# CYTOGENETICS OF INTERGENERIC HYBRIDS BETWEEN DURUM WHEAT (TRITICUM TURGIDUM L.) WITH THINOPYRUM INTERMEDIUM AND SUB-SPECIES ACUTUM, GLAUCUM, PULCHERRIMUM, TRICHOPHORUM, VARNENSE

## A. MUJEEB-KAZI<sup>1</sup>, ALVINA GUL<sup>1</sup>, SUMAIRA RIZWAN<sup>2</sup>, MUHAMMAD FAROOQ<sup>2</sup>, HADI BUX<sup>2</sup>, IFTIKHAR AHMAD<sup>2</sup>, JAVED IQBAL MIRZA<sup>2</sup>, ROMAN DELGADO<sup>3</sup>, VICTOR ROSAS<sup>3</sup> AND ALEJANDRO CORTES<sup>3</sup>

<sup>1</sup>National Institute of Biotechnology and Genetic Engineering (NIBGE), Faisalabad, Pakistan, <sup>2</sup>National Agricultural Research Center (NARC), Park Road, Islamabad, Pakistan and <sup>3</sup>International Maize and Wheat Improvement Center (CIMMYT), Apartado Postal 6-641, Mexico D. F. 06600, Mexico

#### Abstract

Towards diversifying the germplasm base available for durum wheat (Triticum turgidum) the production and morpho-cytogenetic categorization of  $F_1$  hybrid combinations between durum wheat cultivars and *Thinopyrum intermedium* and its sub-species (acutum, glaucum, pulcherrimum, trichophorum, varnense) are reported. All  $F_1$  hybrids were mitotically stable with 2n=5x=35chromosomes, expressed a co-dominant phenotype and exhibited mean meiotic metaphase I chromosomal associations, that in general, do not support alien genetic introgression into the A and / or B genomes of durum wheat via recombinational exchange. This F<sub>1</sub> perennial germplasm has formed the basis to generate backcross derivatives, amphiploids, and shall enable the application of genetic manipulation strategies for transferring useful genes from select combinations for durum wheat improvement. Backcross 1 derivatives ( $F_1$  / durum wheat) were cytologically stable with 2n=7x=49 but exhibited poor self-fertility hampering their maintenance as genetic stocks. Amphiploids however had good stability at the C-0 stage with 2n=10x=70 chromosomes, were self-fertile and set healthy seed progeny. Subsequent generations derived from C-0 seed gave derivatives that were closely true to the C-0 type in composition, as well as combinations that indicated genomic loss where all combinations possessed 56 chromosomal progeny. Aneuploidy, both hyper- and hypo-ploidy, was rampant across both the 56 and 70 chromosome progenies.

#### Introduction

Sustaining agricultural productivity has been associated with genetic diversity and durability of stress resistances (Mujeeb-Kazi, 1998). For wheat improvement the genetic diversity resides in various species of the primary, secondary and tertiary Triticeae gene pools (Jiang *et al.*, 1994). The utilization of this diversity has selective protocols associated with intergeneric or interspecific methodologies. Genes from these species when introgressed and pyramided together can be anticipated to provide crop yield stability by complementing the durability resistance contributed by conventional major and/or minor genes. Numerous wide hybrids have been described during the last decade and a half (Sharma & Gill, 1983; Mujeeb-Kazi & Bernard, 1985; Wang, 1989; Jiang *et al.*, 1994. Mujeeb-Kazi & Hettel, 1995; Sharma 1995) essentially due to the circumvention of species crossability barriers coupled with embryo rescue and plantlet regeneration protocols. The hybridization emphasis however, has mainly been with bread wheat x alien species, due to the fact that bread wheat is the major cereal crop across several global mega-environments. Its cultivated hectarage significantly surpasses that of

durum wheat. Though less, durum wheat cultivation does have its special utilization significance, but faces vulnerability to various biotic/abiotic stresses. Some notable stress constraints are lack of, or further need for resistances/tolerances to *Fusarium graminearum, Helminthosporium sativum*, barley yellow dwarf virus, *Septoria nodorum*, drought and salinity tolerance. These constraints can be addressed by exploiting the genetic diversity within the Triticeae gene pools. The first step however, is generation of durum wheat x alien species hybrids, validation and characterization of the germplasm, with ultimate exploitation of the hybrid combinations for agricultural practicality. We describe here the production of intergeneric hybrids of the *Th. intermedium* sub-species group with various durum cultivars, validate them with morphological-cytogenetic documentation and emphasize their potential contribution to practical sustainable crop production outputs exemplified by advancing the  $F_1$  combinations by backcrossing and amphiploid induction.

#### **Materials and Methods**

**Germplasm:** Seeds of the alien Triticeae species used in this hybrid production study were obtained from late Dr. D. R. Dewey, USDA / ARS Logan, Utah and germinated in Jiffy-7 peat pellets. After 6-weeks of juvenile growth, the seedlings were vernalized in a growth chamber under environmental regimes of 8h diffuse light for 8 weeks at 8°C. Following vernalization, the seedlings were transplanted into 20 cm plastic pots filled with a 2:1:1 (soil: sand: peat) steam-sterilized mix and maintained under greenhouse conditions of 16h of natural daylight and 24°C/14°C day/night temperatures. In the same greenhouse, five plantings in pots (four plants/pot) of several *Triticum turgidum* cultivars were made, 15 days apart. The durum cultivar seeds were obtained from CIMMYT's wheat germplasm bank at El Batan, Mexico, the location at which the major part of this study was conducted.

**Hybrid production:** Spikes of the durum wheat cultivars were emasculated, pollinated by the perennial species pollen 1 to 3 days after emasculation and treated once daily for 3 days with 75ppm gibberellic acid. From the seed set, the embryos were excised 13 to 15 days after pollination and cultured on a special medium for small embryos (Taira & Larter, 1978). These, and subsequent procedures associated with embryo differentiation, plantlet growth, transfer to Jiffy-7 pots, and transplanting to a potted soil mix in the greenhouse, were similar to those reported by Mujeeb-Kazi *et al.*, (1987, 1989). The environmental growth regimes were identical to those maintained for the growth of the parental germplasms in this study.

**F**<sub>1</sub> cloning, somatic/meiotic sampling and colchicine treatment: After assuming vigorous growth, each  $F_1$  hybrid was physically divided into 4 plants, and the clones allowed to grow into vigorous plants. From each clone of each  $F_1$  hybrid, root-tips were collected for somatic cytology and C-banding. The cytological procedures were essentially similar to those described by Mujeeb-Kazi & Miranda, (1985) and Jahan *et al.*, (1990).

Spikes for meiotic analyses were collected in early morning hours (8:00-9:00 a.m.), fixed in Carnoy's (6:3:1, absolute alcohol : chloroform : acetic acid) for 48-72h, and stored under refrigeration ( $4^{\circ}$ C) in 70% alcohol until use. Anthers at metaphase 1 were

1218

stained in alcoholic-acid-carmine for several days, and squashed in 45% acetic acid with a drop of 2% aceto-carmine to enhance the coloration. Meiotic chromosome associations were analyzed at metaphase I. Cytological photography was done of quality representative cells on a black and white high contrast film using a special green/yellow filter combination. One clone of each combination was treated with a colchicine (0.05%) and 2.0% di-methyl-sulfoxide (DMSO) solution using the aerated root treatment protocol of Mujeeb-Kazi *et al.*, (1987) in order to produce amphiploids.

**Spike categorization and backcross-I seed production:** Five fully emerged spikes from each  $F_1$  hybrid and its corresponding durum parent were characterized for spike morphology. Between 5 to 10 self-sterile  $F_1$  spikes were further pollinated with *T. turgidum* cultivars to produce the equivalent of a backcross-I (BC<sub>1</sub>) progeny.

### **Results and Discussion**

**Hybrid production and spike morphology:** Crossing between durum cultivars and perennial Triticeae species (Table 1) leading to seed set and putative hybrid embryo excision ranged from 30.0 to 45.5%. All of these high frequency hybrid recovery combinations possessed well-defined embryos, copious endosperm, and produced rapidly growing vigorous regenerants. Percentages of these hybrids were 37.2 (ssp. *acutum*), 44.0 (ssp. *glaucum*), 45.5 (ssp. *intermedium*), 43.8 (ssp. *pulcherrinum*), 30.0 (ssp. *trichophorum*) and 41.9 (ssp. *varnense*). Several cultivars produced hybrids with *Th. intermedium*, its five sub-species. The durum cultivars combined with the above species (Table 1) were high yielding durums, hence desirable alien introgressions may be anticipated to yield practical out-puts in a relatively short time frame.

<b>*</b> *	Alien	Florets	Seeds	No. of embryos	Plants
<i>T. turgidum</i> cultivar	species	pollinated	set	excised	obtained
Cocorit 71	spp. acutum	48	33	29	17
Yavaros	spp. acutum	52	38	33	18
Arlin	spp. acutum	48	31	21	13
Cappelli	spp. acutum	51	43	39	26
Chen	spp. glaucum	100	74	55	44
Cocorit 71	spp. intermedium	128	98	68	56
Yavaros 79	spp. intermedium	88	70	59	43
Cappelli	spp. intermedium	20	14	10	8
Cocorit 71	spp. pulcherrimum	54	32	21	15
Mexicali 75	spp. pulcherrimum	96	73	62	40
Yavaros 79	spp. pulcherrimum	74	62	55	43
Mexicali 75	spp. trichophorum	24	12	7	3
Croc	spp. trichophorum	22	14	9	5
Dvergand	spp. trichophorum	46	33	26	18
Laru	spp. trichophorum	24	18	11	7
Cappelli	spp. trichophorum	24	16	12	9
Altar 84	Spp. varnense	46	40	32	24
Cappelli	spp. varnense	54	38	28	19
Laru	spp. varnense	30	22	17	13
Mexicali 75	spp. varnense	30	20	15	11

 Table 1. Hybridization details of successful combinations of *Triticum turgidum* L., cultivars with *Thinopyrum intermedium* and its sub-species under greenhouse conditions.

Crossing success could be further enhanced by incorporating procedural manipulations involving early (bud-) and multiple pollinations, pre- and post-pollination hormonal applications (gibberellic acid and 2,4-di-chlorophenoxy-acetic acid), coupled with culture-media formulations (e.g. Taira & Larter, 1978; media for small embryos) where all may as a package significantly modify hybridization outputs.

In bread wheat/alien species hybridization crossability frequencies have received significant mention. Some cultivars like Chinese Spring have been favored since they hybridize readily (Falk & Kasha, 1981) due to the presence of crossability genes kr1, kr2, kr3 on homoeologous group 5 (Fedak & Jui, 1982; Riley & Chapman 1967) and subsequently the contribution of the kr4 loci towards improved crossability (Luo *et al.*, 1992; Yen *et al.*, 1988). Cultivars Asakazekomugi and Fukuhokomugi have better agronomic type than Chinese Spring and are also highly crossable (Jauhar, 1995a, b). In durums, cultivar variability for crossability with rye is present (Immonen *et al.*, 1993). Current observations support this durum crossability diversity but for the durum/alien species combinations attempted by us all have been successful. In BW however, some crosses never set seed leading us to infer that crossability control in BW is more stringent than that of durum wheat cultivars.

**Phenotype of F**<sub>1</sub> **hybrids:** All hybrids were perennial, possessed a vigorous growth habit and tillered profusely. Each hybrid was self-sterile, but female fertile and set various frequencies of backcross I seed (BC<sub>1</sub>) when pollinated by durum cultivars.

A co-dominant wheat/alien species phenotype was a common characteristic (Table 2, Fig. 1) of all hybrids, substantiating alien genetic expressivity in the durum background. The modified  $F_1$  phenotype is advantageous to observe, since this permits the possible selection of beneficial alien characteristics in durum wheats requiring such improvements. The phenotypic parameters generally affected included spike length, spike size, spike width, reduced awn length to even awn absence, lax heads with greater internodal distance and an occasional presence of pubescence (Fig. 1). An intermediate phenotype has been a common observation for several intergeneric hybrids within the Triticeae and has been considered a valid morphological indicator of alien genetic expressivity in a wheat background (Mujeeb-Kazi *et al.*, 1995).

**Somatic and meiotic cytology:** Two satellited chromosomes, (1B and 6B) present as pairs in euploid *T. turgidum* were identified for each  $F_1$  hybrid. Satellites of the alien species were not observed in some hybrid combinations as a consequence of amphiplasty. A mere chromosome count and satellite detail is often enough to initially validate hybridity (Figs. 2a-f). However, when alien species with similar ploidy levels are involved in crosses, additional diagnostics like chromosome banding or *in situ* hybridization become necessary. Unequivocal proof of hybridity however, comes from meiotic analyses that further provide an accurate index of the introgression methodology to be adopted for affecting alien genetic transfers to durum wheats. The meiotic data of Table 3 elucidates the constraints of homoeologous transfers *via* recombination. The *Th. intermedium* species and its sub-species possess two closely related genomes with a distinctly different third genome. Hence, the meiotic associations require careful evaluation before any conclusion is reached to suggest a wheat alien chromosomal union to facilitate a genetic transfer interpretation leading to the crops improvement (Figs. 3a-f).

1220

Cross combination Durum (4X) / Alien species	1	2	3	4	5	9	7	8	6	10	11	12	13
Cappelli / Th. acutum	24.9	0.7	22.5	1.7	1.8	0.5	22.5	5.0	1.1	0.1	1.1	0.1	0.3
Cocorit 71 / Th. acutum	16.5	0.6	21.0	1.0	1.5	0.5	21.0	4.5	0.8	0.1	1.0	0.5	0.3
Yavaros 79 / Th. acutum	17.9	0.5	16.0	1.2	1.8	0.5	16.0	3.5	1.3	0.0	1.3	0.3	0.2
Chen / Th. glaucum	18.4	0.6	22.0	0.9	1.9	0.6	22.0	5.5	1.1	0.1	1.3	0.2	0.3
Cappelli / Th. intermedium	11.9	0.5	13.5	1.1	1.7	0.6	13.5	4.8	1.1	0.1	1.3	0.2	0.3
Cocorit 71 / Th. intermedium	12.0	0.5	15.0	0.8	1.9	0.4	15.0	4.5	0.8	0.1	1.0	0.7	0.2
Yavaros 79 / Th. intermedium	15.6	0.7	16.5	0.9	2.2	10.6	16.5	4.0	0.8	0.0	1.3	1.0	0.3
Cocorit 71 / Th. Pulcherrimum	13.6	0.6	13.0	1.4	1.7	0.5	13.0	5.8	1.0	0.1	1.1	0.1	0.3
Mexicali 75 / <i>Th. pulcherrimum</i>	12.5	0.5	12.5	1.1	2.0	0.7	12.5	7.3	1.0	0.3	1.2	0.1	0.3
Dvergand / Th. trichophorum	17.8	0.7	13.0	1.5	2.6	0.7	13.0	8.5	1.3	0.0	1.3	0.1	0.4
Laru / Th. trichophorum	17.1	0.7	14.5	1.4	2.1	0.7	14.5	7.5	1.2	0.0	1.3	0.5	0.4
Mexicali/ Th. trichophorum	16.0	0.5	13.5	1.5	1.6	0.4	13.5	5.3	1.0	0.0	1.2	0.0	0.3
Altar 84 / <i>Th. varnense</i>	14.6	0.7	22.5	0.8	1.9	0.4	22.5	4.8	0.9	0.1	1.1	0.1	0.3
Cappelli / Th. varnense	15.3	0.6	20.5	0.9	1.8	0.5	20.5	4.5	0.9	0.0	1.1	0.1	3.2
Mexicali 75 / <i>varnense</i>	11.9	0.5	13.5	1.1	1.7	0.6	13.5	4.8	1.1	0.1	1.3	0.2	0.3
Laru / <i>Th. varnense</i>	20.5	0.7	18.0	1.3	2.5	0.7	18.0	7.5	1.0	0.1	1.3	0.7	3.7

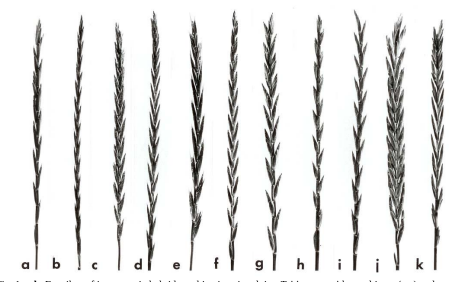


Fig. 1a -  $\mathbf{k}$ . F<sub>1</sub> spikes of intergeneric hybrid combinations involving *Triticum turgidum* cultivars (cv.) and perennial Triticeae sub-species (ssp) of the genus *Thinopyrum (Th) intermedium* showing in  $\mathbf{a}$ . cv. Arlin / ssp *acutum*,  $\mathbf{b}$ . cv. Cocorit 71 / ssp *acutum*,  $\mathbf{c}$ . cv. Croc / ssp *glaucum*,  $\mathbf{d}$ . cv. Cocorit 71 / *Th. intermedium*,  $\mathbf{e}$ . cv. Yavaros / *Th. intermedium*,  $\mathbf{f}$ . cv. Cappelli / *Th. intermedium*,  $\mathbf{g}$ . cv. Mexicali 75 / ssp *pulcherrimum*,  $\mathbf{h}$ . cv. Dvergand / ssp *trichophorum*,  $\mathbf{i}$ . cv. Laru / ssp *trichophorum*,  $\mathbf{j}$ . cv. Altar 84 / ssp *varnense*,  $\mathbf{k}$ . cv. Cappelli / ssp. *Varnense*.

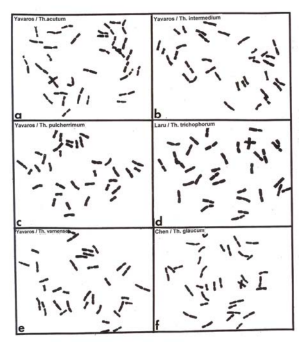


Fig. 2a to **f.** Somatic chromosomes of the hybrid cross combinations between durum and perennial Triticeae sub-species (ssp) of the genus *Thinopyrum (Th) intermedium* showing in **a**.Yavaros / *Thinopyrum acutum* (2n=5x=35), **b.** Yavaros / *Th. intermedium* (2n=5x=35), **c.** Yavaros / *Th. pulcherrimum* (2n=5x=35), **d.** Laru / *Th. trichophorum* (2n=5x=35), **e.** Yavaros / *Th. varnense* (2n=5x=35), and **f.** Chen / *Th. glaucum* (2n=5x=35).

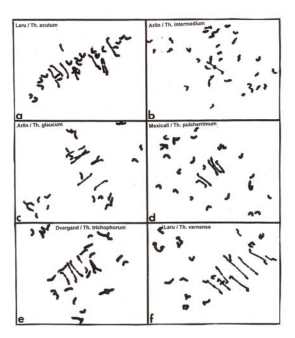


Fig. 3a to **f**. Mean meiotic metaphase I chromosomal relationships in some hybrids amongst durum and perennial Triticeae sub-species (ssp) of the genus *Thinopyrum (Th) intermedium* showing in **a**. Laru / *Th. acutum* (19<sub>1</sub> + 5<sub>11</sub> + 2<sub>111</sub>), **b**. Arlin / *Th. intermedium* (31<sub>1</sub> + 2<sub>11</sub> rods), **c**. Arlin / *Th. glaucum* (22<sub>1</sub> + 5<sub>11</sub> + 1<sub>111</sub>), **d**. Arlin / *Th. pulcherrimum* (25<sub>1</sub> + 5<sub>11</sub>), **e**. Dvergand / *Th. trichophorum* (24<sub>1</sub> + 4<sub>11</sub> + 1<sub>111</sub>), **f**. Laru / *Th. varnense* (19<sub>1</sub> + 8<sub>11</sub>)

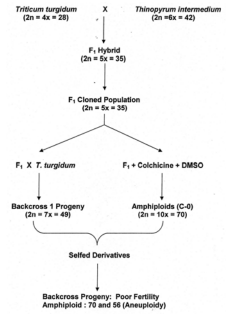


Fig. 4. Schematic elucidating the advance of F1 hybrids to serve maintenance as a clonal population, backcross 1 and amphiploid progeny.

	Chromosome	Mean meiotic chromosomal associations					
Cross combination	number	Ι	II	II	Total	III	IV
Cocorit 71 / Th. acutum	2n=5x=35	20.3 ( <b>15-26</b> )	0.6 ( <b>0-2</b> )	4.9 ( <b>2-8</b> )	5.5	1.1 ( <b>0-2</b> )	0.1 ( <b>0-1</b> )
Yavaros / Th. acutum	2n=5x=35	25.0 ( <b>22-27</b> )	0.4 ( <b>0-2</b> )	4.0 ( <b>2-5</b> )	4.4	0.4 ( <b>0-1</b> )	0 -
Chen / Th. glaucum	2n=5x=35	25.0 ( <b>19-31</b> )	0.1 ( <b>0-1</b> )	4.0 ( <b>2-7</b> )	4.1	0.6 ( <b>0-2</b> )	0 -
Altar 84 / Th. intermedium	2n=5x=35	29.9 ( <b>25-35</b> )	0	2.4 ( <b>0-5</b> )	2.4	0.1 ( <b>0-1</b> )	0 -
Yavaros / Th. intermedium	2n=5x=35	29.3 ( <b>25-33</b> )	0 -	2.5 ( <b>0-5</b> )	2.5	0.1 ( <b>0-1</b> )	0.1
Cocorit 71 / Th. intermedium	2n=5x=35	32.0 ( <b>31-33</b> )	0	1.2 ( <b>0-2</b> )	1.2	0.2 (0-1)	0
Memo / Mexicali // Th. glaucum	2n=5x=35	23.1 ( <b>13-30</b> )	0.3 ( <b>0-1</b> )	5.1 ( <b>1-8</b> )	5.4	0.1 ( <b>0-1</b> )	0.2 ( <b>0-2</b> )
Cocorit / Th. pulcherrimum	2n=5x=35	20.9 ( <b>16-27</b> )	1.4 ( <b>0-3</b> )	4.7 ( <b>2-7</b> )	6.1	0.5 ( <b>0-1</b> )	0.1 ( <b>0-1</b> )
Mexical i/ Th. pulcherrimum	2n=5x=35	24.7 ( <b>19-29</b> )	0.5 ( <b>0-1</b> )	3.9 ( <b>2-6</b> )	4.4	0.5 ( <b>0-2</b> )	0
Dvergand / Th. trichophorum	2n=5x=35	26.1 ( <b>23-33</b> )	0.6 ( <b>0-2</b> )	3.2 ( <b>1-6</b> )	3.8	0.4 ( <b>0-1</b> )	0 -
Laru / Th. trichophorum	2n=5x=35	27.9 ( <b>23-33</b> )	0.3 ( <b>0-2</b> )	3.1 ( <b>1-6</b> )	3.4	0.1 ( <b>0-1</b> )	0.1 ( <b>0-1</b> )
Mexicali / Th. varnense	2n=5x=35	30.6 ( <b>27-33</b> )	0	2.2 (1-4)	2.2	0	0
Altar / Th. varnense	2n=5x=35	31.4 ( <b>29-33</b> )	0	1.8 ( <b>1-3</b> )	1.8	0	0

 Table 3. Mean and range of meiotic associations at metaphase I in intergeneric hybrids of some

 *Triticum turgidum* L. (2n=4x=28, AABB) cultivars with *Thinopyrum intermedium* and its sub-species.

 Table 4. Details of amphiploids derived from various F1 hybrids between

 Triticum turgidum X Thinopyrum intermedium sub-species and its sub-species.

Cross combination	Expected chromosome	Chromosomal range
	number in C–0	in C-n derivatives
T, turgidum* / Th. acutum	2n=10x=70	55 to 70
T. turgidum / Th. glaucum	2n=10x=70	64 to 71
T. turgidum / Th. intermedium	2n=10x=70	55 to 72
T. turgidum / Th. pulcherrimum	2n=10x=70	58 to 71
T. turgidum / Th. trichophorum	2n=10x=70	65 to 71
T. turgidum / Th. varnense	2n=10x=70	58 to 71

\*: T. turgidum combinations involve several cultivars (Table 1) with each Th. intermedium sub-species

**Production and maintenance of backcross 1 derivatives and amphiploids.** The schematic of Fig. 4 elucidates how from the  $F_1$  hybrid production between durum wheat and *Th. intermedium* its maintenance is done with derivatives serving as a living herbarium. From these  $F_1$ s that are male-sterile but female-fertile, BC<sub>1</sub> derivatives are obtained upon crossing with durum cultivars and upon colchicine treatment the  $F_1$ s yield amphiploids. Table 4 provides the details of the BC and amphiploid derivative stocks that are available with their brief mitotic characteristics. The expected mitotic counts of all BC<sub>1</sub>'s expected would be 2n=5x=35 that upon fertilization by the male n=2x=14, AB gamete generate the BC derivative with 49 chromosomes (Fig. 5a). Such derivatives were readily obtained, but contrary to the results seen with similar bread wheat based BC<sub>1</sub>'s, these durum based BC<sub>1</sub>s showed little to no self-fertility. Hence, their best usage would be to immediately advance selected BC<sub>1</sub>'s by further backcrossing leading to alien chromosomal addition line development, agricultural trait categorization and then enforcing alien gene transfers *via* cytogenetic manipulation procedures.

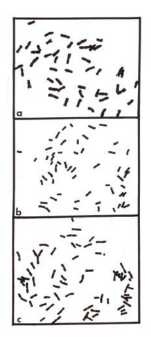


Fig. 5a to **c**. Somatic composition of F1 derived progenies showing in **a**. backcross 1 derivative with 2n=7x=49 chromosomes, **b**. a selfed C-0 derivative with 56 chromosomes indicative of possible genomic elimination, and **c**. a selfed C-0 with the expected near 70 chromosomes (68 with 2 telocentrics)

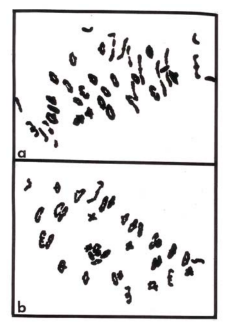


Fig. 6a and **b**. Meiotic association in C-0 selfed derivatives showing in **a**. 4 univalents + 9 rod bivalents + 22 ring bivalents + 1 chain quadrivalent (70 chromosomes), and in **b**. 70 chromosome meiocyte associated as 2 univalents + 3 rod bivalents+32ring bivalents.

All the  $F_1$  hybrids upon colchicine treatment set seed (C-0) that were validated by somatic and meiotic cytology (Figs. 5 and 6). The expected chromosome numbers in the various amphiploids at C-0 generation would be 2n=10x=70 that meiotically associate upto a maximum of 35 bivalents ensuring good fertility and generating healthy seed upon selfing of the various C-0s. This was observed across all amphiploid combinations but aneuploidy was rampant (Fig. 6a, b). More drastic variations from the expected 70 chromosome derivatives were around selfed progeny that had presumably undergone spontaneous genomic loss yielding derivatives with chromosome numbers around 56 (Fig. 5b). These two forms of selfed individuals are classified as complete and partial amphiploids, possess a healthy seed type and are useful genetic stocks for agricultural applications to address biotic and abiotic stress crop production constraints.

#### References

- Falk, D.E. and K.J. Kasha. 1981. Comparison of the crossability of rye (*Secale cereale*) and *Hordeum bulbosum* onto wheat (*Triticum aestivum*). *Canadian J. Genetics* and *Cytology*, 23: 81-88.
- Fedak, G. and P.Y. Jui. 1982. Chromosomes of Chinese Spring wheat carrying genes for crossability with Betzes barley. *Canadian J. Genetics* and *Cytology*, 24: 227-233.
- Immonen, A.S.T., G. Varughese, W.H. Pfeiffer and A. Mujeeb-Kazi. 1993. Crossability of tetraploid and hexaploid wheats with ryes for primary triticale production. *Euphytica*, 65: 203-210.
- Jahan, Q., N. Ter-Kuile, N. Hashmi, M. Aslam, A.A. Vahidy and A. Mujeeb-Kazi. 1990. The status of the 1B/1R translocation chromosome in some released wheat varieties and the 1989 candidate varieties of Pakistan. *Pakistan J. Botany*, 22: 1-10.
- Jauhar, P.P. 1995a. Meiosis and fertility of F1 hybrids between hexaploid bread wheat and decaploid tall wheatgrass (*Thinopyrum ponticum*). *Theoretical* and *Applied Genetics*, 90: 865-871.
- Jauhar, P.P. 1995b. Morphological and cytological characteristics of some wheat X barley hybrids. *Theoretical and Applied Genetics*, 90: 872-877.
- Jiang, J., B. Friebe and B.S. Gill. 1994. Recent advances in alien gene transfer in wheat. *Euphytica*, 73: 199-212.
- Luo, M.C., C. Yen and J.L. Yang. 1992. Crossability percentages of bread wheat landraces from Shaanxi and Henan provinces, China with rye. *Euphytica*, 67: 1-8.
- Mujeeb-Kazi, A. 1998. Evolutionary relationships and gene transfer in the Triticeae. *Triticeae* III, (Ed.): A.A. Jaradat. ICARDA, Syria, pp. 59-65.
- Mujeeb-Kazi, A. and G.P. Hettel. 1995. Utilizing wild grass biodiversity in wheat improvement: 15 years of Wide Cross Research at CIMMYT. *CIMMYT Research Report* No. 2, Mexico, D.F., CIMMYT. pp 1-140.
- Mujeeb-Kazi, A. and J.L. Miranda. 1985. Enhanced resolution of somatic chromosome constrictions as an aid to identifying intergeneric hybrids among some Triticeae. *Cytologia*, 50: 701-709.
- Mujeeb-Kazi, A. and M. Bernard. 1985. Intergeneric hybridization to induce alien genetic transfers into *Triticum aestivum*. *Pakistan J. Botany*, 17: 271-289.
- Mujeeb-Kazi, A., S. Roldan, D.Y. Suh, L.A. Sitch and S. Farooq. 1987 Production and cytogenetic analysis of hybrids between *Triticum aestivum* and some caespitose *Agropyron* species. *Genome*, 29: 537-553.
- Mujeeb-Kazi, A., S. Roldan, D.Y. Suh, N. Ter-Kuile and S. Farooq. 1989. Production and cytogenetics of *Triticum aestivum* L., hybrids with some rhizomatous *Agropyron* species. *Theoretical and Applied Genetics*, 77: 162-168.
- Riley, R. and V. Chapman. 1967. The inheritance in wheat of crossability with rye. *Genetics Research*, 9: 259-267.

Sharma, H.C. 1995. How wide can a wide cross be?. Euphytica, 82: 43-64.

Sharma, H.C. and B.S. Gill. 1983. Current status of wide hybridization in wheat. *Euphytica*, 32: 17-31.

#### 1226

- Taira, T. and E.N. Larter.1978 Factors influencing development of wheat-rye hybrid embryos in vitro. *Crop Science*, 18: 348-350.
- Wang, R.R-C. 1989. Intergeneric hybrids involving perennial Triticeae. *Genet. (Life Sci. Adv.)* 8: 57-64.
- Yen, C., M.C. Luo and J.L. Yang. 1988. The origin of the Tibetan weedrace of hexaploid wheat, Chinese Spring, Chengdu-guang-tou and other landraces of the white wheat complex from China. In: *Proc.* 7<sup>th</sup> *International Wheat Genetics Symposium, Cambridge, England*, 13-19 July 1988, (Eds.): T.E. Miller and R.M.D. Koebner, pp. 175-179.

(Received for publication 28 June 2007)