Salinity induced physiological and biochemical changes in chickpea (Cicer arietinum L.) genotypes

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Abstract: Plant growth and development are adversely affected by salinity- a major environmental stress that limits agricultural production. Chickpea (Cicer arietinum L.) is sensitive to salinity that affects its yield and there is need to identify the tolerant genotypes. The present study was conducted to evaluate the effect of salinity on chickpea genotypes with specific physiological and biochemical attributes contributing to their adaptability to salinity stress. Seven chickpea genotypes both desi (ICC8950, ICCV10, ICC15868, GL26054) and kabuli (BG1053, L550, L552) were evaluated for salinity tolerance. Maximum decrease in relative leaf water content and chlorophyll content was observed with ICC15868 and GL26054 among the desi and L552 from the kabuli genotypes. The photosynthetic pigments, activity of nitrate reductase and relative leaf water content was also reduced in response to salt application with effect being more pronounced in ICC15868, GL26054 and L552 as compared to ICC8950, ICCV10, BG1053 and L550. Lipid peroxidation increases with the increase in NaCl concentration, maximum increment was observed in genotypes ICC15868, GL26054 and L552. Accumulation of proline in response to environmental stresses seems to be widespread among plants. Higher protein fractions were observed with tolerant genotypes in contrast to sensitive genotypes. Salt imposed stress finally caused a higher decline in number of filled pods. On the basis of physiological and biochemical parameters genotypes ICC8950 and ICCV10 from the desi genotypes and BG1053 and L550 from kabuli were identified as the tolerant while ICC15868, GL26054 as the sensitive ones and L552 as the moderately tolerant genotypes. These genotypes could be used as a source of tolerance in breeding programme to develop salt tolerant genotypes.

Keywords: Biochemical, Chickpea, Physiological, Salinity

INTRODUCTION

Plants are exposed to wide range of environmental stresses like high temperature, cold, drought, salinity, alkalinity, UV and pathogen infection. Abiotic stress is the primary cause of crop loss worldwide by more than 50 % (Rasool et al. 2013). Salinity is one of the major abiotic stresses causing severe impact on crop production worldwide as out of the world’s 1.5 billion hectares of cultivable land, 77 million hectares (5 %) do not favor good yields due to high salt content and 20 % of the irrigated agricultural land is adversely affected by salinity. Salt-affected soils are increasing steadily in all the continents, in particular in arid and semiarid areas which cover more than 7 % of the total land surface on earth (Selvakumar et al., 2014). More than 800 M ha of arable lands worldwide are affected by salinity, and this area is expanding and is posing as an ever threat to food production (Munns and Tester, 2008).

Soil salinity is becoming more problematic due to the increase in irrigation around the world. Salinity not only decreases the agricultural production of most crops, but also, as a result of its effect on soil physicochemical properties, adversely affects the associated ecological balance of the area. The harmful impacts of salinity include low agricultural production, low economic returns due to high cost of cultivation, reclamation, management, soil erosion due to high dispersibility of soil, ecological imbalance due to halophytes and marine life forms from fresh water to brackish water, poor human health due to toxic effects of accumulated elements (Hu and Schmidhalter, 2002). Resistance to salt stress does not rely on a single trait but, on the contrary, it has a very complex nature as it depends upon various morphological and biochemical traits. The availability of water to the growing tissue becomes a limiting factor under saline conditions even in the presence of moisture in the soil resulting in what is termed as “Physiological Drought”. Water uptake by plants hence, attains importance under saline conditions. The reduction in photosynthesis, respiration and protein synthesis due to salinity reduced crop yield (Meloni et al., 2003). The primary effect of salinity stress is hyperionic and hyperosmotic stress and in severe cases cause oxidative stress in plants (Ahmad et al., 2012). Oxidative stress is responsible for the
generation of reactive oxygen species (ROS) which are deleterious to plants (Azooz et al., 2011, Ahmad et al., 2012). ROS are highly reactive and cause damage to biomolecules such as lipids, proteins and nucleic acids (Tuteja et al., 2009). Proline is considered as the only osmoregulate which has been shown to scavenge singlet oxygen and free radicals including hydroxyl ions. It also serves as redox potential regulator and protects macromolecules such as proteins, DNA and reduces enzyme denaturation caused by heat, NaCl and other stresses (Kumar et al., 2010). Lipid molecules, in general and unsaturated lipids, in particular are sensitive to oxidative damage to cell membranes causing severe oxidative peroxidation (TBARS), a product of lipid presence of elevated levels of thiobarbituric acid reactive species (TBARS), a product of lipid peroxidation (chickpea, Cicer arietinum L.). genotypes will be identified on the basis of physiological and biochemical indices.

MATERIALS AND METHODS

Seven genotypes of chickpea C. arietinum both kabuli (BG1053, L550, L552) and desi (ICC8950, ICCV10, ICC15868, GL26054) were raised in pots with/without holes in six replications. These genotypes having contrasting behaviour for salinity tolerance were selected out of 20 genotypes evaluated earlier by us for seedling growth and germination behaviour under salt stress (30 mM NaCl) in pots. Plants were grown under saline and non saline conditions in 27cm diameter plastic pots containing 10 kg of soil fertilized with fertilizers as per recommended package of practices for chickpea. Pots with holes were used for control conditions and the pots without holes were used for saline treatments. The experiment was carried out in open-air facility equipped with a rainout shelter. The saline treatment at 20 and 30 mM NaCl was applied in split dose: at the time of sowing and 15 days after sowing (DAS) in a sufficient volume to wet the soil to field capacity. Non saline treated controls were irrigated with tap water. In all the treatments seven seeds were sown in each pot and later thinned to 4 plants per pot at 30 DAS. The physiological (chlorophyll content, relative leaf water content, lipid peroxidation) and biochemical parameters (nitrate reductase, proline content) were estimated in leaves of control and salt treated plants at vegetative (65DAS), flower initiation (90DAS) and at pod initiation stages (110DAS). Characterization of proteins and protein profiling was carried out from dry seeds at maturity.

Estimation of chlorophyll: The chlorophyll content from fresh leaves (100 mg) was extracted with dimethyl sulfoxide (DMSO) (Hiscox and Israelstam, 1979). The tubes were kept in water bath at 65°C for 30 min. 1ml of aliquot was mixed with 2ml DMSO and vortexed. Absorbance was determined photometrically at 645 and 663nm using DMSO as blank.

Estimation of relative leaf water content (RLWC): Fresh leaf (100 mg) sample was submerged in 10 ml distilled water till saturation and kept overnight. Surface water of the leaves was blotted off without putting any pressure and then were weighed to obtain saturated weight. After drying the leaves at 70°C for 72 h their dry weight was obtained. From these data RLWC was calculated by Barr and Weatherley method (1962).

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RLWC = \frac{(\text{Fresh weight-Dry weight}) \times 100}{\text{Saturated weight-Dry weight}}
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Estimation of lipid peroxidation: Lipid peroxidation was determined by measuring the amount of...
malondialdehyde (MDA) produced by the thiobarbituric acid (TBA) reaction (Heath and Packer, 1968). Fresh leaves (100 mg) were ground in 1% TCA (10 ml/g fresh weight) with a mortar and pestle and then was centrifuged at 10,000 rpm for 5 min. 1.0 ml of supernatant was taken in a separate test tube, to which 4.0 ml of 0.5% TBA was added. The mixture was heated at 95°C for 30 min and after cooling in an ice bath it was centrifuged at 5,000 rpm for 5 min for clarification. Absorbance was recorded at 532 nm and corrected for unspecified turbidity by subtracting the value at 600 nm. The concentration of MDA was calculated using an extinction coefficient of 155 mM⁻¹cm⁻¹.

**Extraction and estimation of nitrate reductase:**
Nitrate reductase was estimated from the leaves by using the method of Jaworski (1971). Leaves were cut into small pieces (approximately 1mm size) and 0.2 g sample was suspended in 5 ml of 0.1M KNO₃ (in 0.1M phosphate buffer pH 7.5) solution in 25 ml test tube. The tubes were thereafter, incubated in the dark in a water bath for 30 minutes. At the end of incubation period, the tubes were placed in the boiling water bath for 10 min to stop the enzymatic reaction. After boiling, the tubes were cooled and filtered to remove small pieces of leaflets. The clear supernatant thus obtained was used for estimating nitrate reductase. 0.5 ml of sample extract was took out and to it 1 ml of 1% sulphanilamide solution was added to it. After 5 min, 1 ml of 0.02% N-(1-naphthyl) ethylene diamine dihydrochloride was added and pink colored solution obtained after 30 minutes was measured at 540 nm against reagent blank.

**Estimation of proline content:**
The leaf samples (100 mg) after extraction with methanol:chloroform:water (12:5:1) were centrifuged for 10 minutes and the supernatant was collected. The contents were recentrifuged after adding 4 ml of methanol: chloroform : water. Supernatant were pooled and final volume was made upto 10 ml with same solvent. To this 6 ml of chloroform and 4 ml of distilled water was added. After stirring it was allowed to stand for 15 min in separating funnel till the two layers get separated. Lower layer containing pigments was rejected and upper layer was collected. The final volume of upper layer was made 10 ml by adding distilled water and used to estimate proline content by using ninhydrin reagent (Bates et al., 1973). 5 ml of sample solution was taken and 2.5 ml of acid ninhydrin (125 mg of ninhydrin mixed in 3 ml of acetic acid and 2 ml of orthophosphoric acid, and then kept in oven at 70°C till a clear solution was formed) reagent was added to it. The mixture was boiled for 45 min. The tubes were immediately cooled by keeping in ice bucket. Then 5 ml of benzene was added and vortexed in cycloemixture. Two layers were formed and absorbance of upper layer was recorded at 515 nm by using benzene as blank. Proline was used as standard to make standard curve.

**Characterization of protein in seeds:**
The seed samples taken from treated and control plants were powdered in a hand grinder. The four fractions were separated by using Pant and Tulsiani method (1969).

**Albumins:**
The powdered seed sample was extracted in 5ml of distilled water and centrifuged at 10,000 rpm for 30 min. The supernatant obtained contained albumins.

**Globulins:**
The residue obtained was washed, dried and powdered and treated with 5 ml of 0.5 N sodium chloride, then centrifuged at 10,000 rpm for 30 min. The supernatant obtained contained globulins.

**Prolamins:**
The residue obtained after extraction with 0.5N NaCl was washed, dried, powdered and then treated with 5 ml of isopropanol (50%) and centrifuged at 10,000 rpm for 30 min. The supernatant obtained upon this extraction contained alcohol soluble proteins called prolamins.

**Glutelins:**
Extracted with 5ml of KOH(1N) solution followed by centrifugation at 10,000 rpm for 30 minutes. This supernatant contained glutelins.

To the fractions extracted from (1-4), 1ml of 20% trichloroacetic acid (TCA) was added and kept for 24 hours. This extract was centrifuged for 20 min at 5000 rpm and the precipitates so obtained were dissolved in 0.1 N NaOH. Estimation of proteins was done by Lowry et al. (1951) method. SDS-PAGE electrophoresis of proteins was carried out by the method by Laemmli (1970).

**No. of filled pods:**
The total number of filled pods of six randomly selected plants was counted at harvest.

**RESULTS**

**Total chlorophyll content:** It was observed that salt stress (20 and 30mM NaCl) decreased the total chlorophyll content in chickpea genotypes at vegetative (65DAS), flower initiation (90DAS) and pod initiation (110DAS) stages (Fig. 1). The decrease was more pronounced with 30 mM NaCl concentration. The chickpea genotypes showed a variation of total chlorophyll content even under normal conditions. The decrease in chlorophyll content was more pronounced at vegetative stage in ICC15868 and GL26054 (desi) and L552 (kabuli) chickpea genotypes both at 20 and 30 mM NaCl. Similar trend was observed at flower and pod initiation stages. Maximum chlorophyll content was recorded at pod initiation stage. ICC8950 and ICCV10 (desi) and BG1053 and L550 (kabuli) showed minimum chlorophyll degradation both at 20 and 30 mM NaCl salt stress respectively at pod initiation stage. Maximum degradation of chlorophyll content due to salt stress was recorded with GL26054 and ICC15868. It was observed that salt stress (30 mM NaCl) caused more than 50% reduction in chlorophyll content in ICC15868 and GL26054 at pod initiation stage whereas corresponding decrease in ICC8950, ICCV10, BG1053 and L550 was found to be minimum and
L552 showed moderate decrease.

**Relative leaf water content:** Relative leaf water content of chickpea genotypes was significantly reduced with increasing salinity and the magnitude of the reduction varied among genotypes at all the three stages of growth as shown in Fig 2. Salinity-imposed osmotic stress leads to cell turgor loss and cell volume change. The minimum percentage reduction in relative leaf water content at vegetative stage was recorded in ICC8950 and ICCV10 and the maximum percentage reduction was recorded in GL26054 and ICC15868 desi genotypes both at 20 and 30 mM NaCl.
respectively. Among the kabuli genotypes, the minimum percentage reduction was recorded with BG1053 and L550 while the genotype L552 showed the moderate reduction under salt stress both at 20 and 30 mM NaCl respectively. Similar trend was observed in all the seven genotypes at flower initiation and pod initiation stages. It was observed that RLWC in IC8950, ICCV10, BG1053 and L550 was decreased by 11 to 13% and 12 to 14% with 20 and 30 mM salt application respectively at flower initiation stage. The reduction in RLWC was found to be higher in ICC15868 and GL26054 i.e. 38 to 46% in desi genotypes as compared to IC8950 and ICCV10 where reduction was 11 to 14% at pod initiation stage.
Lipid peroxidation: Lipid peroxidation was determined by evaluating malondialdehyde (MDA) contents of the leaf tissues. In our study, MDA concentration increased at all levels of NaCl treatments when compared to control indicating that salinity induces oxidative stress (Fig 3). The maximum increase in MDA level was observed at 30mM NaCl in all the genotypes but less increment was noticed in ICC8950 and ICCV10 among the desi, while among the kabuli genotypes BG1053 and L550 showed the similar results at all stages of growth and under the salt stress also. The maximum increment in MDA level was noted in genotypes ICC15868 and GL26054 among the desi, and L550 among the kabuli genotypes under both 20 and 30mM NaCl application.

Nitrate reductase: The enzyme activity was measured at the three growth stages viz. vegetative, flower initiation and pod initiation stage. The maximum enzyme activity was observed at 65 DAS (Fig 4). BG1053 and L550 in kabuli genotypes and ICC8950 and ICCV10 in desi genotypes had the highest enzyme activity was while GL26054 and ICC15868 marked the lowest activity in control. Salt stress decreased the activity in all the cultivars at all the stages but the decrease was less in BG1053, L550, ICC8950 and ICCV10 whereas the maximum reduction was observed in GL26054 and ICC15868 as compared to other genotypes. Similar trend was observed for all the genotypes at flower and pod initiation stages.

Proline content: The content of proline increased up to flower initiation stage thereafter decreased (Fig. 5). The proline content was found to be maximum in BG1053 (2.79 mg g⁻¹ DW), L550 (2.60 mg g⁻¹ DW), ICC8950 (2.67 mg g⁻¹ DW) and ICCV10 (2.70 mg g⁻¹ DW) while the minimum proline content was recorded with GL26054 (1.04 mg g⁻¹ DW) and
Fig. 9. Comparative SDS-PAGE protein profiles of total proteins in tolerant chickpea genotypes (ICC 8950 and ICCV 10) under control and salt stress. Lanes: M- Page Ruler TM (Prestained protein ladder); Lane: 1- ICC 8950 (Control), Lane: 2-ICC 8950 (30mM), Lane: 4- ICCV 10 (Control) Lane: 5-ICCV 10 (20mM); Lane: 6-ICCV 10 (30mM).

Fig. 10. Comparative SDS-PAGE protein profiles of total proteins in sensitive chickpea genotypes (ICC15868 and GL26054) under control and salt stress. Lanes: M- Page Ruler TM (Prestained Protein Ladder); Lane 1:ICC15868 (Control), Lane 2: ICC15868 (20mM), Lane 3: ICC15868 (30mM), Lane 4: GL26054 (Control), Lane 5: GL26054 (20mM), Lane 6: GL26054 (30mM).

ICC15868 (1.10 mg g⁻¹DW) in control at vegetative stage. Salt stress at 20 and 30mM NaCl, increased the proline content in BG1053 (4.23, 4.78 mg g⁻¹DW), L550 (3.80, 4.25 mg g⁻¹DW), ICC8950 (4.14, 4.67 mg g⁻¹DW), ICCV10 (4.13, 4.56 mg g⁻¹DW) respectively. The minimum increase in proline content was observed with GL26054 (1.15, 1.20 mg g⁻¹DW) and ICC15868 (1.22, 1.31 mg g⁻¹DW). During flower initiation and pod initiation stages, the maximum proline accumulation was recorded with BG1053, L550, ICC8950 and ICCV10 under non saline conditions. Under the salt stress of 20 mM and 30 mM NaCl the maximum value of proline accumulation was recorded with BG1053 and L550 from the kabuli chickpea genotypes, from the desi genotypes ICC8950 and ICCV10 and the minimum accumulation was noticed in GL26054 and ICC15868 respectively.

Protein characterization: The data on effect of salt stress on total protein and different protein fractions (albumins, globulins, prolamins and glutelins) in seeds of chickpea genotypes are shown in Figs. 6 and 7. The perusal of data reveals that salt stress increased the total protein content as well as different protein fractions in all chickpea genotypes. Maximum increase in protein content was observed in ICC8950, ICCV10, BG1053 and L550 with 30 mM NaCl. Protein content in ICC15868, GL26054 and L550 was increased to a very small amount with salt stress. Globulins content
was found to be maximum and prolamin content was found to be minimum in chickpea genotypes.

**Number of filled pods per plant:** Number of filled pods is one of the important yield contributing attribute and the data pertaining to its significant percentage reduction under salinity is presented in Fig. 8. Higher salinity level (30 mM NaCl) caused 14% decrease in number of filled pods of in BG1053, L550, ICC8950 and ICCV10 and about 46% decline in GL26054, ICC15868.

**Protein profiling (SDS-PAGE):** Fig. 9 represents the comparative SDS-PAGE protein profiles of total proteins in tolerant chickpea genotypes (ICC8950 and ICCV10) under control and stress. Bands of molecular weight greater than 96 KDa are present in ICCV10 and absent in ICC8950 under control and salt stress. Bands of molecular weight 16 KDa are slightly more dense under salt stress than control. No significant variation in protein bands was observed under stress in genotypes ICC8950 and ICCV10 at both the stress concentrations. Fig. 10 represents the comparative SDS-PAGE protein profiles of sensitive genotypes i.e. ICC15868 and GL26054 under control and salt stress. Results were found to be non significant in both the genotypes under salt stress. Fig 11 revealed the SDS-PAGE protein profiles of total proteins in *kabuli* chickpea genotypes viz. BG1053, L552 and L550 under control and salt stress. BG1053 and L552 showed upregulation of proteins under salt stress as compared to their control. L550 showed a non significant result i.e no variation among the different bands of proteins was observed.

**DISCUSSION**

Chickpea genotypes subjected to different concentrations of salt stress showed a decrease in photosynthetic pigments. Salinity stress had negative impact on chlorophyll content. It was observed that BG1053 and L550 (*kabuli*) and ICC8950 and ICCV10 (desi) showed the minimum decrease in chlorophyll content under salt stress whereas ICC15868 and GL26054 showed maximum decrease in chlorophyll content under both applications of salt stress. The minimum degradation of chlorophyll content in chickpea genotypes under salt stress indicates their better photosynthetic ability. The decline in photosynthesis due to salinity stress could be due to lower stomata conductance, depression in carbon uptake and metabolism, inhibition of photochemical capacity, or a combination of all these factors (Mundree et al. 2002).

Photosynthetic pigments decreased in chickpea grown under salt stress (Beltagi 2008) and photosynthesis was reduced to 60% (Murumkar and Chavan 1993). The inhibitory effects of salt stress on chlorophyll pigments could be due to suppression of specific enzymes responsible for the synthesis of the green pigments, or due to increased chlorophyllase activity in wheat, *Catharanthus roseus* and mustard respectably (Kiani et al. 2005; Mishra et al. 2006).

Water uptake by plants hence, attains importance under saline conditions. To overcome the external stress plant cells tend to re-adjust their osmotic potential to prevent water losses that can be achieved either by uptake of inorganic ions or by *de novo* synthesis of compatible solutes. In the present investigation genotypes having minimum chlorophyll degradation were found to have minimum reduction in RLWC. This indicates that these cultivars had the capacity to synthesize photo-assimilates under stressed conditions that could be directed for pod and seed setting under stressed conditions. Kabir et al. (2004) also reported that salinity decreased RLWC and water retention capacity, while increased water saturation deficit and water uptake capacity in mung bean.
The level of lipid peroxidation was measured by estimating malondialdehyde content (MDA). MDA has been widely used as a selection criterion to assess salt injury in various plants (Katsuhara et al., 2005; Jaleel et al., 2007). Results showed that the maximum increase in MDA level was observed in ICC15868 and GL26054 genotypes having maximum chlorophyll degradation and reduction in RLWC under the salt stress indicating that they are more prone to oxidative stress, which is responsible for the generation of ROS that are deleterious to plants and cause damage to biomolecules such as lipids, proteins and nucleic acids (Apel and Hirt, 2004; Tuteja et al., 2009). Salt stress negatively affect the lipid peroxidation and has been reported in many plants, Brassinica juncea (Ahmad et al., 2012), Vicia faba (Azooz et al., 2011) and Solanum tuberosum (Queiros et al., 2011).

Nitrate reductase being an important enzyme of nitrate assimilation and plays an important role in amino acid biosynthesis and protein synthesis. It was observed that salinity decreased the NR activity in ICC15868 and GL26054 genotypes. Decline in nitrate reductase activity under conditions of salinity may be due to enhanced degradation of nitrogen, the inhibition of nitrate reductase activity due to salt ions, or a reduced rate of enzyme synthesis due to the salinity. Results showed increased accumulation of proline under salt stress in chickpea genotypes. The maximum proline was accumulation was observed in ICC8950, ICCV10, BG1053 and L550. Proline is a compatible osmolyte, and perform multiple functions in stress adaptation, recovery and signaling, stabilization of proteins and protein complexes in the chloroplast and cytosol and protection of the photosynthetic apparatus in plants (Szabados, and Savoure, 2009). Ashraf and Foolad (2007) suggested that the application of proline successfully improved stress tolerance in plants. Salinity decreased the number of filled pods per plant, however the reduction in number of filled pods was associated with an increase in pod abortion in salt sensitive chickpea genotypes. However, pollen viability, *in vitro* germination of pollen, or pollen tube growth were not affected by salinity in salt tolerant and sensitive chickpea genotypes under 40mM NaCl (Turner et al., 2013). The increase in protein content may be due to the fact that salinity enhances protein synthesis (Langdale et al., 1973) and promotes conversion of nitrogen into protein (Helal et al., 1975). Significant increase in seed protein content under saline conditions has also been reported by Ashraf (1989) in mashbean cultivars. It can be inferred that salinity activates the metabolism of plants and hence the accumulation of more immediate metabolites, by the cells. A higher content of soluble proteins has been observed in salt-tolerant than in salt-sensitive cultivars of wheat and rice (Maleki et al., 2014; Hakim et al., 2014). Protein accumulation is important for cell survival under salt stress as it causes membrane stabilizations. In our study, salt-tolerant genotypes had the highest proteins than salt-sensitive genotypes. In response to salt stress, plants make new proteins that help them to grow and develop under saline conditions. Salt-tolerant genotypes producing higher protein concentrations is due to higher efficiency of osmotic regulation mechanism in these plants which in turn causes decreasing sodium toxicity in cytoplasm compared to susceptible ones and the result is to prevent protein reduction under salt stress (Flowers and Yeo, 1995).

**Conclusion**

Seven chickpea genotypes both desi (ICC8950, ICCV10, ICC15868, GL26054) and kabuli (BG1053, L550, L552) were evaluated for salinity tolerance on the basis of physiological and biochemical parameters. Salinity inhibited photosynthetic efficiency by chlorophyll degradation and nitrogen assimilation in different genotypes of chickpea. On the basis of physiological and biochemical parameters genotypes ICC8950 and ICCV10 from the desi and BG1053 and L550 from kabuli were identified as the tolerant while ICC15868, GL26054 as the sensitive ones and L552 as the moderately tolerant genotype. These tolerant genotypes had minimum reduction in chlorophyll content, relative leaf water content and MDA level accompanied by higher accumulation of proline and total proteins as compared to susceptible cultivars and the cumulative affect of these changes lead to amelioration of salinity stress tolerance in them.

**REFERENCES**


