EXOGENOUS APPLICATION OF ABSCISIC ACID FOR DROUGHT TOLERANCE IN SUNFLOWER (*HELIANTHUS ANNUUS* L.): A REVIEW

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ABSTRACT

Sunflower has been successfully cultivated over a widely scattered geographical area in the world and also emerged as a crop, which has ability to adopt in a variety of environmental conditions. Sunflower is high yielding oilseed crop and has the potential to bridge up the gap existing between consumption and domestic production of edible oil. Water is essential at every stage of plant growth and development. Water deficit reduces crop yield regardless of the growth stage at which it occurs in field crops including sunflower. The productivity and spatial distribution of agronomic and horticultural crop plants of commercial importance are severely restricted by a variety of environmental factors. Among these factors, drought and salt play very significant role in reducing agricultural production. Water deficit effect is much pronounced at the vegetative and flowering stage. In sunflower water shortage at the flowering stage reduced yield by 29%. Dwarf cultivars of sunflower have higher water potential i.e. least water stress as compared to intermediate and long stature cultivars. This is due to difference in canopy architecture and root penetration depth among different stature sunflower cultivars. Tall stature cultivars produced more leaf area, aerial biomass and deeper root system so they transpired more water. Major effect of drought in plant is reduction in photosynthesis, which is due to decrease in leaf expansion, impaired photosynthetic machinery, leaf senescence and finally reduction in assimilates production and partitioning. Sunflower exhibits a large varietal difference for osmotic adjustment in response to water shortage. Drought tolerance has been observed in all plant species, but its extent varies from species to species. One way to ensure future food needs of the increasing world populations should involve better use of water by the development of crop varieties those require lesser amounts of water and more tolerant to water shortage. Water deficit in root zone causes an increase in rate of root respiration which leads to an imbalance in the utilization of carbon resources, decrease in production of adenosine triphosphate (ATP) and an increase in production of reactive oxygen species (ROS). Genotypic variation for osmotic adjustment has also been reported in sunflower. The dwarf sunflower cultivars have initiated osmotic adjustment earlier than intermediate and large stature cultivars which indicated better osmotic response of dwarf cultivars. The delay in osmotic adjustment in intermediate and tall stature sunflower genotypes is due to faster root penetration and more water extraction from deeper layer of soil. So, tall stature sunflower cultivars face water stress later as compared to short stature. Three growth stages of sunflower as heading, flowering and milking are sensitive to water shortage. In limited irrigation application study, water applied at different growth stages of sunflower significantly decreased seed yield, particularly during three growth periods: heading, flowering and milking. The role of abscisic acid (ABA) as plant stress hormone is well established. Under drought condition, ABA is synthesized in plant tissue and sent to the guard cell as a stress signal. Here ABA causes stomatal closure, which improves the water relations of plant. ABA entering a leaf can be metabolized rapidly. In response of water deficit, ABA begins to increase markedly in plant leaf tissues and to a lesser extent, in other tissues including roots. This leads to stomatal closure and decreased transpiration. It also inhibits shoot growth and root growth appears to be promoted which increased the water supply. It helps in promoting drought tolerance, both from the use of exogenous application to intact plants and from the measurement of the endogenous ABA concentration. In response to exogenous application of ABA in drought tolerant line of sunflower dehydrin protein accumulated in vegetative tissue. Dehydrin protects cytosolic structures from the deleterious effects of cellular dehydration. ABA plays a critical role in regulating plant water status through guard cells and growth as well as by induction of genes that encode enzymes and other proteins which create cellular dehydration tolerance. In plants stomata response to soil drying is mediated by ABA, originating from roots and transported to the shoot via the transpiration stream. ABA under drought is produced in dehydrated roots, transported to the xylem and regulates stomatal opening and leaf growth in the shoots. Stomata respond to the concentration of ABA in the guard cell apoplast. Screening of sunflower cultivars under drought applied at different crop growth stages and exogenous application of ABA will defiantly help in saving water for successful crop production.

Key Words: Sunflower cultivars, drought, growth stages, ABA application.
**INTRODUCTION**

Pakistan is chronically deficient in the production of edible oil and situation is getting worse day by day with alarming increase in population growth rate along with preferences of people for edible oils and deficient local production (Asif et al., 2001). Pakistan is the third largest importe of edible oil in the world and spends huge foreign exchange on the import of edible oil, which is second to that spent on the import fuel oil. The consumer’s demand has steadily increased from 0.3 million tons to 2.764 tons during the last two and half decades with almost stagnant domestic production of 0.857 million tons. At present indigenous oil seed production meets only 27% of domestic requirement, while the remaining 73% is met through imports (Govt. of Pakistan, 2009a).

Domestic oil production during 2007-08 (July–March) was 0.833 million tons, while 2.23 million tons of edible oil was imported and 0.349 million tons recovered from imported oil-seeds. The total availability of edible oil from all sources amounted to be 3.066 million tons. The import of vegetable oils in increasing amounts is imposing a severe drain on national economy. The import bill of edible oil was Rs. 50 billion in 2006-07 and has risen up to Rs. 84 billions 2008-09 (Govt. of Pakistan, 2009b). All this necessitates concentrating on more efforts to increase domestic edible oil production to reduce drain on the foreign exchange reserves.

Edible oilseed crops in Pakistan can be classified as traditional (rapeseed, mustard, groundnut, sesame, cotton and non traditional (sunflower, soybean and safflower). Rapeseed and mustard oil is not regular cooking oil due to the presence of higher concentrations of erucic acid and thioglucoside could be more than 5% in oil blending for ghee manufacturing. Cottonseed contributes about 70-75% to the total domestic oil production only as a by-product. Cotton is mainly grown for fiber purpose and its oil content and fiber are negatively correlated with each other (Govt. of Pakistan, 2009b). Among the non-traditional oilseed crops sunflower can play an important role narrowing the wide gap between production of edible oils in the country and its import (Khan et al, 2003). In Pakistan, area, production and yield per unit area of sunflower have increased tremendously with the passage of time. During 2007-08 area under sunflower cultivation was 1130 thousand acres with 683 thousand tons of seed yield and 264 thousand tons of oil production. During 2008-09 area of sunflower was provisionally estimated as 1250 thousand acres with 755 thousand tons seed and 287 thousand tons oil production (Govt. of Pakistan, 2009b).

As a plant of economic importance, sunflower has been successfully cultivated over a widely scattered geographical area in the world and also emerged as a crop, which has ability to adopt in a variety of environmental conditions (Beard and Geng, 1982). Sunflower is high yielding oilseed crop and has the potential to bridge up the gap existing between consumption and domestic production of edible oil. Furthermore, it is a short duration crop (90-120 days) and can be grown twice a year. It fits well in existing cropping system and can be grown without replacing any major crop. Water is essential at every stage of plant growth and development. Water deficit reduces crop yield regardless of the growth stage at which it occurs in field crops including sunflower (Jensen and Mogenson, 1984; Hussain, 2010). The productivity and spatial distribution of agronomic and horticultural crop plants of commercial importance are severely restricted by a variety of environmental factors. Among these factors, drought and salt play very significant role in reducing agricultural production (Boyer, 1982). In the face of a global scarcity of water resources, drought has already become a primary factor in limiting crop production. Sunflower genotypes have performed differently in various environmental conditions. Different environmental conditions are required to evaluate the good yielding, better adaptive and stable crops varieties (Luquez et al., 2002; Prusti et al., 1999). Under drought sunflower head diameter, achene’s yield, seed weight per head, 1000-seed weight, oil content and oil yield are significantly decreased (Jasinkas, 1999; Kazi et al., 2002). At present, around 18% of the global farmland is irrigated (more than 240 million hectares) and up to 40% of the global food supply is produced on this land (Somerville and Briscoe, 2001).

Diminishing water resources in the world emphasize to limit irrigation for field crops. In some areas of the world the available water supply is not sufficient to produce the maximum yield on the irrigable area. While in other regions, the available moisture for irrigation is already regulated and requires deficit irrigation. For many surface water conservation projects, the annual supply of irrigation water is limited by reservoir capacity and the annual reservoir inflow. These adverse water available conditions highlight the need for deficit irrigation management for different crops (Martin et al., 1989).

The role of abscisic acid (ABA) as plant stress hormone is well established. Under drought condition, ABA is synthesized in plant tissue and sent to the guard cell as a stress signal. Here ABA causes stomatal closure,
which improves the water relations of plant. ABA entering a leaf can be metabolized rapidly (Loveys, 1984; Gowing et al., 1993; Jia et al., 1996). In sunflower, stomatal control depends on the concentration of ABA in xylem sap (Tardieu et al., 1996). Stomatal closure due to response of water shortage is one of the drought-adaptation mechanisms in plant. Physiological observations associated with the varietals differences in stress tolerance in field crops have been reported (Moons et al., 1995; Pelah et al., 1997) and ABA also produced genetic difference in drought tolerant and drought sensitive plants (Ouvrard et al., 1996). Varietal differences in tolerance may be associated with increase of ABA in response to various environmental stresses. This includes drought tolerance of maize (Pekic and Quarrie, 1987), chilling tolerance of rice seedlings (Lee et al., 1993) and salt tolerance of rice (Moons et al., 1995). In wheat growth of dwarf stature cultivars (Dariel) is less sensitive to constant exogenous application of ABA than that of tall isolines (Bet Hashita) (Blum et al., 1997).

**Sunflower commercial cultivation:** Sunflower (*Helianthus annuus* L.) is one of the important oil seed crops in the world. It originated in North America. Sunflower was “camp follower” of western Native American tribes who had domesticated the crop (possibly 1000 BC) and then took it eastward and southward of North America (Weiss, 2000). Firstly Europeans watched its cultivation at many places from Southern Canada to Mexico. It was first introduced in Europe through Spain, and spread in Europe as a curiosity until it approached to Russia where it was readily adapted. After the World War II high oil containing sunflower lines from Russia were introduced in America. It was the production of male sterile and restorer gene system that made hybrid cultivation feasible and increased commercial interest in sunflower and its use as an oil birdseed crop and human snack food crop.

**Sunflower morphology:** It is an annual, erect, broad leaf plant having strong, prolific and laterally spread tap root system. Stem is round in early season and becomes angular and woody later in the season. It normally remains un-branched. Leaves of sunflower are phototropic and follow the sun rays with a lag of 12° behind the sun azimuth. This feature has been contributed to enhance light interception and possibly increased photosynthesis (Putnam et al., 1990).

The Sunflower head is made up of 1,000 to 2,000 individual flowers joined at a common receptacle. It has ligulate ray flowers around circumference which are without stamens and pistils, while the remaining flowers are perfect. Anthesis starts at the periphery and approach to the centre of the head.

In temperate regions, sunflower needs approximately 11 days from planting to emergence, 33 days from emergence to head formation, 27 days from head visible to first anther formation, 8 days from first to last anther and 30 days from last anther to crop maturity. Varietals differences in maturity have been associated with changes in vegetative period before the head become visible.

**Drought:** Water is very essential for plant growth and makes up 75 to 95 percent of plant tissue. A vast amount of water moves throughout the plant daily. Plants use water and carbon dioxide to form sugars and complex carbohydrates. Water acts as a carrier of nutrients and also a cooling agent. It also provides an element of support through turgor and as an intercellular reaction medium (Ashraf and Harris, 2005).

Drought can be defined as an extended period of deficient rainfall relative to the statistical mean for a region. Drought can also be defined according to meteorological, hydrological or agricultural criteria. Meteorological drought is qualified by any significant deficit of precipitation, hydrological drought is manifested as reduced river and stream flow and critically deep ground water tables, while an agricultural drought indicates extended dry period that results in crop stress and harvest reduction. The impact of drought on agriculture is due to the shortage of moisture in the soil, when the moisture in the soil is no longer sufficient to meet the needs of growing crops. It results from a lack of input of moisture in soil from rainfall or irrigation to an extended period.

Among the environmental stress factors one of the most widely limiting for crop production on global basis is water. According to one estimate, around 28 percent of the world’s land is too dry to support vegetation (Kramer and Boyer, 1995). On global basis, water is a paramount factor to determine the distribution of species, and the responses and adaptation of species to water stress are critical for their success in any environmental niche and for their use and productivity in agricultural ecosystem. Severe droughts occurred periodically in several major food producing countries, having far-reaching impacts on global food production and supply. The global production of grain has, in some years, been reduced by 5 percent or more as a result of severe droughts in key countries (Wisner and Chase, 1984). It has been estimated that droughts cause an average annual yield loss of 17 percent in the tropics (Edmeades et al., 1999), but losses can be much more severe and total crop failure is also possible.

The world human population is expected to reach 8.0 billion by 2025 and 8.9 billion by 2050 (FAO, 2006). About 80 million people are being added to the population total each year and 97 percent of the predicted population growth will take place in the developing countries. It is projected that there is need to double world food production in order to feed 8.0 billion people by 2025. This will certainly place more pressure on the
environment. The developing countries, in particular, are confronted with severe food security challenges. With a severe limit to the amount of unused land available to bring in to cultivation, improving crop yields in both normal and less productive soils under limited water supply is an utter requirement to satisfy future world food needs.

Environmental stresses represent a major constraint to meet the world food demand. There are relatively few stress free areas where crops may approach their potential yields. For instance, up to 45% of the world agricultural lands are subject to continuous or frequent drought, wherein 38% of the world human population resides (Bot et al., 2000). In the face of a global scarcity of water resources, drought has already become a primary factor in limiting crop production worldwide. Water-limited crop production depends on the intensity and the pattern of drought which vary from year to year. In some sub-tropical countries like Pakistan, however, there is a high probability that crop water deficits increase in severity as the season progresses, due to lack of rainfall and to the high evaporative demand (Ashraf and Foolad, 2006).

The severity of water shortage imposed on field crops also depends on the susceptibility of crops during different stages of their development. The general effects of drought on plant growth are well known. When soil moisture is deficient, crop establishment may be reduced, plant growth limited, normal development patterns disrupted and eventually, final yield is lowered (Manivannan et al., 2007; Sankar et al., 2007).

Abiotic stresses are considered to be the main source (71%) of yield reductions (Boyer, 1982). The estimation of potential yield losses by individual abiotic stresses are estimated at 17% by drought, 20% by salinity, 40% by high temperature, 15% by low temperature and 8% by other factors (Ashraf and Harris, 2005).

Water deficit plays a key role in reducing yields of field crops (Jaleel et al., 2007a). Limitation in crop production due to water shortage depends on the intensity and the pattern of drought, which vary from year to year. The efficient use of limited water resources and better growth under limited moisture supply are desirable traits for crops in drought conditions. Crop production is severely constrained by water shortage during the growing period (Jaleel et al., 2007b). In agriculture, water use productivity is associated with the amount of available water (Jaleel et al., 2007c; Jaleel et al., 2007d). Growth parameters showed reduction under salt and water stress condition. Salt stress together with moisture stress also adversely affected the growth rate (Hussain and Ismail, 1994)

Drought tolerance has been observed in all plant species, but its extent varies from species to species. One way to ensure future food needs of the increasing world populations should involve better use of water by the development of crop varieties those require lesser amounts of water and more tolerant to water shortage (Jaleel et al., 2007e).

Plant acclimation to drought has developed certain mechanisms necessary to sustain key physiological processes in plant. Maintenance of plant leaf turgor under deficit soil moisture is an important adaptation trait that attribute to drought tolerance (Hsiao et al., 1976). Plant metabolic processes are more sensitive to turgor and cell volume than to absolute water potential (Jones and Corlett, 1992). The major physiological mechanism to maintain leaf turgor pressure by decreasing osmotic potential is osmotic adjustment (Jones and Turner, 1980). Sunflower exhibits a large varietal difference for osmotic adjustment in response to water shortage (Conroy et al., 1988; Chimienti and Hall, 1994).

Drought has detrimental effects on plant growth and development. Cellular growth appears to be the most sensitive to water deficit. Decreasing the external water potential by only -0.1 M Pa or less might result in a perceptible decrease in cellular growth. The response of cellular growth to water deficit appears as a slowing of shoot and root growth. This is usually followed by a reduction in cell wall synthesis. Protein synthesis in the cell may be almost equally sensitive to drought. The activities of certain enzymes especially nitrate reductase, phenylalanine ammonia lyase decrease quite sharply as water deficit increase. A few enzymes, such as α-amylase and ribonuclease show increase in activities. Such hydrolytic enzymes might break down starches and other material to make the osmotic potential more negative, thereby resisting the drought. Nitrogen fixation and reduction also drop with water deficit, as nitrate reductase activity is reduced. At levels of water deficit that cause observable changes in enzyme activities, cell division is also inhibited. Stomata begin to close leading to reduction in transpiration and photosynthesis (Bradford and Hsiao, 1982).

Reduction in relative water content (RWC) and leaf water potential caused decrease in photosynthesis of higher plants (Lawlor and Cornic, 2002). Drought mainly limits photosynthesis by stomatal closure and through metabolic impairment (Tezara et al., 1999; Lawson et al., 2003). Drought can also affect growth of plant organs, which results in the alteration of the morphology of the plants (French and Turner, 1991). Plants have different mechanisms to cope drought stress such as drought avoidance and drought tolerance. The ability of plants to delay harmful decrease in the water potential of the protoplasm is considered as avoidance of desiccation (Hussain, 2010). Desiccation tolerance means that plants have ability to maintain their normal functions even at low tissue water potentials. Both morphological and physiological diversity have
been developed in plants to tolerate the drought (Blum, 1997).

Sunflower is an adaptive species to available soil moisture. In water deficit conditions, high achene yield can be obtained by increasing total water use and harvest index while reducing soil evaporation and finally placing the crop at low vapor pressure deficit. Water is mostly localized deep in the profile in dry region, so the deep rooted species and cultivars with deep rooting should be used for maximum water utilization (Hoad et al., 2001). Sunflower cultivars deplete available soil moisture differently. Mahal et al. (1998) conducted a trial about the effect of depletion of available soil moisture and nitrogen levels at Ludhiana, Indian Punjab. They reported that application of water at 40% depletion of available soil moisture (ASMD), saved 8.2 cm (12.2%) more water as compared to frequent irrigation (ASMD 20%) with out any adverse effect on growth and yield.

Water deficit in sunflower has adverse effect on yield and its components while oil content is least affected. Seed yield has positive correlation with days to flowering, days to physiological maturity, head diameter, plant height, 1000-seed weight and oil yield under limited and normal irrigation (Razi and Asad, 1999). Water shortage also reduced total dry matter production, leaf area, relative water content, transpiration rate, total number of seeds per head, seed set percentage, grain yield and water use efficiency. Water deficit effect is much pronounced at the vegetative and flowering stage. In sunflower water shortage at the flowering stage reduced yield by 29% (Velue and Palanisami, 2001).

There is evidence that ABA normally plays a role in resistance of mesophytes to water stress. Most studies have been done with drought sensitive cultivar of crop plants. Resistant cultivars have higher levels of ABA when they are exposed to water stress, and sensitive cultivars can be converted to resistant types by exogenous application of ABA (Quarrie, 1980; Cellier et al., 1998). In response to water deficit, ABA begins to increase markedly in plant leaf tissues and, to a lesser extent, in other tissues including roots (Bradford and Hsiao, 1982; Salisbury and Marinos, 1985; Walton, 1980). This leads to stomatal closure and decreased transpiration. It also inhibits shoot growth, and root growth appears to be promoted which increased the water supply. Most of these adaptations involving ABA are best observed in mesophytes; xerophytes often have other adaptations (Kriedemann and loveys, 1974).

Drought, photosynthesis and respiration: Photosynthesis is an important metabolic pathway which is responsible for growth and development. However, photosynthetic efficiency of particular plant is influenced by surrounding environmental conditions (Wise, 1995). The potential of plants to absorb light energy and funnel it to reaction center does not get altered, instead, the capacity of plants to utilize the light absorbed by them declines significantly when they are exposed to environmental stresses such as drought, salinity, low temperature and high light intensity (Dubey, 1997; He et al., 1995). Major effect of drought in plant is reduction in photosynthesis, which is due to decrease in leaf expansion, impaired photosynthetic machinery, leaf senescence and finally reduction in assimilates production (Wahid and Rasul, 2005). Drought stress reduced the crop yield due to changes in photosynthetic pigments (Anjum et al., 2003) and diminished activities of calvin cycle enzymes (Monakhova and Chernyadev, 2002).

The earliest response to drought is closure of stomata at mild to moderate drought. In parallel, a progressive down-regulation or inhibition of metabolic processes leads to decreased RuBP content in leaf under severe drought, and finally inhibits photosynthetic CO₂ assimilation (Flexas and Medrano, 2002). The decreases in CO₂ assimilation and stomatal conductance are due to reduction in carboxylation and RuBP regeneration capacity. Although Rubisco specific activity decreases in severely stressed plant leaves, its content increase under prolonged drought. The increase of Rubisco content in the leaves of the drought-tolerant sunflower hybrid indicating that it could be the one factor which conferring better acclimation and good drought tolerance (Pankovic et al., 1999; Hussain, 2010). Reaction of RuBP to 3-PGA decreased with reducing leaf relative water content (RWC), indicated regeneration of RuBP inhibited under drought stress. Water stressed leaves of sunflower has low RWC and ATP synthesis was known to limit photosynthesis rate because of the inhibition of photophosphorylation (Lawlor, 2002; Tezara et al., 1999).

Relative water content, leaf water potential and osmotic potential of sunflower decreased with water shortage but there were no effects of elevated CO₂ on them. Increased Pn (net CO₂ assimilation rate) led to acclimation, with decreased Rubisco content and increased its activity; drought decreased Pn with metabolic inhibition involving loss of Rubisco activity (Tezara et al., 2002).

Photosynthetic efficiency is reduced by drought stress which leads to photo-inhibition. Most prominent conditions of this phenomena are low and high temperature, osmotic and high light stress (Wise, 1995; Huner et al., 1993). It may also be the result of activity of Reactive Oxygen Species (ROS) such as super-oxide anion radicals (O₂⁻), hydrogen peroxide (H₂O₂), hydroxyl radicals (OH⁻), alkoxy radicals (RO) and singlet oxygen (O₂). When the electrons transferred from the PS- II reaction center molecules in the presence of light exceed the number of electrons accepted by NADP⁺, these are promptly accepted by O₂ through PS- I and lead to the formation of toxic oxygen species such as O₂ and H₂O₂.
Osmotic adjustment under drought: Osmotic adjustment (OA) is a common stress tolerance mechanism in plants accomplished by the production of different compatible organic solutes (Serraj and Sinclair, 2002). Osmotic adjustment (OA) has been considered as an important drought tolerance mechanism in plants which is helpful in postponement of dehydration stress (Morgan, 1990).

These compatible solutes including soluble sugars, sugar alcohols, proline and glycinebetaine are low molecular weight and highly soluble which are nontoxic for plant even at higher cytosolic concentration. They protect plants from cellular dehydration through detoxification of reactive oxygen species (ROS), stabilization of membranes, and structures of enzymes and proteins. Under drought stress, accumulation of compatible solutes occurs in the cell which lowered the osmotic potential and attracts water molecules into the cell and ultimately maintains the cell turgor. Due to OA plant organelles and cytoplasmic activities take place at about normal rate which help plants to perform well in terms of growth, photosynthesis and assimilate partitioning to grain filling (Ludlow and Muchow, 1990; Subbarao et al., 2000).

Glycinebetaine (GB) is accumulated in diverse organisms under environmental stress such as drought, salt, or cold. Glycinebetaine, a naturally occurring quaternary ammonium compound has received great attention as a compatible solute because under water stress it may aid in drought tolerance by maintaining turgor pressure (Agboma et al., 1997). GB has been shown to protect functional proteins, enzymes (e.g. Rubisco), and lipids of the photosynthetic apparatus and to maintain electron flow through thylakoid membranes (Xing and Rajashekar, 1999; Allakhverdiev et al., 2003). Plants with greater ability of glycinebetaine accumulation are more tolerant to water shortage (Monyo et al., 1992).

In field crops accumulation of compatible solutes has been considered as a parameter of selection for stress tolerance. Accumulation of proline can be occurring in plants under stresses such as temperature, drought and starvation. Higher levels of proline enabled the plant to maintain low water potentials (Jalil et al., 2007; Sankar et al., 2007). Plant responses to drought stress include accumulation of ABA and proline which results in speculation that ABA may trigger proline increase in plant cells (Aspinall, 1980; Stewart, 1980).

Water relations of sunflower cultivars: The success of semi-dwarf genes in cereals has prompted scientist to consider height reduction in sunflower. Agronomic benefits of dwarf sunflowers have been reported (Miller, 1992; Johnston et al., 1995). In sunflowers differential water relations response has been demonstrated by different stature genotypes. Dwarf cultivars of sunflower have higher water potential i.e. least water stress as compared to intermediate and long stature cultivars. This is due to difference in canopy architecture and root penetration depth among different stature sunflower cultivars (Angadi and Entz, 2002a). Tall stature cultivars produced more leaf area, aerial biomass and deeper root system so they transpired more water (Angadi and Entz, 2002b). Gimenez and Fereres (1986) have reported a negative correlation between leaf area and leaf water potential. Greater leaf overlap due to compact canopy in dwarf versus standard height sunflower has also been reported by Sadras et al. (1991). Compact crop canopies affect microclimate by reducing wind speed and light penetration by increasing relative humidity (Nobel, 1999), which in turn reduces transpiration loss from the canopy (Pataki et al., 1998). An open canopy, with greater light penetration and better air circulation has potential to increase the plant water status (Nobel, 1999).

Genotypic variation for osmotic adjustment has also been reported in sunflower (Chimenti and Hall, 1994. The dwarf sunflower cultivars have initiated osmotic adjustment earlier than intermediate and large stature cultivars which indicated better osmotic response of dwarf cultivars. The delay in osmotic adjustment in intermediate and tall stature sunflower genotypes is due to faster root penetration and more water extraction from deeper layer of soil. So, tall stature sunflower cultivars
face water stress later as compared to short stature (Angadi and Entz, 2002b). At any rate, dwarf sunflower cultivars are quicker and more efficient in using osmotic adjustment to ameliorate the effects of drought stress (Angadi and Entz, 2002a). The superior osmotic adjustment and lower stress level enabled the dwarf sunflower cultivars to maintain the higher leaf pressure potential (Angadi and Entz, 2002b).

Genotypic variation in rooting depth has been noted in sunflower and deep root system of sunflower cultivars is usually associated with longer growth duration (Fereres et al., 1986; Schneiter, 1992). Short stature cultivars are gaining popularity among farmers due to ease in cultivation and shorter growth duration (Johnston et al., 1995). Standard height sunflower hybrids have shown greater water depletion from soil as compared to dwarf open pollinated cultivars (Schneiter, 1992).

**Drought in relation to crop growth stages:** Sunflower is commonly grown as a rain fed crop in the world but it also responds well to irrigation. Three growth stages of sunflower as heading, flowering and milking are sensitive to water shortage (Doorenbos and Kassam, 1979). In limited irrigation application study, water applied at different growth stages of sunflower significantly decreased seed yield (Stone et al., 1996), particularly at three growth periods of heading, flowering and milking (Demiroren, 1978), at three growth stages (heading, beginning of flowering and end of flowering) (Unger, 1983); and at 50 % ray flower stage (Alessi et al., 1977). In rape seed response of drought stress mostly depend upon genotypes, weather conditions, intensity and duration of drought stress, growth and developmental stages (Robertson and Holland, 2004). Timing of water stress is most crucial than its intensity (Korte et al., 1983).

Normal irrigations are essential for bumper crop production, but when there is scarcity of water, it becomes imperative to differentiate the critical growth stages of the crop, where irrigation could be missed, without reducing the grain yield significantly. In sunflower, irrigation missing at flowering drastically reduced achene yield (Demir et al., 2006) and biological yield (Petchu et al., 2003) as compared to achene yield (El-Tayeb, 2005) and biological yield (De-Guang et al., 2001) obtained by irrigation missing at vegetative stage (Laghrab et al., 2003) on account of reduction in number of achenes head^{-1} (Kadayifei and Yildirim, 2000).

Sunflower irrigated at budding were 19cm taller than those irrigated only at flowering or late flowering. Irrigation at budding stage also favored leaf and stem dry matter production. Irrigation at flowering or late flowering was important for head and seed development. Highest seed and total dry matter yield were obtained with the full irrigation treatment, but they were not always significantly higher than those resulting from less well timed irrigation, namely those at flowering and late flowering (Hussain, 2010). Irrigation treatments affected the rate of oil accumulation in seed and had a relatively small but significant effect on final oil percentage. Linoleic and oleic acid concentrations of oil were little affected by irrigation treatments (Unger, 1983).

Amount and timing of applied irrigation has pronounced effect on growth and yield of sunflower. Limited irrigations as one irrigation, at the “beginning flower stage” (BF) with 686 m^3 ha^{-1} of water or of two irrigations at “flower bud stage” (BF) and at “beginning flower stage” (BF) with 956 m^3 ha^{-1} or at “beginning flower stage” (BF) and at “middle seed growth stage” (MG) with 1155 m^3 ha^{-1}, gave yields of 3.1, 3.3 and 3.5 t ha^{-1} respectively. The achene yield (3.7 t ha^{-1}) obtained with three irrigations (volume 1509 m^3 ha^{-1}), distributed in “FB”, “BF” and “MG” stages were higher than the yield obtained with the lesser irrigations volumes (D’Amato et al., 1999).

Early sunflower growth was limited when the water availability was reduced during budding. Full irrigation was effective for maximum grain production. The corresponding yield increases were 1.4, 1.8 and 2.2 t ha^{-1} as compared with medium inputs or moderately irrigated crop. The best yields (up to 4.0 t ha^{-1}) were observed in dry years under full irrigation. When drought occurred before anthesis, grain yield and achene number m^{-2} were reduced. When drought occurred during the vegetative and flowering period the achene numbers per head decreased. In the absence of stressful environmental conditions during grain filling 3000-12000 achenes m^-2 and potential 1000-seed weight can be obtained. Oil concentration and grain yield increased together while oil and protein concentration were negatively related. So it was concluded that sufficient water is necessary during grain filling to achieve a high oil concentration (Debake et al., 1998).

Supplemental water treatment applied to sunflower during vegetative, reproductive, or both growth stages has pronounced effect on seed yield. Seed yield ranged from 2100 to 2700 kg ha^{-1}, supplemental irrigation increased seed yields by 480 kg ha^{-1} each year (Aiken and Stockton, 2001).

In sunflower irrigation application at heading, flowering and milking (HFM) is the best choice for maximum yield. Goksoy et al. (2004) applied irrigation at 60% heading, flowering and milking (H_{0.60}FM), 40% heading, flowering and milking (H_{0.40}FM), heading, 60% flowering and milking (H_{0.60}FM), heading, 40% flowering and milking (H_{0.40}FM), heading, flowering and 60% milking (HFM_{0.60}) and heading, flowering and 40% milking (HFM_{0.40}). They reported that seed yield and oil yield increases for the limited irrigation treatments were; 78.7 % and 77.4% for H_{0.60}FM; 77.4 % and 78.9% for H_{0.40}FM; 72.2 % and 75% for HFM_{0.60}; 76.4 % and 79.2%
Water deficit has significant effect on development stages of rapeseed. Among six water deficit levels/stages viz. normal irrigation, no irrigation from germination to roset, no irrigation at stem elongation, no irrigation at flowering, no irrigation at siliqua formation and no irrigation at seed filling, the siliqua number was decreased significantly when plants were exposed to water deficit at flowering stage (Rad et al., 2005).

Daneshian et al. (2005a) conducted an irrigation control experiment based on 60mm evaporation from an evaporation basin. Second experiment irrigated when 180mm water was evaporated from evaporation basin. Drought stress started before the reproductive stage. Results showed that water stress reduced seed yield due to reduced head diameter, seed number, seed weight and plant height. The effective seed filling period, harvest index, leaf area index, and seed oil content were also decreased by drought. A reduction in leaf area of sunflower has been observed under continuous 30 day drought period, ending at anthesis than full irrigation (Hammadeh et al., 2005).

Sunflower genotypes performed differently against drought at different growth stages. Maruthi et al. (1998) evaluated six genotypes of different growth period and their performance to rainfall distribution. All cultivars were subjected to moisture deficit at grain filling stage. They found that medium duration cultivars, despite encountering terminal drought, performed better as compared to other maturity groups. When water shortage in sunflower occurred for longer than 12 days at grain filling and flowering stage it reduced seed yield as compared with non-stress plants (Reddy et al., 1998). Seed yield in sunflower is also reduced in high oleic acid hybrids under limited water availability when irrigation was terminated at 100% flowering or 10, 20 and 30 days later. Seed yield was adversely affected when irrigation was terminated at flowering than on later dates (Salera and Baldini, 1998; Singh et al., 1999).

Irrigation interval has prominent effect on growth, development and yield of sunflower. Al-Ghamdi et al. (1991) irrigated sunflower with (a) 5 (b) 10 or (c) 15-days intervals corresponding to 40, 60 and 80% depletion of available soil moisture, respectively. Plant height and seed yield were significantly lower in (c) than in (a) or (b).

The influence of limited irrigation on growth, seed yield and yield components has well documented. Five irrigation treatments were applied, designated as T1 full irrigation and T2, T3, T4 and T5 received 75, 50, 25 and 0% of applications of the fully irrigated treatment on the same day. Seed yields averaged the highest with full irrigation treatment (T1) and differences between full irrigation and other treatments were significant (Erdem et al., 2001). In sunflower when drought occurs during seed filling it reduced seed yield, seed weight per head, 1000-seed weight and head diameter. Seed oil content and oleic acid also reduced but palmitic acid increased under water deficit conditions. (Jasinkas, 1999; Mekki et al., 1999).

In canola 33.3% and 20.7% seed yield reduction were observed when drought stress applied at flowering and siliqua development stage, respectively. This yield reduction due to short term drought during flowering and siliqua development was associated with few siliques per plant and a reduction in seed weight (Sinaki et al., 2007). Drought stress during vegetative or early reproductive stages in soybean reduces seed yield due to reduction in number of seeds per unit area, while water shortage during seed filling reduces seed size (Korte et al., 1983; Brevedan and Egli, 2003).

**Sunflower quality and yield under drought stress:**

Sunflower genotypes have shown differential yield response under drought. Water stress increased the net assimilation rate while it decreased the leaf area ratio in sunflower. Water stress had no effect on relative growth rate. An inverse and negative correlation was found between net assimilation rate, achene yield and oil yield where as relative growth rate had a very low and positive correlation with both achene yield and oil yield. Leaf area ratio had positive and significant correlation with achene and oil yield (Hussain et al., 2008). A decreasing trend in head diameter, number of achenes per head, 100-achene weight and achene yield per plant has been observed as the level of water stress increased from 100% to 25% of field capacity. Seed oil content was sensitive to even mild water stress but showed stability under increasing stress conditions (Khan et al., 2000). Water deficit reduced harvest Index (HI). Excellent positive correlations were found between HI and grain yield under dry land conditions. The reduction in HI due to water deficits was mostly due to adjustments in seed number with less variation in individual seed weight. In turn, the reduction in seed number was due to a combination of reduced head size and of the area having viable seeds (Fereres et al., 1986). Water stress significantly decreased yield and its components; however, oil content did not differ significantly. Genotype and phenotype correlation revealed that seed yield had significant positive correlations with head diameter, plant height, 1000-seed weight and oil yield under normal and limited irrigation. Seed yield had significant correlations with oil content under normal irrigation regime, while these correlations were largely reduced under stress conditions and were not significant. The highest direct effect was exhibited by 1000-seed weight in normal and limited irrigation environments. Direct effect of oil was quite lower than its correlation with seed yield. The lowest direct effect was found for plant height and oil content under normal and stress environments, respectively (Razi and Asad, 1999).
A significant negative correlation of head diameter with fresh shoot weight was observed under water stress. Positive and significant correlation existed between dry shoot weight and seed yield per plant under water stress conditions (Tahir and Mehdi, 2001). Tahir et al. (2002) evaluated inbred lines of sunflower under water stress and normal irrigation. The maximum (34.13%) decrease in yield per plant was observed under water stress as compared with that of under normal irrigation condition followed by 25.56% decrease in leaf area and 22.63% decrease in 100-achene weight. Chimenti et al. (2002) also recorded a significant effect of drought on biomass at the end of anthesis, at physiological maturity, on grain yield and harvest index. Ardakani et al. (2005a) showed non-significant differences for biological yield, seed oil percent among water treatments. Water stress has significant effect on total dry matter, net assimilation rate, relative growth rate, crop growth rate, and leaf area index. Ardakani et al. (2005b) and Nasri (2005) also reported that water stress decreased seed number, seed weight; head diameter, harvest index, oil percent, oil yield and growth indices. Daneshian et al. (2005b) investigated the irrigated treatment as normal condition (60 mm evaporation from pan) and irrigated as stress condition (180 mm evaporation from pan). Normal condition showed the highest yield, high seeds per head and seed weight while oil seed content decreased under irrigated as stress condition.

Sunflower oil quality is affected under different water regimes. The major effect of irrigation was an increase of linoleic acid content and reduction in oleic acid. However oleic/linoleic acid ratio increased under water stress, especially when plants were subjected to soil water stress during elongation stage in standard genotypes (Flagella et al., 2002).

Abscisic Acid (ABA): ABA has been associated as a stress hormone in vascular plants. It has been detected in mosses but absent in liverworts. Several genera of fungi make ABA as a secondary metabolite (Milborrow, 2001). ABA has been detected in every major organ of plant from the root cap to the apical bud and synthesized in almost all cells that contain chloroplast and amyloplasts. It is a 15-carbon compound that resembles the terminal portion of some carotenoid molecules. The orientation of carboxyl group at carbon 2 determines the cis and trans isomers of ABA. Nearly all the naturally occurring ABA is in the cis form, and by convention the name abscisic acid refers to that isomer (Taiz and Zeiger, 2003).

Biosynthesis, metabolism and transport of ABA: ABA biosynthesis takes place in chloroplast and other plastids via the pathway as depicted in figure 2.2. The pathway begins with isopentenyl diphosphate (IPP), the biological isoprene unit, and leads to the synthesis of the violaxanthin. Synthesis of violaxanthin is catalyzed by zeaxanthin epoxidase (ZEP), converted to neoxanthin, which is then cleaved to form the C15 compound xanthenoxal. The cleavage is catalyzed by 9-cis-epoxy-carotenoid dioxygenase (NCED). Synthesis of NCED is rapidly induced by water stress, suggesting that the reaction it catalyzes is a key regulatory step for ABA synthesis. The enzyme is localized in on the thylakoid, where the carotenoid substrate is located. Finally, xanthenoxal is converted to ABA via oxidative steps involved the intermediate(s) ABA-aldehyde and or
possibly xanthoxic acid. This final step is catalyzed by a family of enzymes like aldehyde oxidases that synthesize ABA. Its biosynthesis is not the only factor that regulates ABA concentration in the tissue. The concentration of free ABA in the cell cytosol is also regulated by degradation, compartmentation, conjugation, and transport (Hussain, 2010). For example, cytosolic ABA increases during water stress as a result of synthesis in the leaf, redistribution within the mesophyll cell, and import from the roots and recirculation from other leaves. The concentration of ABA declines after rewatering because of degradation and export from the leaf as well as a decrease in the rate of synthesis.

ABA is inactivated through oxidation, yielding the unstable intermediate 6-hydroxymethyl ABA, which is rapidly converted to Phaseic acid (PA) and dihydrophaseic acid (DPA). Free ABA is also inactivated by covalent conjugation to another molecule, such as a monosaccharide. A common example of an ABA conjugate is ABA-β-D-glucosyl ester (ABA-GE). Conjugation is not only renders ABA inactive as a hormone but also alters its polarity and cellular distribution. Whereas free ABA is localized in the cytosol, ABA-GE accumulates in vacuoles and thus could theoretically serve as a storage form of the hormone. When plants were subjected to a series of stress and re-watering cycles, suggesting that the conjugated form is not broken down during water stress (Taiz and Zeiger, 2003).

ABA is transported by both the xylem and the phloem, but it is normally much more abundant in phloem sap. When radioactive ABA is applied to a leaf, it is transported both up the stem and down toward the roots. Most of the radioactive ABA is found in the roots within 24 hours. Destruction of the phloem by a stem girdle prevents ABA accumulation in the roots, indicating that the hormone is transported in the phloem sap. ABA synthesized in the roots can also be transported to the shoot via the xylem. Whereas the concentration of ABA in the xylem sap of well watered sunflower plants is between 1.0 and 15.0 nM, while in water stressed sunflower plant it increases to as much as 3000 nM (3.0µM) (Schurr et al., 1992).
The magnitude of stress induced change in xylem ABA content varies widely among species and it has been suggested that ABA also is transported in a conjugated form, and then released by hydrolysis in leaves although this has yet to be identified. As water stress begins, some of the ABA carried by the xylem stream is synthesized in roots that are in direct contact with the drying soil. Because this transport can occur before the low water potential the soil cause any measurable change in the water status of the leaves, ABA is believed to be a root signal that helps to reduce the transpiration rate by closing stomata in leaves (Davies and Zhang, 1991).

The 3.0µM ABA concentration in the apoplast is sufficient to close stomata but all of the ABA in the xylem stream does not reach the guard cells. Much of the ABA in the transpiration stream is taken up and metabolized by the mesophyll cells. During the early stages of water stress, however, the pH of the xylem sap becomes more alkaline, increasing from pH 6.3 to about pH 7.2 (Wilkinson and Davies, 1997). Drought induced alkalization of the apoplast favors formation of the dissociated form of abscisic acid, ABA•, which does not readily cross membranes. Hence, less ABA enters to the mesophyll cells and more reaches to the guard cells via the transpiration stream. In this way ABA is redistributed in the leaf without any increase in the total ABA level. This increase in xylem sap pH may function as a root signal that promoted early closure of the stomata (Taiz and Zeiger, 2003).

Abscisic Acid and drought: Plants are under periodic water stress due to unpredictable rainfall and limited availability of irrigation water. They have certain mechanisms which allow them to perceive the incoming stresses and rapidly regulate their physiology and metabolism to cope with these stresses. A good example of such mechanisms is the ability of plants to regulate their water loss through partial closure of stomata, reduced leaf development. It occurs long before loss in leaf turgor and irreversible damage to inner membrane systems (Jones, 1980; Cowan, 1982; Davies and Zhang, 1991). The plant hormone ABA is produced under water deficit conditions and plays a major role in response and tolerance to dehydration (Shinozaki and Yamaguchi Shinozaki, 1999).

ABA plays a critical role in regulating plant water status through guard cells and growth as well as by induction of genes that encode enzymes and other proteins which create cellular dehydration tolerance (Luan, 2002; Zhu, 2002). In plants stomata response to soil drying is mediated by ABA, originating from roots and transported to the shoot via transpiration stream (Davies and Zhang, 1991). ABA under drought is produced in dehydrated roots, transported to the xylem and regulates stomatal opening and leaf growth in the shoots (Zhang et al., 1987; Zhang and Davies, 1990). Stomata respond to the concentration of ABA in the guard cell apoplast (Harris and Outlaw, 1991; Hornberg and Weiler, 1984; Anderson et al. 1994).

In gradual soil drying wilting of plant leaves and gradual stomatal closure are common phenomena. When soil drying prolonged ABA comes from two sources initially from the roots and later from the older plant leaves, which wilted earlier than the younger leaves (Zhang and Davies, 1989b). It appears that plants have evolved two responses to soil drying. Initial soil drying may be sensed by part of the root system and ABA in root xylem regulates stomatal conductance such that water loss in the shoot may be reduced and water deficit can be avoided (Zhang and Davies, 1989a). In severe soil
drying shoot water deficit becomes unavoidable and some older leaves may wilt, due to weak hydraulic link with the main stem or weaker control of the stomatal conductance. This wilting accounts for the accelerated ABA concentration in the xylem and much severe stomatal inhibition in the young leaves (Zhang and Davies, 1989b).

It has been shown that leaf conductance is closely related to xylem ABA concentration (Jia and Zhang, 1999). Closure of plant stomata depends upon the amount of xylem ABA that enters in leaves per unit time i.e. ABA flux (Raschke, 1975; Gowing et al., 1993; Jarvis and Davies, 1997). This implies that ABA is helpful to regulate stomatal movement under normal conditions due to changes in transpiration under non stressed conditions, or that ABA’s role is magnified after the soil drying has led to the very low transpiration. However, studies have strongly suggested that stomatal movement responds to xylem ABA concentration rather than its flux, like water stressed conditions when leaf conductance has decreased to low value (Zhang and Davies, 1991; Jackson et al., 1995; Trejo et al., 1995).

The rapid metabolism of xylem derived ABA in the leaves is essential to prevent ABA accumulation and keep the stomata sensitivity regulated. ABA acts as stress hormone in root systems as it facilitates penetration of roots in compacted soil layers and increase desiccation tolerance in maize (Hartung et al., 1994). Plants growing in arid and semi-arid climate often face some degree of drought stress. Due to which, plants have developed various mechanisms to enhance their tolerance including well-developed root systems, osmotic adjustments and certain leaf morphologies (Levitt, 1980; Kramer, 1983; Jones, 1992; Larcher, 1995; Blum, 1997; Kozlowski and Pallardy, 2002). ABA, a well-known stress induced plant hormone has long been studied as potential mediator for induction of drought tolerance in plants (Davies and Jones, 1991; Zeevart, 1999). It helps in promoting drought tolerance, both from the use of exogenous application to intact plants (Gibson et al., 1991; Heschel and Hausmann, 2001; Xu et al., 2002; Wang et al., 2003) and from the measurement of the endogenous ABA concentration (Van Rensburg and Kruger, 1994; Li and Wang, 2003). Exogenous application of 5 μM ABA to cotton under drought significantly increased seed number and lint mass per plant (Pandey et al., 2003). Jia and Zhang (1999) has also shown 50% decrease in leaf conductance in sunflower detached shoot fed with 5 μM ABA solution for 30 minutes. In soybean, reproductive potential reduced considerably due to less pod set under drought stress. This problem has been alleviated by exogenous application of 0.1 mM abscisic acid (Liu et al., 2004). Population variation due to exogenous application of ABA have been observed in well watered and stressed plants (Amdt et al., 2001; Tsialtas et al., 2001; Ponton, et al., 2001). Exogenous application of ABA affected water use efficiency more in droughty plant than well-watered plants of jewel weed (Heschel and Hausmann, 2001). Plants would grow more rapidly by using a finite water supply more efficiently when water is limited, in this case high water use efficiency would positively affect plant productivity (Li, 1999).

The involvement of drought induced ABA and ethylene in shoot and root growth is still a controversial subject (Robert and LeNoble, 2002). Under drought-stressed conditions, stomata close in response to either a decline in leaf turgor and/or water potential, indicating that stomatal responses are closely linked to soil moisture content and leaf water status. Much is known about the role of ABA in closing the stomata as well as ABA production in dehydrating roots and ABA circulation in the plant (Wilkinson and Davies, 2002).

ABA-induced stomatal closure causes depression in net CO₂ uptake, which involves mechanisms at both the stomatal and chloroplast levels. The mid-day decline in stomatal conductance in several plant species under drought conditions may be due to increased sensitivity to xylem-carried ABA, which is induced by low leaf water potentials (Wilkinson and Davies, 2002). Also, the decline in intercellular CO₂ following stomatal closure apparently induces a down regulation of photosynthetic machinery to match the available carbon substrate. The amount of ABA in xylem sap can increase substantially as a function of reduced water availability in the soil and this might result in an increased ABA concentration in different compartments of the leaf. Zhang and Outlaw (2001) reported that stressing Vicia faba roots could change ABA concentrations at the guard cell apoplast and that the apoplastic guard cell ABA concentration correlated with changes in stomatal aperture more effectively than did the guard cell symplastic fraction. These studies indicate that apoplastically facing guard cell ABA receptors seem to be important in the responses to stress signals experienced by plants. Increases in the xylem sap ABA and leaf ABA were correlated with reduced stomatal conductance under partial root drying conditions in grape vines (Stoll et al., 2000).

Abscisic Acid, drought and crop cultivars: Varietal differences in drought tolerance may be associated with changes in ABA in response to various environmental stresses. This includes drought tolerance of maize (Pekic and Quarrie, 1987), chilling tolerance of rice seedlings (Lee et al., 1993) and salt tolerance of rice (Moons et al., 1995). In wheat, growth of dwarf stature cultivars (Dariel) are less sensitive to constant exogenous application of ABA than that of tall isolines (Bet Hashita) (Blum et al., 1997). Similarly higher accumulation of ABA in drought tolerant wheat cultivars C-306 and HW-24 than susceptible cultivar Hira have been reported (Chandrasekar et al., 2000).
Cellier et al. (1998) studied a model system consisting of a drought tolerant line and a drought sensitive line of sunflower subjected to progressive drought. They described that in both lines exogenous application of 10 μM ABA caused stomata closure within 6 hours. It displayed that stomata of both lines showed similar sensitivity to exogenous application of ABA. It is widely accepted that ABA mediates general adaptive responses to drought. There is also evidence to suggest that additional signals are involved in this process (Munns and King, 1988; Trejo and Davies, 1991; Munns et al., 1993; Griffiths and Bray, 1996). In response to exogenous application of ABA in drought tolerant line of sunflower dehydrin protein accumulated in vegetative tissue (Labhili et al., 1995) and in wheat and poplar (Pelah et al., 1997). Dehydrin protects cytosolic structures from the deleterious effects of cellular dehydration (Baker et al., 1988; Dure et al., 1989; Close, 1996).

ABA has pronounced effects on proline content and water status of different genotypes of sunflower. Unyayar et al. (2004) studied the effect of exogenous application of 10^{-5} M ABA on two sunflower genotypes (Nantio F1 and Ozdemirbey) exposed to drought and water logging. Leaf area and relative leaf water content decreased under drought stress but in ABA treated leaves relative water content had lower values than in non treated leaves of Nantio F1. Dry weight decreased under drought stress while it increased in ABA treated genotype Nantio F1. ABA and proline accumulation increased under water deficit conditions. Exogenous application of ABA increased proline and ABA content in treated leaves as compared to non treated leaves of Ozdemirbey. Genotypic response to ABA in drought is correlated with variation in ABA concentrations (Trejo et al., 1995; Borel et al., 1997; Blum and Sinmena, 1995; Cellier et al., 1998). In Cassava under drought genotypic variation in accumulation of ABA among young, older, expanding and folded leaves have been observed which ultimately affect leaf expansion, growth and transpiration. Under both water shortage and well watered condition young leaves accumulated more ABA than mature leaves (Alfredo and Setter, 2000). In *Euphorbia Lathyris* water stress increased 10-fold ABA concentration in expanding leaves and 5-fold in mature leaves (Sivakumaran and Hall, 1978), while in castor bean many fold increase in mature leaves and 11-fold increase in xylem sap ABA has been observed by Jokhan et al. (1996). This variation in accumulation of ABA in mature and young leaves under water stress is due to increase in rate of catabolism of ABA into biologically inactive products in mature leaves and slow ABA transport in them (Cornish and Radin, 1990). In droughty leaves an accumulation of high level of ABA soon comes to normal level after one day of re-watering (Alfredo and Setter, 2000; Hussain, 2010). This rapid return to normal level of ABA results in recovery of plant growth and development. This behavior contributes to adoption of plants to cope periodic drought followed by renewed water application / rainfall (Connor and Cock, 1981; Palta, 1984). Genotypic variation in accumulation of ABA in field due to drought and other stresses have been observed which reflects gradual depletion of soil water. This genotypic variation also depends upon the extent to which leaf growth is arrested in response to mild and incipient water deficit and ability of crops to retain leaves during drought (Connor and Cock, 1981; Palta, 1984). Crop species like Cassava has ability to partially close the stomata and maintain well photosynthetic rate in low water potential. It is due to rapid accumulation of ABA and halting of leaf growth on the onset of drought and rapid recovery of growth after re-watering (Alfredo and Setter, 2000).

**Conclusion:** Drought is a serious menace to agriculture productivity. Drought stress at divergent growth stages of sunflower hybrids pounced severe damage to growth, development, oil quality and yield. ABA is a well-known stress hormone which improves drought tolerance in sunflower hybrids through osmotic adjustment and improving water relations. It is therefore suggested that screening of sunflower cultivars against drought should be done and exogenous application of ABA will defiantly help in conserving water within plants which is necessary for successful crop production under water deficit conditions.

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