A conserved HSF:miR169:NF-YA loop involved in tomato and Arabidopsis heat stress tolerance

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Abstract

Heat stress transcription factors (HSFs) and miRNAs regulate different stress and developmental networks in plants. Regulatory feedbacks are at the basis of these networks. Here, we report that plants improve their heat stress tolerance through HSF-mediated transcriptional regulation of \textit{MIR169} and post-transcriptional regulation of \textit{NF-YA} transcription factors. We show that HSFs recognize tomato and Arabidopsis \textit{MIR169} promoters using yeast-one-hybrid/ChIP-qPCR. Silencing tomato HSFs using virus induced gene silencing (VIGS) reduced \textit{Sly-MIR169} levels and enhanced \textit{Sly-NF-YA9/A10} target expression. Further, \textit{Sly-NF-YA9/A10-VIGS} knock-down tomato plants and Arabidopsis plants overexpressing \textit{At-MIR169d} or \textit{At-nf-ya2} mutants showed a link with increased heat tolerance. In contrast, Arabidopsis plants overexpressing \textit{At-NF-YA2}, or those expressing a non-cleavable \textit{At-NF-YA2} form (miR169d-resistant \textit{At-NF-YA2}) as well as plants inhibited for \textit{At-miRNA169d} regulation (miR169d mimic plants) were more sensitive to heat stress, highlighting \textit{NF-YA} as negative regulator of heat tolerance. Furthermore, post-transcriptional cleavage of \textit{NF-YA} by elevated miR169 levels resulted in alleviating the repression of heat stress effectors HSFA7a/b in tomato and Arabidopsis revealing a retroactive control of HSFs by the miR169:NF-YA node. Hence, a regulatory feedback loop involving HSFs, miR169s and NF-YAs plays a critical role in the regulation of heat stress response in tomato and Arabidopsis plants.

Introduction

Heat stress (HS) adversely affects the distribution and productivity of agriculturally important plants worldwide by affecting all aspects of plant processes like germination, growth, development, reproduction and yield (Mittler et al. 2010; Lobell et al. 2011; Hasanuzzaman et al. 2013). In nature, such HS conditions may be chronic or recurring, or both (Bäurle 2016); therefore, plants have evolved a variety of responses to elevated temperatures that minimize damage and ensure protection of cellular homeostasis.

HSFs are important regulators that play critical roles in signal transduction processes to mediate gene expression in response to multiple abiotic stresses, including cold,
drought, salt, and heat (Collins et al. 1995; Nover et al. 2001; Guo et al. 2016; Jacob et al. 2017). HSFs recognize heat stress elements (HSE: 5′-GAA[n]TTC-3′) that are conserved in promoters of heat stress-responsive (HSR) genes including HSP genes (Busch et al. 2005). During the initial stages of HS, HSFs are the primary molecules responsible for relaying signals of cellular stress to the transcriptional apparatus and reprogram gene expression to repair protein damage through elevated synthesis of molecular chaperones and proteases (Morimoto 2002; Scharf et al. 2012; Guo et al. 2016). In plants, there are multiple genes coding for HSFs for example, Arabidopsis has 21 HSF family members (von Koskull-Döring et al. 2007) and tomato has 26 HSFs (Yang et al. 2016). Another class of regulatory molecules, the microRNAs (miRNAs) have emerged as a major class of small non-coding RNAs with roles in plant growth and development as well as under stress (Sunkar and Zhu, 2004; Lin et al. 2018; Ravichandran et al. 2019). HSF-mediated induction of MIR398 has been shown to trigger a regulatory loop that is critical for thermotolerance in Arabidopsis (Guan et al. 2013). Yan et al. (2012) have shown that stress-induced alternative splicing of MIR400 provides a mechanism for the regulation of miRNA processing in Arabidopsis during HS. Lin et al. (2018) have shown that miR160 modulates plant development and heat shock protein gene expression to mediate heat tolerance in Arabidopsis. Heat-responsive miRNAs in *Apium graveolens* (Li et al. 2014), switch grass (Hivrale et al. 2015), banana (Vidya et al. 2018) and apple (Niu et al. 2019) have also been reported.

The miR169 family members have significant roles in plant abiotic and biotic stress responses and are involved in various aspects of plant development including root architecture, nodule formation etc. by targeting *Nuclear factor-Y A* (NF-YA) gene family members (Zhao et al. 2011; Ni et al. 2013; Sorin et al. 2014; Zhang et al. 2015; Li et al. 2017). Tomato plants overexpressing miR169 show improved resistance to drought stress (Zhang et al. 2011) while in Arabidopsis enhanced miR169 levels promote leaf dehydration (Li et al. 2008; Ni et al. 2013). Over-accumulation of miR169 is correlated with a reduction of *NF-YA* target transcripts in Arabidopsis seedlings under cold stress (Zhou et al. 2008; Lee et al. 2010). Role of miR169 is also established in long-distance signalling and nitrogen/phosphorus starvation (Pant et al. 2009; Buhtz et al. 2010; Zhao et al. 2011). In a previous publication from our laboratory, we have reported 18 *MIR169*
family members in tomato (Rao et al. 2020). Assessment of Sly-MIR169 precursors in different heat stress (HS) regimes highlighted that fifteen of the sixteen precursors (two of the loci are bicistronic) are differentially regulated in response to HS. Heat-mediated up-regulation of MIR169s has also been reported in Arabidopsis (Li et al. 2010; Guan et al. 2013) however, a detailed mechanism was lacking. Here, we have dissected the transcriptional regulation of MIR169 family by HSFs in tomato and Arabidopsis HSR.

Our laboratory has previously established miR169-mediated transcript cleavage of five classical targets belonging to the NF-YA class of transcription factors (Sly-NF-YA1, Sly-NF-YA3, Sly-NF-YA8, Sly-NF-YA9 and Sly-NF-YA10) (Rao et al. 2020). Here, we show that miR169 mediated post-transcriptional down-regulation of At-NF-YA2 in Arabidopsis and Sly-NF-YA9 and Sly-NF-YA10 in tomato enhances HS tolerance and impacts HSF expression. NF-YA transcriptionally regulates a suite of HSR genes including HSFs which exert a feedback regulation on NF-YAs through the control of miR169 expression. This is a new feedback loop involved in the regulation of plant tolerance to HS.

**Results**

**MIR169 members are transcriptionally regulated by HSFs in tomato**

The transcriptional activation of heat stress genes is regulated by HSFs that recognize HSE cis-elements in promoters of HSR genes (Busch et al. 2005). All the 15 heat up-regulated Sly-MIR169 promoters (Rao et al. 2020), contain one or more HSEs indicating possible involvement of HSFs in their heat inducibility (Supplementary figure S1, Supplementary table S1). This prompted us to investigate the transcriptional regulation of MIR169 promoters by HSFs using yeast-one-hybrid (Y1H) assays. Out of 322 interactions studied between tomato HSFs and 14 heat-responsive Sly-MIR169 promoters (one promoter could not be assessed as a minimal inhibitory Aureobasidin A antibiotic concentration could not be achieved), we found specific binding of seven HSFs on five Sly-MIR169 promoters (Figure 1A, Supplementary table S2, Supplementary figure S2). Promoters of Sly-MIR169a-1 and Sly-MIR169d-1 are regulated by a single HSF viz., Sly-HSF1a and Sly-HSFA1a, respectively (Figure 1A, Supplementary table S2). Sly-MIR169b promoter interacts with Sly-HSFA1a and Sly-
HSFA7a while *Sly-MIR169c* promoter shows Sly-HSFA4a and Sly-HSFA6b binding. Transcription of *Sly-MIR169d* is regulated by Sly-HSFA8a and Sly-HSFB1a (Figure 1A).

Further, to confirm the binding of these Sly-HSFs on *Sly-MIR169* promoters, we performed chromatin immunoprecipitation (ChIP) assays by co-expressing specific *Sly-MIR169* promoters and *P35S:HSF:GFP* constructs in *Nicotiana benthamiana* leaves. ChIP-qPCR was then performed on the HSE-specific regions in the *Sly-MIR169* promoters. Fold enrichment was calculated by comparing the Ct values of anti-GFP immunoprecipitated samples with IgG immunoprecipitation control (Figure 1B-I). ChIP-qPCR analysis of all HSFs with *Sly-MIR169* promoters matched the Y1H results except for HSFA4a and HSFA8a on *Sly-MIR169c* and *Sly-MIR169d* promoters (Figure 1E, G).

The HSF-mediated regulation of *MIR169* promoters was also assessed in *Nicotiana benthamiana* leaves using agro-infiltration (Supplementary Figure S3). For this, *Sly-MIR169* promoters were fused with *GUS* reporter gene and co-transformed with effector plasmids containing HSFs under CaMV35S promoter. All the five HSFs showing promoter enrichment in the ChIP assays were transformed individually or in combinations (as the case may be) to check their ability to drive *GUS* gene expression under respective *Sly-MIR169* promoters. qRT-PCR based estimation of *GUS* transcripts in presence of HSFs confirmed HSF-mediated activation of *Sly-MIR169* promoters for all the HSFs (Supplementary Figure S3) except Sly-HSFA1a (Supplementary Figure S3B). However, co-transformation of Sly-HSFA1a with Sly-HSFA7a (Supplementary Figure S3B) leads to enhanced levels of *GUS* transcripts as compared to individual HSFs suggesting a cooperative action in regulating *Sly-MIR169b* promoter.

**HSFs regulate the orchestration of miR169:NF-YA module during heat stress in tomato**

To assess whether the HSF:miR169:NF-YA module is functional in tomato HS, we quantified the transcript abundance of all components of this module in wild type tomato plants upon HS. Expression profiles of HSFs reveal the heat-mediated up-regulation of *Sly-HSFA6b*, *Sly-HSFA7a* and *Sly-HSFB1a* while *Sly-HSFA1a* and *Sly-HSFA1e* are not HS responsive (Figure 2A, Supplementary figure S4A). HSFA1s, the master regulators of HSR response, are present in bound forms with HSPs in cytoplasm. Upon HS, HSFs
dissociate from HSPs and move to nucleus to transcriptionally regulate HSR genes (Kotak et al. 2004; Chan-Schaminet et al. 2009) hence, HSFAs do not exhibit transcriptional up-regulation upon HS. However, we note that in non-stressed conditions the Sly-HSFA1s are more abundant in comparison to other HSFs (by comparing Ct values of these HSFs), suggesting their sufficient levels to orchestrate gene transcription upon HS (Supplementary table S3A). In agreement with our previous results (Rao et al. 2020), four miR169 mature forms (Figure 2B, Supplementary figure S4B) exhibit heat-mediated induction in comparison to non-stressed WT tomato plants. Comparison of their Ct values in control conditions revealed the abundant presence of miR169c in non-stressed plants, while imposition of heat stress led to strong induction of miR169a (Supplementary table S3B). Furthermore, expression analysis of miR169s targets (NF-YAs) shows reduction of only Sly-NF-YA9 and Sly-NF-YA10 transcripts in HS in WT plants (Figure 2C, Supplementary figure S4C).

To investigate the direct role of HSF-mediated regulation of the MIR169:NF-YA module in tomato, we utilized TRV-based virus induced gene silencing (VIGS) of HSFs. The VIGS silencing assays were performed 3-weeks-post agro-infiltration. Setting up of efficient silencing for TRV-HSFs was assessed by the reduction in the endogenous HSFs transcripts in silenced plants (Figure 2D-G). To ensure the specific downregulation of selected HSFs, we have examined the expression profiles of putative off targets HSFs predicted by SGN-VIGS tool (Supplementary table S4) by qRT-PCR. The unaltered expression of putative off targets in different HSFs-VIGS silenced plants confirms the specificity of the silencing (Supplementary figure S5A-C). Subsequently, downstream components in the pathway like MIR169 precursor levels and their heat responsive NF-YA targets (Sly-NF-YA9 and Sly-NF-YA10) were quantified by qRT-PCR after these plants were given a HS (Figure 2D-G, Supplementary figure S6A-D). Silencing of Sly-HSFA1a and Sly-HSFA7a reduces the transcription of Sly-MIR169b by 3 and 5 fold respectively, and leads to more than 2 fold enhanced expression of Sly-NF-YA9 and Sly-NF-YA10 transcripts upon HS (Figure 2D, E). Transcription of Sly-MIR169c is reduced by 2.6 folds in Sly-HSFA6b silenced plants, this in turn leads to up-regulation of Sly-NF-YA10 upon HS (Figure 2F). Silencing of Sly-HSFB1a (Figure 2G) reduces the expression of Sly-MIR169d by 2.9 folds, resulting in enhanced
accumulation of Sly-NF-YA9 and Sly-NF-YA10 transcripts upon HS. Sly-HSFA6b silencing also reduces the levels of Sly-MIR169d-1 by 4.9 folds that in-turn enhances the transcript levels of Sly-NF-YA10 upon HS (Figure 2F). Depressed expression of Sly-MIR169s with concomitant enhanced expression of Sly-NF-YA9/10 in Sly-HSF-silenced backgrounds upon HS suggests the HSF-mediated regulation of Sly-MIR169:Sly-NF-YA9/10 module in HS.

The miR169: NF-YA module is critical for high temperature tolerance in tomato

To establish the functions of Sly-NF-YA9 and Sly-NF-YA10 in heat tolerance, we used the VIGS-silencing approach to knock-down their expression in tomato plants and assessed their thermotolerance post HS. Three-weeks-after agro-infiltration, a silencing efficiency of ~80% was achieved for both Sly-NF-YA9 and Sly-NF-YA10 transcripts as compared to the TRV-vector negative control plants (Supplementary Figure S7A). Expression analysis of Sly-NF-YA9 or Sly-NF-YA10 in WT control (non-stressed) and TRV-EV infiltrated plants revealed no significant fluctuation in their expression levels (Supplementary figure S7B), suggesting the specificity of the VIGS silencing constructs. The TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 plants were given a HS and scored for heat tolerance six-days-after HS. qRT-PCR based expression analysis revealed that a strong downregulation of Sly-NF-YA9 and Sly-NF-YA10 is maintained in VIGS silenced plants post HS (Supplementary figure S7C-D), ruling out any time point and treatment biased alteration in their expression levels. Phenotypic evaluation post-HS showed that the leaves in vector control plants are highly damaged in these conditions, in contrast to the Sly-NF-YA9 and Sly-NF-YA10 silenced plants which are robust with green, healthy, erect leaves and also show emergence of new leaves (Figure 3A). The Sly-NF-YA9 and Sly-NF-YA10 knock-down plants show ~55% and 80% survival rate (judged by phenotype of 7th to 9th leaves and emergence of new leaves) in comparison to ~33% for vector-control plants (Figure 3B). Further, we gauged the physiological performance of TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 silenced plants, six days after HS using a Li-Cor 6400 photosynthesis measuring system. Both the NF-YA silenced plants show significant enhancement in water use efficiency and net photosynthetic rates as compared to TRV-control plants under HS (Figure 3C). To rule out that the post HS
phenotyping and survival scoring is not because of an effect of silencing of off targets, we predicted any possible off targets of Sly-NF-YA9 and Sly-NF-YA10 genes. Unaltered expression patterns of these putative off target genes in Sly-NF-YA9 and Sly-NF-YA10 VIGS silenced plants confirmed that the phenotype observed is specific to the knock-down of Sly-NF-YA9 and Sly-NF-YA10 genes (Supplementary table S4, Supplementary figure S8A-B).

To further delineate the molecular events driving the improved thermotolerance of TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 knock-down plants, we checked whether the expression of conserved HSR genes like HSFs (Sly-HSFA3a and Sly-HSFA7a), Sly-shSP and Sly-APX (ascorbate peroxidase) is significantly modulated upon heat stress. First, to ascertain that these genes are reliable HSR genes in the VIGS conditions, we show that they all are upregulated in TRV-EV plants under heat stress in comparison to non-stressed control TRV-EV plants (Figure 3D-H, Supplementary figure S9A-E). Furthermore, these HSR genes are inducible in TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 knock-down plants during heat stress as compared to non-stressed conditions (Figure 3D-H, Supplementary figure S9A-E). The strong upregulation of HSR genes in heat stressed TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 knock-down plants in comparison to TRV-EV HS plants confirms the direct involvement of these HSR genes in regulating the thermotolerance of tomato plants (Supplementary figure S9F). Further, since all these HSR genes (except Sly-APX) exhibit upregulation in TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 knock-down plants in comparison to TRV-EV plants in non-stressed conditions (Supplementary figure S9G), this indicates that these HSR genes (except Sly-APX) are downstream targets of Sly-NF-YA9 and Sly-NF-YA10. Furthermore, promoter analysis (1kb) of these HSR genes revealed the presence of CCAAT boxes in Sly-HSFA3a and Sly-HSFA7a (Supplementary table S5), suggesting a direct Sly-NF-YA9/10 mediated regulation on these genes and an indirect regulation on other HSR genes. Since the HSR genes are upregulated in TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 knock-down plants, these results suggest that Sly-NF-YA9 and Sly-NF-YA10 act as negative regulators of thermotolerance.*

*MIR169s are regulated by HSFs during heat stress in Arabidopsis*
To assess whether the HSF:miR169:NF-YA module under HS is active in other plants too, we decided to evaluate this module in the model plant Arabidopsis. Guan et al. (2013) have reported heat-mediated induction of miR169 and miR398 by Northern hybridization in Arabidopsis. However, the regulation of miR169 in the heat responses of Arabidopsis remains unknown. Using taqman based qRT-PCR analysis of WT control and heat stress challenged Arabidopsis plants, we show that all four mature miR169 forms are strongly upregulated (≥5 fold) under HS (Figure 4A, Supplementary figure S10). Further, to support a probable mechanism of their heat inducibility, promoters of all 14 At-MIR169 genes (Li et al. 2010) were screened for the presence of HSEs and we found two to multiple HSEs in all MIR169 promoters (Supplementary table S6). To delineate this speculated interplay among Arabidopsis HSFs and At-MIR169 promoters, Y1H assays were performed (Supplementary table S7) with 21 At-HSFs and 11 At-MIR169 promoters (as six precursors are transcribed as three bi-cistronic units viz., Pro-At-MIR169i/j, Pro-At-MIR169k/l, Pro-At-MIR169m/n; Li et al. 2010). This analysis identified 5 HSFs interacting with 3 At-MIR169 promoters (Figure 4B). Three HSFs (At-HSFB3a, At-HSFA7b, At-HSFB2b) regulate At-MIR169b; At-HSFA2a controls At-MIR169d transcription and transcription of At-MIR169h is regulated by At-HSFA1d. At-HSFA2 is considered as the key regulator of Arabidopsis HS-response and is required for the maintenance of acquired thermotolerance (Chang et al. 2007). Therefore, we decided to dissect the At-HSFA2 mediated At-MIR169d module for characterization of HS-response and thermotolerance in Arabidopsis. In wild type Arabidopsis plants, we find strong induction of At-MIR169d precursor (≥15 fold) upon HS (Figure 4C, Supplementary figure S11). Transgenic promoter-At-MIR169d:GUS plants support this heat-dependent transcriptional acceleration (Figure 4D, E). Moreover, GUS aided reporter analysis of promoter-At-MIR169d:GUS in WT and in the At-hsfa2 knockout mutant (Supplementary figure S12A, B, C) background in HS highlights minimal GUS expression in the At-hsfa2 mutant as compared to WT plants (Figure 4F, G). Further, there is strong reduction of At-MIR169d precursor levels in At-hsfa2 mutant as compared to WT plants (Figure 4G) confirming At-HSFA2-mediated regulation of At-MIR169d in Arabidopsis HS and suggesting a general regulatory loop in plants.
The miR169d: NF-YA2 regulatory node is heat responsive and spatially co-expressed in Arabidopsis

To explore the functional significance of heat-mediated regulation of MIR169s, we assessed the expression of the eight canonical NF-YA targets (Sorin et al. 2014) in HS in Arabidopsis. qRT-PCR based profiling shows significant reduction of At-NF-YA2 (-3.5 fold), At-NF-YA3 (-2.2 fold), At-NF-YA6 (-2.1 fold) and At-NF-YA10 (-2.1 fold) transcripts in WT plants when exposed to HS (Supplementary figure S13A, B). Further, to assign the functional relevance of the selected At-MIR169d isoform in particular, we analyzed target mimicry plants generated by overexpressing MIM169defg constructs under CaMV35S, using constructs from Todesco et al. (2010). The expression of 4 mature forms of At-miR169s in these MIM169defg lines revealed that miR169defg is the most reduced form (Supplementary figure S14A). The target mimic lines sequester the specific mature miRNAs by acting as non-cleavable targets; thereby they protect the target genes from miRNA cleavage (Todesco et al. 2010). The expression of At-NF-YA2, At-NF-YA3, At-NF-YA6, At-NF-YA8 and At-NF-YA10 is significantly increased in MIM169defg lines as compared to WT plants, with At-NF-YA2 transcript levels being highest in the MIM169defg lines (Supplementary figure S14B). Moreover, overexpressing At-MIR169d (At-MIR169d-OE) precursor under constitutive CaMV35S promoter in Arabidopsis strongly represses (-4 to -11-fold in three different At-MIR169d-OE lines) At-NF-YA2 transcripts, establishing it as the major miR169d target (Supplementary figure S14C). Moreover, promoter-At-NF-YA2 lines do not show HS-mediated up-regulation specifying a pure post-transcriptional regulation (Supplementary figure S15A, B).

To delineate the existence and localization of miR169d:At-NF-YA2 functional module in plant tissues and development stages, promoter reporter (GUS) lines were generated by transforming WT Arabidopsis plants with constructs that contained the 2 kb genomic sequences upstream of the At-MIR169d precursor. Parallelly, 2 kb region ahead of the translation start site of At-NF-YA2 was similarly assessed. The At-MIR169d and At-NF-YA2 promoters show prominent GUS localization in the vasculature of roots and aerial parts of seedling, mature rosette/cauline leaves and sepals in flowers of Arabidopsis,
suggesting functional co-existence of the miR169d:NF-YA2 module (Supplementary figure S16).

**Arabidopsis plants with enhanced miR169d or reduced NF-YA2 expression are thermotolerant**

To further investigate the miR169d:NF-YA2 module in HS-response in Arabidopsis, we assayed 2-weeks-old plants of **At-MIR169d-OE**, **At-NF-YA2-OE** (NF-YA2 overexpressed under 35SCaMV promoter, Supplementary figure S17), **At-nf-ya2 knockout mutant** (Supplementary figure S18), an miR169d resistant version of **At-NF-YA2** under native promoter (**pNF-YA2:NF-YA2**-r) and MIM169defg lines for their ability to survive HS. The survival rate of plants was gauged 6-days-after exposure to 2 h of HS at 42 °C. Plants overexpressing **At-MIR169d** (**At-MIR169d-OE**) that have reduced target **At-NF-YA2** levels and plants that lack **At-NF-YA2** transcripts (**At-nf-ya2** mutant) exhibit twice the survival rate than WT plants, in contrast, plants with high abundance of **At-NF-YA2** transcripts viz. **At-NF-YA2-OE**, **MIM169defg** and the **pNF-YA2:NF-YA2-r** (a variant **At-NF-YA2** transcript that allows miRNA169 binding but escapes miRNA-mediated cleavage) are nearly 3-folds more HS sensitive (Figure 5A, B). While only ~10% plants survive in transgenic plants overexpressing **At-NF-YA2** mRNAs, as much as 80-90% plants survive the HS in **At-MIR169d-OE** and **At-nf-ya2** mutant plants. This establishes that HS-tolerance in Arabidopsis by At-miR169d is mediated by down-regulation of **At-NF-YA2** functions wherein, **At-NF-YA2** acts as negative regulator of heat stress-response.

To further explore the mechanism underlying the better survival of **At-MIR169d-OE** and **At-nf-ya2** lines as well as the sensitivity of **At-NF-YA2-OE**, MIM169defg and **pNF-YA2:NF-YA2-r** lines, we analyzed the expression of several heat responsive genes reported in literature (Larkindale and Vierling 2008) in all these lines. Expression profiles of 11 HSR genes in response to HS (Supplementary table S8) revealed that the improved thermotolerance in **At-MIR169d-OE** and **At-nf-ya2** mutant plants is correlated with enhanced expression of **At-Ascorbate peroxidase2** (**At-APX2**), **At-HSFA3a**, **At-HSFA7a**, **At-HSFA7b** and **At-HSP17.ACI** genes (Figure 5C, Supplementary figure S19). In contrast, transgenic plants **At-NF-YA2-OE**, **pNF-YA2:NF-YA2-r**, MIM169defg that are
defective in heat-responsive gene regulation, exhibit reduced At-APX2, At-HSFA3a, At-HSFA7a, At-HSFA7b, and At-HSP17.ACI transcripts (Figure 5C, Supplementary figure S19). These results ascertain At-NF-YA2 as negative regulator of many HSR genes and thermotolerance in Arabidopsis.

It is well established that imposition of HS can give rise to excess levels of reactive oxygen species (ROS), resulting in cellular oxidative damage. APX plays crucial role as H$_2$O$_2$-scavenging enzyme in plant cells. Therefore, a higher level of At-APX2 transcripts in At-MIR169d-OE and At-nf-ya2 mutant plants should effectively maintain the antioxidants that protect plants. Indeed, these plants have low H$_2$O$_2$ levels as judged by DAB staining (Figure 5D). In contrast, the overexpression of At-NF-YA2 gene (At-NF-YA2-OE) and chelation of At-miR169d (MIM169defg lines) and pNF-YA2:NF-YA2-r show enhanced DAB staining due to higher H$_2$O$_2$ levels (Figure 5D). Further, cellular death estimation as a consequence of this in At-MIR169d-OE and At-nf-ya2 lines is negligible while it is much pronounced in the At-NF-YA2-OE, MIM169defg and pNF-YA2:NF-YA2-r lines, as assessed by trypan blue staining (Figure 5E).

**Arabidopsis At-NF-YA2 negatively regulates HSR genes leading to HS sensitivity**

Repressed expression of HSR genes in At-NF-YA2 abundant lines and enhanced expression of HSR genes in At-NF-YA2 down-regulated lines confirms that At-NF-YA2 is a negative regulator of these HSR genes (Figure 5C, Supplementary figure S19). At-HSP17.ACI, At-HSFA3a, At-HSFA7a, At-HSFA7b and At-APX2 promoters have NF-YA binding sites (CCAAT-box), making them probable downstream targets of At-NF-YA2 transcription factor (Figure 6A). Indeed, ChIP-qPCR and Y1H assay shows that the At-NF-YA2 protein interacts with At-HSFA3a and At-HSFA7b promoters, but not with At-HSP17.ACI, At-HSFA7a and At-APX2 promoter (Figure 6B, Supplementary figure S20). To further confirm the repression of At-HSFA3a and At-HSFA7b transcription by At-NF-YA2 in planta, GUS promoter reporter constructs Pro-At-HSFA3a:GUS and Pro-At-HSFA7b:GUS were introduced in WT, At-NF-YA2-OE and At-nfy-a2 mutant backgrounds. Expression analysis and quantification of GUS transcription using these lines confirm At-NF-YA2-mediated repression of At-HSFA3a and At-HSFA7b as GUS accumulation is reduced in At-NF-YA2-OE line and strongly expressed in At-nf-ya2
mutant background as compared to WT (Figure 6C, D). Thus, At-NF-YA2 regulates the transcription of At-HSFA3a and At-HSFA7b directly. Thermotolerance assay using At-hsfa3a and At-hsfa7a mutants show high sensitivity to HS (Figure 6E, Supplementary figure S21A, B), only 5% At-hsfa3a and 1.7% At-hsfa7a plants are able to recover as compared to 45% WT plants six-days-after HS imposition (Figure 6F, Supplementary figure S21B). This confirms the positive role of At-HSFA3a and At-HSFA7a genes in maintaining thermotolerance. Furthermore, role of At-HSFA7b gene in salt and heat stress tolerance via an E-box-like motif was recently reported by Zang et al. (2019). Thus, in Arabidopsis, HS induces At-HSFA2 that increases the expression of At-miR169d which in turn post-transcriptionally down-regulates At-NF-YA2 transcriptional repressor. At-NF-YA2 in turn enhances transcription of HSR genes like At-HSFA3a and At-HSFA7b. This regulatory feedback loop plays a critical role in thermotolerance.

Discussion

Previously we have shown heat-responsiveness of MIR169 members in tomato (Rao et al. 2020). Moreover, heat-mediated induction of miR169 in Arabidopsis (Guan et al. 2013; Li et al. 2010) and in rice (Zhao et al. 2007; Zhao et al. 2009) is also reported. This study focuses on defining the underlying upstream regulatory mechanism viz., the HSF-mediated transcriptional control of MIR169 genes as well as the post-transcriptional regulation of downstream targets of miR169s i.e., the NF-YA transcription factors, which in turn transcriptionally regulate HSR genes in both tomato and Arabidopsis.

Cooperative HSFs binding regulate MIR169 transcription in HS

At-HSFA1b and At-HSFA7b mediated induction of At-MIR398b during heat stress has been reported earlier in Arabidopsis (Guan et al. 2013). Presence of multiple HSEs in promoters foster cooperative interactions between multiple HSF trimers (Topol et al. 1985; Xiao et al. 1991; Bonner et al. 1994). The promoters of HS-responsive MIR169s exhibit multiple HSFs binding making these promoters suitable targets for cooperative binding of HSFs as evidenced by Y1H/ChIP-qPCR assays in both tomato and Arabidopsis (Figures 1A-I, 4B and 6B). The binding ability of Sly-HSFs on Sly-MIR169 promoters in tomato was further confirmed by transient assays in N. benthamiana,
where effector plasmids containing different Sly-HSFs were co-transformed with specific 
*Sly-MIR169* promoters hooked with *GUS* reporter gene. Sly-HSF-mediated activation of 
*Sly-MIR169a-1*, *Sly-MIR169b*, *Sly-MIR169c*, *Sly-MIR169d* and *Sly-MIR169d-1* 
promoters was confirmed for all Sly-HSFs except Sly-HSFA1a. Individual co-
transfection of Sly-HSFA1a with Pro-*Sly-MIR169b* was unable to induce *GUS* 
transcription. Co-infiltration of Sly-HSFA1a with Sly-HSFA7a (the other HSF binding on 
Pro-*Sly-MIR169b*) leads to strong induction of *GUS* transcripts, suggesting the 
cooperative interactions between Sly-HSFA1a and Sly-HSFA7a. The *Sly-HSFA1a-VIGS* 
study suggests that threshold level Sly-HSFA1a is sufficient to regulate transcription of 
its target genes, as portrayed by strong reduction of *Sly-MIR169b* transcription (Figure 
2D). Some studies have also reported the stress-mediated hetero-oligomeric complex 
formation of HSFA1a with other A-class HSFs to synergistically regulate the expression 
of a number of small HSPs, DnaJ/Hsp40, and HSP70 (Chan-Schaminet et al. 2009; Liu 
et al. 2011; Hahn et al. 2011; Li et al. 2014).

The HSF ‘B-class’ is known primarily as a repressor of transcription (Czarnecka-verner 
et al. 2000; Ikeda et al. 2009). We find Sly-HSFB1a-mediated transactivation of Pro-*Sly-
MIR169d* (Supplementary figure S3D). Sly-HSFB1a has been shown to activate 
transcription of Myc-HSP17.6, alone and synergistically with Sly-HSFA1a as well as 
with acidic activators like HAC1/CBP (Bharti et al. 2004). The class-A1 HSFs are 
considered as master regulators of HS response (Liu et al. 2011; Yoshida et al. 2011). 
In our study we find ‘Class-A’ HSFs regulate expression of all *MIR169* either singularly 
or in combination with other HSFs belonging to class-A/-B/-C in both tomato and 
Arabidopsis. The ‘Class-A1’ HSFs regulate 2 of 5 and 1 of 3 *MIR169* promoters in 
tomato and Arabidopsis, respectively reiterating the significance of this class in HS 
tolerance.

**Alleviation of *Sly-NF-YA9*/*Sly-NF-YA10* and *At-NF-YA2* enhances thermotolerance**

Further, analysis of thermotolerance of plants with reduced *NF-YA* levels (TRV-*Sly-NF-
YA9* and TRV-*Sly-NF-YA10* in tomato and *At-MIR169d*OE and *At-nf-ya2* knockout 
mutant in Arabidopsis) and those with higher *NF-YA* expression (At-*NF-YA2*OX, pNF-
*YA2*:NF-YA2-r and *MIM169defg*) revealed that reduced *NF-YA* levels increase
thermotolerance (higher survival rate) and there is elevated expression of HSR genes including HSF genes, APX and several HSP genes (Figure 3D-H, Supplementary figure S9 and Figure 6C, Supplementary figure S19). Thus, establishing Sly-NF-YA9 and Sly-NF-Y10 as negative regulators of HSR gene expression and thermotolerance in tomato and At-NF-YA2 in Arabidopsis. Other reports by Ceribelli et al. (2008) and Leyva-González et al. (2012) have also shown NF-YAs as repressors. Further, we have demonstrated in both tomato and Arabidopsis that the diminished expression of Sly-NF-YA10 (Rao et al. 2020) and At-NF-YA2 (Figure 2C, Supplementary figure S13A, B and supplementary figure S15A, B) upon HS is a post-transcriptional regulation which is a result of miR169-mediated cleavage and not a transcriptional regulation.

**At-NF-YA2 binds to HSR gene promoters**

Our study establishes direct At-NF-YA2 mediated transcriptional regulation of At-HSFA3a and At-HSFA7b by Y1H/ChIP-qPCR assays. Further, loss of Pro-HSFA3a:GUS, Pro-HSFA7a:GUS and Pro-HSFA7b:GUS activity in At-NF-YA2-OE lines and high GUS expression in At-nf-ya2 mutant background confirms regulation of these promoters by At-NF-YA2 as a transcriptional repressor (Figure 6C and D). Studies in literature support the involvement of HSFA3a in governing thermotolerance in Arabidopsis and tomato (Li et al. 2013; Wu et al. 2018). Larkindale and Vierling (2008) have reported At-HSFA3a as a universal candidate gene commonly induced in heat acclimated plants in different regimes of heat stress treatment. We also observe high thermo-sensitivity in At-hsfa3a mutant plants (Figure 6E). It is known that the NF-YA subunit forms functional trimer with NF-YB and NF-YC subunits to regulate transcription of a large number of genes (Warpeha et al. 2007; Kumimoto et al. 2010; Sato et al. 2014). Sato et al. (2014) have shown synergistic interaction of At-DREB2A with trimer of NF-YA2, NF-YB3 and NF-YC10 to form a transcriptional complex that activates At-HSFA3 expression in Arabidopsis protoplasts. DREB2A is also a HS inducible gene and its overexpression significantly increases HS tolerance (Sakuma et al. 2006). Moreover, Sato et al. (2014) have found heat-inducible expression of NF-YB3 and NF-YC10 but relatively stable expression of At-NF-YA2 transcripts upon HS. Here, we conclusively show that there is heat-mediated down-regulation of At-NF-YA2 (Supplementary figure
S13A, B) that is a consequence of miR169-mediated cleavage and also that At-NF-YA2 acts as a repressor of At-HSFA3a transcription (Figures 5C, 6C and 6D). Thus, we speculate that the reduced availability of NF-YA2 subunits during HS (due to degradation by miR169d) may favor functional DREB2A:NF-YC-10:NF-YB3 complex formation, which in turn up-regulates At-HSFA3a expression during HS (Figure 7).

**An HSF-mediated feedback loop regulates MIR169 in heat stress**

Furthermore, under HS, *Sly-HSFA7a* is up-regulated in *Sly-NF-YA9/A10* silenced tomato plants and there is enhanced expression of *At-HSFA7b* in *At-nf-ya2* mutant plants (Figures 3D-H, 5C, and supplementary figure S19 and table S8). We have shown that these HSFs interact with *MIR169* promoters. This suggests the existence of an HSF-mediated feedback module regulating HS-mediated transcriptional regulation of *MIR169s* directly or indirectly that may be crucial for thermotolerance in both tomato and Arabidopsis (Figures 1A, 3D-H, 4B, 5C and 7). While our study demonstrates a direct role of NF-YA class to regulate HSFs, one cannot rule out the possibility that NF-YAs regulate HSFs indirectly. It is known that NF-YA genes regulate abiotic/biotic stress response mainly via ABA pathway (Zhao et al. 2009; Luan et al. 2015; Ding et al. 2016). Similarly, HSFs are also induced in response to ABA in Arabidopsis (Huang et al. 2016). Thus, alteration in HSFs levels in *Sly-NF-YA9/10* down-regulated tomato plants or *At-nf-ya2* knockdown Arabidopsis plants could also be a result of the fluctuations in the ABA levels of the silenced plants. Moreover, the HSF perturbation during HS could also be a result of ROS-mediated HSF induction (Volkov et al. 2006; Guan et al. 2013), thus feeding to the HSF:*MIR169:NF-YA* loop. We find that heat stress induces At-HSFA2a that in turn transcriptionally activates *At-MIR169d*. At-HSFA2a has also been shown to be positively regulated by NF-YC10 together with HSFA1s (Yoshida et al. 2011; Sato et al. 2014). Overexpression of miR156 and the resultant down-regulation of target SPL genes also induce HSR genes including *At-HSFA2a* (Stief et al. 2014) (Figure 7). Moreover, our data shows other heat responsive miR169s may also act parallely to regulate NF-YAs and contribute towards heat tolerance of Arabidopsis (Figure 7). Thus, multiple pathways work in tandem, interacting among themselves to regulate cellular homeostasis during HS.
A conserved thermotolerance mechanism in tomato and Arabidopsis

It is also noteworthy, that tomato Sly-NF-YA9/A10 are the best orthologues of At-NF-YA2 (Supplementary figure S22). Also, we show that Sly-HSFA7a and At-HSFA7b regulate the transcription of MIR169 in both tomato and Arabidopsis, respectively. Interestingly, the heat-responsive At-MIR398 is also shown to be regulated by an HSFA7 class member (Guan et al. 2013). This suggests that the Sly-NF-YA9/10 and At-NF-YA2 orthologues and Sly-HSFA7-class members have conserved roles in heat tolerance in both plant species.

Our study also highlights the existence of HSF-independent pathways in regulating MIR169s transcription during HSR as HSF-mediated regulation could be established for only 5 out of 15 and 3 out of 11 MIR169 promoters of tomato and Arabidopsis, respectively (Figures 1A, 4B). Though these MIR169 loci may be indirectly controlled by other cascades regulated by HSFs like the DREB2-mediated HSFA3 regulation or NF-YA-mediated regulation as we have shown. The present study establishes a conserved strategy that involves HSF-mediated induction of miR169 in both tomato and Arabidopsis that leads to reduction of target NF-YA genes that in-turn regulate HSF transcription which feeds back to the miR169:NF-YA pathway. Down-regulation of At-NF-YA2 homologues may be a viable strategy for improving the thermotolerance and yield stability of crop plants.

Materials and Methods

Plant materials and growth conditions

Tomato cultivar CLN seeds (Balyan et al. 2020) were germinated on filter paper soaked with deionized water at 26 °C. Three-days-old seedlings of uniform growth were transplanted in plastic pots, filled with soilrite and placed in a plant growth chamber (CMP6050, Conviron, Canada) maintained at 26 °C/21 °C (day/night: 16/8 h) relative humidity 60%, light intensity 300 µM per m² per sec.

Arabidopsis thaliana (ecotype Columbia) was used as the wild-type in this study. Seeds of the MIR169d-OE, NF-YA2-OE, Pro-NF-YA2:NF-YA2-r, Pro-At-MIR169d: GUS, Pro-At-NF-YA2: GUS were from Sorin et al. (2014) study. Seeds of At-nf-ya2
(SALK_021228) were obtained from the Arabidopsis Biological Resource Center (Columbus, OH). Homozygous plants were identified by diagnostic PCR analysis using the primers listed in Table S6. Arabidopsis plants were grown in soilrite containing plastic pots and kept under cool white light of 100 μmol m−2 sec−1 intensity, with a 16 h light/8 h dark photoperiod at 22 °C/20 °C (day/night) in a plant growth chamber (CMP6050, Conviron, Canada).

One-month old tomato plants were initially exposed for 4 h at a gradual increasing range of temperature from 26 °C to 45 °C then exposed additionally for 4.5 h at 45 °C for HSFs and miR169s expression profiling study. VIGS silenced tomato plants were exposed to heat stress for 4.5 h at 45 °C while 2-weeks old Arabidopsis plants were exposed to 42 °C for 2 h heat stress. For phenotype scoring and survival assays, HS treated VIGS silenced plants and Arabidopsis plants were allowed to recover for six days at 26 °C/21 °C (day/night: 16/8 h) and at 22 °C/20 °C (day/night) in a plant growth chamber (CMP6050, Conviron, Canada). Leaves from these experimental plants were harvested immediately after HS for expression profiling of HSR genes.

Real-time quantitative RT–PCR analysis

Four (expression studies of HSFs and miR169s) and five weeks (VIGS silenced) old tomato plants and two-weeks-old seedlings of Arabidopsis were used for total RNA extraction with Trizol reagent (Invitrogen, USA). Total RNA was treated with DNase I (Ambion, USA) to remove genomic DNA contamination. For real-time RT–PCR (qRT–PCR) analysis, 2 μg of total RNA was used for cDNA synthesis by using high capacity cDNA reverse transcription kit (Applied Biosystem, USA) following the manufacturer’s protocol. The cDNA reaction mixture was diluted 2-fold, a 0.5 μl aliquot was used as a template in a 10 μl PCR reaction. PCR reactions included a pre-incubation at 95°C for 2 min, followed by 40 cycles of denaturation at 95°C for 15 sec, annealing at 60°C for 40 sec, and extension at 72°C for 45 sec. All the reactions were performed in a CFX-Connect real-time PCR detection system (Bio–Rad, www.bio-rad.com/) using GO Taq qPCR master mix (Promega, USA). Each experiment was repeated at least three times. The relative fold change was calculated by following the $2^{-ΔΔCT}$ method using Actin and Tubulin gene as endogenous control (Paul et al. 2016). The stability and similarity in the
Ct range of *Actin* and *Tubulin* were assessed across the experimental conditions (Supplementary table S9). For calculation of fold change in the gene expression the control value was set as one for all qRT-PCR expression analysis. The primers used in this study are listed in Supplementary table S10 and all the qRT-PCR primers having a good amplification efficiency ranging between 90-110% were used (Supplementary table S11).

For miRNA expression analysis 2 µg of the small RNA was polyadenylated using the Poly(A) Tailing kit (epicenter- An Illumina company) following the manufacturer's protocol. The above poly-A tailed small RNA reaction mixture was then reverse transcribed with a special 100 bp primer i.e. miR_oligodT_RTQ (Mutum et al. 2013) which is composed of two variable bases at its end followed by the poly(T)25 and then with a random adapter sequence using the Superscript III Reverse Transcriptase (Invitrogen, USA). The taqman based q-RT-PCR was performed using the TaqMan Fast Universal PCR Master Mix (2X, Applied Biosystems) and ‘Fam (Fluorescein) dye’ and BHQ labeled taqman universal probe specific to the adapter region of the miR_oligodT_RTQ (Supplementary table S8) which enable the use of a common reverse primer (Mutum et al. 2013) and miRNA specific forward primer in the expression analysis. For each well a total reaction volume of 7 µl was prepared and for each miRNA two technical with 3 biological replicates were analyzed, *5S rRNA* and *U6snRNA* (Supplementary table S8) were used as the endogenous control for data normalization. For calculation of fold change in the gene expression the control value was set as one for all qRT-PCR expression analysis.

**Promoter sequences analysis**

The 2 kb sequences, upstream of the precursor start site of *MIR169* precursors and translational start codon (ATG) of Arabidopsis HSF genes were downloaded from Sol Genomic Network database (http://sgn.cornell.edu) and TAIR, respectively and analyzed for the presence of putative *cis*-regulatory elements by PlantPAN 3.0 (Chow et al. 2019). Positions of heat stress elements (HSE) were also searched manually in miRNA169 promoters. We searched for both perfect and imperfect HSE types manually,
as these imperfect variants also have been validated for HSFs binding by different in vitro and in vivo assays (Santoro et al. 1998; Mittal et al. 2011; Guo et al. 2008).

**Yeast one-hybrid assays**

The coding sequences for 23 tomato HSFs and 21 Arabidopsis HSFs were cloned into the pGADT7 vector (Clontech) as a prey (Supplementary table S8). DNA fragments corresponding to the promoters (250 to 1500 bp) of heat responsive *MIR169* genes from tomato and Arabidopsis (Supplemental tables S7, S3) were cloned into the pABAI plasmid (Clontech) as baits. For positive control, sHSP promoter previously reported to have HSFA3a binding (Li et al. 2013) was cloned. Promoter-specific bait yeast strains were generated by transforming Y1HGold yeast strain with 1 µg of linearized bait plasmids and plated on SD-ura medium. Positive bait strains having insertion at the URA3 locus were selected by performing colony PCR using primers specific to the bait sequences (Supplementary table S10). To negate this bait specific background *AbAr* expression, minimal inhibitory concentrations for all 27 baits (miRNA promoters) were determined by spotting yeast cultures at OD$_{600}$ ranging from 0.1 to 0.0001 on SD/-Ura medium supplemented with different concentration of aureobasidin A (100- 1000 ng/ml; Supplementary tables S6, S12). For one-on-one bait/ prey interaction, bait strains were transformed with pGADT7 vector containing CDS of tomato HSFs. Positive bait/prey containing yeast colonies were selected by plating the transformed yeast cells on SD/-Leu medium. To study the interaction between different tomato HSFs and miRNA promoters, specific bait/prey containing yeast strains were spotted on SD/-Leu/AbA medium having bait specific aureobasidin A concentration.

**Chromatin immunoprecipitation (ChIP)-qPCR assays**

Promoters for tomato *Sly-MIR169a-1, Sly-MIR169b, Sly-MIR169c, Sly-MIR169d* and *SlyMIR169d-1* were amplified from the genomic DNA and cloned into pbi101 vector upstream of the *GUS* reporter gene. All transcription factors (*Sly-HSFA1a, Sly-HSFA7a, Sly-HSFA8a, Sly-HSFB1a, Sly-HSFA4a, Sly-HSFA6b* and *Sly-HSFA1e* CDS) were cloned in pCAMBIA1302 in fusion with the *GFP* reporter. The binary constructs were co-infiltrated as per the combinations obtained from Y1H assay into 4-weeks-old *N. benthamiana* leaves via *Agrobacterium tumefaciens* (strain EHA105). These leaves
were harvested 2 days after the infiltration and ChIP assays was performed using four infiltrated leaves from five independent plants for each co-infiltration. To identify the binding of At-NF-YA2 on HSR gene promoters, NF-YA2:GFP overexpressing transgenic lines were used. Leaves from *N. benthamiana* plants (for tomato ChIP) and 3-week-old NF-YA2:GFP overexpressing Arabidopsis plants were cross-linked with 1% formaldehyde. Chromatin was sheared to an average length of 500bp by sonication, and immunoprecipitated with GFP tag-specific monoclonal antibody (abcam; catalogue no. ab290). The fold enrichment was calculated for different promoters by performing real-time quantitative PCR analysis on immunoprecipitated DNA using a Bio–Rad CFX96 real-time PCR detection system. For each promoter pair used for ChIP-qPCR the primer amplification was first tested on input DNA and the fold enrichment was checked by keeping no-antibody, IgG and input DNA controls. Data for enrichment of Sly-HSFA6b on Sly-MIR169d-1 promoter is presented as an example in Supplementary figure S23. All the primers used for ChIP-qPCR are listed in Supplementary Table S10.

**Transient promoter reporter assay to confirm HSFs-mediated regulation of Sly-MIR169 promoters in *N. benthamiana***

Agro-infiltration was performed in 4-weeks-old *N. benthamiana* leaves. For promoter reporter assays Pro-Sly-MIR169s:GUS and p35S:Sly-HSFs were transformed in *Agrobacterium tumefaciens*. For co-infiltrations, overnight cultures of individual constructs were harvested and suspended at OD 1 in infiltration buffer (10 mM MgCl2, 10 mM MES, pH 5.8, 0.5% glucose and 150 μM acetosyringone). After 3 h of incubation at room temperature, various combinations were mixed at a 1:1:1 ratio; these mixes were used to infiltrate leaves. These leaves were harvested 2 days after the infiltration and used for RNA extraction and qRT-PCR as described above. The expression of GUS transcript is measured by qRT-PCR and presented as fold-change values calculated using the 2−ΔΔCt method. The values were normalized with NPTII that is co-expressed in the promoter: GUS construct. The GUS expression from the reporter (pro-Sly-MIR169:GUS) infiltrated samples was set to one to normalize the background signal.

**Virus induced gene silencing**
Transient silencing assays following VIGS was performed in tomato plants as described by Senthil and Mysore (2014). For silencing of selected genes, 300 to 400 bp cDNA fragments designed using VIGS tool (Sol Genomics Network) were amplified and cloned into the Eco RI–Bam HI site of the TRV2 vector. After confirmation by Sanger sequencing, the TRV2-gene vectors were transformed into A. tumefaciens strain GV3101. About 15-days-old tomato plants were inoculated with a mixture of A. tumefaciens strains containing the TRV2 vector constructs and TRV1 (a helper plasmid, contains RNA-dependent RNA polymerase (RdRp), movement protein (MP), and a 16 kDa cysteine-rich protein) into the first and second true leaf. Empty TRV2 vector was used as a control. Infected plants were kept in dark and moist conditions for 3 days and then transferred to light; HS was imposed after 3 weeks (the time required for appearance of chlorophyll bleaching phenotype in the leaves of TRV-Phytoene desaturase-infiltrated plants used as silencing control) (Supplementary figure S24A, B) at 45 °C for 4.5 h. Leaves were frozen for qRT-PCR analysis. Off targets of the selected VIGS constructs were also predicted at VIGS tool (Supplementary table S4) and validated by qRT-PCR in the silenced plants. To confirm the maintenance of silencing throughout the phenotype scoring and biochemical assays time period, the expression of Sly-NF-YA9 and Sly-NF-YA10 was assessed further during the post heat stress period. Survival and cell death was assayed after 6 days’ recovery. The experiment was performed in four biological replicates with 8 plants per replicate for each gene.

Gas-exchange parameters measurements
Gas-exchange parameters, like water use efficiency (WUEi), net photosynthetic rate (A), and transpiration rate (E) were measured simultaneously by using a portable Licor 6400 photosynthesis system (LI-6400, Li-Cor Inc., Lincoln NE, USA). WUEi was calculated using the formula A/E. For all measurements, sixth and seventh fully expanded tomato leaves of 6 plants for each VIGS construct were used and the experiment was repeated thrice keeping similar parameters. The measurement conditions were as follows: leaf temperature 27°C, leaf-air vapour pressure deficit 1.5±0.5 kPa, photosynthetic photon flux 300 μmol m⁻² s⁻¹, relative air humidity 60% and ambient CO2 concentration 400±5 μmolmol⁻¹.

Histochemical detection of H₂O₂ accumulation and cell death in HS
Hydrogen peroxide (H$_2$O$_2$) levels and cell death were estimated by using 3,3'-diaminobenzidine (DAB) and trypan blue stain, respectively on the 6th and 7th leaves of 6-week-old TRV-Vector and TRV-gene plants immediately after heat stress (45 °C for 6 h). DAB staining was performed as previously described (Daudi et al. 2012). The viability of leaf cells was measured using the trypan blue exclusion method (Koch and Slusarenko 1990). Experiment was repeated twice with similar parameters having a sample size of 8 plants per replicate.

**GUS assays**

To analyze the histochemical localization of GUS reporter protein, tissue samples were incubated for 12 h at 37 °C with the substrate solution (1 mM 5-bromo-4-chloro-3-indolyl-β-D-glucuronide, pH 7.0, 100 mM sodium phosphate buffer, 10 mM Na$_2$EDTA, 0.5 mM potassium ferricyanide, 0.5 mM potassium ferrocyanide, and 0.1% Triton X-100). Subsequently, seedlings and leaves were washed with de-staining solution containing ethanol:acetone:glycerol (3:1:1) to eliminate chlorophyll. They were photographed with a Nikon SMZ1000 Stereomicroscope (Tokyo, Japan) and OLYMPUS SZX16 Stereomicroscope (India). GUS transcripts were quantified using qRT-PCR. The assays were performed with three biological replicates.

**Constructs generation**

For yeast one hybrid assay, all the promoters and CDS of transcription factors (HSFs and NF-YA) were cloned in pABAI and PGADT7 vector, respectively. For promoter reporter assays, all promoters were cloned in pbi101 vector and all transcription factors (Sly-HSFA1a, Sly-HSFA7a, Sly-HSFA8a, Sly-HSFB1a, Sly-HSFA4a, Sly-HSFA6b and Sly-HSFA1e CDS) were cloned in pCAMBIA1302. Promoters were amplified as approximately 2000 bp region upstream of the precursor start site for MIR169s (Pro-Sly-MIR169a-1:GUS, Pro-Sly-MIR169b:GUS, Pro-Sly-MIR169c:GUS, Pro-Sly-MIR169d:GUS, Pro-Sly-MIR169d-1:GUS, Pro-At-MIR169d:GUS) and translation start site for HSFs (Pro-At-HSFA3a:GUS and Pro-At-HSFA7b:GUS) and Pro-NF-YA2:GUS from genomic DNA. For overexpressing miR169d (p35S:At-MIR169d-OE) a 300 bp fragment surrounding the miRNA sequence was amplified from genomic DNA, cloned in pbi121 and transformed into Arabidopsis WT plants by floral dip method. For VIGS,
300-400 bp fragment of *Sly-HSFA1a, Sly-HSFA7a, Sly-HSFB1a, Sly-HSFA6b* and *Sly-NF-YA9, Sly-NF-YA10* were cloned in pTRV-2 vector.

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**Author contributions**

SM designed and supervised the study; SR designed and performed all the experiments and analyzed the data; CB did mutant genotyping for hsfa2 and nf-ya2; CS generated MIR169, NF-YA2 and NF-YA2-r transgenic lines of Arabidopsis. SR and SM wrote the article; CS and MC complemented the writing and critically reviewed the manuscript. SM agrees to serve as the author responsible for contact and ensures communication.

**References:**


specifically enhances heat stress-induced gene expression by forming a heat stress-
specific transcriptional complex with NF-Y subunits. The Plant Cell, 26 4954-4973.

transcription factor (Hsf) family: structure, function and evolution. Biochimica et


Sorin C, Declerck M, Christ A, Blein T, Ma L, Lelandais-Brière C, Njo MF, Beeckman T,
Crespi M and Hartmann C (2014) A miR169 isoform regulates specific NF-YA targets

miR156 regulates tolerance to recurring environmental stress through SPL transcription
factors. The Plant Cell, 26:1792-1807.


for comprehensive analysis of microRNA function in Arabidopsis thaliana. PLoS
Genetics, 6: e1001031.


microRNAs in banana during acquired thermo tolerance. Journal of Horticultural

required for effective expression of heat shock genes in Arabidopsis. Plant Molecular
Biology, 61, 733-746.


**Figure Legends**

**Main Figures:**

**Figure 1:** Yeast one hybrid and Chromatin immunoprecipitation (ChIP-qPCR) assay of Sly-MIR169 promoters and HSFs. (A) Yeast-one-hybrid assay to assess transcriptional regulation of heat-responsive *MIR169* promoters by HSFs. Only the
positive Y1H interactions between different HSFs and Sly-MIR169 promoters have been shown. The Sly-MIR169 promoters were cloned in the reporter vector pABAi and the full-length cDNA of HSFs were cloned in the pGADT7 vector. After the co-transformation into Y1H gold yeast strain, serial dilutions (O.D. 0.1 to 0.0001) of yeast cultures were spotted onto plates lacking URA and LEU with specific Aureobasidin A concentration and incubated for 3 days. Promoter of sHSP has been used as a positive control to show binding of HSFA3a on sHSP promoter (Li et al. 2013) and empty vectors as negative controls. The Aureobasidin A concentrations for all Sly-MIR169 promoters is presented in supplementary table S7. (B-I) Chromatin immunoprecipitation (ChIP-qPCR) assay of Sly-MIR169 promoters and HSFs in Nicotiana benthamiana. Fold enrichment of Sly-MIR169a-1 promoter in HSFA1e immunoprecipitated sample (B); Fold enrichment of Sly-MIR169b promoter in HSFA1a immunoprecipitated sample (C); in HSFA7a immunoprecipitated sample (D); Fold enrichment of Sly-MIR169C promoter in HSFA4a immunoprecipitated sample (E); Fold enrichment of Sly-MIR169C promoter in HSFA6b immunoprecipitated sample (F); Fold enrichment of Sly-MIR169d promoter in HSFA8a immunoprecipitated sample (G); in HSFB1a immunoprecipitated samples (H); Fold enrichment of Sly-MIR169d-1 promoter in HSFA6b immunoprecipitated samples (I). These experiments were repeated 3-4 times; the error bars represent standard deviation between the replicates. The values for different promoters in IgG immunoprecipitated sample were set as one for normalization.

Figure 2: Validation of HSF:miR169:NF-YA node components in tomato upon heat stress. (A) Expression profiles of HSF genes that regulate Sly-MIR169 transcription in WT tomato leaves during control and heat stress condition by qRT-PCR using the 2−ΔΔCt method. (B) Expression of mature miR169s in response to heat stress as determined by taqman based qRT-PCR in control and heat challenged WT tomato leaves. (C) qRT-PCR based expression profiles of target NF-YA transcripts in WT control and heat stressed tomato leaves using the 2−ΔΔCt method. (D-G) The expression analysis of HSF genes, Sly-MIR169 precursors and NF-YA transcripts in VIGS plants silenced for Sly-HSFA1a (D), Sly-HSFA7a (E), Sly-HSFA6b (F) and Sly-HSFB1a (G). In (D-G) 15-days-old tomato plants were agro-infiltrated with empty vector (TRV-1, TRV-2 VC) or TRV-Sly-HSF VIGS constructs. The VIGS established plants
were subjected to heat stress after 3-weeks-of agro-infiltration, and used for expression studies of different genes. Graphical data represents mean values of expression of three to four biological sets. Error bars show standard deviation. To represent the negative fold-change on y axis in (C-G), the relative expression values of down-regulated genes were transformed using the formula \[-(1 / RQ \leq 0.5)\]. Actin was used as endogenous control in (A), (C) and (D-G), while 5SrRNA was used as reference control in (B). The fold change normalization was done by setting the control value as one for all qRT-PCR expression analysis.

**Figure 3. Functional validation of the role of Sly-NF-A9 and Sly-NF-A10 in HS response in tomato.** (A) Phenotype of TRV-VC (Vector control), TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 silenced plants after heat stress. Fifteen-days-old tomato plants were agro-infiltrated with empty vector (TRV-VC), TRV-Sly-NF-YA9 or TRV-Sly-NF-YA10 silencing constructs. Plants were given HS 3-weeks-after infiltration. Phenotype of HS-treated silenced plants was scored six-days-post HS. Experiments were repeated 3-4 times with 8 plants per replicate. (B) Percent survival of TRV-EV, TRV-Sly-NF-YA9 and TRV-Sly-NFYA10 silenced plants after HS. (C) Estimation of water use efficiency (mmol mol⁻¹), net photosynthesis rate (μmol m⁻²s⁻¹), and transpiration rate of TRV-EV, TRV-Sly-NF-YA9 and TRV-Sly-NFYA10 silenced plants after HS. (D-E) Expression profiles of HSR genes in control and heat treated TRV-EV, TRV-Sly-NF-YA9 and TRV-Sly-NFYA10 silenced plants. The expression levels of genes were calculated using the 2−ΔΔCt method and presented using fold-change values. The fold change normalization was done by setting the control value as one in (D-H). Actin was used as endogenous control. Error bars represent the standard deviation of two independent biological replicates. *p<0.05, **p<0.01 and ***p<0.001 vs. wild type, by two-tailed Student’s t-test.

**Figure 4. Transcriptional regulation of MIR169s promoters by HSFs in Arabidopsis.** (A) miR169s expression in response to heat stress as determined by Taqman-based qRT-PCR of control and heat challenged plants. 5S rRNA was used as the normalization control. Experiment was repeated three times and average values are plotted as bars, error bars depict standard deviation between the replicates. (B) Y1H
assay showing the positive interaction of HSFs with the promoters of different At-MIR169s, Y1H assay shows binding of At-HSFA7b, At-HSFB2b, At-HSFB3a, At-HSFA2 and At-HSFA1b on At-MIR169b, d and h promoters. This experiment was repeated at least three times with similar results, and data from one representative experiment is shown. Serial dilutions (O.D. 0.1 to 0.0001) of yeast cultures were spotted onto plates lacking URA and LEU with specific Aureobasidin A concentration and incubated for 3 days. The Aureobasidin A concentrations for all At-MIR169 promoters is presented in supplementary table S3. (C) Expression levels of At-MIR169d in wild-type plants subjected to 0 (control) and 2 h heat stress at 42 °C. Data are shown as means ± SD of three biological replicates and normalized to actin reference gene using (2−ΔΔCT) method. (D) Expression patterns of Pro-At-MIR169d:GUS lines of transgenic Arabidopsis plants subjected to 0 (control) or 2 h of heat stress at 42°C. (E) Levels of GUS transcripts in Pro-At-MIR169d:GUS lines, measured by qRT-PCR after 0 or 2 h of heat stress at 42°C. (F) Transcriptional activity of Pro-At-MIR169d:GUS reporter construct in WT and At-hsfa2 mutant background during 2 h of heat stress at 42°C. (G) Expression of GUS transcript driven by Pro-At-MIR169d and At-MIR169d precursor in soil-grown wild-type plants of Arabidopsis and At-hsfa2 mutant. To represent the negative fold-change on y axis in figure (G), the relative expression values of down-regulated genes were transformed using the formula [-1 divided by RQ ≤ 0.5]). The fold change normalization was done by setting the control value as one in Figure (A, C, E and G). Error bars represent the standard deviation. These experiments were repeated at least three times with similar results, and data from one representative experiment are shown in D and F.

**Figure 5: Assaying the miR169d:At-NF-YA2 node in heat stress in Arabidopsis.** (A) Thermotolerance was gauged in At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NF-YA2-r and MIM169defg transgenic plants in comparison to wild-type (WT) plants. Two-weeks-old soil-grown plants were subjected to 0 (control) or 2h heat stress at 42°C, and damage was recorded 6 days later. (B) Estimation of survival (percentage) of HS treated At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NF-YA2-r and MIM169defg plants. Plants were assayed for heat stress tolerance after 1 week of HS. These experiments were repeated at least five times with similar results, and average
data of all experiments are shown. Error bars, ± SD of five repeats. (C) Expression patterns of heat stress-responsive genes in WT, At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NYA2-r and MIM169defg plants. Data are shown as means ± SD of biological replicates and normalized to Actin using \(2^{-\Delta\Delta CT}\) method. Log2 transformation was applied to the RQ data to obtained negative fold change. (D-E) DAB and Trypan blue staining of HS treated At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NYA2-r and MIM169defg plants to estimate the production of ROS and cell death, respectively. *p<0.05, **p<0.01 and ***p<0.001 vs. wild type, by two-tailed Student’s t-test. Three independent lines of MIR169d-OE were used and similar results were obtained for all studied parameters, a single representative picture was shown in the figure A, B, D and E. The fold change normalization was done by setting the control value as one for figure (C).

**Figure 6: Transcriptional regulation of At-HSFA3a and At-HSFA7b by At-NF-YA2 during heat stress in Arabidopsis.** (A) Graphical representation of CCAAT cis-elements in At-HSR gene promoters HSFA3a, HSFA7a, HSFA7b, APX and sHSP17.6A. The 2 kb upstream sequences of start codon were analysed by plant care and Plant CAN promoter analysis tools. The positions of CCAAT boxes are depicted by violet coloured tabs. (B) Chromatin immunoprecipitation (ChIP)-qPCR assay of At-NF-YA2 on At-HSR gene promoters. (C-D) In-planta estimation of transcriptional regulation of At-HSFA3a and At-HSFA7b transcription by At-NF-YA2 during heat stress. (C) Pro-At-HSFA3a:GUS reporter construct was agro-infiltrated in WT, At-NF-YA2-OE and At-nf-ya2 backgrounds. GUS was visualized 2-days post-infiltration by histochemical staining. (D). qRT-PCR based quantification of GUS transcripts in promoter reporter constructs infiltrated plants, 2-days-post infiltration by using GUS specific primers. The NPTII gene present in the same vector backbone was used as reference control. To represent the negative fold-change on y axis in, the relative expression values of down-regulated genes were transformed using the formula \(-\left(1 \text{ divided by RQ} \leq 0.5\right)\). The fold change normalization was done by setting the control value as one. (E) Thermotolerance assay of WT and At-hsfa3a mutant plants. Two-weeks-old soil-grown plants were subjected to 0 (control) or 2h heat stress at 42°C, and damage was recorded 6 days post-stress. (F) Estimation of survival (percentage) of WT and At-
hsfa3a plants 6 days post heat stress. The experiments were repeated at least three times with similar results, and average data of all experiments are shown. Error bars, ± SD of three repeats. *p<0.05, **p<0.01 and ***p<0.001 vs. wild type, by two-tailed Student’s t-test.

**Figure 7:** Model for HSF-induced miR169:NF-YA module in mediating thermotolerance in tomato and Arabidopsis. Heat stress induces the expression of HSFs by altering the cellular ROS levels as well as other cues independent of ROS signaling. HSFA1a, HSFA2 and HSFA7a are the key player of plant HSR response. These HSFs bind to the promoters of MIR169 leading to transcriptional enhancement of miR169s in both tomato and Arabidopsis. Enhanced accumulation of miR169s reduces the levels of Sly-NF-YA9/A10 in tomato and At-NF-YA2 in Arabidopsis that leads to enhancement of HSR genes expression like HSFA2, HSFA3 and HSFA7s, possibly via altering ABA levels. In Arabidopsis, At-miR169-mediated down-regulation of At-NF-YA2 during HS may favor functional DREB2A:NF-YC-10:NF-YB3 complex formation to up-regulate At-HSFA3a expression during HS. Enhanced levels of HSFAs in turn feed the transcription of MIR169 genes. The heat-mediated orchestration of parallel miRNA pathways (miR169:NY-YA, miR398:CSD and miR156:SPL) alters the levels of HSR genes and regulate plant thermotolerance. Solid lines denote a confirmed interaction and dotted lines are predictive. Red and black lines denote findings of this study and existing literature, respectively. Black colored numerals refer to the studies that established the particular connection between different components of the model.


**Supplementary Figure legends:**

**Supplementary figure S1:** Promoter analysis of MIR169 genes in tomato. Distribution of perfect and imperfect heat stress elements (HSE) in MIR169 promoters.
of tomato. HSEs were identified manually by curating a list from published literature and marked as pink (perfect HSE) and turquoise blue (imperfect HSE) colored shapes on black colored lines representing MIR169 promoters. The red right-handed arrow represents precursor start site. The HSE sequence variants for perfect and imperfect HSEs is provided in supplementary table S1.

Supplementary figure S2: Yeast-one-hybrid assay of 24 tomato HSFs on Sly-MIR169d-1 promoter. Interaction of only Sly-HSFA6b on Sly-MIR169d-1 promoter highlights the specificity of the Y1H assay. The co-transformed Y1H gold yeast strain cultures were spotted (O.D. 0.1 to 0.0001) onto plates lacking URA and LEU with and without specific Aureobasidin A concentration and incubated for 3 days. Promoter of sHSP has been used as a positive control to show binding of HSFA3a on sHSP promoter (Li et al. 2013) and empty vectors as negative controls. The positive binding of HSFA6b on Sly-MIR169d-1 promoter was assessed at 200 ng/ml Aureobasidin A concentration.

Supplementary figure S3: HSF-mediated transcriptional activation of tomato MIR169 promoters. In-planta transient assays showing HSF-mediated transcriptional activation of Sly-MIR169:GUS promoters in Nicotiana benthamiana as assessed by qRT-PCR. The fold-change of GUS transcripts were calculated using the 2−ΔΔCt method in (A) to (E). (A) Sly-HSFA1e with pro-Sly-MIR169a-1:GUS. (B) Sly-HSFA7a and Sly-HSFA1a with pro-Sly-MIR169b:GUS. (C) Sly-HSFA6b with pro-Sly-MIR169c:GUS. (D) Sly-HSFB1a with pro-Sly-MIR169d:GUS. (E) Sly-HSFA6b with pro-Sly-MIR169d-1:GUS. The NPTII gene co-expressed in the pro-Sly-MIR169:GUS construct was taken as the reference gene for normalization. The GUS expression from the reporter (pro-Sly-MIR169:GUS) infiltrated samples was set to one for normalization (marked as green bars). These experiments were repeated at least six times; the error bars represent standard deviation between the biological replicates.

Supplementary figure S4: Expression profiling of HSFs:miRNAs:NF-YAs during heat stress in tomato leaves. (A) Expression profiles of HSF genes that regulate Sly-MIR169 transcription in WT tomato leaves during control and heat stress condition by...
qRT-PCR using the 2−ΔΔCt method. (B) Expression of mature miR169s in response to heat stress as determined by taqman based qRT-PCR in control and heat challenged WT tomato leaves. (C) qRT-PCR based expression profiles of target NF-YA transcripts in WT control and heat stressed tomato leaves using the 2−ΔΔCt method. Data represents mean values and standard deviation of biological replicates. Tubulin was used as endogenous control in (A) and (C), while Sly-U6snRNA was used as normalization control in (B). Error bars in (A), (B) and (C) represent the standard deviation of biological replicates. The fold change normalization was done by setting the control value as one for all qRT-PCR expression analysis.

Supplementary figure S5: Determining the specificity of HSFs silencing in VIGS silenced tomato plants. (A-C) qRT-PCR based expression profiles of putative off target genes using the 2−ΔΔCt method. (A) Relative expression of Sly-HSFA1b and Sly-HSFA1c off targets in TRV-Sly-HSFA1a silenced plants. (B) Relative expression of Sly-HSFA6a off target in TRV-Sly-HSFA6b silenced plants. (C) Relative expression of Sly-HSFA4c and Sly-HSFB4a off targets in TRV-Sly-HSFA7a silenced plants. Error bars in (A-C) represent the standard deviation of biological replicates. ACTIN was used as endogenous control in A-C. Data normalization and fold change values with TUBULIN were presented in Supplementary table S3. The fold change normalization was done by setting the TRV-EV expression as one for all qRT-PCR expression analysis.

Supplementary figure S6: In planta validation of heat governed HSFs mediated transcriptional regulation of MIR169s and NF-YAs in tomato leaves.

The expression analysis of HSF genes, Sly-MIR169 precursors and NF-YA transcripts in VIGS plants silenced for Sly-HSFA1a (A), Sly-HSFA7a (B), Sly-HSFA6b (C) and Sly-HSFB1a (D). 15-days-old tomato plants were agro-infiltrated with empty vector (TRV-EV) or TRV-Sly-HSF VIGS constructs. The VIGS established plants were subjected to heat stress after 3-weeks-of agro-infiltration, and used for expression studies of different genes. Graphical data represents mean values of expression of three to four biological sets. Error bars show standard deviation. To represent the negative fold-change on y axis, the relative expression values of down-regulated genes were transformed using the formula \[-(1 / RQ ≤0.5)]]. Tubulin was used as endogenous reference control. The
fold change normalization was done by setting the control value as one for all qRT-PCR expression analysis.

**Supplementary figure S7: Virus induced silencing of Sly-NF-YA9 and Sly-NF-YA10.** (A) qRT-PCR analysis of Sly-NF-YA9 and Sly-NF-YA10 gene in TRV-EV, TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 plants in control (non-stressed) conditions confirming their silencing. The fold change normalization was done by setting the TRV-EV expression as one for all qRT-PCR expression analysis. (B) Expression of Sly-NF-YA9 and Sly-NF-YA10 in WT tomato and TRV-EV infiltrated plants in control (non-stressed) conditions. The fold change normalization was done by setting the WT control value as one for all qRT-PCR expression analysis (C-D) Expression profiles of Sly-NF-YA9 (C) and Sly-NFY-A10 (D) in TRV-Sly-NF-YA9/A10 VIGS silenced plants in heat stress (HS) in comparison to control non-stressed conditions. The fold change normalization was done by setting the control value as one for all qRT-PCR expression analysis. The expression levels of genes were calculated using the 2−ΔΔCt method and presented using fold-change values. Actin was used for normalising expression in (A-D). Fold change values for (A-D) with TUBULIN normalisation were presented in Supplementary table S3. EV: vector control. Error bars represent the standard deviation of biological replicates.

**Supplementary figure S8: Determining the specificity of NF-YA9/A10 silencing in VIGS silenced tomato plants.** (A-B) qRT-PCR based expression profiles of putative off target genes using the 2−ΔΔCt method. (A) Relative expression of Sly-NF-YA1, Sly-NF-YA3 and Sly-NF-YA10 off targets in TRV-Sly-NF-YA9 silenced plants. (B) Relative expression of Sly-NF-YA9 (no off targets were predicted for TRV-Sly-NF-YA10, we used Sly-NF-YA9 to show specificity of silencing) as in TRV-Sly-NF-YA10-VIGS silenced plants. Error bars in (A-B) represent the standard deviation of biological replicates. ACTIN was used as endogenous control in A-B. Fold change values for (A-B) with Tubulin normalisation were presented in Supplementary table S3. The fold change normalization was done by setting the TRV-EV expression as one for all qRT-PCR expression analysis.
Supplementary figure S9: Expression analysis of heat stress responsive genes in control and heat treated TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 silenced tomato plants.

(A-E) Expression profiles of HSR genes in control and heat treated TRV-EV, TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 silenced plants. qRT-PCR based fold change in expression in control and heat stressed plants of Sly-HSFA2 (A); Sly-HSFA7a (B); Sly-HSFA3a (C); Sly-APX (D) and Sly-HSP17.6 CII. (E). Relative expression of HSR genes in TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 silenced plants in control conditions as compared to TRV-EV plants. (G) Expression profiles of HSR genes in control and heat treated TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 silenced plants. The expression levels of genes were calculated using the 2−ΔΔCt method and presented using fold-change values. Tubulin was used as reference gene for A-E and Actin was used as endogenous control for F-G. Fold change values for (F-G) with TUBULIN normalisation were presented in Supplementary table S3. The fold change normalization was done by setting the expression in control non-stressed as one for all qRT-PCR expression analysis in (A-E). The fold change normalization was done by setting the TRV-EV heat stressed expression as one for all qRT-PCR expression analysis in (F). The fold change normalization was done by setting the expression of TRV-EV in control non-stressed as one for all qRT-PCR expression analysis in (G). Error bars represent the standard deviation of independent biological replicates.

Supplementary figure S10: Expression profiling of At-miR169s in heat stress.

Expression of mature At-miR169s in response to heat stress as determined by Taqman-based qRT-PCR. U6snRNA was used as the endogenous normalization control. The average values of multiple biological replicates are plotted as bars, error bars depict standard deviation between three replicates. The fold change normalization was done by setting the control value as one for all qRT-PCR expression analysis.

Supplementary figure S11: Expression analysis of At-MIR169d in heat stress in Arabidopsis.

Expression levels of At-MIR169d in wild-type plants subjected to 0
(control) and 2 h heat stress at 42 °C. Data are shown as means ± SE of three biological replicates and normalized to At-EF-α reference gene using \(2^{-\Delta\Delta CT}\) method. The fold change normalization was done by setting the control value as one for the qRT-PCR expression analysis.

**Supplementary figure S12: Schematic structure of At-HSFA2 gene carrying the T-DNA insertion and lack of At-HSFA2 expression in the At-hsfA2 mutant (Salk_008978).** (A) The two exons of At-HSFA2 genomic DNA are shown according to the gene features annotated at TAIR database. T-DNA insertion site is shown as inverted triangle in the second exon of At-HSFA2. The exon sequences are depicted by red and the UTRs are represented by green color. (B) Genotyping PCR of At-hsfa2 mutants using genomic DNA, presence of border specific amplification in At-hsfa2 confirms their homozygosity, while presence of only gene specific amplification in control Columbia plants confirms their wild type nature. (C) Expression of At-HSFA2 was determined by RT-PCR, using full length cDNA from WT and mutant At-hsfa2 plants. Absence of At-HSFA2 specific band in mutant plants confirms the knockout nature of At-hsfa2 homozygous mutants. RNA was isolated from detached mature leaves of the wild-type (wt) or At-hsfa2 plants. Expression of actin gene is shown as a loading control.

**Supplementary figure S13: Expression pattern of miRNA169d:NF-YA in Arabidopsis.** (A) Expression levels of mature miR169d and eight miR169 target genes in wild-type plants subjected to 0 (control) and 2 h heat stress at 42°C using qRT-PCR. Data are shown as means ± SD of biological replicates and normalized to actin and 5SrRNA reference genes, for mRNA and miRNA respectively, using \(2^{-\Delta\Delta CT}\) method for A and with At-EF-α reference gene and U6snRNA for mRNA and miRNA respectively, using \(2^{-\Delta\Delta CT}\) method for B. The fold change normalization was done by setting the control value as one for all qRT-PCR expression analysis.

**Supplementary figure S14: Establishing miRNA169:target modules operating in Arabidopsis.** (A) Taqmann based qRT- qPCR analysis of the accumulation of mature miR169abc, miR169defg, miR169hijklmn and miR171bc (used as negative control) in
Arabidopsis MIM169defg lines. Average values of expression for different miR169s in three independent lines of MIM169defg transgenic plants were plotted as bars and error bars depict the standard deviation between the replicates. (B) qRT-PCR analysis of the accumulation of the At-NF-YA transcripts in MIM169defg Arabidopsis lines. Data are presented as average fold induction relative to control where induction is the ratio of transcript levels in two independent MIM169defg transgenic lines to those in wild-type plants (WT). Error bars, ± SD of three repeats. (C) qRT-PCR analysis of the reduction of At-NF-YA2 transcripts in three independent miR169d over expressing (OE) lines. Data are presented as average fold in two independent MIM169defg lines relative to WT control, that was set as one.

Supplementary figure S15: Heat stress mediated transcriptional regulation of At-NF-YA2. (A) Histochemical GUS expression of Pro-At-NF-YA2:GUS transgenic Arabidopsis plants during control and heat stress. (B) qRT-PCR based determination of GUS transcripts of Pro-At-NF-YA2:GUS transgenic Arabidopsis plants during control and heat stress. These experiments were repeated 3 times with 20 seedling per replicate. Error bars depict standard deviation between three replicates. The fold change normalization was done by setting the control value as one for qRT-PCR expression analysis.

Supplementary figure S16: Delineating the existence and localization of miR169d:At-NF-YA2 functional module in Arabidopsis. Histochemical GUS expression patterns of Pro-At-MIR169d:GUS and Pro-At-NF-YA2:GUS promoter reporter transgenic Arabidopsis plants in: seedling (2 weeks old), complete rosette (4 weeks), mature rosette leaf, cauline leaf and flower

Supplementary figure S17: Expression levels of At-NF-YA2 in At-NF-YA2 overexpressing transgenic plants. Relative expression of At-NF-YA2 in transgenic Arabidopsis plants overexpressing At-NF-YA2 under constitutive CaMV35S promoter. Bar represents the average data of three biological replicates and error bar represents standard deviation of the replicates.
Supplementary figure S18: Schematic structure of At-NF-YA2 gene carrying the T-DNA insertion and lack of At-NF-YA2 expression in the At-nf-ya2 mutant (SALK_021228). (A) The four exons of At-NF-YA2 genomic DNA are shown according to the gene features annotated at TAIR database. T-DNA insertion site is shown by inverted triangle in the fourth exon of At-NF-A2. The exon sequences are depicted by red and the UTR are represented by green color. (B) Genotyping PCR of At-nf-ya2 mutants, presence of border specific amplification in At-nf-ya2 plants genomic DNA confirms their homozygosity, while presence of only gene specific amplification in control Columbia plants confirms their wild type nature. (C) Expression of At-NF-YA2 was determined by RT-PCR, using full length cDNA from WT and mutant At-nf-ya2 plants. Absence of At-NF-YA2 specific band in mutant plants confirms the knockout nature of At-nf-ya2 homozygous mutants. RNA was isolated from detached mature leaves of the wild-type (WT) or At-nf-ya2 plants. Expression of actin is shown as a loading control.

Supplementary figure S19: Expression profiling of HSR genes in different transgenic lines of Arabidopsis. Expression patterns of five heat stress-responsive genes in WT, At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NF-YA2-r and MIM169defg plants. Data are shown as means ± SD of biological replicates and normalized to At-EF-α reference gene using (2^{-ΔΔCT}) method. The fold change normalization was done by setting the control value as one for qRT-PCR expression analysis.

Supplementary figure S20: Yeast-one-hybrid assay based transcriptional regulation of At-HSFA3a, At-HSFA7a and At-HSFA7b promoters by At-NF-YA2. Positive interaction of At-NF-YA2 with the promoters of At-HSFA3a and At-HSFA7b were obtained. Serial dilutions (O.D. 0.1 to 0.0001) of yeast cultures were spotted onto plates lacking LEU with specific Aureobasidin A concentration and incubated for 3 days.
Supplementary figure S21: Thermotolerance assay of WT and At-hsfa7a plants.
(A) Two-weeks-old soil-grown WT and At-hsfa7a mutant plants were subjected to 0 (control) or 2h heat stress at 42°C, and damage was recorded 6 days later. (B) Estimation of survival (percentage) of heat stress treated WT and At-hsfa7a plants. Plants were assayed for heat stress tolerance after 6 days of HS. These experiments were repeated at least three times with similar results, and average data of all experiments are shown. Error bars, ± SD of three repeats. ***p<0.001 vs. wild type, by two-tailed Student’s t-test.

Supplementary figure S22: At-NF-YA2, Arabidopsis orthologue of tomato Sly-NF-YA9 and Sly-NF-YA10. (A-B) Protein sequence alignment of tomato Sly-NF-YA9 (A) and Sly-NF-YA10 with At-NF-YA2. Protein sequences were aligned with Arabidopsis proteome at TAIR database by using Blastp tool.

Supplementary figure S23: Binding of Sly-HSFA6b to Sly-MIR169d-1 promoter. Absence of any amplification in no-Ab (no antibody) and slight amplification in IgG control confirms the specificity of immunoprecipitation. Presence of strong amplification on input DNA confirms the good amplification efficiency of the primer used for analysis. Presence of amplification in Anti-GFP lane confirms the enrichment of Sly-MIR169d-1 promoter in Sly-HSFA6b:GFP CHIP. L denotes the 100 bp ladder.

Supplementary figure 24: Virus induced silencing of tomato Phytoene desaturase (PDS) gene. PDS silencing was used as a control for assessing establishment of gene silencing by VIGS. (A) Silencing of the PDS control gene causing photobleaching in tomato plants 3 weeks after agroinfection. (B) Enlarged individual leaves of TRV-PDS and control WT leaves.

Supplementary Table legends:

Supplementary table S1: Distribution of perfect and imperfect heat stress elements (HSE) in Sly-MIR169 promoters.
**Supplementary table S2:** Interaction summary of *Sly-MIR169* promoters and Sly-HSFs. Positive HSF interaction is marked in red colored boxes.

**Supplementary table S3:** All Qrt-PCR raw data, calculations and fold change normalizations with different endogenous controls.

**Supplementary table S4:** Off targets of candidate VIGS constructs used in the study.

**Supplementary table S5:** Promoter analysis of HSR genes in tomato.

**Supplementary table S6:** Number of HSE elements and minimal inhibitory concentration of Aureobasidin (Aba) for all At-MIR169 promoters.

**Supplementary table S7:** Interaction summary of At-MIR169 promoters and At-HSFs. Positive HSF interaction is marked in red colored boxes.

**Supplementary table S8:** qRT-PCR based expression profiles of HSR genes in *At-MIR169d OE, At-NF-YA2 OE, At-nf-ya2, At-pNF-YA2:NF-YA2-r* and *MIM169defg* transgenic lines.

**Supplementary table S9:** Analysis of Ct similarities across different conditions for the reference genes used in the study.

**Supplementary table S10:** List of Primers used in the study.

**Supplementary table S11:** Primer efficiency of all the primers used in the study.

**Supplementary table S12:** Minimal inhibitory concentration of Aureobasidin (Aba) for all *Sly-MIR169* promoters.
Figure 1: Yeast one hybrid and Chromatin immunoprecipitation (ChIP-qPCR) assay of Sly-MIR169 promoters and HSFs. (A) Yeast-one-hybrid assay to assess transcriptional regulation of heat-responsive MIR169 promoters by HSFs. Only the positive Y1H interactions between different HSFs and Sly-MIR169 promoters have been shown. The Sly-MIR169 promoters were cloned in the reporter vector pABAi and the full-length cDNA of HSFs were cloned in the pGADT7 vector. After the co-transformation into Y1H gold yeast strain, serial dilutions (O.D. 0.1 to 0.0001) of yeast cultures were spotted onto plates lacking URA and LEU with specific Aureobasidin A concentration and incubated for 3 days. Promoter of sHSP has been used as a positive control to show binding of HSF3a on sHSP promoter (Li et al. 2013) and empty vectors as negative controls. The Aureobasidin A concentrations for all Sly-MIR169 promoters is presented in supplementary table S7. (B-I) Chromatin immunoprecipitation (ChIP-qPCR) assay of Sly-MIR169 promoters and HSFs in Nicotiana benthamiana. Fold enrichment of Sly-MIR169a-1 promoter in HSFA1e immunoprecipitated sample (B); Fold enrichment of Sly-MIR169b promoter in HSFA1a immunoprecipitated sample (C); in HSFA7a immunoprecipitated sample (D); Fold enrichment of Sly-MIR169C promoter in HSFA4a immunoprecipitated sample (E); Fold enrichment of Sly-MIR169b promoter in HSFA6b immunoprecipitated sample (F); Fold enrichment of Sly-MIR169d promoter in HSFA8a immunoprecipitated sample (G); in HSFB1a immunoprecipitated samples (H); Fold enrichment of Sly-MIR169d-1 promoter in HSFA6b immunoprecipitated samples (I). These experiments were repeated 3-4 times; the error bars represent standard deviation between the replicates. The values for different promoters in IgG immunoprecipitated sample were set as one for normalization.
Figure 2: Validation of HSF:miR169:NF-YA node components in tomato upon heat stress. (A) Expression profiles of HSF genes that regulate Sly-MIR169 transcription in WT tomato leaves during control and heat stress condition by qRT-PCR using the 2−ΔΔCt method. (B) Expression of mature miR169s in response to heat stress as determined by taqman based qRT-PCR in control and heat challenged WT tomato leaves. (C) qRT-PCR based expression profiles of target NF-YA transcripts in WT control and heat stressed tomato leaves using the 2−ΔΔCt method. (D-G) The expression analysis of HSF genes, Sly-MIR169 precursors and NF-YA transcripts in VIGS plants silenced for Sly-HSFA1a (D), Sly-HSFA7a (E), Sly-HSFA6b (F) and Sly-HSFB1a (G). In (D-G) 15-days-old tomato plants were agro-infiltrated with empty vector (TRV-1, TRV-2 VC) or TRV-Sly-HSF VIGS constructs. The VIGS established plants were subjected to heat stress after 3-weeks of agro-infiltration, and used for expression studies of different genes. Graphical data represents mean values of expression of three to four biological sets. Error bars show standard deviation. To represent the negative fold change on y axis in (C-G), the relative expression values of down-regulated genes were transformed using the formula \[-(1 / RQ ≤ 0.5)]. Actin was used as endogenous control in (A), (C) and (D-G), while 5SrRNA was used as reference control in (B). The fold change normalization was done by setting the control value as one for all qRT-PCR expression analysis.
Figure 3. Functional validation of the role of Sly-NF-A9 and Sly-NF-A10 in HS response in tomato. (A) Phenotype of TRV-VC (Vector control), TRV-Sly-NF-YA9 and TRV-Sly-NF-YA10 silenced plants after heat stress. Fifteen-days-old tomato plants were agro-infiltrated with empty vector (TRV-VC), TRV-Sly-NF-YA9 or TRV-Sly-NF-YA10 silencing constructs. Plants were given HS 3-weeks-after infiltration. Phenotype of HS-treated silenced plants was scored six-days-post HS. Experiments were repeated 3-4 times with 8 plants per replicate. (B) Percent survival of TRV-EV, TRV-Sly-NF-YA9 and TRV-Sly-NFYA10 silenced plants after HS. (C) Estimation of water use efficiency (mmol mol⁻¹), net photosynthesis rate (μmol m⁻²s⁻¹), and transpiration rate of TRV-EV, TRV-Sly-NF-YA9 and TRV-Sly-NFYA10 silenced plants after HS. (D-E) Expression profiles of HSR genes in control and heat treated TRV-EV, TRV-Sly-NF-YA9 and TRV-Sly-NFYA10 silenced plants. The expression levels of genes were calculated using the 2−ΔΔCt method and presented using fold-change values. The fold change normalization was done by setting the control value as one in (D-H). Actin was used as endogenous control. Error bars represent the standard deviation of two independent biological replicates. *p<0.05, **p<0.01 and ***p<0.001 vs. wild type, by two-tailed Student’s t-test.
Figure 4. Transcriptional regulation of MIR169s promoters by HSFs in Arabidopsis. (A) miR169s expression in response to heat stress as determined by Taqman-based qRT-PCR of control and heat challenged plants. 5S rRNA was used as the normalization control. Experiment was repeated three times and average values are plotted as bars, error bars depict standard deviation between the replicates. (B) Y1H assay showing the positive interaction of HSFs with the promoters of different At-MIR169s, Y1H assay shows binding of At-HSFA7b, At-HSFB2b, At-HSFB3a, At-HSFA2 and At-HSFA1b on At-MIR169b, d and h promoters. This experiment was repeated at least three times with similar results, and data from one representative experiment is shown. Serial dilutions (O.D. 0.1 to 0.0001) of yeast cultures were spotted onto plates lacking URA and LEU with specific Aureobasidin A concentration and incubated for 3 days. The Aureobasidin A concentrations for all At-MIR169 promoters is presented in supplementary table S3. (C) Expression levels of At-MIR169d in wild-type plants subjected to 0 (control) and 2 h heat stress at 42 °C. Data are shown as means ± SD of three biological replicates and normalized to actin reference gene using (2−ΔΔCT) method. (D) Expression patterns of Pro-At-MIR169d:GUS lines of transgenic Arabidopsis plants subjected to 0 (control) or 2 h of heat stress at 42°C. (E) Levels of GUS transcripts in Pro-At-MIR169d:GUS lines, measured by qRT-PCR after 0 or 2 h of heat stress at 42°C. (F) Transcriptional activity of Pro-At-MIR169d:GUS reporter construct in WT and At-hsfa2 mutant background during 2 h of heat stress at 42°C. (G) Expression of GUS transcript driven by Pro-At-MIR169d and At-MIR169d precursor in soil-grown wild-type plants of Arabidopsis and At-hsfa2 mutant. To represent the negative fold-change on y axis in figure (G), the relative expression values of down-regulated genes were transformed using the formula [-1 divided by RQ ≤0.5)]. The fold change normalization was done by setting the control value as one in Figure (A, C, E and G). Error bars represent the standard deviation. These experiments were repeated at least three times with similar results, and data from one representative experiment are shown in D and F.
Figure 5: Assaying the miR169d:At-NF-YA2 node in heat stress in Arabidopsis. (A) Thermotolerance was gauged in At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NF-YA2-r and MIM169defg transgenic plants in comparison to wild-type (WT) plants. Two-weeks-old soil-grown plants were subjected to 0 (control) or 2h heat stress at 42°C, and damage was recorded 6 days later. (B) Estimation of survival (percentage) of HS treated At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NF-YA2-r and MIM169defg plants. Plants were assayed for heat stress tolerance after 1 week of HS. These experiments were repeated at least five times with similar results, and average data of all experiments are shown. Error bars, ± SD of five repeats. (C) Expression patterns of heat stress-responsive genes in WT, At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NF-YA2-r and MIM169defg plants. Data are shown as means ± SD of biological replicates and normalized to Actin using (2^{−ΔΔCT}) method. Log2 transformation was applied to the RQ data to obtained negative fold change. (D-E) DAB and Trypan blue staining of HS treated At-MIR169d-OE, At-NF-YA2-OE, At-nf-ya2, pNF-YA2:NF-YA2-r and MIM169defg plants to estimate the production of ROS and cell death, respectively. *p<0.05, **p<0.01 and ***p<0.001 vs. wild type, by two-tailed Student’s t-test. Three independent lines of MIR169d-OE were used and similar results were obtained for all studied parameters, a single representative picture was shown in the figure A, B, D and E. The fold change normalization was done by setting the control value as one for figure (C).
Figure 6: Transcriptional regulation of At-HSFA3a and At-HSFA7b by At-NF-YA2 during heat stress in Arabidopsis. (A) Graphical representation of CCAAT cis-elements in At-HSR gene promoters HSFA3a, HSFA7a, HSFA7b, APX and shSP17.6A. The 2 kb upstream sequences of start codon were analysed by plant care and Plant PAN promoter analysis tools. The positions of CCAAT boxes are depicted by violet coloured tabs. (B) Chromatin immunoprecipitation (ChIP)-qPCR assay of At-NF-YA2 on At-HSR gene promoters. (C-D) In planta estimation of transcriptional regulation of At-HSFA3a and At-HSFA7b transcription by At-NF-YA2 during heat stress. (C) Pro-At-HSFA3a:GUS reporter construct was agro-infiltrated in WT, At-NF-YA2-OE and At-nf-ya2 backgrounds. GUS was visualized 2-days post-infiltration by histochemical staining. (D). qRT-PCR based quantification of GUS transcripts in promoter reporter constructs infiltrated plants, 2-days-post infiltration by using GUS specific primers. The NPTII gene present in the same vector backbone was used as reference control. To represent the negative fold-change on y axis in, the relative expression values of down-regulated genes were transformed using the formula [-1 divided by RQ ≤0.5)]. The fold change normalization was done by setting the control value as one. (E) Thermotolerance assay of WT and At-hsfa3a mutant plants. Two-weeks-old soil-grown plants were subjected to 0 (control) or 2h heat stress at 42°C, and damage was recorded 6 days post-stress. (F) Estimation of survival (percentage) of WT and At-hsfa3a plants 6 days post heat stress. The experiments were repeated at least three times with similar results, and average data of all experiments are shown. Error bars, ± SD of three repeats. *p<0.05, **p<0.01 and ***p<0.001 vs. wild type, by two-tailed Student’s t-test.
Figure 7: Model for HSF-induced miR169:NF-YA module in mediating thermotolerance in tomato and Arabidopsis. Heat stress induces the expression of HSFs by altering the cellular ROS levels as well as other cues independent of ROS signaling. HSFA1a, HSFA2 and HSFA7a are the key player of plant HSR response. These HSFs bind to the promoters of MIR169 leading to transcriptional enhancement of miR169s in both tomato and Arabidopsis. Enhanced accumulation of miR169s reduces the levels of Sly-NF-YA9/A10 in tomato and At-NF-YA2 in Arabidopsis that leads to enhancement of HSR genes expression like HSFA2, HSFA3 and HSFA7s, possibly via altering ABA levels. In Arabidopsis, At-miR169-mediated down-regulation of At-NF-YA2 during HS may favor functional DREB2A:NYC-10:NYB3 complex formation to up-regulate At-HSFA3a expression during HS. Enhanced levels of HSFAs in turn feed the transcription of MIR169 genes. The heat-mediated orchestration of parallel miRNA pathways (miR169:NYC-10, miR398:CSD and miR156:SPL) alters the levels of HSR genes and regulate plant thermotolerance. Solid lines denote a confirmed interaction and dotted lines are predictive. Red and black lines denote findings of this study and existing literature, respectively. Black colored numerals refer to the studies that established the particular connection between different components of the model. Reference key- 1: Guan et al. 2013; 2: Stief et al. 2014; 3: Sato et al. 2014; 4: Zhao et al. 2009, Luan et al. 2015, Ding et al. 2016; 5: Sakuma et al. 2006. CSD: Copper superoxide dismutase; SPL: SQUAMOSA-promoter binding like; ABA: Abscisic acid; ROS: Reactive oxygen species; DREB: Dehydration responsive element binding protein.