## Can Tritipyrum, a New Salt Tolerant Potential Amphiploid, Be a Successful Cereal Like Triticale?

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#### ABSTRACT

Soil affected by salt (NaCl) is a major problem worldwide and in areas with potential agriculture; lands in many countries are not enough to support crop production. The development of salt tolerant cultivars would be enhanced by better understanding of the genetic control of tolerance to salt stress. A new cereal, tritipyrum, a range of amphiploids between Triticum spp. and Thinopyrum spp. offers such a new chance. Those with the 6x construction (2n=6x=42, AABBE<sup>b</sup>E<sup>b</sup>) derived from *Triticum durum* (2n=4x=28, AABB) and Thinopyrum bessarabicum (2n=2x=14, E<sup>b</sup>E<sup>b</sup>) are of the potential to become a new high salt tolerant cereal crop. Tritipyrum is prone to problems similar to those exhibited by early triticales, e.g. chromosome instability and low fertility, which in that crop were eventually overcome by breeding. Other problems could be overcome through substitution of  $\mathbf{E}^{\mathbf{b}}$  genome chromosomes by D genome ones, and the feasibility of this has been assessed in the progenies of (6x tritipyrum) x (6x wheat) hybrids with the aid of fluorescent in situ hybridization (FISH). The cytological, morphological and agronomic studies of existing tritipyrum lines, including the effect of vernalization, were carried out, too. A novel multiple-pistil/seed characteristic of one original tritipyrum line has also been investigated and its genetic basis established. The results have shown that, first creation of substituted lines is feasible, and thus it could be a route for the elimination of undesirable traits. Second, improvement should be possible via selection for chromosomally stable lines, with increased fertility and yield. Third, it may also be possible to exploit the perennial habit and multi-tillering traits in a dual-purpose forage/grain crop. Fourth, the multiple-pistil/seed trait may be controlled by two recessive genes. Fifth, there is a high probability of having established the seven possible monosomic additions of Th. bessarabicum to T.durum for the first time.

Keywords: Amphiploid, Tritipyrum, Fluorescent *in situ* hybridization (FISH), Multiplepistil/seed, *Triticum durum, Thinopyrum bessarabicum*, Salt tolerance.

#### INTRODUCTION

Based on different reports, it has been estimated that in the range of 3.4-9 million km<sup>2</sup> of lands are salt affected (Flowers & Yeo, 1988); more than one third of which can be found in Asian countries. In many of these countries, soil salinity and alkalinity have spread to such a degree that they have brought about severe problems for the national economies (Szabolcs, 1989). In Asia, the greatest extension of salt affected soils occurs in the former USSR, China, India, Pakistan and Iran (Dewan & Famouri, 1942). The genetic studies of salt tolerance in various species including sorghum, soy-

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bean and rice have shown that salt tolerance is controlled polygenetically and is quantitatively inherited (Azhar & McNeilly, 1988). Variation for salt tolerance in wheat has been reported (Quershi et al. 1980; Kingsbury & Epstein, 1984; Sayed, 1985) and there is some potential for improving the salt tolerance of wheat by conventional breeding. As wheat is the most important source of human nutrition, the study and ultimate improvement of salt tolerance in this crop serves the high priority. The development of crop varieties that combine high yield potential with high salt tolerance will not be an easy task. Triticeae contains several halophytic and perennial species, which are naturally adapted to saline environments. A number of these species, particularly members of the genus Thinopyrum, have been hybridized with wheat (Gorham et al. 1985, 1986) and some of the resulting progenies have been tested for salt tolerance (Storey et al., 1985) but successful incorporation of genes imparting salt tolerance to T. aestivum background has not been established yet, either due to the failure in obtaining fertile  $F_1$  hybrids or due to difficulty in getting direct gene introgression by homoeologous recombination (Tomar et al., 1995). One of the most notable of these species is Thinopyrum bessarabicum, a littoral diploid grass native to the Crimea, Ukraine. It has been identified as a useful source of salt tolerance genes for transfer to wheat for the following reasons; 1-It is a diploid species and highly tolerant to prolonged exposure to 250 mM NaCl (Gorham et al., 1985), 2-The genes conferring tolerance to salt are expressed in a wheat genetic background (Gorham et al., 1986; Forster et al., 1987), 3-Its chromosomes pair and presumably can combine with wheat homoeologous in the absence of chromosome 5B (Forster & Miller, 1985; King et al., 1993).

Amphiploids between wheat and its relatives have been of interest for many years as potential sources for wheat improvement and as potential crops in their own right, but only triticale (the amphiploid between wheat and rye) has so far become a new cereal crop

(Gupta & Priyadarshan, 1982; Gregory, 1987). Tritordeum, the amphiploid between Triticum durum and Hordeum chilense is considered to possess potential as a novel crop species (Martin et al., 1996). A similar amphiploid between T. durum and Th. bessarabicum might also be of potential, especially as Th. bessarabicum is known to exhibit tolerance to high levels of salt (Gorham et al., 1985). The assessment of today's tritipyrum situation in comparison with triticale and the possible problems facing breeders in future is not an easy undertaking. The quantum of work done in triticale is evident from several reviews written in recent years on different aspects of triticale and shows the importance of this first man-made cereal. There is no reason that the same success should not be achieved in the case of tritipyrum. Tritipyrum exhibits salt tolerance (King et al., 1997) and data on the salt stress tolerance in an incomplete set of wheat-Th bessarabicum addition lines (Forster et al., 1988) and a set of wheat-H. chilense addition lines (Forster et al., 1990) revealed analogous effects of group 2 and 5 chromosomes. It is interesting to note that homoeologous group 5 chromosomes in the Triticeae appear to be responsible for tolerance to several abiotic stresses (Forster, 1990). Chromosome  $5E^{b}$  was found to carry a dominant gene for salt tolerance (Forster et al., 1988).

The gene(s) conferring salt tolerance located on chromosome 5E<sup>b</sup> from Th. bessarabicum has (have) greater effect when substituted for homoeologous wheat group 5 chromosomes than when present as an addition chromosome, as in the Chinese Spring  $5E^{b}$  addition line. This characteristic of  $5E^{b}$ chromosome demonstrates the potential for production of new salt-tolerant wheat varieties (King et al., 1996). The benefits of the R genome in triticale (AABBRR) over D genome in wheat (AABBDD), are that the R genome is superior to the D genome with respect to yield potential, disease resistance, tolerance to minor element deficiencies and toxicities and to phosphorus uptake efficiency (Varughese, 1996). Similarly in tritipyrum the  $E^{b}$  genome is superior to the D genome in terms of salt tolerance potential. Further breeding and selection may modify some of the negative traits. In the light of the studies of the substitution of D genome chromosomes into triticale it seems likely that similar substitution into tritipyrum could prove valuable. There are a number of detrimental characters in tritipyrum and beneficial ones in wheat that are obvious candidates for removal or insertion, respectively. The brittle rachis, which is almost certainly carried by 3E<sup>b</sup>, could be removed by substitution of 3E<sup>b</sup> by 3D. Similarly the blue aleurone colour carried by 4E<sup>b</sup> could be replaced by substitution with 4D; at the same time the Rht2 semi-dwarfing gene could be inserted with the 4D. Bread making quality could be introduced by replacing 1E<sup>b</sup> with 1D. On the other hand, substitution of 5E<sup>b</sup> by 5D would be detrimental, as this would delete the salt tolerance. This could be a case for partial substitution in the form of Robertsonian translocation where only one arm of a chromosome is replaced (Hassani, 1998).

Alien chromosome addition lines, in which single pairs of homologous chromosomes from a related species are added to wheat, were first produced by O'Mara (1948), who succeeded in adding rye (Secale cereal) chromosomes to wheat by backcrossing a wheat-rye amphiploid (triticale) to wheat. Since that time chromosome addition lines have been obtained from a range of alien species (Sears, 1956; Kimber, 1967; Islam et al., 1978; Miller et al., 1982). The multiple pistil/seed characteristic occurs in rice and has been variously termed as multiple pistil (Parthasarathi, 1935), polygynous rice (Shen, 1933), or multiple seeded rice (Wickramasekera, 1939). This malformation proved to be under the control of three recessive genes in the progenies of a cross of two rice cultivars (Butany & Bhattacharyya, 1962) and monogenic recessive in crosses investigated by Ghose & Butany, 1952. This paper reports the recent preliminary investigations of tritipyrum potential, including the possibility of improvement by the substitution of D genome chromosomes for E<sup>b</sup> ones,

characterization of the alien E<sup>b</sup> chromosomes in the possible (1-7, DD)E<sup>b</sup>E<sup>b</sup> substitution lines by FISH, morphological study of the existing tritipyrum lines in glass house, agronomic behavior in the first ever field trial and cytological studies for determination of the source of the chromosomes involved in aneuploidy. The genetic heritability of the novel multiple-pistil/seed characteristic of one original Tritipyrum line (Creso/Th. bessarabicum) and primary studies of the chromosomal location of the genes are discussed. To achieve the latter, the first ever attempt of conceiving and constructing a novel set of alien chromosome addition lines of Th. bessarabicum chromosomes to T. durum cv. Creso is also reported.

#### **MATERIALS AND METHODS**

1-Eight hexaploid tritipyrum (2n=6x=42, AABBE<sup>b</sup>E<sup>b</sup>) and two octaploid tritipyrum AABBDDE<sup>b</sup>E<sup>b</sup> (2n=8x=56,lines and  $AABBE_1E_1E_2E_2$ ). All Tritipyrum lines were produced in the Cereal Research Department at John Innes Center, UK (King et al., 1997, Fig. 2b). 2-Triticum aestivum cvs. Chinese Spring (CS), Wembley, Axona and two breeding lines  $S_{2-4}$  and  $S_{6-4}$  (Fig. 2c). 3-T. durum cvs. Aziziah; Creso; Karim Langdon (4B) 4D, Macoun, Neodure, Stewart and Th. bessarabicum (Fig. 2a). 4-Genomic DNA's of tetraploid wheat cv. Creso (AABB), T. aestivum cv.CS and Th. bessarabicum.

#### Cytology and Cytogenetic Survey

In chromosome engineering studies both conventional Feulgen stained (Table 1) and FISH at meiotic metaphase I of Tritipyrum lines (Table 2) were utilized. The *in situ* hybridization followed the technique of Reader *et al.* (1994) with the addition of preblocking hybridization stage in which wheat DNA was hybridized to the template DNA for one hour prior to the normal hybridization.



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**Figure 2.** a) From left to right: *T. durum* cvs. Aziziah; Creso; Karim; Langdon (4B)4D, Macoun, Neodure, Stewart and *Th. Bessarabicum.* b) From left to right 6x Tritipyrum Az/b, Cr/b, La/b, La (4B)4D, Ka/b, Ma/b, Ne/b, St/b and 8x Tritipyrum, CS/b, St/*Th. junceiforme.*c) From right to left *T. aestivum* cvs. Chinese Spring, Wembley, Axona and two breeding lines  $S_{2-4}$  and  $S_{6-4}$  d) The Tritipyrum section of the field trail, note the lateness of the lines as compared with the surrounding wheat crop. e) Brittle rachis of Tritipyrum. f) Perennial habit of Tritipyrum.

Tritipyrum lines	PMC	Univ.	E Rods	ivalents Rings	Total	Thy.	Quadriv.	Arm assoc.	Pairing frequency
Az/b	30	1.20	6.97	13.43	20.40	-	-	33.83	0.81
	25	(0-4) 1.36 (0-6)	(0-17) 5.96 (0-14)	(4-20) 14.36 (11-20)	(17-21) 20.32 (18-21)	-	-	34.68	0.83
Cr/b	30	2.20	4.47	15.43	19.90	-	-	33.33	0.79
	37	(0-16) 4.00	(1-11) 5.78	(7-17) 13.22	(13-21) 19.00	-	-	32.22	0.77
La/b	27	(0-12)	(1-10)	(13-21)	(14-21)	0.04	-	33.60	0.80
	24	(0-14)	(0-12)	(7-21)	(11-21) 20.48	(0-1)	_	36.44	0.87
La (4B)4D/b	30	(0-4) 2.17	(2-9) 4 69	(11-19)	(17-21) 19-71	- 0.06	- 0.06	33.73	0.80
	30	(0-8) 3.23	(0-12) 6.00	(8-18) 13.37	(17-21) 19.37	(0-1)	(0-1)	32.74	0.78
Ka/b	30	(0-8) 2.60	(0-10) 4.92	(9-17) 15.78	(16-21) 20.70	-	-	36.48	0.87
Ma/b	23	(0-8) 3.82	(3-11) 6.26	(7-17) 12.83	(15-21) 19.09	-	-	31.92	0.76
Ne/b	30	(0-14) 2.83	(1-14) 5.14	(4-9) 14.36	(14-21) 19.50	0.04	-	33.86	0.81
	27	(0-6) 3.89	(1-8) 5.89	(7-20) 13.16	(14-21) 19.05	(0-1) -	-	32.21	0.77
St/b	30	(0-9)	(2-11)	(10-18)	(15-21)	0.06	-	33.32	0.79
	20	3.30 (0-7)	5.20 (2-10)	14.06 (11-17)	19.26 (18-21)	(0-1)		22.22	0.70
CS/h	30	3.43 (0-8)	5.26 (2-10)	(10-18)	19.29 (16-21)	-	-	33.32	0.79
0.5/0	23	5.16	8.38	16.68	25.06	0.03	0.16	41.74	0.75
	30	(2-7) 2.60 (0-6)	(4-12) 8.15 (4-13)	(12-21) 18.37 (12-20)	(22-28) 26.52 (22-28)	(0-1) 0.03 (0-1)	(0-2) 0.07 (0-2)	44.89	0.80
St/Th.j	19	4.89	7.12	18.14	2526	(0 - 1) (0 - 1)	(0.2) 0.11 (0-1)	43.4	0.78
		(4-8)	(3-14)	(10-25)	(19-28)	(3-1)	()		
Analysis of variance Ms between 6x lines Ms within 6x lines Ms between all lines Ms within all lines	7df 7df 9df 9df	1.30 <sup>ns</sup> 0.64 1.73 <sup>ns</sup> 1.02	$0.52^{ns}$ 0.55 2.06* 0.47	0.87 <sup>ns</sup> 1.17 4.11 1.21	0.36 <sup>ns</sup> 0.15 10.15 <sup>**</sup> 0.28	  	  	2.61 <sup>ns</sup> 1.15 27.61 1.69	$0.0016^{ns}$ 0.0006 $0.0014^{ns}$ 0.0007

**Table 1.** Mean chromosome pairing of individual plants of Tritipyrum genotypes by the Feulgen staining technique (ranges in parenthesis).

ns,\*,\*\* Non.significant, significant at P=0.05 and P=0.01, respectively.

#### **Greenbouse Test**

Six 42-chromosome seedlings of each line were selected from among 20 seedlings and transplanted into plastic pots. Three were

	Univa	alents	Row	Rod bi	Row	
Genotype	E <sup>b</sup>	AB	total	E <sup>b</sup>	AB	total
Cr/b	1.12 (1.27)	0.81 (0.66)	1.93	1.35 (1.44)	2.38 (2.25)	3.73
La/b	1.53 (1.51)	0.77 (0.79)	2.30	2.20 (2.05)	3.12 (3.21)	5.32
La(4B) 4D/b	1.31 (1.23)	0.56 (0.64)	1.87	1.62 (1.63)	2.60 (2.58)	4.26
Ka/b	1.54 (1.56)	0.81 (0.80)	2.35	1.59 (1.65)	2.68 (2.57)	4.27
St/b	2.30 (2.24)	1.11 (1.17)	3.41	1.86 (1.86)	2.96 (2.90)	4.82
Column total	7.80	4.06		8.62	13.47	
Grand total			11.86			22.36
$\chi^2$			$0.058^{**}$		.006**	

Table 2. Univalents and rod bivalents of E<sup>b</sup> and AB genomes of Tritipyrum lines

Expected values in parenthesis

\*\*Significant P=0.01

grown under natural day length ranging from 12-17 hours during spring/summer 1995 in the greenhouse. At the same time, the other three seedlings were vernalized at  $7.5^{\circ}$ C with an eight-hour photoperiod for six weeks with normal light density. After vernalization, they were transferred to the greenhouse and grown under the same conditions as the unvernalized plants for the duration of the experiment, using a completely randomized design consisting of three replicates. Each set of three vernalized and unvernalized plants were scored for a number of traits; percent fertility, spikelet number and total grain of the first and second floret of each spikelet were scored and analysis of variance was carried out (Table 3).

#### **Field Trial**

A small-scale trial field was sown and divided into sections, one section containing the tritipyrum and the other containing the parents as a randomized block design with four replications. A range of morphological and agronomical traits were measured or counted on each plot (Table 4). For protein content and grain hardiness 5-10 g of mixed grain from each plot was milled by a Cyclotec 1093 Sample Mill, 3.5g of the flour of each sample being analyzed by an Oxford QN near infrared analyzer. Analysis of variance was applied.

#### The Production of Substituted lines

To produce substituted Tritipyrum, spikes of each 42-chromosome (6x) line of Tritipyrum were emasculated, then pollinated with pollen from the 6x wheat cultivars (Fig.lb). For selecting, the 42-chromosome  $F_2$  seedlings from the  $F_1$  hybrids, the mitotic chromosome preparation technique was made. To identify the number of  $E^{\bar{b}}$  chromosomes of F<sub>2</sub> and BC<sub>1</sub> plants via FISH, the meiotic chromosome preparations were made following the method of Hutchinson et al. (1980). Total genomic DNA in situ hybridization (GISH) was carried out as described by Schwarzacher et al. (1992) and Reader et al. (1994), with the addition of a preblocking step (Hassani, 1998). The chromosome constitution of possible substituted Tritipyrum plants of the  $F_2$  and the BC<sub>1</sub> progenies were assessed by scoring the number of E<sup>b</sup> chromosomes present in meiotic cells at metaphase I for individual fertile plants (Table 5).

#### Novel Multiple-pistil/seed

#### **Genetic Inheritance of Multiple Pistils**

Tritipyrum lines	Height (cm) Tiller no.		er no.	Matu	rity <sup>a</sup>	Bagged f	ertility %	Unbagged fertility %		
	Unver.	Ver.	Unver.	Ver.	Unver.	Ver.	Unver.	Ver.	Unver.	Ver.
Az/b	119.3±32	118.7±4.6	25±2.94	9.7±1.70	54.7±11.9	61.3±3.7	45.5±7.93	55.4±14.3	63.4±14.8	54.3±20.7
Cr/b	103±10.9	89.7±1.5	26.3±2.4	20.3±1.7	64.0±4.7	54.0±1.0	14.2±3.06	18.8±12.7	34.8±7.14	48.3±9.21
La/b	145±4.59	16.3±13.8	23.3±3.2	24±4.71	66.3±10.1	60.3±1.3	39.1±7.53	13.2±11.3	42.9±14.6	20.6±2.63
La (4B)4D/b	$149 \pm 2.56$	14.8±2.2	21.7±2.7	17.7±1.5	67.3±7.5	48.0±1.0	$17.8 \pm 16.5$	$39.2{\pm}6.34$	35.6±21.2	62.1±14.4
Ka/b	72±7.69	91.0±2.56	$11\pm3.97$	9.0±0.59	$65.0{\pm}2.1$	57.3±43	29.5±15.4	35.7±11.3	35.2±25.9	19.3±11.6
Ma/b	115±14.1	126±2.35	26.7±4.1	13.7±2.8	52.0±1.6	58.3±1.7	$30.0{\pm}15.6$	59.1±1.06	$36.2 \pm 25.0$	50.9±6.94
Ne/b	161±5.79	143±3.24	27.7±3.8	13±2.04	56.7±5.9	52.7±1.3	49.9±19.9	63.1±1.80	20.7±9.31	67.0±5.16
St/b	148±5.89	127.3±7.5	25±2.56	11.7±2.1	65.0±5.1	52.7±1.3	25.1±13.1	48.4±9.13	39.4±15.8	18.1±2.69
CS/b	131±2.70	116.3±6.01	26.7±2.8	14.7±2.7	54.3±2.7	42.7±2.7	37.0±13.3	20.4±18.9	$37.0{\pm}1.88$	26.5±12.0
St/Th.j.	130±23.9	94.0±7.71	28.3±3.4	21.3±2.6	54.0±3.3	53.7±4.7	16.4±8.76	13.3±11.4	29.1±20.3	13.5±13.5
Mean	127.5	117.4	24.17	15.50	59.93	54.93	30.44	35.59	42.18	38.29
Grand Mean	12	2.45	19.84		57.43		33.02		40.24	
LSD5% for Genotype	6.9		2.37		4.07		9.66		11.94	
LSD5% for (Treatment)	1.38		0.47		0.81		1.93		2.39	
LSD5% for (Genotype x Treatment)	13	3.81	4.73		8.14		19.32		23.89	

**Table 3.** Mean height, tiller number, maturity and fertility of tritipyrum under unvernalized and vernalized conditions

<sup>a</sup> Days from ear emergence to ripening.

The presence or absence of multiple pistils and seeds were screened in the florets of immature and mature spikes of Creso/Th. bessarabicum (Cr/b), all the durum wheat parents of the Tritipyrum lines and Th. bessarabicum. Segregation of the characters was studied in the progenies of (Cr/b) x *bessarabicum=Ka/b)*, (Karim/Th. Macoun/Th. bessarabicum (Ma/b) and Stewart/Th. bessarabicum (St/b) crosses (Table 6). All individual plants at flowering and maturity were carefully checked for multiple pistils or seeds. The Chi-Square analysis for the fixed-ratio dihybrid hypothesis, i.e. one multiple pistil: fifteen single pistils, was performed (Gomez & Gomez, 1984).

## Production of Monosomic *T. durum-Th. bessarabicum* addition lines

After emasculation, the selected 42chromosome seedlings of Cr/b were pollinated with the pollen of Creso (Fig. 1a). The resulting amphiploid F'<sub>1</sub>s (2n=5x=35; AAB-BE<sup>b</sup>) were grown. The mitotic and meiotic studies of (Cr/b)/Creso progenies and FISH were carried out. The 29-chromosome plants were selected from the self-pollinated progenies and checked for the presence of multiple pistils and seeds in the vegetative and mature stages of growth, respectively. The 29-chromosome plants in F<sub>3</sub> (Fig. 5g) and F<sub>4</sub> (Fig. 5d) progenies were classified into seven morphologically distinct groups.

#### **RESULTS AND DISCUSSION**

Tritipyrum is a new amphiploid combination between wheat (*Triticum* spp.) and *Thinopyrum* spp., and to date, there hav only been three scientific publications, two of which are the work of the author of this paper. In the first paper, all produced Triti-

Genotype	Survival	Height	Tiller	Spikelet	Fertility	Fmgrgmce	1000-Grain	Protein %	Grain
Genotype	%	(cm)	number	number	%	(day)	weight (g)	1 lotelli 70	Hardness
Az/b	58.8±5.5	49.3±1.6	1.3±0.3	$6.5 \pm 0.7$	$57.2 \pm 3.0$	238.5±0.7	22.1±1.7	15.4±0.3	59.6±2.7
Crr/b	61.3±6.3	58.0±3.3	$6.8 \pm 2.8$	$10.0 \pm 1.6$	60.9±6.1	239.0±1.6	$38.4 \pm 4.8$	15.7±02	60.4±2.0
La/b	58.3±7.7	99.3±2.5	13.3±1.6	$14.8\pm0.9$	75.3±4.7	240.0±0.4	46.4±1.0	16.3±0.3	62.1±0.8
La(4B)4D/b	76.3±8.3	92.3±1.9	$13.8 \pm 2.3$	$15.5\pm0.4$	69.3±3.5	238.3±1.1	48.2±1.0	17.8±0.5	53.0±3.9
Ka/b	51.3±2.4	59.3±2.3	$14.8 \pm 3.3$	12.7±1.5	$63.5 \pm 5.2$	242.2±1.4	59.4±7.2	15.3±0.3	$62.2 \pm 0.7$
Ma//b	61.3±4.3	81.5±1.9	$10.0\pm0.9$	$15.0\pm0.3$	$61.2\pm7.7$	240.3±0.3	41.9±0.5	16.6±0.3	57.0±2.5
Ne/b	62.5±7.2	$101.8 \pm 2.1$	13.7±1.1	16.5±0.0	59.1±12	$240.0\pm0.4$	46.3±0.8	16.7±0.2	60.2±1.5
St/b	65.0±5.4	99.8±3.1	$14.0\pm0.6$	$15.5\pm0.7$	$60.2 \pm 8.4$	239.7±1.7	43.5±1.5	17.8±0.3	56.7±2.6
Cs/b	53.8±5.2	77.8±1.3	$10.5 \pm 2.2$	19.0±1.1	63.2±2.9	236.5±0.7	37.1±0.6	16.7±0.3	$58.2 \pm 2.6$
St/Th.j	63.8±4.3	99.0±1.2	$13.0\pm0.9$	$17.0 \pm 1.7$	35.3±2.4	250.3±2.5	49.1±4.5	18.0±0.3	54.1±1.9
Az	31.3±2.4	65.3±2.5	$5.5 \pm 0.9$	14.3±1.0	$84.5 \pm 8.9$	236.5±1.6	42.7±2.6	16.5±0.4	34.6±32
Cr	41.3±3.8	$60.8\pm\pm4.2$	7.5±5.3	14.5±0.5	90.1±4.4	238.3±0.3	$50.8 \pm 0.7$	16.0±0.4	38.7±3.5
La	43.8±4.3	$101.8 \pm 2.0$	$18.5 \pm 2.0$	20.5±1.0	90.6±3.2	238.0±0.9	49.8±2.1	16.3±0.2	38.5±1.6
Ka	32.5±3.3	$55.0 \pm 2.6$	$5.0 \pm 1.2$	12.5±0.9	78.2±14	237.5±1.2	46.1±5.5	15.5±0.3	39.8±2.4
Ma	47.5±6.3	81.5±2.5	$7.8\pm0.9$	$17.8\pm0.3$	$85.2 \pm 4.0$	236.8±0.8	51.6±1.0	$17.2\pm0.1$	39.2±1.8
Ne	40.0±2.0	62.8±0.9	$10.0\pm1$	17.5±1.1	92.1±1.4	239.7±0.6	59.3±5.4	$17.2\pm0.4$	37.7±2.1
St	41.3±2.4	101.5±2.3	13.5±2.6	19.0±0.3	93.4±0.6	237.3±1.1	52.4±0.5	16.8±0.4	39.5±2.8
CS	57.5±12.0	75.3±2.3	9.8±2.2	19.5±0.6	90.6±1.8	233.8±1.8	38.2±0.9	13.9±0.4	60.5±0.6
Grand Mean	52.6	78.3	10.5	15.44	72.78	239	45.72	16.41	50.67
CV%	21.0	5.95	34.4	11.99	17.47	0.99	12.85	3.83	9.45
LSD (Block)	7.40	3.12	2.41	1.24	8.52	1.57	3.94	0.42	3.21
LSD (Genotype)	15.7	6.62	5.11	2.61	18.07	3.33	8.35	0.89	6.81

Table 4. Mean of morphgical characters of tritipyrum and wheat cultivars.

pyrum lines, which were tested in the hydroponic culture, exhibited good salt tolerance (King *et al.*, 1997). For example, the Neodur/*Th. bessarabicum* and Langdon/*Th. bessarabicum* lines exhibited 27.8 and 38.9%, survival rates, respectively, in 250 mM NaCI, whereas both Neodure and Langdon wheat parents died. This paper reports the results in some aspects of this new third man-made cereal as follows.

#### Cytology and Cytogenetic Survey

Chromosome counts of progenies of 42chromosome hexaploid Tritipyrums exhibited a considerable amount of instability in the form of aneuploidy (45.3 %) with a proportion of plants having lost chromosomes (40-41), and some having addition chromosomes (43-44). In the 56-chromosome octaploid genotypes, the aneuploidy was even higher (62.5 %) with the chromosome number ranging from 52 to 60 (Table 1). King *et al.* (1997) reported an overall mean of 17.80 bivalents for 6x tritipyrum in the early generation of tritipyrum, but in this study it was

19.05. This suggests that after relatively few selfing generations the pairing behavior has already improved. A very low rate of multivalents was found in some of the lines. This could be due to homoeologous pairing between A, B and  $E^{b}$  suggesting that the  $E^{b}$ genome may have the capacity to partly suppress the activity of the Ph1 locus of chromosome 5B of wheat, which normally prevents homoeologous pairings (Miller et al., 1998). FISH, using total genomic DNA of the alien Th. bessarabicum species as a probe (GISH), was a valuable technique for identifying the E<sup>b</sup> chromosomes in tritipyrum genotypes (Table 2, Fig. 4a-b). Although the random univalent and rod bivalent ratio was expected to be 1 ( $E^b$ ): 2 (A+B) and I (E<sup>b</sup>E<sup>b</sup>): 2 (AA+BB) respectively, but ratios of 2 ( $E^{b}$ ): 1 (A+B) and 1 ( $E^{b}E^{b}$ ): 1.5 (AA+BB), respectively, were found and confirmed by Chi-Square analysis. These results show that although the E<sup>b</sup> chromosomes play a major part in the pairing failure and meiotic instability of tritipyrums, the A and B genome chromosomes, must not be ignored, as univalent and rod bivalent chromosomes occur in both E<sup>b</sup> and AB genomes

E Dianta	No. of E	E <sup>b</sup> chrom	nosomes	Back	No. of E <sup>b</sup> chromosomes			
1 <sup>2</sup> 1 Iants	Univ.	Biv.	Total	cross plants	Univ.	Biv.	Total	
1	5	3	11	1	3	4	11	
2	4	3	10	2	3	4	11	
3	1	4	9	3	3	3	9	
4	3	3	9	4	4	2	8	
5	6	1	8	5	2	3	8	
6	4	2	8	6	6	1	8	
7	2	3	8	7	4	2	8	
8	2	3	8	8	5	1	7	
9	3	2	7	9	3	2	7	
10	1	3	7	10	4	1	6	
11	3	2	7	11	3	1	5	
12	4	1	6	12	5	0	5	
13	2	2	6	13	3	1	5	
14	3	1	5	14	3	1	5	
15	5	0	5	-	-	-	-	
16	2	1	4	-	-	-	-	
17	1	1	3	-	-	-	-	
18	2	0	2	-	-	-	-	
Mean	3	2	6	-	3.8	1.7	8	

**Table 5.** The  $E^b$  chromosome constitution at meiosis of the Tritipyrum x 6x wheat  $F_2$  and Tritipyrum/6x wheat x Tritipyrum backcrosses.

(Table 2). The level of *in situ* cross hybridization also indicated that there is a relatively close genetic relationship between the, A, B and  $E^{b}$  genomes of Tritipyrum.

#### **Greenhouse Test**

The morphologies of the glass-house grown Tritipyrum lines were predominantly wheat-like. They had relatively low fertility and showed perennial habit (Fig. 2f), ears with a brittle rachis (Fig. 2e), with the exception of CS/*Th. bessarabicum* line, a range of tiller production and variation in plant and spike morphology in both vernalized and unvernalized conditions (Table 3), which are in agreement with King *et al.* (1997). Despite the involvement of a single accession of *Th. bessarabicum*, these observations suggest that improvement is possible by breeding and selection within the A and

**Field Trial** The results of the Tritipyrum lines alongde their wheat parents demonstrated that

late maturity of Tritipyrum.

side their wheat parents demonstrated that the Langdon (4B)4D/*Th. bessarabicum* [La (4B/4D)/b] line showed the best overall performance. It had the greatest survival rate of plants, the second highest fertility level, and was the first to produce ears. Only Chinese Spring/*Th. bessarabicum* was earlier. It had the third highest 1000-grain weight, the third highest tiller number, and the second highest protein content, in comparison with the other Tritipyrums including the normal Langdon/*Th. bessarabicum* (La/b) line (Ta-

B genome complements. All lines had rela-

tively the same range of late maturity under

both conditions (Table 3), therefore; the lack

of vernalization is not a major factor in the

	1 1								
Progeny	Observed plants rogeny withpistil		Expected plants withpistil on 3:1		$\chi^2$	Expect withpis	ed plants til on 15:1	$\chi^2$	Р
	Single	Multiple	Single	Multiple		Single	Multiple		value
а	13	2	11.25	3.75	$0.56^{**}$	14.063	0.938	0.36**	0.01
b	92	5	72.75	24.25	19.3 <sup>ns</sup>	90.938	6.063	$0.056^{**}$	0.01
a+b	105	7	84	28	20.1 <sup>ns</sup>	105	7	0.038**	0.01

**Table 6.** Segregation ratio for multiple pistil in F<sub>2</sub> progeny. a) Small population of Cr/b x Ks/b. b) Large population of Cr/b x Ks/b, Ma/b and Sub.

ns, \*\* Non-significant, significant p=0.01 respectively.

ble 4). Therefore, it could be concluded that the removal of chromosome 4B and/or the substitution of 4D has a pronounced beneficial effect and could be an important factor in future Tritipyrum breeding.Variation in survival (Table4) indicated that in contrast to the wheat parents, which were all of spring habits, the Tritipyrums had greater winter hardiness and possessed a potential as a winter-sown crop, presumably as a result of genes of the E<sup>b</sup> genome. All Tritipyrums showed relatively low levels of fertility in comparison with the wheat parents (Table 4). This may be due to chromosome instability at meiosis, although there is no evidence of a correlation between fertility and variation in chromosome pairing (King et al., 1997). Therefore, other factors in E<sup>b</sup> genome may be responsible for this phenomenon. Ear emergence can be used as an indicator of the maturity behavior of Tritipyrum in comparison with the wheat parents. All Tritipyrum lines had later ear emergence than their wheat parents (Fig. 2d). The 8x St/Th. junceiforme showed the greatest delay in ear emergence, presumably due to the presence of the two Thinopyrum genomes (Hassani et al., 1998).

The highest protein amount result, that of the 8x Tritipyrum involving *Th. junceiforme* probably partly results from the generally accepted negative correlation between yield and protein content, but may also indicate a potential of this genotype for introducing into Tritipyrum variation for this character from within the E genomes. The large variation for all the characters studied shows there is a considerable potential for improvement of Tritipyrum as a new cereal. In this respect, the development of Tritipyrum has parallel aspects with the development of triticale, which in early stages, it also showed similar variations, but breeding and continuing selection, especially within the wheat components produced a successful new cereal to the world farming system.

#### The Production of Substituted Tritipyrum

In spite of considerable variation between the Tritipyrum lines, the morphology of  $F_1$ hybrids was predominantly wheat-like. Pollen sterility was observed in all hybrid combinations ranging from 1.9% (Cr/b x Wembley) to 63.1% (La/b x Wembley). Seven of the unpaired chromosomes could be identified as E<sup>b</sup> chromosomes by FISH on meiotic preparations in  $F_1$  progeny (Fig. 4c-d). The in situ results on mitotic preparations also confirmed the presence of seven E<sup>b</sup> chromosomes (Fig. 4e). Due to the large number of univalents in F'<sub>1</sub>s a high range of aneuploidy and cytological segregation was, as expected, observed in the F<sub>2</sub> generation. A considerable range of segregation was also observed in terms of plant morphology (Fig. 3a-b). There was also a high level of sterility, production of back tillers and late maturity. From the large population of F2 selfgeneration the seed production was mainly from those involving [La (4B)4D/b, Ma/b and Ne/b] x 6x Wembley and La (4B)4D/b x S<sub>6-4</sub> combinations. This suggests that the choice of wheat parent, both primary



tetraploid and secondary hexaploid, is important if the problem of hybrid necrosis is to be avoided (Fig. 3e). Wembley apparently is a better parent than  $S_{6-4}$ . Lines with specific *T. durum* backgrounds, especially La (4B)4D, showed higher performance, indicating that the initial 4x wheat parent is also important. These effects suggest that variation in both 4x and 6x wheat could play a part in improving Tritipyrum (Hassani *et al.*, 1998).

The mean ratio of fertile to sterile plants in  $F_3(1.5:1)$  and the BC<sub>1</sub>- $F_2(1.83:1)$  progenies in comparison with  $F_2$  (0.41:1) and  $BC_1$ (0.55:1) generations were very much higher, respectively. This is presumably the result of higher chromosomal stability in F<sub>3</sub> and BC<sub>1</sub>- $F_2$  in comparison with  $F_2$  and  $BC_1$  generations, respectively (Hassani, 1998). The GISH on meiotic slides from fertile F<sub>2</sub> and BC<sub>1</sub> plants ranging (2-11) and (5-11) of  $E^{b}$ chromosomes show that it is possible to produce Tritipyrum with variable numbers of  $E^{b}$  and D genome chromosomes (Table 5, Fig. 4f-g), and that FISH is a useful technique for determining the number of E<sup>b</sup> chromosomes present. To recognize E<sup>b</sup> chromosomes from wheat ones (A, B and D genome chromosomes) various probe, hybridization and prehybridization mixtures with different fluorochrome labeled nucleotides were tried. The best differentiation results were achieved when dCTP nucleotide labeled with Texas Red fluorochrome was used following preblocking by total DNA of Chinese Spring wheat. GISH also showed the presence of T. durum-Th. bessarabicum Robertsonian translocated chromosomes (Fig. 4h). The 1BL.1RS wheat/rye translocated chromosome is an example of this type of chromosome and is widespread in European cultivated wheat varieties, where it initially conferred resistance to certain foliar diseases. Although this type of chromosome may have a far from ideal, large amount of unwanted and possibly detrimental alien genetic material, it may nevertheless, be useful for Tritipyrum, by permitting the substitution of a single chromosome arm. Such translocated chromosomes may also

have value as a means of introducing *Th.* bessarabicum characters into wheat. The low level of fertility and crossability means that a considerable effort will be required to produce sufficient progeny to produce specific combinations of  $E^b$  and D genome chromosomes. Further investigations by molecular cytogenetics and molecular markers along with morphological selection will be required in subsequent generations to establish stable genotypes with varying  $E^b$  and D genome chromosome substitutions and desirable phenotypes. However, this study supports the feasibility of substituted Tritipyrum.

#### Novel Multiple pistil/seed

#### **Genetic Inheritance of Multiple Pistils**

King et al. (1997) reported the occurrence of multiple seeds in the Creso/Th. bessarabicum Tritipyrum line. This appears to be the first report of this phenomenon in the Triticeae with no indication of the mechanism. Pistillody in which the anthers become converted or partially converted to pistils is known in wheat (Sears, 1954). This condition is produced by the absence of a gene on chromosome 6B, and rarely gives rise to multiple seeds within a floret. In the Creso/Th.bessarabicum, the number of pistils (gynoecia) in each floret varies from one to three with single pistil florets predominating in each spike of individual plants. The number of stamens remains unchanged. Observation of both Th. bessarabicum and T. durum showed that neither had multiple pistils and the phenomenon is not a postfertilization event as Cr/b has multiple pistils in pre-anthesis (Fig.5a-b). The possibility that plants from multiple seed florets were haploid was considered, but no baploid plants were found from pairs of seeds from multiple seed florets of Cr/b and Cr/b x Ka/b. However, aneuploid differences did occur within their florets, which prove the pre-pollination mechanism of multiple seed formation if the multiple seeds arose from a post-pollination event, seed within a single floret would be expected to have identical chromosome numbers (Hassam, 1998). If the multiple pistil trait is the result of two recessive genes (Table 6), one in each parent, then it follows that there is variation for the genes in the tetraploid wheats as the trait only occurred in the Creso based line, and possibly in the Ma/b line which had a few multiple seeds in the early generation (Miller, Pers. Comm.). The variation is less likely to be from Th. bessarabicum as a single accession was used as the parent for all of the lines (King et al., 1997). The trait does not appear to be environmentally affected because it occurs both under glass house and field conditions. The possibility of a specific Creso cytoplasmic effect may also warrant consideration, although in general all T. durum cultivars would be expected to have similar cytoplasm (Hassani et al., 1998).

# Production of Monosomic *T. durum-Th. bessarabicum* addition lines

In order to determine the chromosomal location of the trait a novel set of addition lines of the chromosomes of Th. bessarabicum to T. durum Creso were produced (Fig. 1a). Some of the  $F_1$  plants had 35 chromosomes but the majority did not, presumably as a result of aneuploid female gametes in the Cr/b. The 35-chromosome plants exhibited 14 bivalents and seven univalents (Fig. 5e). The FISH study showed that, as expected, the 14 bivalents were from A and B genomes and the seven univalents from the  $E^{b}$  genome (Fig. 5f). The 29-chromosome  $F_{2}$ (Fig. 5c) and F<sub>3</sub> plants (Fig. 5h) were selected, by chromosome counting, as potential T. durum-Th. bessarabicum addition lines. At maturity, these were classified into seven morphologically distinct groups (Fig. 5g&d). These results indicate a high probability of having established the seven possible monosomic additions of Th. bessarabicum to T. durum. Although the multiple pistil trait showed heritability through several

generations of the Cr/b Tritipyrum and in the hybrid progenies, but there was no evidence for its presence in any of the 29chromosome monosomic additions. This could be due to the need for the gene(s) to be present in two doses before expression occurs. If this is the case, it will be necessary to obtain 30-chromosome disomic additions to identify its chromosomal location. Alternatively, if the trait results from genes on more than one Th. bessarabicum chromosomes, e.g. two chromosomes, then single additions will not locate it. In this case multiple addition lines will be required to find the cbromosomal location of the genes (Hassani, 1998). Although seven individual addition lines have not been categorically identified, it seems likely that the seven morphological groups equate with the seven different chromosomes. Chromosome 4E<sup>b</sup> is certainly present in group B on the evidence of the presence of seeds with blue aleurone, a character carried by chromosome 4E<sup>b</sup> and group F, on the basis of its narrow ears, probably carries 2E<sup>b</sup> (Forster et al., 1988). Further characterization, however, including RFLP analysis is required before it can be claimed with certainty that a complete set of seven distinct addition lines has been established.

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**Figure. 3.** Morphological variation of Tritipyrum x wheat  $F_2$  plants a) Vegetative stage. b) At maturity. c and d) A range of (Tritipyrum x wheat) x Tritipyrum BC<sub>1</sub>-F<sub>1</sub> and BC<sub>1</sub>-F<sub>2</sub> plants at maturity. e) Mature plants of Tritipyrum x wheat  $F_1$  hybrids. Left: four necrotic plants of Tritipyrum genotypes x 6x wheat Axona. Right: four normal plants of Tritipyrum genotypes x 6x wheat Wembley.

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**Figure. 4.** a) A metaphase I cell of meiotic preparation of Creso/*Th. bessarabicum* stained with DAPI, showing two rod bivalent and nineteen ring bivalent, undifferentiated chromosomes. b) Meiotic preparation of *Creso/Th. bessarabicum* showing one  $E^b$  rod bivalent and six  $E^b$  ring bivalents. c) GISH on a meiotic preparation of a Tritipyrum x wheat  $F_1$  hybrid. A metaphase I cell stained with DAPI, showing 14 undifferentiated univalent chromosomes of the  $E^b$  and D genomes. d) The same cell, the seven  $E^b$  univalent fluoresce (bright red). e) GISH on mitotic cells of Tritipyrum x wheat  $F_1$  hybrids. A mitotic preparation of the St/b x S  $_{6-4}$  hybrid showing seven univalent chromosomes of the  $E^b$  genome (Single filter). f) A pollen mother cell at Metaphase I of a Tritipyrum x Wembley  $F_2$  plant stained with DAPI showing 42 chromosomes, 13 ring bivalents, four rod bivalents and eight univalents. g) A pollen mother cell at Metaphase I of a Tritipyrum x Wembley  $F_2$  plant stained with DAPI showing 40 for univalents. h) A pollen mother cell at metaphase I of a BC<sub>1</sub> plant showing five  $E^b$  chromosomes, two ring bivalents, one univalent and one rod bivalent consisting of a wheat-*Th. bessarabicum* Robertsonian translocation chromosome (arrowed).



**Figure 5.** a) Multiple pistils in a single floret of the Creso/*Th. bessarabicum* line. b) Multiple seeds of individual florets (photo, King *et al.* 1997). c) A PMC at Metaphase I of (Creso/*Th besarabicum*) x Creso  $F_2$  plant shows one univalent  $E^b$  chromosome. d) Seven morphological distinct 29-chromosome potential tetraploid-*Th. bessarabicum* addition lines from (Cr/b) x Cr  $F_4$  progeny. e) A meiotic preparation at Metaphase I of Cr/b x Cr  $F_1$  hybrid with 35 chromosomes, 14 pairs of A and B and seven univalents of  $E^b$  genomes stained with DAPI. f) *In situ* hybridization on a meiotic preparation at Metaphase I Cr/b x Cr  $F_1$  hybrid with 35 chromosomes, 14 of A and B and 7 of  $E^b$  genomes under bright red filter. g) Potential addition line groupings of Cr/b x Cr  $F_3$  progenies. h) A PMC at Metaphase I of (Creso/*Th. bessarabicum*) x Creso  $F_1$  plant showing one univalent  $E^b$  chromosome.

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### آیا تریتیپیرم (TRITIPYRUM) ، آمفی پلوئیدی جدید با پتانسیل تحمل به شوری، می تواند غلهای موفق مانند تریتیکاله باشد؟

### چکیدہ

خاکهای متاثر از شوری مشکل عمده و فراگیر جهانی است که سطح اراضی قابل توجهی را در بسیاری از کشورها از تولید محصولات زراعی خارج نموده است. اصلاح واریته های ژنتیکی متحمل به شوری مستلزم دانش و آگاهی

ما در درک بهتر ماهیت و کنترل ژنتیکی پیچیده این صفت خواهد بود. تریتیپیرم آمفی پلوئید جدید متحمل به شوری حاصل از تلاقی بین گونه های دو جنس گندم نان ( پایه مادری) و تینوپیرم وحشی (پایه پدری) می تواند نوید و امیدی تازه در این زمینه باشد. اگر چه در این غله جدید بویژه ارقام هگزایلوئید، تلاقی بین ارقام گندم دورم و یک گونه علف شورپسند دیپلوئید، شواهدی محکم دال بر دارا بودن پتانسیلی بالقوه مبنی بر ظهو به عنوان یک گیاه زراعي جديد و مقاوم به تنش شوري ديده مي شود وليكن اين غله مصنوعي نوظهور داراي معايب ناپايداري جزئي کروموزومی و باروری کم همانند تریتیکاله در اوان پیدایش خود به عنوان یک غله مصنوعی می باشد. همانگونه که صفات نامطلوب در آن گیاه به تدریج با کارهای اصلاحی مرتفع گردید به نظر می رسد که غلبه بر این صفات در این گیاه نیز با جایگزین نمودن کروموزومهای ژنوم D گندم نان با کروموزمهای ژنوم وحشی پایه پدری آن (E<sup>b</sup>) میسر گردد. در راستای تحقق عملی این فرضیه در آزمایشاتی نتایج حاصل از تلاقی آن با ارقام مختلف گندم هگزاپلوئید با تکنیک هیبریداسیون DNA فلورسنت در محل طبیعی خود (FISH) مطالعه گردید. علاوه بر آن خصوصیات مرفولوژیکی، سیتولوژیکی، و رفتار زراعی این گیاه برای اولین بار در مزرعه بررسی گردید. همچنین صفت جدید چند بذری در یک لاین بررسی و اساس ژنتیکی – کروموزومی آن تا حدود زیادی پایه ریزی شد. نتايج اوليه و اميدوار كننده حاصل از تحقيقات فوق الذكر نشان داد اولا" توليد لاين هاى ثانويه تريتيپيرم حاوى کروموزومهای جایگزین شده از ژنومD به جای ژنومE<sup>b</sup> جهت حذف برخی صفات نامطلوب و دستیابی به لاینهای با ثبات كروموزومي توام با عملكرد و باروري بيش از لاينهاي موجود از طريق روشهاي اصلاحي امكان پذير مي باشد. ثانیا" وجود خواص پنجه زنی مستمر و چند ساله بودن این گیاه امکان اصلاح یک نبات زراعی دو منظوره (تولید علوفه و دانه) امری دور از انتظار و دسترس نیست. ثالثا" به احتمال بسیار قوی صفت چند بذری تحت کنترل دو ژن مغلوب یکی در رقم وحشی و دیگری در ارقام گونه زراعی گندم دوروم بوده و شواهد موجود در بخش مطالعه اساس کروموزومی یدید ه نادر چند بذری در خانواده گندمیان به احتمال زیاد دلالت بر ایجاد یک مجموعه هفت تایی از یک رقم گندم تتراپلوئید (Creso)، هر یک حاوی یک کروموزوم اضافه متفاوت از گونه وحشی والد پدری، برای اولین بار در سطح دنیا دارد.