High temperature induced changes in quality and yield parameters of tomato (*Solanum lycopersicum* L.) and similarity coefficients among genotypes using SSR markers

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**ABSTRACT**

High temperature induced by climatic fluctuations are an important threat for plant growth, development and quality of agricultural produces. Adaptableness to environmental changes generally derives from a large set of genetic traits affecting physio-morphological, biochemical and agronomic parameters. Therefore, the identification of genotypes with higher yield and good quality parameters at high temperatures is becoming increasingly necessary for future breeding programs. Here, we analyzed the performance of different tomato genotypes grown under elevated temperatures in terms of yield and nutritional quality of the fruit. High temperature stress was induced from flower initiation to maturity stage by keeping the pots in a temperature controlled green house facility for 45 days. The quality and yield parameters were taken at the harvesting stage. Starch and soluble sugar concentration in the leaves of tomato genotypes showed significant reduction in its amount under heat stress. Titrable acidity (TA), total soluble solids (TSS) and ascorbic acid content of tomato fruits were highest under high temperature conditions compared to ambient condition but lycopene content decreased with rise in temperature. The yield attributes viz., number of fruits/plant, fruit set %, average fruit weight (g), yield per plant (g/plant) were significantly lower for Arka Saurabh, Arka Rakshak and Pusa Rohini when compared to other genotypes under study. Molecular characterization of selected 22 tomato genotypes were assessed using 25 simple sequence repeat (SSR) markers. Phylogenetic tree was constructed by the unweighted neighbour-joining method (UPGMA) using NTSYSpc cluster analysis software. The Jaccard’s similarity matrix was constructed using the SIMQUAL method using UPGMA algorithm in NTSYSpc. Jaccard’s similarity matrix among these tomato genotypes ranged from a minimum of 0.22 to a maximum of 1 with an average genetic similarity of 0.67. Hence this study has importance in identifying genotypes that could maintain good quality and higher yield under high temperature condition.

1. Introduction

High temperature is one of the major abiotic stress affecting plants, having adverse effects on both growth and reproduction (Beena, 2013). Its impact on agriculture is severe by affecting the productivity of crop negatively. The global average temperature has increased by 0.6 °C over the past 100 years and is projected to increase at a rapid rate in future (Root et al., 2003). The average increase is expected to be 0.5–2.8 °C by the end of the 21st century (Meibl et al., 2005; Vuuren et al., 2008; Beena et al., 2018b). Tomato (*Solanum lycopersicum* L.) is considered as an important vegetable crop native to South America. The genus *Solanum* includes annual or short-lived perennial herbaceous plants. It is a typical day neutral plant and is mostly self-pollinated crop. It is an excellent source of carotenoids, vitamins, antioxidants, lycopene and lutein. The limited caloric supply, relatively high fibre content and presence of minerals, vitamins and phenols such as flavonoids make the tomato fruit an excellent “functional food” providing many physiological benefits and basic nutritional requirements. The recent scenario of global warming affected agricultural production and productivity (Ainsworth and Ort, 2010) and the most essential goal of plant breeders should be to develop high yielding varieties that are resistant to biotic and abiotic stress factors. Studies are conducted to evaluate the performance of tomato cultivars for heat tolerance at reproductive phase. Under high temperature condition, reproductive phase is particularly sensitive to...
continuous mild heat (CMH; Kinet and Peet, 1997). Male sterility and the position of the stigma relative to the anther cone seem to be major factors limiting fruit and seed set (Dane et al., 1991; Levy et al., 1978). Tolerant genotypes were identified under high temperature condition (Levy et al., 1978; Dane et al., 1991; Sato et al., 2000, 2004; Bhattarai et al., 2016). High temperature stress changes the physiological and biochemical responses in plants (Camejo et al., 2005; Min et al., 2014) which later on decreases crop quality and its yield. However, the susceptibility of plants to high temperature differs according to genotypes and also the developmental stages (Wahid et al., 2007). Variation in the response of cultivars to high temperature stress is not only in the vegetative organs (Camejo et al., 2006) but also in the reproductive organs (Firon et al., 2006).

Genetic diversity analysis exclusively based on phenotypic traits may not be a reliable measure of genetic differences as they are influenced by environmental factors (Shehzad et al., 2009). Thus, DNA based molecular markers such as RAPD (Random Amplified Polymorphic DNA), SSR (Simple Sequence repeat), AFLP (Amplified fragment length polymorphism) and SNP (Single Nucleotide Polymorphism) have been routinely used to assess the genetic divergence among the genotypes as they are not influenced by environmental factors. Multi-allelic nature and high polymorphism of SSR markers help to establish the relationship among the individuals even with less number of markers. SSR markers are preferred as they are abundant in the genome, well-distributed throughout the genome, hyper-variable, multi-allelic and co-dominant nature, ease of assaying, highly reproducible and highly informative markers are immensely valuable in studies of variation detection, diversity analysis, phylogeny, population structure, gene mapping and association studies (Beena et al., 2012b; Ditta et al., 2018). The knowledge of the extent of genetic variation, diversity and genetic relationships between genotypes of the crop is vital and foundation for developing an improved cultivar possessing high yield, good fruit quality and adapted to various abiotic and biotic stresses situations (Sheshshayee et al., 2011). Knowledge of the genetic diversity of tomato genotypes is useful for core collection development and effective conservation strategy. The Jaccard’s similarity matrix can be constructed using the SIMQUAL method using UPGMA algorithm in NTSYSpc. UPGMA is perhaps the most widely used techniques which is the only method to be suggested if group averages are obtained (Aldenderfer and Blashfield, 1978). Even though the unweighted pair group approach using arithmetic averages (UPGMA) and neighbour-joining (NJ) algorithms is intended to generate single trees, depending mostly on order of data entry, they can derive more than one topology from a single matrix (Stefan Van Dongen and

Winnepenningcx, 1996). The UPGMA dendrogram was designed on the basis of similarity indices that demonstrated distinct clustering into groups of different genotypes (Punia et al., 2009). Thus in this study an attempt has been made to (a) evaluate the performance of 22 tomato genotypes for changes in quality and yield traits under high temperature condition. (b) calculate the similarity coefficients among genotypes using SSR markers.

2. Materials and methods

2.1. Planting material

Planting material used in this study is the cultivated tomato varieties released from various states of India and germplasm collections from Indian Council of Agricultural Research-Indian Institute of Horticultural Research, Bangalore (Table 1).

2.2. Methodology

This experiment was conducted at College of Agriculture, Vellayani, Kerala Agricultural University, 8.4316° N, 76.9860° E. Tomato seeds were obtained from NBPGR (substation), Thrissur. Tomato seeds were sown in germination tray (60 cells in one tray of size 54 × 35 × 5 (L x W x H) in cm) and filled with potting mixture (coir pith compost and vermicompost @ 2:1 ratio) and labelling was done properly. Irrigation was provided regularly using a rose can. The one month old seedlings were transplanted to pots (30cm height, 20 cm diameter) with potting mixture made from loamy soil of pH 5.8, sand and cow dung on equal volume by volume basis. Six replications were maintained for a single variety. The plants were grown in natural, outdoor environment conditions in a wired enclosure (32.1/24 ± 1 °C mean day/night temperature, 1350–1550 μmol m⁻² s⁻¹ light intensity, 60–65% relative humidity) until flower initiation. Five plants per each replication were maintained. The experiment was laid out in completely randomized design with two treatment levels i.e. control and high temperature stress (36±2 °C) with three replications each. 20 days after transplanting, a set of 22 genotypes with three replicates were transferred to temperature controlled greenhouse for heat stress induction. The average maximum and minimum air temperatures for control condition during crop growth period was 32.1 °C and 24 °C and the average maximum and minimum relative humidity (RH) of air was 90.6% and 59.2% respectively. The daily temperatures including maximum and minimum temperatures were recorded under control as well as heat stress conditions using digital thermo-hygrometer throughout the experiment. Quality parameters and yield parameters were taken at harvesting stage.

2.3. Quality parameters

The carbohydrate content in plants was estimated by Phenol-sulphuric acid method (Dubois et al., 2002). The starch content in plant leaves was estimated by Anthrone method (Hodge and Hofreiter, 1962). Glucose content in the sample was calculated using the standard graph.

The lycopene content in the fruit was quantified by the method explained by Rangana (1976). Optical density (OD) of the extract was measured at 503 nm in UV-VIS-spectrophotometer (Elico SL-160) using petroleum ether as a blank. Lycopene content of the sample was calculated by using the following formula:

\[
\text{Absorbance (1 unit)} = 3.1206 \times \text{O.D. of sample} \times \text{volume made up} \times \text{dilution} \times 100 \div \text{weight of sample} \times 1000
\]

Lycopene (mg 100g⁻¹) = (3.1206 × O.D. of sample × volume made up × dilution × 100) / (weight of sample × 1000) (1)

Titration method was used to estimate titrable acidity (AOAC, 2000). Five tomatoes from each genotype were homogenized in a mixer to a fine puree. Five grams of homogenized tomato puree was extracted with distilled water and made up the volume to 50 mL. Ten mL of filtrate was titrated against 0.01 N NaOH using a drop of phenolphthalein indicator. Acidity was calculated as using citric acid as standard equivalents and expressed as percent of acidity.
Titratable acidity = Volume of NaOH used (mL) / Volume of juice taken (mL) 
\times 0.0064 \times 100 \tag{2}

TSS in terms of Brix units was measured in fresh tomato juice using a digital refractometer (Model DG-NXT, ARKO India Ltd).

The ascorbic acid content in plants was estimated volumetrically by the method explained by Sadasivam and Manickam (2008). Working standard solution of 5ml containing 100 μg/ml of ascorbic acid was pipetted out into a 100 ml conical flask. 4% oxalic acid was added to it and titrated against 2, 6- dichlorophenol indophenol dye (V1 mL). End point was noted on appearance of pink colour which persisted for a few minutes. The sample (0.5g) was weighed and ground in a mortar with pestle using 15ml 4% oxalic acid. The homogenate was filtered through a double layered cheese cloth. The filtrate was made up to a known volume and centrifuged at 10,000 rpm for 10 min. The supernatant was collected and made up to 25ml using oxalic acid. 5.0 ml aliquot was pipetted into a conical flask to which

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**Table 1. List of twenty-two tomato genotypes used for the study.**

| Sl. No. | Varieties            | Released from                     |
|---------|----------------------|-----------------------------------|
| 1       | Nandi                | UAS & AVRDC                       |
| 2       | IC-45                | IIHR collections                  |
| 3       | Pusa Rohini          | IARI                              |
| 4       | Pusa Ruby            | IARI                              |
| 5       | IIHR-2200            | IIHR collections                  |
| 6       | Anagha               | KAU                               |
| 7       | Akshaya              | KAU                               |
| 8       | Vellayani Vijay      | KAU                               |
| 9       | Arka Vikas           | IIHR                              |
| 10      | Kasvi Vishesh        | ICAR-IIHR                         |
| 11      | Vaibhav              | UAS & AVRDC                       |
| 12      | IIHR-26372           | IIHR collections                  |
| 13      | Palam Pride          | HPAU                              |
| 14      | Arka Abha            | IIHR                              |
| 15      | Arka Alok            | IIHR                              |
| 16      | Manulakshmi          | KAU                               |
| 17      | Sakti                | KAU                               |
| 18      | Manuprabha           | KAU                               |
| 19      | Arka Samrat          | IIHR                              |
| 20      | Arka Sourabh         | IIHR                              |
| 21      | PKM-1                | TNAU                              |
| 22      | Arka Rakshak         | IIHR                              |

UAS - University of Agricultural Sciences, Bangalore.
AVRDC - Asian Vegetable Research and Development Center.
ICAR-IIHR - Indian Institute of Horticultural Research, Bangalore.
ICAR-IARI – Indian Agricultural Research Institute, New Delhi.
KAU – Kerala Agricultural University, Thrissur.
ICAR-IIHR – ICAR- Indian Institute of Vegetable Research, UP.
HPAU – Himachal Pradesh Agricultural University, Solan.
TNAU - Tamil Nadu Agricultural University, Coimbatore.

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Figure 1. Gel profile with DNA bands of tomato (Lane 1-Manuprabha, Lane 2-Akshaya, Lane 3-Pusa Ruby, Lane 4-IC 45, Lane 5- Nandi, Lane 6-IIHR 2200, lane 7-IIHR 26372, lane 8-Palam Pride, lane 9-PKM 1, lane 10-Manulakshmi, lane 11-Arka Sambrat, lane 12-Rakshak, lane 13-Arka Vikas, lane 14-Pusa Rohini, lane 15-Arka alok, lane 16-Sakthi, lane 17-Vaibhav, lane 18- Vellayani Vijay, lane 19-Anagha, lane 20-Kashi Vishesh, lane 21- Arka Sourabh, lane 22-Arka Abha).
10ml of 4% oxalic acid was added. This was titrated against dichlorophenol indophenol (DCPIP) solution until the appearance of pink colour (V2 mL). The amount of ascorbic acid is calculated as follows:

\[
\text{Ascorbic acid} = \frac{0.5 \text{mg}}{V_1 \text{ml}} \times \frac{V_2 \text{ml}}{5 \text{ml}} \times \frac{100}{\text{weight of sample}}
\]

(3)

2.4. Yield parameters

The height of plants was measured from the base of the stem to the tip of the shoot at harvest stage, and the average height was calculated on per plant basis and expressed in cm.

Fruit set was also expressed in percentage by counting the total number of flowers as well as total number of fruits per plant.

Table 2. List of primers along with their sequence, chromosome number and expected product size.

| Sl. no. | Chromosome | Primer | Sequence | Expected product size | Reference |
|--------|------------|--------|----------|-----------------------|-----------|
| 1      | 1          | SSR 134| F: CCCTCTTGCTAAACATCCA R: CGTTGGAATCAGGATGTG | 171       | Wen et al. (2019) |
| 2      | 1          | SSR 270| F: AGCTCAAGGCTGTGTTG R: AACACACGTGCAGCTCAT | 231       | Wen et al. (2019) |
| 3      | 2          | SSR 96 | F: GGTTTATGAGCAATGGTGG R: CCTTTATGTCAGCCGGTGT | 222       | Wen et al. (2019) |
| 4      | 2          | SSR 47 | F: CCTCTCAAGAATGAAAGCTCGTG R: CCTTGGGATACCAACACCAA | 191       | Wen et al. (2019) and Khan et al. (2020), Kumar et al. (2016) |
| 5      | 2          | SSR 276| F: TCCGGCAAGAGTGAACATT R: CGAGGGAGTAACTGGGATT | 148       | Wen et al. (2019) |
| 6      | 3          | SSR 304| F: CCTTCGCGGTTACTCGCAC R: TTGCAGCTTCAGCCGGTCG | 186       | Wen et al. (2019) |
| 7      | 3          | SSR 63 | F: CCACAAACATTCATCCTCA R: GTCTGGGATACCAACACCAA | 250       | Wen et al. (2019) and Khan et al. (2020), Kumar et al. (2016) |
| 8      | 4          | SSR 4  | F: TTCTTGGGAGCAGAGGGTTA R: CCAATACAGCAGACCCGAT | 166       | Kaushal et al. (2017) |
| 9      | 4          | SSR 293| F: GAAAGAGTGGAGTCTTGCA R: TTCTGACACTGGGATAC | 129       | Wen et al. (2019) |
| 10     | 5          | SSR 10 | F: TCTTGGCAGAGCGTTTGC R: GGGAGATGCCACATCACATA | 249       | Khan et al. (2020), Kumar et al. (2016) |
| 11     | 5          | SSR 115| F: CACTTCTTTTATCTCAAGAGC R: ATGGAGGTGATGCAACAGCC | 211       | Kumar et al. (2016) |
| 12     | 6          | SSR 19 | F: GCTTCCCGACACCACTCATC R: GAGGAGTGGAGTCTTGCA | 188       | Khan et al. (2019) and Khan et al. (2020), Kumar et al. (2016) |
| 13     | 6          | SSR 293| F: GAAAGAGTGGAGTCTTGCA R: TTCTGACACTGGGATAC | 129       | Wen et al. (2019) |
| 14     | 7          | SSR 248| F: GATCGGTTGATGTCGGTTT R: GGGAGATGCCACATCACATA | 249       | Khan et al. (2020), Kumar et al. (2016) |
| 15     | 7          | SSR 124| F: TCAATCCCATAGACCGGTTA R: GAGGAGAAGACCCAGGCAA | 131       | Dhaliwal et al. (2011) |
| 16     | 8          | SSR 70 | F: TCTTCTGTGTTGTTGTTG R: GGGTGGATGATGGGATAG | 120       | Dhaliwal et al. (2011) |
| 17     | 8          | SSR 111| F: TTCTTCTTTTATCTCATGCT R: TTGAGTGGAGTCTTGCA | 188       | Khan et al. (2020) and Kaushal et al. (2017), Kumar et al. (2016) |
| 18     | 8          | SSR 20 | F: GAGGAGGCGAACAACACAGGA R: GACATGGCGATTAGTACCAA | 157       | Khan et al. (2020), Kumar et al. (2016) |
| 19     | 9          | SSR 602| F: GGGTACATGACACATCTAAGGA R: GCCAATGATACCGGATCGT | 299       | Kwon et al. (2009) |
| 20     | 9          | SSR 450| F: AAATGAAAGACCGAATCAGC R: CATAGGCGCCAGAATGAC | 265       | Kwon et al. (2009) |
| 21     | 10         | SSR 341| F: TTCTCTGGGTTGGGCAAT R: AAGCCCGGATCTTGGTACG | 292       | Khan et al. (2020) |
| 22     | 10         | SSR 331| F: GCATTGATGACACCATCAGC R: ATGATGCTTTGGTCG | 178       | Khan et al. (2020) |
| 23     | 11         | SSR 80 | F: GCAAATGCTTAAGAAGGATG R: AGGCTGATCTGTGGTACGA | 180       | Benor et al. (2008) |
Fruit setting% = (Total number of fruits / Total number of flowers) × 100

(4) Average fruit weight was calculated by adding the weight of fruits from each of three replication plants at harvest and divided it by total number of fruits and expressed in grams per fruit.

The weight of all the fruits collected per plant was taken and the total yield was calculated at the harvesting stage.

2.5. DNA isolation

Twenty-two genotypes of tomatoes were used for the present study. The leaf samples were obtained from one month old plant samples, and genomic DNA was isolated using CTAB method as defined in the procedure by (Murray and Thompson, 1980). The genomic DNA isolated from 22 varieties of tomatoes were validated using agarose gel electrophoresis.

Quality was assessed by using gel electrophoresis with 5μl of crude DNA sample on agarose gel (0.8%) and stained with ethidium bromide. After electrophoresis, the gel was visualized under UV trans-illuminator and photographed with gel documentation system. The observations on the intactness of bands of DNA samples were taken which revealed the quality of the DNA (Figure 1).

2.6. PCR amplification using SSR primers

Twenty five SSR primers were randomly selected and tested on isolated genomic DNA of Solanum lycopersicum L. (Table 2). These primers are associated with high temperature stress traits and sequences were obtained from the Sol Genomics Network (SGN, http://solgenomics.net/) database. PCR reaction was performed in a 20μl reaction mixture which consisted of, 2.0 μl of genomic DNA with quantity 25ng/μl, 2.0 μl of 10X Taq assay buffer A, dNTPs mix (10mm each) of about 1.5 μl, 0.3 μl of Taq DNA Polymerase (1U). 0.75 μl of Forward and reverse primer (10pM) respectively and 12.7 μl of autoclaved distilled water (Kaushal et al., 2017).

2.7. Detection of polymorphism among tomato genotypes using SSR primers and analysis of similarity coefficient

Twenty five primer combinations were screened. The documented SSR profiles were carefully examined for the polymorphism in banding pattern among the genotypes. Markers were scored according to the standard protocol using binary codes. Banding patterns were scored for absence (0) and presence (1) of bands.

The amplified gel pictures obtained from twenty five SSR markers were scored. The binary data generated for all the varieties for the polymorphic markers was entered in the NTedit program of NTSYSpc version 2.10 software (Jhamshidi and Jhamshidi, 2011).

2.8. Statistical analysis

The overall effects of treatment and cultivar and their interaction were analysed by means of two-way ANOVA with heat treatment and genotypes taken as fixed factors. Genotypes were treated as fixed factors because we were interested in the response of the specific genotypes used in this experiment. Twenty two varieties were analysed with 3 replicates each for the treatment levels. The statistical analysis were done using OPSTAT software (Two factorial CRD).

3. Result

Significant genotypic differences for starch content was observed among tomato varieties under high temperature. Among the genotypes, Vaibhav (312.97 mg g⁻¹ fresh weight) recorded the maximum starch accumulation followed by Manulakshmi (304.45 mg g⁻¹ fresh weight) under control condition, while the minimum starch content was recorded in Arka Rakshak (214.06 mg g⁻¹ fresh weight). In heat stress condition, the highest starch content was observed in Anagha (235.67 mg g⁻¹ fresh weight), while the lowest was observed in Arka Sourabh (84.37 mg g⁻¹ fresh weight). The percent decrease in starch content was more in Arka Sourabh and less in IISR-2200. The average starch content of the tomato genotypes at flowering stage was 170.71 mg g⁻¹ fresh weight and 262.86 mg g⁻¹ fresh weight under heat stress and control condition respectively (Table 3).

Table 3. Significant genotypic differences for soluble sugar content was observed in different genotypes under high temperature. Highest soluble sugar concentration was observed in Nandi (77.73 mg g⁻¹ fresh weight) and lowest concentration in Arka Rakshak (51.92 mg g⁻¹ fresh weight) under control condition, whereas under stress condition Vellayani Vijay (59.6 mg g⁻¹ fresh weight) showed the highest soluble sugar content and minimum in Arka Rakshak (35.73 mg g⁻¹ fresh weight). The percent decrease in starch content was more in Arka Rakshak and less in Arka Abha.

The lycopene content decreased with a rise in temperature and ambient condition recorded the highest lycopene content in fruits (2.03 mg g⁻¹ fresh weight). The highest lycopene content was recorded in IISR-2200 (5.49 mg g⁻¹ fresh weight) and lowest was observed in Arka Alok (0.36 mg g⁻¹ fresh weight) under the control conditions whereas, maximum lycopene content was recorded for Nandi (2.94 mg g⁻¹ fresh weight) and minimum was recorded for Arka Vikas (0.35 mg g⁻¹ fresh weight) under high temperature conditions (Figure 2). The percent reduction in lycopene content under stress condition was maximum for IISR-2200 (52%) and minimum for Kashi Vishesh (3%).

Titratable acidity of tomato fruits was found to be significantly different among different genotypes. Highest concentration recorded at high temperature condition when compared to low temperature regimes (control). The highest titratable acidity was recorded for Kashi Vishesh (0.76%) which is at par with Vaibhav (0.75%) and Nandi (0.71%), minimum was recorded for IC-45 (0.33%) under control condition and maximum for Kashi Vishesh (0.86%) which is at par with Vaibhav (0.80%) and Nandi (0.81%), and minimum for IC-45 (0.37%) under high temperature condition (Figure 3). The average titratable acidity under control condition was 0.52% and 0.60% under high temperature condition. The percent increase in titrable acidity under heat stress was highest for Arka Alok (27%) and minimum for Pusa Rohini (2%).

In our study, TSS increased in all the genotypes under temperature stress condition compared to control (Figure 4). Highest TSS was recorded for Arka Samrat (5.72%) and lowest for IC-45 (2.32%) under control ambient condition. But under high temperature condition highest TSS was recorded for Kashi Vishesh (6.23%) and lowest for IC-45 (2.57%). The percent increase in TSS was highest for IISR-2200 (53%) and lowest for Arka Vikas (1%) under stress condition.

Vitamin C content showed significant differences among the genotypes, all tolerant genotypes showed higher vitamin C under temperature stress conditions compared to control (Figure 5). Under high temperature condition, highest concentration of vitamin C was observed for Nandi (32.71 mg g⁻¹ fresh weight) and lowest for Arka Sourabh (9.67 mg g⁻¹ fresh weight) whereas, ascorbic acid was found highest in Palam Pride (40 mg g⁻¹ fresh weight) and lowest in Arka Samrat (9.39 mg g⁻¹ fresh weight) for control conditions. The percent increase in vitamin C content was maximum for IISR-2200 (36%) and minimum for Pusa Ruby (1%).

Under high temperature stress in polyhouse condition, all the genotypes showed an increment in the plant height because of the shading effect of polyhouse (Table 5) and elevated CO₂ (570 μmol mol⁻¹). Maximum plant height was observed for Nandi (143.97 cm) and minimum height for Vellayani Vijay (51.9 cm) under control condition and for high temperature condition, highest value of plant height was observed for IC-45 (219.33 cm) and lowest for Arka Sourabh (128.33 cm). The average value of plant height under control and temperature
Table 3. Effect of high temperature on starch content in leaves of tomato varieties expressed in mg g⁻¹ fresh weight.

| Sl. No. | Genotypes | Control | Treatment | Mean |
|---------|------------|---------|-----------|------|
| 1       | Nandi      | 281.33  | 222.76    | 252.04 |
| 2       | IC-45      | 250.06  | 208.07    | 229.06 |
| 3       | Pusa Rohini| 245.63  | 114.08    | 179.85 |
| 4       | Pusa Ruby  | 252.98  | 212.82    | 232.90 |
| 5       | IIHR-2200  | 250.73  | 127.44    | 189.09 |
| 6       | Anagha     | 271.17  | 235.67    | 253.42 |
| 7       | Akshaya    | 271.11  | 211.80    | 241.46 |
| 8       | Vellayani Vijay | 288.52 | 191.91    | 240.22 |
| 9       | Arka Vikas | 209.70  | 92.33     | 151.02 |
| 10      | Kashi Vishesh | 279.95 | 204.87    | 242.41 |
| 11      | Vaibhav    | 312.97  | 200.21    | 256.59 |
| 12      | IIHR-26372 | 239.47  | 168.27    | 203.87 |
| 13      | Palam Pride| 264.15  | 189.25    | 226.70 |
| 14      | Arka Abha  | 221.26  | 165.21    | 193.23 |
| 15      | Arka Alok  | 280.82  | 171.93    | 226.38 |
| 16      | Manulakshmi| 304.45  | 191.52    | 247.99 |
| 17      | Sakthi     | 245.40  | 162.70    | 204.05 |
| 18      | Manuprabha | 283.79  | 219.24    | 251.52 |
| 19      | Arka Samrat| 283.12  | 183.69    | 233.41 |
| 20      | Arka Sourabh| 262.54  | 84.37     | 173.46 |
| 21      | PKM-1      | 269.61  | 103.21    | 186.41 |
| 22      | Arka Rakshak| 214.06  | 94.20     | 154.13 |

Mean 262.86 170.71

Factors SE(m) C.D. (0.5%)
Varieties 3.65 10.29
Treatments 1.10 3.10
Factor (V X T) 5.17 14.55

Table 4. Effect of high temperature on soluble sugar content in leaves of tomato varieties expressed in mg g⁻¹ fresh weight.

| Sl. No. | Genotypes | Control | Treatment | Mean |
|---------|------------|---------|-----------|------|
| 1       | Nandi      | 77.73   | 53.65     | 65.69 |
| 2       | IC-45      | 57.52   | 41.54     | 49.53 |
| 3       | Pusa Rohini| 56.03   | 41.35     | 48.69 |
| 4       | Pusa Ruby  | 60.65   | 49.38     | 55.02 |
| 5       | IIHR-2200  | 57.23   | 55.53     | 56.38 |
| 6       | Anagha     | 66.43   | 49.74     | 58.09 |
| 7       | Akshaya    | 65.88   | 56.82     | 61.35 |
| 8       | Vellayani Vijay | 67.79 | 59.60     | 63.70 |
| 9       | Arka Vikas | 53.41   | 46.41     | 49.91 |
| 10      | Kashi Vishesh | 65.99 | 53.30     | 59.65 |
| 11      | Vaibhav    | 63.67   | 51.06     | 57.36 |
| 12      | IIHR-26372 | 54.58   | 42.33     | 48.46 |
| 13      | Palam Pride| 61.80   | 42.80     | 52.30 |
| 14      | Arka Abha  | 55.91   | 50.07     | 52.99 |
| 15      | Arka Alok  | 57.40   | 53.46     | 55.43 |
| 16      | Manulakshmi| 72.03   | 56.81     | 64.42 |
| 17      | Sakthi     | 56.96   | 45.80     | 51.38 |
| 18      | Manuprabha | 71.83   | 52.91     | 62.37 |
| 19      | Arka Samrat| 62.56   | 43.98     | 53.27 |
| 20      | Arka Sourabh| 54.94  | 46.30     | 50.62 |
| 21      | PKM-1      | 53.83   | 45.61     | 49.72 |
| 22      | Arka Rakshak| 51.92  | 35.73     | 43.83 |

Mean 61.19 48.83

Factors SE(m) C.D. (0.5%)
Varieties 0.976 2.748
Treatments 0.294 0.829
Factor (V X T) 1.380 3.886
stress condition were 96.79 cm and 162.21 cm respectively. The percent increase in plant height was maximum for Vellayani Vijay (70%) and minimum for Arka Vikas (14%).

Fruit set significantly decreased at high temperature in all the tomato genotypes as compared to control temperature (Table 6). Highest fruit set % under control condition was recorded in Vellayani Vijay (53.68%) and lowest in Pusa Rohini (13.56%) whereas, highest fruit set % for high temperature stress condition was recorded in IC-45 (7.69%) and lowest for Palam Pride (1.23%). The average fruit set percentage under control and high temperature stress was 33.52% and 2.87% respectively. The percent decrease in fruit set % was maximum for Palam Pride (96.42%) and minimum for Arka Rakshak (86.17%). Significant decrease was observed in average fruit weight of tomato genotypes at high temperature. There was decrease in number of fruits per plant, percent fruit set and fruit yield per plant in all tomato genotypes under high temperature. Nandi, Kashi Vishesh, Anagha showed lesser magnitude of reduction for parameters like lycopene content and fruit set % as compared to Arka Sourabh, Pusa Rohini, PKM-1.

Average fruit weight was significantly decreased at high temperature in all the tomato genotypes as compared to control temperature (Table 7). The maximum average fruit weight was observed for Arka Vikas (37.23g) and minimum for IC-45 (3.91g) under control conditions whereas, it is maximum for Kashi Vishesh (6.61g) which is on par with Nandi (6.30g) and minimum for Arka Rakshak, Arka Samrat, Arka Sourabh, PKM-1 (0.12g). The maximum percent decrease under heat stress was recorded for Pusa Rohini, Pusa Ruby, Arka Rakshak, Arka Samrat, Arka Sourabh, PKM-1 (susceptible varieties - > 95%) and minimum for IC-45 (77%) as compared to ambient condition.
Yield per plant significantly decreased at high temperature in all tomato genotypes as compared to control temperature. Nandi (213.12g/plant) gave the maximum yield per plant under control condition whereas, Arka Rakshak (22.41g/plant) showed minimum yield per plant (Figure 6). Under heat stress condition, genotypes Nandi, Anagha, Akshaya, IIHR-2200, Vellayani Vijay, Kashi Vishesh, Arka Abha, Arka Alok, Vaibhav, Manuprabha, Manulakshmi, IC-45 and IIHR-26372 recorded higher fruit yield per plant. Varieties like Arka Saurabh, Arka Rakshak, PKM-1, Sakthi, Palam Pride, Arka Samrat recorded the maximum percent reduction in yield per plant (99%) and minimum was recorded in Kashi Vishesh (69%).

3.1. Marker analysis

Twenty-five SSR markers were used for PCR screening, and the sequence was taken from the Sol Genomics Network database. In 3 percent agarose gel electrophoresis, 7 out of the 25 primers displayed polymorphism (Figure 7) and the other primers were monomorphic. Therefore seven markers were used for determining the coefficient of similarity. The temperatures of these reactions were optimized using gradient PCR technique. Different annealing temperatures (Tm ± 5°C) were set between each block in this process.

3.2. Similarity coefficient analysis

Based on the DNA banding pattern of twenty two tomato genotypes using 25 SSR markers, Jaccard’s similarity coefficient were developed and displayed in Figure 8. The genetic similarity coefficients of these tomato genotypes ranged from a minimum of 0.22 to maximum of 1. The average genetic similarity range was 0.67.

Maximum genetic similarity (1) was shown by; Pusa Ruby with IC-45, IIHR-26372, Manulakshmi, Arka Samrat, Arka Alok, Sakthi and Arka Abha with Nandi. Quality and yield parameters also showed similarities among these genotypes. Titrable acidity content of Arka Abha and Nandi ranged from 0.6 to 0.7 and for Pusa Ruby with IC-45, IIHR-26372, Manulakshmi, Arka Samrat, Arka Alok, Sakthi ranged from 0.4 to 0.6. Fruit set percentage of Nandi and Arka Abha ranged from 36 to 44% while it ranged from 30 to 36% in case of Pusa Ruby with IC-45, IIHR-
Table 5. Effect of high temperature on plant height of tomato expressed in cm.

| Varieties         | Control | Treatment | Mean  |
|-------------------|---------|-----------|-------|
| Nandi             | 143.97  | 161.33    | 152.65|
| IC-45             | 85.67   | 219.33    | 152.50|
| Pusa Rohini       | 101.70  | 154.67    | 128.18|
| Pusa Ruby         | 104.17  | 172.67    | 138.42|
| IIHR-2200         | 113.00  | 166.33    | 139.67|
| Anagha            | 96.40   | 146.67    | 121.53|
| Akshaya           | 101.83  | 165.33    | 133.58|
| Veilayani Vijay   | 51.90   | 176.67    | 114.28|
| Arka Vikas        | 127.57  | 148.67    | 138.12|
| Kasvi Vishesh     | 84.33   | 147.00    | 115.67|
| Vaibhav           | 91.87   | 167.67    | 129.77|
| IIHR-26372        | 109.97  | 174.67    | 142.32|
| Falam Pride       | 101.23  | 164.00    | 132.62|
| Arka Abha         | 93.33   | 183.00    | 138.17|
| Arka Alok         | 68.83   | 163.67    | 116.25|
| Manulakshmi       | 80.33   | 172.67    | 126.50|
| Sakti             | 90.17   | 130.33    | 110.25|
| Manuprabha        | 106.37  | 142.67    | 124.52|
| Arka Samrat       | 96.93   | 159.67    | 128.30|
| Arka Sourabh      | 73.50   | 128.33    | 100.92|
| PKM-1             | 109.67  | 174.33    | 142.00|
| Arka Rakshak      | 96.77   | 149.00    | 122.88|
| **Mean**          | 96.80   | 162.21    |       |

Factors SE(m) C.D. (0.5%)

| Varieties         | 6.75    | 19.00    |
| Treatment         | 2.04    | 5.73     |
| Factor (V X T)    | 9.55    | 26.87    |

Table 6. Effect of high temperature on fruit set percentage of tomato genotypes expressed in %.

| Varieties         | Control | Treatment | Mean  |
|-------------------|---------|-----------|-------|
| Nandi             | 44.55   | 5.56      | 25.05 |
| IC-45             | 30.56   | 7.69      | 19.13 |
| Pusa Rohini       | 13.56   | 1.59      | 7.57  |
| Pusa Ruby         | 32.63   | 2.38      | 17.51 |
| IIHR-2200         | 31.79   | 2.08      | 16.94 |
| Anagha            | 40.66   | 4.17      | 22.42 |
| Akshaya           | 42.81   | 2.73      | 22.77 |
| Veilayani Vijay   | 53.68   | 2.30      | 27.99 |
| Arka Vikas        | 15.27   | 1.96      | 8.61  |
| Kasvi Vishesh     | 48.72   | 5.13      | 26.92 |
| Vaibhav           | 38.10   | 2.38      | 20.24 |
| IIHR-26372        | 31.64   | 2.90      | 17.27 |
| Falam Pride       | 34.43   | 1.23      | 17.83 |
| Arka Abha         | 36.33   | 2.86      | 19.59 |
| Arka Alok         | 35.80   | 3.03      | 19.42 |
| Manulakshmi       | 35.82   | 1.90      | 18.86 |
| Sakti             | 29.43   | 2.22      | 15.83 |
| Manuprabha        | 35.68   | 2.15      | 18.41 |
| Arka Samrat       | 36.70   | 2.56      | 19.63 |
| Arka Sourabh      | 26.16   | 2.56      | 14.36 |
| PKM-1             | 24.60   | 2.22      | 13.41 |
| Arka Rakshak      | 18.54   | 2.56      | 10.55 |
| **Mean**          | 35.52   | 2.87      |       |

Factors SE(m) C.D. (0.5%)

| Varieties         | 3.18    | 8.96     |
| Treatment         | 0.96    | 2.70     |
| Factor (V X T)    | 4.50    | 12.67    |
26372, Manulakshmi, Arka Samrat, Arka Alok, Sakthi. Yield of Nandi and Arka Abha showed a range between 180-213 g/plant and that of Pusa Ruby with IC-45, IIHR-26372, Manulakshmi, Arka Samrat, Arka Alok, Sakthi ranged 50–80 g/plant. Minimum genetic similarity coefficient (0.22) was shown by two pairs of genotypes viz. Pusa Rohini with Akshaya and Kashi Vishesh. Since they have low similarities, they shown differences in the yield, physiological data and in molecular characterization. In case of yield, Pusa Rohini with Akshaya and Kashi Vishesh shown wide range differences in lycopene content (1.58–3.53 mg plant⁻¹), fruit set percentage (13–48%) and yield (80–135g/plant).

4. Discussion

Heat stress reduces the sucrose transport and its accumulation in the leaves of both heat tolerant and heat-sensitive tomato genotypes,

### Table 7. Effect of high temperature on average fruit weight content of tomato genotypes expressed in g.

| Varieties   | Control | Treatment | Mean    |
|-------------|---------|-----------|---------|
| Nandi       | 26.91   | 6.30      | 16.61   |
| IC-45       | 3.91    | 0.96      | 2.43    |
| Pusa Rohini | 34.78   | 0.27      | 17.53   |
| Pusa Ruby   | 32.41   | 0.15      | 16.28   |
| IIHR-2200   | 15.00   | 1.14      | 8.07    |
| Anagha      | 19.84   | 3.46      | 11.65   |
| Akshaya     | 25.01   | 1.45      | 12.23   |
| Vellayani Viji | 17.08 | 3.25      | 10.16   |
| Arka Vikas  | 37.23   | 0.14      | 18.68   |
| Kashi Vishesh| 17.24 | 6.61      | 11.92   |
| Vaibhav     | 25.99   | 1.49      | 12.74   |
| IIHR-26372  | 20.86   | 1.28      | 11.07   |
| Falam Pride | 31.11   | 0.16      | 15.64   |
| Arka Abha   | 35.18   | 1.21      | 18.19   |
| Arka Alok   | 16.04   | 0.97      | 8.51    |
| Manulakshmi | 19.88   | 1.03      | 10.46   |
| Sakthi      | 16.18   | 0.34      | 8.26    |
| Manuprabha  | 29.66   | 1.02      | 15.34   |
| Arka Samrat | 31.10   | 0.12      | 15.61   |
| Arka Sourahi| 18.14   | 0.12      | 9.13    |
| PKM-1       | 14.76   | 0.12      | 7.44    |
| Arka Rakshak| 11.21   | 0.12      | 5.66    |
| Mean        | 22.52   | 1.44      |         |
| Factors     | SR(m)   | C.D. (0.9%)|
| Varieties   | 1.91    | 5.37      |
| Treatments  | 0.58    | 1.62      |
| Factor (V X T) | 2.69 | 7.59      |

**Figure 6.** Clustered column graph showing yield obtained from tomato genotypes maintained at control and high temperature conditions.
Figure 7. Amplification profile of 22 genotypes with a) SSR 96, b) SSR 63, c) SSR 13, d) SSR 270, e) SSR 356, f) SSR 605. 1-Marker 100 bp ladder. Lane 2–23 tomato genotypes in the same order of Table 1.

Figure 8. Jaccard’s similarity coefficient matrix for 22 tomato genotypes based on SSR data. Where, G1 = Manuprabha, G2 = Akshaya, G3 = Pusa Ruby, G4 = IC-45, G5 = Nandi, G6 = IIHR-2200,G7 = IIHR-26372, G8 = Palam Pride, G9 = PKM-1, G10 = Manulakshmi,G11 = Arka Samrat, G12 = Arka Rakshak, G13 = Arka Vikas, G14 = Pusa Rohini, G15 = Arka Alok, G16 = Sakthi,G17 = Vaibhav, G18 = Vellayani Vijay, G19 = Anagha,G20 = Kashi Vishesh, G21 = Arka Sourabh, G22 = Arka Abha.
indicating that the carbohydrate translocation and partitioning to other plant parts are negatively affected under high temperatures, similar to the results obtained from wheat (Wahid et al., 2007; Shanmugam et al., 2013). A decrease in the starch content was observed under different exposure of abiotic stress (Vinocur and Altman, 2005). In our study also a drastic change in the starch content was observed for different varieties. In heat stress condition, the highest starch content was observed in Anagha, while the lowest was observed in Arka Sourabh. Under heat stress, the concentration of starch and soluble sugar in the pollen grains was lower than that under control conditions (Kumar et al., 2015). These findings are similar to those obtained with that from rice (Sheoran and Saini, 1996) and wheat (Durion et al., 1996). It has been suggested that carbohydrates starvation in those grains are not responsible for the stress-induced pollen sterility. Pollen of heat tolerant varieties have high amount of glucose rather than sucrose and fructose and it can also retain high amount of carbohydrates (Firon et al., 2006). Xu et al. (2017) revealed that fruit set directly influenced the number of fruits and yield in tomato crop moreover, there is no significant correlation between vegetative and reproductive traits. Flower number per inflorescence and membrane thermo-stability are also relevant characteristics and might be used as indicators of reproductive heat tolerance. Several workers also reported that high temperatures cause significant loss in tomato productivity due to reduced fruit set, number of fruits and poor-quality fruit (Zinn et al., 2010; Akhtar et al., 2012; Nahar and Ullah, 2012; Solankey et al., 2017).

Shi and Le Maguer (2000) reported the inhibition of lycopene production at higher temperatures (38 °C). The relatively heat tolerant genotypes showed lesser decrease in lycopene content in the fruit at high temperature as compared to susceptible genotypes (Sharma and Le Maguer, 1996). Lycopene constitutes 80-90 % of the total carotenoids in tomato fruits (Valverde et al., 2002). The result obtained from the present study also pointed the fact that lycopene production under heat stress was severely affected. Under high temperature condition, maximum lycopene content was recorded for Nandi and minimum was recorded for Arka Vikas. Amrutha and Beena (2020) observed in tomato genotypes for fruit quality parameters at high temperature conditions.

The fruits showed lower content of phenols, flavonoids, ferric reducing antioxidant potential, total soluble solids, and titrable acidity in plants grown at heat stress as compared with the control. The ascorbic acid content was high at stress condition. Carotenoids and lycopene content was low at temperature stress compared to higher content observed at control condition (Mamatha et al., 2014).

Increase in temperature increased TSS and titrable acidity but decreased total sugars, lycopene, and total carotenoids content in tomato (Lokesha et al., 2019). The sugars contribute to the total soluble solids content of tomato fruits (Laxman et al., 2013; Selahle et al., 2014). TSS ranged from 4 to 6 °Brix in tomato fruits. The change in the glucose to fructose ratio and the organic acids content is the main cause for changes in the TSS changes in tomato. For the taste of tomatoes, TSS was reported as a beneficial indicator (Klunklin and Savage, 2017). TSS increased in the genotypes under temperature stress compared to control, which is on par with inferences by Shivashankara et al. (2015). Our study also showed that an increment in TA and TSS value were observed for all the tomato genotypes. Under stress conditions Vellayani Vijay showed the highest soluble sugar content and minimum in Arka Rakshak. The phenolic substances have a protective role on ascorbic acid content (Wang and Zheng, 2001) the presence of phenolics and flavonoids in tomato fruits help to maintain the vitamin C level. A significant increase in total phenolic acids and flavonoids under high-temperature were reported in strawberry (Wang, 2006) and also in other crops (Pervez et al., 2009; Bitra and Gerats, 2013). Vitamin C content showed significant differences among the tolerant genotypes, all tolerant genotypes showed higher vitamin C under temperature stress conditions compared to control. Vitamin C content increased when the heat stress was imposed during flowering and fruit set stages, indicating that the plant metabolism is adapted to high temperature. The synthesis and accumulation of health-promoting metabolites, termed phytochemicals, depends mainly on the genetic material, although the agronomic practices and environmental factors also have an important influence on yield and quality characteristics of fruits and vegetables (Rouphael et al., 2012; Schreiner et al., 2013). Thus, salt and nutritional stresses have been used for the improvement of the nutritional quality of fruits (Colla et al., 2013; Fanciullino et al., 2014; Massareto et al., 2018). Heat stress increased ascorbic acid (ABA) and reduced salicylic acid (SA) content; however, combined application of SA + HA markedly reduced ABA and increased SA. Antioxidant enzymes activities revealed that SA and HA treated plants exhibited increased levels of ascorbate peroxidase (APX), super-oxide dismutase (SOD), and reduced glutathione (GSH) (Hemmati et al., 2015; Gururani et al., 2015). Seed filling parameters recorded at milky and dough stage revealed that high temperature stress condition increased the amount of reducing sugar, carbohydrates, starch, and flavonoids. However, amylase, seed protein, and anthocyanin showed reduction under high temperature stress condition. Activity of invertase was reduced under high temperature condition compared to control in all varieties from 15 to 30 days after 50% flowering (Pravalika et al., 2020).

Fruit setting percentage is affected by changing temperatures during different crop growing seasons (Nahar and Ullah, 2012; Beena et al., 2018a). High temperatures cause significant loss in tomato productivity due to reduced fruit set and poor fruit quality (Mitcham and McDonald, 1992; Khanal, 2012). While shading increased the number of fruits per plant and total fruit yield. The maximum fruit yield was obtained by plants grown under 50% shading in both cultivars under study. Tomato plants grown under shading gave the best physical characteristics of tomato fruits (fruit length and diameter) and TSS %. Leaf concentrations of N, K and Ca were significantly increased with the increased shading levels. The highest content of N, K and Ca was observed with shading with black net at 50% density. On the contrary, plants grown without shading had the highest content of P (El-Bassiony et al., 2014). Alsamir et al. (2017) reported that high temperature in tomato reduced number of fruits, flower to fruit set ratio and fresh fruit weight. These results are supporting the present inferences from our study. The higher pollen viability, high pollen germination and high soluble sugar content in pollen grains at anthesis may be the reason for better number of fruits per plant, percent fruit set and fruit yield in tolerant genotypes at high temperature. Pollen viability and fertility are reported to be reason for better plant productivity during heat stress. The fruit number, fruit set percentage and fruit weight per plant were decreased with increase in temperature. In the present study, the yield attributes viz., number of fruits/plant, fruit set %, average fruit weight (g), yield per plant (g/plant) were significantly lower for varieties like Arka Saurabh, Arka Rakshak and Pusa Rohini.

Under heat stress conditions Nandi, Anagha, Akshaya, IIHR-2200, Vellayani Vijay, Kashi Vishesh, Arka Abha, Arka Alok, Vaibhav, Manuprabha, Manulakshmi, IC-45 and IIHR-26372 produced higher fruit yield per plant. But the varieties like Arka Saurabh, Arka Rakshak, PKM-1, Sakthi, Palam Pride, Arka Samrat recorded the maximum percent reduction in yield per plant and the minimum was recorded in Kashi Vishesh. At high temperature levels, transpire more, and hence yield reduction is caused by the impaired pollen, anther development, and reduced pollen viability. The temperature values higher than 35 °C reduce the fruit set and delay the development of normal fruit colour (Kang et al., 2002; Sato et al., 2006). Reduced allocation of assimilates under high temperature stress compared with control temperature condition (Sang et al., 2005) and reduced supply of phytomethanes and poor production of growth regulators in sink tissues are pointed out to be the reasons for reduced yield related traits (Islam, 2011; Hasanuzzaman et al., 2013). Studies on heat-tolerant tomato genotype demonstrated that high invertase activity and increased sucrose import into young tomato fruits contributed to heat tolerance through increasing sink strength and sugar signalling activities, by regulating a programmed cell death pathway (Li et al., 2012). The genotypes producing higher proline concentrations in plant parts and with higher membrane thermo-stability.
under high temperature produced highest fruit yield, and exhibited higher temperature tolerance (Din et al., 2015). Heat tolerant genotypes maintained higher net photosynthesis \( (P_N) \) and increased stomatal conductance \( (g_s) \) at 38 °C, and better leaf cooling. Sensitive genotypes had lower \( P_N \) and \( g_s \) at 38 °C, and the increased less than in the tolerant group and less leaf cooling. Under controlled conditions, all eight genotypes had the same plant size and pollen viability reduced dramatically in the sensitive group and less leaf cooling. Under controlled conditions, all eight genotypes accumulated more biomass, had a lower heat injury index and higher protective effect of this gene under salinity and drought (Baranova et al., 2011).

**4.1. Similarity coefficient analysis**

A wide range of similarity coefficients between certain genotypes indicated the presence of significant genetic variability between some of the investigated genetic stocks. Among other tomato varieties, Dhalliwal et al. (2011) previously reported similar findings of similarities coefficients among the tomato genotypes. The limited range of coefficient of similarity between these genotypes suggested the existence of limited genetic similarities among the analyzed genotype. A similar study was conducted by Kumar et al. (2016) and reported that the genotypes Arka Vikas and 2012TODVAR-2 are 100% similar.

Based on present study, Akshaya, IIHR-2200, Manuprabha are moderately tolerant varieties showed 89% similarity. Pusa Ruby, IC-26372, Manulakshmi, Arka Samrat, Arka Aloks, Sakti showed 100% similarity. Kaushal et al. (2017) reported a maximum of 96% similarity among tomato. High level of similarity (95%) was revealed among 39 tomato (Al-Abadi, 2007), similarity of 100% was found among tomato (Tam et al., 2005). Kashi Vishesh, Anagha and Vellayani Vijay are tolerant varieties. Kashi Vishesh and Anagha showed 78% similarity. Kashi Vishesh and Vellayani Vijay showed 67% similarity. Vellayani Vijay and Anagha showed 89% similarity. Susceptible varieties are: Arka Sourabh, Pusa Rohini, Palam Pride, Arka Rakshak and observed 78% similarity for three pairs of genotypes viz. Arka Sourabh with Pusa Rohini, Palam Pride and Arka Rakshak. A 68% similarity observed for two pairs of genotypes viz. Pusa Rohini with Palam Pride and Arka Rakshak.

**5. Conclusion**

Significant genotypic differences for starch, soluble sugars, titratable acidity (TA), total soluble solids (TSS), lycopene content, yield attributes viz., number of fruits/plant, fruit set %, average fruit weight (g) and yield per plant (g/plant) were observed among tomato genotypes. Nandi, Anagha, Akshaya, Vellayani Vijay, Kashi Vishesh showed high temperature tolerance. Jaccard's similarity coefficient matrix of these tomato genotypes ranged from a minimum of 0.22 to a maximum of 1. Further study has to be conducted for the confirmation of heat tolerance with respect to different attributes contributing for tolerance mechanisms.

**Declarations**

**Author contribution statement**

Amrutha Vijayakumar: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Wrote the paper.

Shanija Shaji; Sarada, S.: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Beena R.: Conceived and designed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Sajitha Rani, T; Roy Stephen.; Manju, R.V.; Viji, M.M.: Contributed reagents, materials, analysis tools or data.

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**Declaration of interests statement**

The authors declare no conflict of interest.

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