + 14S	2x (4x)
‘Netja Dark’	TTTT × SS	42	28T + 14S	3x (6x)
H07-102-13	TTTT × SS	42	28T + 14S	3x (6x)
H07-III-02	TTTT × SS	41	27T + 14S	3x (6x)
H08-143-03	TTTT × SS	42	28T + 14S	3x (6x)
04-187-23	Unknown × TTTT	63	56T + 7S	4x + 7 (9x)
H08-132-01	TTS × SS	56	28T + 28S	4x (8x)
H08-143-01	Unknown	45	27T + 18S	3x + 4 (6x + 4)
04-122-04	Unknown	44	44T	3x + 2 (6x + 2)

^a T—genome of tuberous Begonia; S—*B. socotrana* genome

^b Ploidy based on $x = 13$ and 14 according to Legro and Haegeman (1971). In brackets ploidy based on $x = 6$ and 7 according to Matsuura and Okuno (1936)

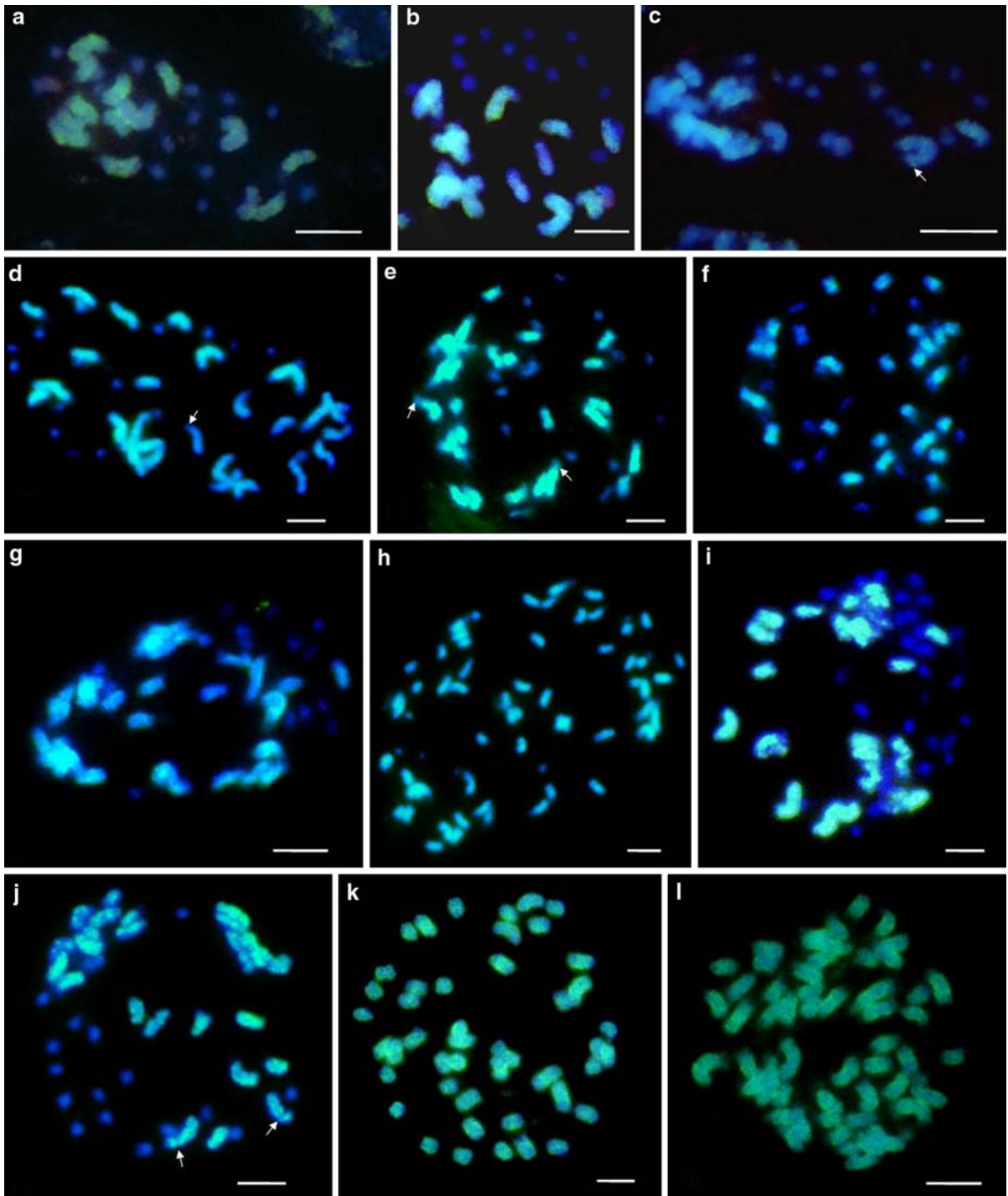


Fig. 2 Chromosome discrimination in *Begonia* hybrids. The digoxigenin-11-dUTP labeled tuberous *Begonia* genomic DNA was detected with anti-Dig FITC (*pale/green fluorescence*) and *B. socotrana* chromosomes were counterstained with DAPI (*dark/blue fluorescence*). **a** HO-7256-3 ($2n = 28$); **b** HO-7256-4 ($2n = 28$); **c** H08-124-01 ($2n = 28$); **d** 'Netja Dark'

($2n = 42$); **e** H07-102-13 ($2n = 42$); **f** H07-III-02 ($2n = 41$); **g** H08-143-03 ($2n = 42$); **h** 04-187-23 ($2n = 63$); **i** H08-132-01 ($2n = 56$); **j** H08-143-01 ($2n = 45$); **k** 04-122-04 ($2n = 44$); **l** 04-117-04 ($2n = 56$). *Arrows* indicate chromosomes carrying satellites. Bars = 5 μm (Color figure online)

were distinct only for a few chromosomes at less condensed stage in prometaphase. Similar conclusions were drawn by Arends (1970) for ‘Elatior’-begonias and Okuno and Nagai (1954) and Legro and Haegeman (1971) for hybrids tuberous Begonias (*B. × tuberhybrida* Voss). The chromosomes of *B. socotrana* differ in size from those of tuberous Begonias, so it seems possible to identify them in ‘Elatior’-begonias hybrids. However, in our study, in a few metaphase plates chromosomes carrying a satellite were observed (Fig. 2, arrows) and the minute chromosomes of *B. socotrana* and satellites were difficult to recognize from each other in crowded metaphases of ‘Elatior’-begonias. Chromosomes carrying satellites were also observed for other species e.g., in *B. evansiana* (Okuno and Nagai 1953) and *B. semperflorens* (Zeilinga 1962).

Genome differentiation by GISH

GISH distinguished chromosomes derived from tuberous Begonia and *B. socotrana* in ‘Elatior’-begonia. The uniform labeling of tuberous Begonia genome was observed when the probe concentration was ~ 3.75 ng/ μ l and the ratio of probe to *B. socotrana* blocking DNA was 1:30. *B. socotrana* DNA used as a probe failed to label *B. socotrana* chromosomes both in presence and lack of the blocking DNA, which may be caused by the very small genome size-0.63 pg/1C (Marie and Brown 1993) and small amount of repetitive DNA. The problem with hybridization of genomic probes has been also observed in other plant species with small genomes. For many species with small genomes having a relatively low proportion of the middle and high repetitive DNA families, GISH signals were limited to the pericentromeric regions for example in *Brassica* (Hasterok et al. 2005), *Oryza* (Li et al. 2001), *Rubus* (Lim et al. 1998) and *Brachypodium distachyon* (Hasterok et al. 2004). However, there are also a few species with small genomes, e.g. in the genera *Musa* (D’Hont et al. 2000), *Solanum* and *Lycopersicon* (Gavrilenko et al. 2001) where genomic probes hybridize to the entire length of the chromosomes. This has been interpreted to be the result of a more even distribution of repetitive DNA families along the chromosomes. According to Raina and Rani (2001) the critical genome size below which it is difficult to paint along entire chromosomes might

be around 0.6 pg/1C. Nevertheless, in *Arabidopsis thaliana* representing the smallest genomes among angiosperms (1C = 0.16 pg; according to Bennett et al. 2003) a uniform labeling of entire chromosomes was achieved by increasing DNA probe concentration up to 7.5–15 μ g per slide or 5 μ g of probe and increasing time of hybridization to 60 h (Ali et al. 2004). Due to satisfying differentiation of genomes in ‘Elatior’-begonia hybrids using single-target GISH with tuberous Begonia probe we did not try to modify the method by increasing concentration of *B. socotrana* DNA.

Chromosome constitutions of ‘Elatior’-begonia hybrids

Most of the progeny of ‘Elatior’-begonias are said to be triploids having 40–42 chromosomes, a few are diploids and occasionally tetraploids with 56 chromosomes (Mikkelsen 1976; Hvoslef-Eide and Munster 2006). In our study, seven different chromosome numbers have been found in eleven genotypes of ‘Elatior’-begonia hybrids (Table 2). The observation of 14 chromosomes of tuberous Begonia and 14 of *B. socotrana* in three accessions of ‘Elatior’-begonia hybrids (Fig. 2a–c) indicates that they are the result of hybridizing at diploid level of *B. socotrana* ($2n = 28$) with tuberhybrids ($2n = 28$). According to Arends (1971), crosses between diploid tuberous Begonia hybrids and *B. socotrana* do not succeed. However, Doorenbos (1973) obtained diploid ‘Elatior’-begonias hybrids from hybridization between diploid cultivar of the ‘Pendula’-type with *B. socotrana*.

Four ‘Elatior’-begonias hybrids investigated in our study (‘Netja Dark’, H07-102-13, H07-III-02, H08-143-03) were triploids (27-28T + 14S) (Fig. 2d–g) whereas genotype H08-143-01 was near triploid ($3x + 4$) comprising of 27 tuberous Begonia and 18 *B. socotrana* chromosomes (Fig. 2j). Similar genome composition has been observed by Arends (1970) in other twenty-one ‘Elatior’-begonias hybrids. Our conclusion on the origin of these hybrids are consistent with those presented by Arends (1970) that they resulted from crosses of tuberhybrids with $2n = 54$ or 56 chromosomes and *B. socotrana* with $2n = 28$ chromosomes.

The hybrid H08-132-01 ($2n = 28T + 28S$) (Fig. 2i) may be the result of backcrossing of

'Elatior'-begonia ($2n = 28T + 14S$) with *B. socotrana* ($2n = 28S$) where the former genotype produced diploid egg cell. Similar genome constitution of 'Elatior'-begonia can be expected from crosses between tetraploid tuberous Begonia hybrids ($2n = 4x = 56$) with the *B. socotrana* where the species provided unreduced gametes. Dewitte et al. (2009) proved that occurrence of viable $2n$ pollen is not a rare phenomenon in *Begonia*. In their study, 14% of investigated plants produced unreduced gametes. They have obtained successful crosses with genotypes producing $2n$ pollen, and showed via flow cytometry analysis the increase of the DNA content in the progenies. Similarly, in *Begonia*, allotetraploid semperflorens begonias have originated with the use of unreduced gametes (Horn 2004). However, there is no data on gamete formation in *B. socotrana*.

In genotype 04-122-04 ($2n = 44$) (Fig. 2k) all chromosomes showed uniform labelling with tuberous Begonia DNA probe. This plant has morphological characters of tuberous Begonia (date not shown) and may have developed apomictically from unfertilised maternal cells. After crosses between *B. socotrana* and *B. × semperflorens-cultorum*, Preil and Lorenz (1983) obtained progenies having the genotype of the male parent which must have originated from androgenesis or by elimination of the maternal chromosomes. Another possible explanation of genome composition in 04-122-04 may be elimination of *B. socotrana* chromosomes. The elimination of *B. socotrana* chromosomes has been suggested before by Arends (1970) for two 'Elatior'-begonias 'Riegers Leuchtfeuer' ($28T + 9S$) and 'Rose Queen' ($28T + 12S$). Chromosome elimination has been reported in cross in other species, e.g. *Nicotiana tabacum* × *N. plumbaginifolia* (Ar-Rushdi 1957). The selective elimination of the chromosomes of one of the parental genotypes may be associated with disturbances in nuclear division (such as non-congressed chromosome at metaphase, lagging chromosomes and bridges at anaphase, chromatin fragments, degraded type of chromatin, multipolar spindles and micronuclei) in hybrid embryos and endosperm (Lange 1971; Bennett et al. 1976). According to Noda and Kasha (1981) the failure of congregation of some chromosomes during prometaphase appears to be main mitotic disturbance

leading to chromosome elimination in hybrids between *Hordeum vulgare* and *H. bulbosum*. Gupta (1969) suggested that significant difference between the S and G2 periods of the parental species might lead to elimination of a chromosome segments or the whole genome of the species having the longer S or G2 phases.

In genotype 04-187-23 (Fig. 2h) 56 chromosomes were derived from tuberous Begonia and 7 chromosomes from *B. socotrana*. The same genome constitution was observed by Arends (1970) in a second generation 'Elatior'-begonia 'Eveleens Orange' which originated as the result of crossing 'Elatior'-begonia 'Flambeau' ($2n = 52T + 14S$) with the tuberous hybrid 'Flamboyant' ($2n = 42T$).

Basic chromosome number

The genome constitutions of 'Elatior'-begonias 04-187-23 and 'Eveleens Orange' having 7 chromosomes of *B. socotrana* are difficult to explain considering the basic chromosome number in *Begonia* $x = 13$ and 14 which have been postulated by Legro and Haegeman (1971). It is also difficult to explain that according to Legro and Haegeman (1971) a triploid tuberous begonia 'Tasso' is propagated by seeds. These data may support suggestion of Matsuura and Okuno (1936, 1943) that the basic chromosome number in the genus *Begonia* is 6, 7 and 13, where the last may be of secondary origin derived from synthesis the former two. Similar conclusions were drawn by Okuno and Nagai (1953, 1954) based on analysis of meiotic chromosome configuration in *B. evansiana* and *B. tuberohybrida*. In our study, taking into consideration 6 and 7 as the basic chromosomes number in genus *Begonia*, *B. socotrana* ($2n = 28$) would be an autopolyploid which would explain genome composition in 04-187-23 and 'Eveleens Orange' ($2n = 56T + 7S$) with 7 chromosomes of *B. socotrana*. In Table 2 the ploidy level of analyzed genotypes were evaluated based on basic chromosome number $x = 13$ and 14 according to Legro and Haegeman (1971) and $x = 6$ and 7 according to Matsuura and Okuno (1936). This knowledge on the ploidy level and genome composition of analyzed 'Elatior'-begonias is essential for further progress in breeding Begonias.

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