Molecular and Cytogenetic Studies on Abiotic Stress tolerance in wheat

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ABSTRACT

In the present experiment, ten Egyptian wheat (Triticum aestivum L.) cultivars were used, i.e. Gemmiza 9, Gemmiza 10, Gemmiza 11, Sids 1, Sids 2, Misr 1, Sakha 93, Giza 168, Shakha 94 and Shakha 95. Morphological, chemical, biochemical, and molecular markers were used to detect the genetic differentiation among the cultivars under different salt levels on seedling performance. The results indicated that there are highly significant variations between the studied wheat cultivars. For RAPD-PCR analysis, a total of 75 bands were detected among the studied cultivars, 53 bands showed polymorphism and for SSR markers a total of 11 bands were detected among the studied genotypes and 8 bands showed polymorphism

Key words: genetic differentiation, wheat, polymorphism, SSR.

INTRODUCTION

Wheat (Triticum aestivum L.) is one of the world’s major cereal crops as the unique molecular makeup of its grain allows its use as a primary structural ingredient of breads, pastas, tortillas, and other products worldwide (Collard et al., 2005). Wheat (Triticum spp.) is a monocot and belongs to tribe Triticaceae of family Poaceae (Gramineae). Other important crops like rice (Oryza sativa L.), maize (Zea mays L.) and bamboo also belong to this family (Shitsukawa et al., 2006).

Wheat is the main cereal grain crop grown in Egypt for thousands of years, serving as the principal source of calories in Egyptian diet. The productivity of Egyptian wheat cultivars has increased dramatically from 2.4 Mg ha⁻¹ in 1958 to 6.8 Mg ha⁻¹ in 2004. Enrichment of genetic diversity plays a crucial role in wheat cultivar improvement (Chao et al., 2007). Genetic markers represent genetic differences between individual organisms or species.

Generally, they do not represent the target genes themselves but act as ‘signs’ or ‘flags’. Genetic markers that are located in close proximity to genes (i.e. tightly linked) may be referred to as gene ‘tags’. Such markers themselves do not affect the phenotype of the trait that interest because they are located only near or ‘linked’ to genes controlling the trait. All genetic markers occupy specific genomic positions within chromosomes (like genes) called ‘loci’ (singular ‘locus’). There are three major types of genetic markers: (1) morphological (also ‘classical’ or ‘visible’) markers which themselves are phenotypic traits or characters; (2) biochemical markers, which include allelic variants of enzymes called isozymes; and (3) DNA (or molecular) markers, which reveal sites of variation in DNA (Jones et al., 1997)

MATERIALS AND METHODS

The present research was carried out at the Faculty of Agriculture, Alexandria University, Egypt during the seasons of 2019 up to 2020 to study the morphological, biochemical, and molecular genetic markers of some selected bread wheat cultivars to detect the genetic differentiation. Ten seeds were sown in each pot from each cultivar in Selica-jel after washing with adequate amount of distill water. Sodium chloride solution was applied after 8 days of germination in four levels (0mM, 50mM, 150mM and 200mM) were applied as foliar spray after three weeks of germination. The following morphological parameters were measured: shoot length (cm), root length (cm), No of leaves/plant, No of roots/plant and proline content. Completely Randomized Design (CRD) with four replicates was used.

Leaves from each cultivar were grounded separately, using a cooled mortar with a pestle, and adding 0.23 M Tris-acetate, pH 5.0. Homogenate was extracted by the solution containing Tris (27.7 g) and citric acid (11.0 g) in one liter volume adjusted with distilled water. Electrophoresis was carried out by the prescriptions recommending 1% agar-starch-polyvinyl-pyrrolidone gel and Tris-orate or Tris- acetate separation buffers. Electrophoresis was conducted at 270 v, 4°C for 100 min. 100 ml of 0.01 M acetate buffer pH 5.0, containing 0.1% benzidine and 0.5% hydrogen peroxide (H₂O₂) were layered over the gel immediately before staining (Sabrah, 1980). Proline was determined according to the method of Bates et al., (1973) by 3% aqueous sulfosalicylic acid, acid ninhydrin: 1.25 gm ninhydrin, 30 ml glacial acetic acid, 20 ml 6M phosphoric acid.

RAPD analysis was carried out using 10 oligonucleotide primers (Table 1) that were selected from the Operon Kit (Operon Technologies Inc., Alabameda, CA). The polymerase chain reaction mixture (25µl) consisted of 13µl master mix (Promega) Tag DNA polymerase; 2µl of genomic DNA, 2µl primer, 8µl deionized water. PCR amplification was performed in a BiometraTI gradient thermal cycler for 35 cycles after initial denaturation for 5min at 94°C. Each cycle consisted of denaturation at 94°C for 1min; annealing at 36°C for 1min; extension at 72°C for 2min and final

DOI: 10.21608/ASEJAIQSJSAE.2020.129275

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Received November 2, 2020, Accepted, December 12, 2020.
extension at 72°C for 5 min (Williams et al., 1980). Amplification products were separated on 2% agarose gels at 100 volts for 1.30 hrs with 1 x TBE buffer. To detect ethidium bromide/DNA complex, agarose gels were examined on ultraviolet transilluminator (302 nm wave length) and photographed. Using 100pb Plus DNA ladder, ready-to-use (Gene Ruler, Fermentas, and Life Sciences), the length of the different DNA fragments were determined. For each sample, the reproducible DNA bands from two runs were scored for their presence or absence.

Table 1. Primers name and their oligonucleotide sequences used in the current study.

| Primer Code | Sequence |
|-------------|----------|
| OPN-04      | 5' - GACCGGACCCA - 3' |
| OPD-05      | 5' - TGAGCGGACA - 3' |
| OPC-05      | 5' - GATGACGGCC - 3' |
| OPM-05      | 5' - GGAGACGTG - 3' |
| OPB-07      | 5' - GAAACGGTG - 3' |
| OPN-10      | 5' - ACAACTGGGG - 3' |
| OPG-12      | 5' - CAGCTCGA - 3' |
| OPQ-12      | 5' - AGTACGGGCAC - 3' |
| OPN-13      | 5' - AGCGTCCTCA - 3' |
| OPQ-14      | 5' - GAGACGCTTCA - 3' |

Simple Sequence Repeats (SSRs), also known as microsatellites, are repeating sequences of nucleotides, such as (AC)n. They have been found in abundance on most eukaryotic chromosomes and are often highly polymorphic (Rafalski and Tingey 1993). Five SSR markers described (Kong et al., 2000) were used for genotyping assays (Table 2). One way ANOVA in completely randomized experiments was used to reveal the significant differences among the samples. The LSD test was conducted to identify the significant differences among the means at 5% level of probability (Siugh, 1994).

Table 2. Sequences of the SSR loci and annealing temperature for PCR reaction used in the current study

| Locus  | Sequence of forward and reveres primers |
|--------|----------------------------------------|
| Wmc661 | F: CCACCATGGTGCTAATAGTGTC             |
|        | R: AGCTCGTAACGTAATGCAACTG             |
| Xtxp-8 | F: ACAT CTACT AC CCT CTACCC           |
|        | R: ACACATCGAGACCCAGTTG                |
| Xtxp-10| F: ATACTATCAAGAGGGGAGC                |
|        | R: AGTACTAGCCACACGTCAC                |
| Xtxp-12| F: ATAT GGAGAGGAAGAGC CG              |
|        | R: AACACAACAT GCAC GCAT G             |
| Xtxp-19| F: ATACTATCAAGAGGGGAGC                |
|        | R: AGTACTAGCCACACGTCAC                |

RESULTS AND DISCUSSION

Morphological markers

Regarding on the morphological markers, results presented in Table 3 indicated significant variations among the Egyptian wheat cultivars in the seedling and root length (cm). Sids 2 was the highest one by average 18.95 cm, followed by misr 1 in average 18.68 cm, while Gemmeiza 10 was the shortest one (15.04 cm). Concerning to the Egyptian wheat cultivars data indicated significant values between Sakha 93, Giza 168 and the other cultivars. Sooht length of Giza 168 was (15.67 cm) followed by Sakha 93 (13.90 cm), however no significant variations were observed between Gemmeiza 9, Sakha 94 and Sakha 95. The range between Egyptian wheat cultivars in relation to shoot length ranged from 18.95 cm (Sids 2) to 13.28 cm (Gemmeiza 9) by value ~5.67 increasing 35% and between the shortest cultivars ranged from 15.05 (GEMMIZA A11) to 13.28 (Gemmeiza 9) in increase by ~ 10% (Table 3).

According to root length, results indicated significant variations among all studied wheat cultivars. The Sakha 93 wheat cultivar recorded the highest value (4.29 cm) followed by Gemmeiza 10 in average 4.09 cm, while Gemmeiza 11 and Sakha 94 was the shortest in average 2.56 and 2.56 cm, in respect. The highest root length was recorded to Sakha 93 in average 4.29 cm followed by Gemmeiza 11 (4.09 cm). The range between Egyptian wheat cultivars in root length ranged from 4.29 cm (Sakha 93) to 2.56 cm (Sids 1) by value = 1.73 increasing 40% (Table 3). Wheat cultivars showed decrease in root length with increasing the salt levels until 200 mM salt, Sakha 93 and followed by Gemmeiza 11 showed the tallest root length (4.33 and 3.83 cm), respectively. While Sakha 95 showed the shortest root (2.33 cm) under 200 mM of sodium chloride.

Results in Table 4 indicated no significant variations were observed among all the studied wheat cultivars in number of leaves/seedling except with Sids 2. The Sids 2 wheat cultivar recorded the highest value (2.44 cm) followed by Gemmeiza 10 in average 4.09 cm, while Gemmeiza 11 and Sakha 94 was the shortest in average 2.56 and 2.56 cm, in respect. The highest root length was recorded to Sakha 93 in average 4.29 cm followed by Gemmeiza 11 (4.09 cm). The range between Egyptian wheat cultivars in root length ranged from 4.29 cm (Sakha 93) to 2.56 cm (Sids 1) by value = 1.73 increasing 40% (Table 3). Wheat cultivars showed decrease in root length with increasing the salt levels until 200 mM salt, Sakha 93 and followed by Gemmeiza 11 showed the tallest root length (4.33 and 3.83 cm), respectively. While Sakha 95 showed the shortest root (2.33 cm) under 200 mM of sodium chloride.
Table 3. Seedling and root length (cm) of Egyptian wheat cultivars as affected by four different salinity concentrations and their interaction during 2019 – 2020 seasons.

| Variety       | Salinity concentration | Mean  | Salinity concentration | Mean  |
|---------------|------------------------|-------|------------------------|-------|
|               | 0          | 50    | 150       | 200  | 0     | 50    | 150       | 200  |
| Gemmiza 11    | 17.67      | 15.09 | 15.10     | 16.10 | 5.40  | 4.07  | 3.07      | 3.83 |
| Gemmiza 10    | 18.19      | 14.46 | 14.66     | 15.04 | 6.18  | 2.93  | 3.23      | 3.23 |
| Sids 1        | 16.13      | 15.37 | 15.11     | 15.72 | 3.17  | 2.55  | 2.55      | 2.67 |
| Sids 2        | 18.37      | 17.20 | 19.67     | 18.95 | 4.47  | 4.37  | 2.90      | 3.40 |
| misr 1        | 17.97      | 18.46 | 18.53     | 18.68 | 2.63  | 2.63  | 2.37      | 2.62 |
| Sakha 93      | 14.27      | 12.80 | 13.72     | 13.90 | 4.53  | 4.50  | 3.80      | 4.33 |
| Geiza 168     | 14.42      | 14.37 | 15.67     | 14.68 | 4.57  | 2.93  | 3.07      | 3.43 |
| Gammeiza 9    | 12.87      | 14.40 | 13.73     | 13.28 | 3.87  | 1.87  | 3.33      | 2.80 |
| Sakha 94      | 13.82      | 12.73 | 14.33     | 13.29 | 3.13  | 2.17  | 2.10      | 3.20 |
| Sakha 95      | 13.97      | 12.66 | 13.50     | 13.44 | 3.86  | 3.20  | 2.87      | 2.33 |
| Mean          | 15.77      | 14.74 | 15.38     | 13.29 | 4.18  | 3.09  | 2.93      | 3.21 |

LSD at 0.05 (V) 0.288
LSD at 0.05 (SC) 0.181
LSD at 0.05 (V x SC) 0.573

Table 4. Number of leaves and roots/seedling of Egyptian wheat cultivars as affected by four different salinity concentrations and their interaction during 2019 – 2020 seasons.

| Variety       | Salinity concentration | Mean  | Salinity concentration | Mean  |
|---------------|------------------------|-------|------------------------|-------|
|               | 0          | 50    | 150       | 200  | 0     | 50    | 150       | 200  |
| Gemmiza 11    | 2.67       | 2.00  | 2.07      | 2.00 | 2.19  | 7.60  | 7.00      | 6.87 |
| Gemmiza 10    | 2.53       | 2.00  | 2.00      | 2.00 | 2.15  | 7.47  | 7.13      | 6.96 |
| Sids 1        | 2.47       | 2.10  | 2.10      | 2.10 | 2.32  | 6.70  | 6.27      | 6.53 |
| Sids 2        | 2.67       | 2.10  | 2.00      | 3.00 | 2.44  | 6.10  | 6.13      | 4.40 |
| misr 1        | 2.67       | 2.00  | 2.00      | 2.00 | 2.17  | 6.80  | 7.50      | 6.00 |
| Sakha 93      | 2.67       | 2.00  | 2.00      | 2.00 | 2.17  | 6.13  | 6.93      | 6.47 |
| Geiza 168     | 2.67       | 2.00  | 2.00      | 2.00 | 2.17  | 6.67  | 6.33      | 6.87 |
| Gammeiza 9    | 2.67       | 2.00  | 2.00      | 2.00 | 2.17  | 6.60  | 5.40      | 6.33 |
| Sakha 94      | 2.67       | 2.00  | 2.00      | 2.00 | 2.17  | 6.60  | 6.10      | 5.40 |
| Sakha 95      | 2.67       | 2.00  | 2.00      | 2.00 | 2.17  | 6.60  | 6.10      | 5.40 |
| Mean          | 2.64       | 2.05b | 2.02b     | 2.11b | 6.65a | 6.39b | 6.18c     | 5.95d |

LSD at 0.05 (V) 0.218
LSD at 0.05 (SC) 0.138
LSD at 0.05 (V x SC) 0.435

Concerning to the wheat cultivars data in Table 4 indicated no significant difference between Sakha 93 and Giza 168, also between Sakha 94 and Sakha 95 cultivars.

Our data are agreed with those reported by (Hamada, 1996) which assessed 13 Aegilops and three wild Triticum originally Turkish species by using morphological, pathological, qualitative and agricultural traits. Also, our result is agreed with Hamada, (1996) used 12 yield parameters and 5 morphological traits of spring wheat to evaluate genetic divergence among 19 durum wheat genotypes. These genotypes were subsequently classified into 7 separate clusters revealing high level of genetic divergence independent of original cultivated zone. The present work is agreement with Hu and Schmidhalter (1998), who reported that wheat growing in 120 mM NaCl reacted with a 25% reduction in growth rate, Na+ in the growing cells of leaves was at maximum only 20 mM, and Cl- only 60 mM. However, Ball (1998) found that the common decrease in leaf expansion is not related to a loss.
in turgor pressure and is most likely a result of a change in hormonal signaling from roots to leaves. In the salt-sensitive genotypes, in which salt is not effectively excluded from the transpiration stream, salt will build up to toxic levels in the leaves, resulting in death of old leaves and new leaves becoming injured and succulent (Munns and James, 2003). Our results are agreement with Ashraf et al., (1986) who reported that root length can be used as selection criteriou under salinity stress.

Heritability estimates were low under 200 and 250 mM NaCl stress indicating that improvement in root length is difficult under stress conditions. Our results are agreement with Hasegawa et al., (2000) who reported that roots play several important roles during plant growth and development and are typically the first part of the plant to encounter salinity. In glycophytes, the root is the primary site of salt stress and the ability to maintain ion homeostasis and redox potential is critical for the normal root growth and function under saline stress.

**Chemical markers**

Results in Table 5 indicated significant variations among all the studied wheat cultivars in chlorophyll. The misr1 wheat cultivar recorded the highest value (30.55) followed by GIZA 168, GEMMIZA 9 and GEMMIZA 11 by average 29.93, 29.24 and 29.04, respectively. Analysis of variance in Table 5 showed significant variation between the wheat cultivars in relation to Chlorophyll by L.S.D.-0.05= 0.825. The reduction in chlorophyll content under salinity agreed with those reported by Iqbal et al., (2006).

Decrease in total chlorophyll content could be due to ion accumulation and functional disorders observed during stomata opening and closing under salinity stress (Nawaz et al., 2010). Another reason for the decrease of chlorophyll content under salt conditions is stated to be the rapid maturing of leaves (Nawaz et al., 2010). Decrease in chlorophyll content under salinity stress is observed more in salt sensitive genotypes in comparison to cultivars with low tolerance (Khan et al., 2009).

Proline content was determined as indicator for salt tolerant in the studied wheat genotypes. Results showed that the proline content was increased by increasing concentration of salt. The regression coefficient was done to determine the relationship between the two variables. Proline was considered as the dependent factor (Y) while the salt concentration was determined as the independent variable (X) for the cultivars. Analysis of variance in Table 5 and fig.1 showed significant variation between wheat cultivars in proline content under different salt concentrations. The highest values with no significant were recorded to misr1 and Gimmelza 11 in average 1.05 and 1.04 in respect with L.S.D. =0.034, while the lowest value was recorded to Sakha 93 with average 0.55.

| Variety | Chlorophyll (μ moles/g/fresh weight) | Proline (μ moles/g/fresh weight) |
|---------|--------------------------------------|----------------------------------|
|         | Mean 0 50 150 200                     | Mean 0 50 150 200                 |
| Gemmiza 11 | 27.73 28.07 26.07 27.13              | 0.453 1.18 1.12 1.38             |
| Gemmiza 10 | 25.33 29.40 31.20 30.33              | 0.488 0.533 0.670 0.995          |
| Sids 1   | 26.93 25.73 27.73 22.07              | 0.727 0.798 0.320 0.592          |
| Sids 2   | 31.67 27.30 25.27 21.87              | 0.436 0.833 0.769 1.27           |
| misr 1   | 27.73 29.13 34.27 31.07              | 0.727 1.09 1.19 1.20            |
| Sakha 93 | 21.67 28.00 24.40 28.13              | 0.530 0.631 0.593 0.426          |
| Geiza 168 | 33.00 28.40 29.93 28.40              | 0.644 0.468 0.543 1.24           |
| Gammeiza 9 | 34.73 26.93 33.70 21.60              | 0.447 0.570 0.441 1.09           |
| Sakha 94 | 26.87 23.93 25.10 24.40              | 0.488 0.589 1.08 1.17            |
| Sakha 95 | 25.93 21.60 29.30 32.93              | 0.757 0.644 0.824 0.886          |
| Mean     | 28.16b 26.85c 28.70a 26.79c           | 0.57 c 0.73b 0.76 b 1.02a        |

LSD at 0.05 (V) 0.825
LSD at 0.05 (SC) 0.522
LSD at 0.05 (V x SC) 1.65
Proline is the only organic cytosolate which able to make the major contribution or osmotic adjustment at sever salinity in roots, while in shoots and spikes the contribution of proline in osmoregulation might be reduced. We concluded that there is no stable situation in usage of organic or inorganic soluble components in osmotic adjustment in the cultivars and lines on different salinity levels. This is happened not only in different cultivars, but also in different organs which conferring the contrasting opinions about the physiological significance of proline which has remained controversial among physiologists. Many reports have pointed out that proline is mostly accumulated when plants growth ceased (Joly et al., 2000).

The overall results of the present study indicated that for all wheat cultivars grown at 50 Mm, 150 Mm, and 200 Mm of salt maintained a higher proline level than those grown at control as Figure 30. The increment of proline level was higher under high salt concentration than that in low salt concentration. Increasing praline level due to high salt can be used to screen wheat genotypes, which is comparable to cell membrane thermo-stability test.

**Biochemical markers**

Peroxidase isozymes exhibited a wide range of variability among the different cultivars at different localities. In control three loci were obtained for all cultivars as two cathodal Pex.1c and Pex. 2a found as common band and Pex 2c was unique for some wheat, while, Pex 3c was unique for some wheat cultivars. With increase of salt levels the results showed increase in number of loci (4 and 5) as shown with 200 Mm salt two anodal (Pex.1A and Pex.2A) were found as common band for all the wheat samples. While (pex.1c) at cathodal was as common band for all the wheat samples. misr1 wheat cultivar gives two bands in cathodal (Pex.2c and Pex.4c).
Our results in a line with Hassanein (1999) who reported that the results showed that band number was exhibited in untreated and treated plants with salt of all cultivars. This band was higher densities and intensities in the salt treated cultivars than grown under control conditions. These results indicated that salt stress increased the accumulation of the esterase enzyme and that encoding gene(s) which accelerated in response to salt stress. Salinity increase esterase isozymes, the highest number of esterase isozymes were detected under the highest NaCl concentration.

**Molecular markers**

A total of 75 bands were detected among the studied cultivars. 53 bands showed polymorphism. Out of these polymorphic bands, 13 unique bands were scored and the number of monomorphic bands was 9 (Table 6). Results indicated clearly that OPC-05 and OPQ-14 showed three unique fragments followed by OPM-05 and OPG-12 by two unique fragments. However, the primer OPC-05, OPM-05 OPN-10 and OPG-12 show 100% polymorphism. While primer OPN-04, OPD-05, OPB-07, OPQ-12, OPN-13 and OPQ-14 showed 60, 88, 80, 75, 60 and 89% polymorphism, respectively. The range of DNA size was between 138 bp and 1825 bp. Molecular markers have also been extensively used to analyze the genetic diversity in crop plants. Based on the data obtained by RAPD analysis, it was possible to discriminate between the ten wheat cultivars used. The genotype-specific markers indicated that the highest number of RAPD specific markers was scored for OPC-05 and OPQ-14 (3 markers), while both OPM-05, and OPG-12 scored two markers each. On the other hand, OPD-05, OPQ-12 and OPN-13 scored one marker each as Table 6 and Figure 2.

For SSR markers a total of 11 bands were detected among the studied genotypes. 8 bands showed polymorphism. Out of these polymorphic bands, 3 bands were monomorphic. However, the primer wmc661 and primer xtxp19 shows 100% polymorphism. While primer XTXP8, and XTXP12 showed 50% polymorphism, respectively. The range of DNA size was between 84bp in primer wmc 661 to 254 bp in primer XTXP12 as Table 7 and Figure 3.

| Primer code | Total amplicons | Monomorphic | polymorphic | unique | Positive unique markers | Polymorphism % |
|-------------|-----------------|-------------|-------------|--------|-------------------------|----------------|
| OPN-04      | 5               | 2           | 3           | 0      | -                       | 60             |
| OPD-05      | 8               | 1           | 6           | 1      | 1096                    | 88             |
| OPC-05      | 7               | 0           | 4           | 3      | 582-292-275             | 100            |
| OPM-05      | 14              | 0           | 12          | 2      | 608-290                 | 100            |
| OPB-07      | 5               | 1           | 4           | 0      | -                       | 80             |
| OPN-10      | 6               | 0           | 6           | 0      | -                       | 100            |
| OPG-12      | 8               | 0           | 6           | 2      | 277-188                 | 100            |
| OPQ-12      | 8               | 2           | 5           | 1      | 199                     | 75             |
| OPN-13      | 5               | 2           | 2           | 1      | 487                     | 60             |
| OPQ-14      | 9               | 1           | 5           | 3      | 417-306-213             | 89             |
| Total       | 75              | 9           | 53          | 13     | 13                      | 85.2           |

Table 6. Polymorphism data as detected by RAPD markers, total number of amplicons, monomorphic and polymorphic amplicons and the percentage of polymorphism among the ten cultivars.

| Primer | Fragment size (bp) | Number of alleles | Monomorphic bands | Polymorphic bands | Polymorphic bands% |
|--------|--------------------|-------------------|-------------------|-------------------|--------------------|
| wmc661 | 84-101             | 2                 | 0                 | 2                 | 100                |
| xtxp8  | 140-244            | 2                 | 1                 | 1                 | 50                 |
| xtxp10 | 272                | 1                 | 1                 | 0                 | 0                  |
| xtxp12 | 221-254            | 2                 | 1                 | 1                 | 50                 |
| xtxp19 | 143-253            | 3                 | 0                 | 3                 | 100                |

Table 7. Number of alleles, fragment size range and polymorphism detected by SSR loci in the ten wheat genotypes.
Fig. 2. DNA polymorphism of the Egyptian wheat cultivars using RAPD DNA (1) Sakha93, (2) Geiza168, (3) Gammeiza9, (4) Sakha94, (5) misr1, (6) Gammeiza10, (7) Gammeiza11, (8) Sids 1, (9) Sids 2, and (10) Sa
REFERENCES

Ashraf, M., T. McNeil and A.D. Bradshaw. 1986. The response of selected salt tolerant and normal lines of four grass species to NaCl in sand culture. New phytologist 104: 453-461.

Ball, M. C. 1988. Salinity tolerance in the mangroves, Aegceras. Geiza168, (3) Gammeiza9, (4) Shakha94, (5) misr1, (6) Gammeiza10, (7) Gammeiza11, (8) Sids 1, (9) Sids 2, and (10) Sakha 95.

Hasegawa, M. 2000. Plant cellular and molecular responses to high salinity. Annual Review of Plant Physiology and Plant Molecular Biology. 51:463-499.

Hassanein, A. 1999. Alternation in protein and esterase wheat in response to salinity stress. Biologia plantarum: 42(2): 241-248.

Hu, Y., U. Schmidhalter. 1998. Spatial distributions and net deposition rates of mineral elements in the elongating wheat (Triticum aestivum L.) leaf under saline soil conditions. Planta. 204: 212-219.

Iqbal, N., M.Y. Ashraf, J. Farrukh, M. Vicente and A. Kafeel. 2006. Nitrate reduction and nutrient accumulation in wheat (Triticum aestivum L.) grown in soil salinization with four different salts. J. Plant Nutrition. 29: 409-421.

Joly, R.J., A. Maggio and M.P. Reddy. 2000. Leaf gas exchange and solute accumulation in the halophyte Salvadora persica grown at moderate salinity. Environmental and Experimental Botany. 44: 31-38.

Jones, N., H. Oughan and H. Thomas. 1997. Markers and mapping: We are all geneticists now. New phytologist. 137: 165-177.

Khan, D.K., M. Tuna, M. Tal, A. Nejidat and A. Golan-Goldhirsh. 2009. Variability in the pattern of random amplified polymorphic DNA. Electrophoresis. 18:2852-2856.

Kong, L., J. Dong and G.E. Hart. 2000. Characterization, linkage map positions, and allelic differentiation of (Sorghum bicolour (L.) Moench) by DNA simple-sequence repeats (SSRs). Theoretical and Applied Genetics. 101:438-448.

Munns, R., James, R.A. 2003. Screening methods for salinity tolerance: a case study with tetraploid wheat. Plant Soil. 59: 1-18.
Mohamed H. Harby: Molecular and Cytogenetic Studies on Abiotic Stress tolerance in wheat

Nawaz, R., P. Thompson, J. McNaught and S. Ananiadou. 2010. Meta Knowledge Annotation of Bio Events. In Proceedings of LREC. 2498 – 2507.

Rafalski, J.A. and S.V. Tingey. 1993. Genetic diagnostics in plant breeding: RAPDs, microsatellites, and machines. Trends in Genetics. 9:275-280.

Sabrah, N.S. 1980. Genetical and cytological studies on maize. Ph.D. Thesis, Faculty of Agriculture, University of Alexandria, Egypt.

Shitsukawa, N., A. Takagishi, C. Ikari, S. Takumi and K. Murai. 2006. WFL, a wheat FLORICAULA/LEAFY ortholog, is associated with spikelet formation as lateral branch of the inflorescence meristem. Genes Genetic System. 81:3–20.

Siugh, P.K. 1994. Genetic diversity in durum wheat germplasm. Annals of Agricultural Research. 15: 418 – 422.

Williams, J.K., A. Kubelik, K.J. Livak, J.A. Rafalski and S.V. Tingey. 1990. DNA polymorphisms amplified by arbitrary primers are useful as genetic markers. Nucleic Acids Research. 18:6531-6535.

الملخص العربي

دراسات جزيئية ووراثية خلوية على القدرة لتحمل الإجهاد البيئي في القمح

محمد حسن حربى

يهدف البحث الحالي لدراسة الإجهاد البيئي مثلاً في القدرة على تحمل الملوحة لأصناف من القمح، ولتحقيق هذا الغرض تم إجراء دراسات جزيئية وكميحيوية ووراثية خلوية.

تم إجراء هذا البحث باستخدام 10 أصناف من القمح وهي:

- جميزه 1
- جميزه 2
- سداس 1
- سداس 2
- مصر
- سخا 7
- جيزه 8
- سخا 9
- سخا 10
- سخا 94

وتم توظيف معلمات جزيئية وكيميحيوية وسيتوغوتية وذلك تحت ظروف الإجهاد بسبب الملوحة.

وأظهرت نتائج البحث الحالي أن هناك فروقاً جوهراً عالية لتحمل بعض الأصناف للملوحة وذلك باستخدام تكنيك تحليل تفاعل البلمرة المتسلسل السريع.

وأظهرت النتائج تحديد 75 حزمة منها 53 حزمة متعددة الشكل المظهري كما أظهر تكنيك ISSR ان هناك 11 حزمة منها 8 حزم متعددة الشكل المظهري.

أهمية هذا البحث لمربى القمح أنه يستطيع في برامج التربية والاختيار أن يحصل على تراكيب وراثية جيدة ومميزة ومفيدة في تربية القمح لتحمل الملوحة حيث أن القمح يعتبر من المحاصير الهامة والاستراتيجية على مستوى العالم.

وقد تم زراعة عشرة حبوب من كل صنف في أرض وذلك بعد غسلها بماء مقشر وقد تم معاملة الحبوب بكوليدين الصوديوم بعد ثمانية أيام من الانتباه وذلك تحت اربعة مستويات مختلفة وهي (صفر، 50، 150، 200) وتم قياس الخواص المورفولوجية في المجموع الخضرى مثل: