Transfer RNA-derived non-coding RNAs (tncRNAs): Hidden regulation of plants’ transcriptional regulatory circuits

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1. Introduction

The profusion of non-coding RNAs (ncRNAs) with diverse regulatory actions discovered over the last decade has been proven to be a crucial breakthrough in molecular biology. They are powerful regulators of gene expression at the epigenetic, transcriptional, and post-transcriptional levels in the living system [1]. Amongst the diverse pool of smaller ncRNAs, microRNAs (miRNAs), and small interfering RNAs (siRNAs) are the most extensively surveyed molecules [2]. They alter gene expression by binding to their target mRNAs [3]. The advancement of high-throughput sequencing technology has tremendously resolved the surveillance of the other classes of small untranslated RNAs beyond miRNAs and siRNAs [4,5]. Amongst them, transfer RNA (tRNA)-derived non-coding RNAs (tncRNAs) have been reported in all three domains of life [6]. This distinct group of regulatory RNAs is derived from the endonucleolytic cleavage of precursor tRNAs (pre-tRNAs) or mature tRNAs [7]. tncRNAs includes well-known shorter tRNA-derived RNA fragments (∼12 to 30 nucleotides [nt]) popularly termed as tRFs [8] or tDRs [9], and longer tRNA halves or tRHs (∼30 to 40 nt) [10]. Precise tRNA cleavage leading to their production is governed by tRNA type, cell type, developmental stage, stress, and tissue [11–13]. It has been revealed that these novel molecules can originate from nuclear as well as organellar tRNAs [14,15]. This class of small RNAs has been particularly well studied in human cancers [16–18]. In plants, the earliest report by Thompson et al. 2008 showed that processed tRNA halves were generated in Arabidopsis during oxidative stress [19]. In another study, phloem-specific tRNA fragments were shown to interfere with protein synthesis in pumpkin (Cucurbita maxima) [20]. For the first time in plants, Loss Morais and his group in 2013 showed the association of 19 nt tRFs with Argonaute under stress conditions [21]. Alves et al. extended the research and highlighted the differential accumulation of specific tRFs under abiotic stresses in Arabidopsis and rice [22]. In addition to abiotic stresses, some studies have shown links to tncRNAs generation in response to biotic stresses, e.g., in...
response to Apple stem grooving virus (ASGV) infection in domesticated apple [23], and Phytophthora capsici infection in black pepper [24].

Depending on the position of cleavage on mature tRNA, tRFs belong to two categories: 5’ end derived tRF-5, and 3’ end derived tRF-3 generated upon cleavage in the D and T region, respectively. Among the tRNA halves, 5’ or 3’ tRHs are produced by cleavage in the anticodon region containing 5’ or 3’ portions of mature tRNA [25]. The pre-tRNA produces 5’U-tRFs/leader tRF, and tRF-1/tSRNA/3’U-tRFs from the 5’ leader and 3’ trailer portion respectively [7,26,27]. By now, it has been established that tncRNAs can be generated in a DICER-dependent or independent manner in plants [28,29]. Besides Dicer-like proteins (DCLs), members of RNases T2, namely RN51, RN52, and RN35 are the key contributors to the biogenesis of tncRNAs originating from mature tRNAs in Arabidopsis [22,30]. Noteworthy, the tncRNA generation from pre-tRNAs in plants has been understudied.

Identification of tncRNAs in small RNA-seq (sRNA-seq) datasets is a very challenging and error-prone process. During the library preparation, extensively modified tRNA nucleosides affect the reverse transcription process, resulting in truncation or misincorporation of nucleotides [31]. This, in turn, increases the probability of mismatch. However, allowing just a single mismatch to over- come sequencing errors, may lead to base misidentification, and hence causes the inflation of the false negatives. This is due to high sequence similarities among 20 major tRNA isotypes covering the whole tRNA family of isoacceptors (tRNAs with different anticodons but charged with the same amino acid), and isodecoders (tRNAs carrying the same anticodon with variations in the body sequence). Researchers have suggested different mapping strategies for tRNA-derived reads [32–34]. Some tools are available for tncRNA detection, e.g. tDrMapper [35], MiNTmap [36], and tRF2Cancer [37], but they are trained and tested on human data- sets, and thereby suit for human data only. As compared to humans, tncRNAs have been less explored in the plant domain. Although, exclusive plant databases, e.g. tRex [38] and PtRfDb [39], harboring information related to the transfer RNA-derived fragments (tRFs), have also been developed in the recent past, yet a convenient methodology for the accurate identification and exploration of tncRNAs is still lacking in plants.

In this study, we developed a computational methodology for accurate tncRNA detection. We have identified a diverse pool of tncRNAs in six different angiosperms viz. Arabidopsis thaliana (model plant), Medicago truncatula (model legume), tomato (Solanum lycopersicum), chickpea (Cicer arietinum), rice (Oryza sativa), and maize (Zea mays) by exploiting publicly available sRNA-seq datasets. With the help of our in-house pipeline, we have identified, classified, and quantified tncRNAs into different subtypes based on their origin and cleavage position viz. tRF-5, tRF-3, tRF-1, leader-tRF, 5’ and 3’ tRNA halves (tRHs). As there is no single standardized nomenclature for tRNA subtypes yet, researchers have followed the classical and most frequently used terms for tRNA subtypes. tRNA modifications play an integral role in tRNA structural and functional integrity of fragments generated from the 3’ end. We also observed a remarkable fraction of fragments generated from internal portions of mature tRNA. These fragments constitute nearly 45% of the total tncRNA sequences identified in each plant (Supplementary Fig. 1A–F). Longer tRHs occupy less than 10% of the tncRNAs. Compared to 3’tRHs, the 5’tRHs were more abundant. The remaining fraction (<1%) consisted of tncRNAs generated from pre-tRNAs. The length diversity of tncRNAs indicates their prospective roles besides canonical miRNA-like gene expression

2. Results

2.1. In silico tncRNA identification across six plants

By utilizing sRNA-seq data available at Sequence Read Archive (https://www.ncbi.nlm.nih.gov/sra), from the National Center for Biotechnology Information (NCBI), we have identified diverse groups of tncRNAs from 2,448 samples in six angiosperms viz. A. thaliana (1676), S. lycopersicum (1 6 0), C. arietinum (21), M. truncatula (1 2 7), O. sativa (2 4 3), and Z. mays (2 2 1). After processing and quality control, filtered reads were provided as input to the in-house pipeline (Fig. 1A). Due to the huge number of tRNA gene copies predicted by tRNAscanSE [44], only high-quality tRNAs with a score equal to or above 50 were selected for mapping. To avoid ambiguous reads from non-tRNA regions, we created an artificial genome by masking the genuine tRNA gene region with 50 nt upstream and downstream in the genome (masked genome) and adding them as artificial chromosomes (Fig. 1A). We classified only those reads as tncRNAs that mapped exclusively to the artificial chromosomes (i.e. tRNA set consisting of mature tRNA, 50 nt leader, and 50 nt trailer). The identified tncRNAs were categorized into tRF-5, tRF-3, tRF-1, leader-tRF, 5’tRH, 3’tRH, and other tRF (Fig. 1B). The sample-wise tncRNA analysis results containing various information related to each tncRNA, i.e. tncRNA type, parent tRNA locus information, position (start–end), sequence, length, read count, RPM, nature, and position of modified nucleosides, alignment of each tncRNA on its progenitor tRNA, and the count of different tncRNA subclasses utilized in this study can be downloaded from “Data download” module at http://www.nipgr.ac.in/tncRNA. The sample number, total, and unique tncRNA counts for individual plants have been summarised in Table 1 (the same tncRNA sequences were grouped and termed as unique).

2.2. tncRNA classification according to origin and site of cleavage

We have computed the relative percentages of sRNA-seq samples, tncRNA abundance (total and unique), and tncRNA classes (total and unique) for each plant (Fig. 2A). The total tncRNAs in Arabidopsis showed higher abundance due to greater sample size, but the relative abundance of unique sequences in Arabidopsis was fairly comparable to other plants (removing duplicate sequences from several samples reduces the bias of sample number). The unique count of reads per tncRNA category for individual samples has been provided in Supplementary sheets 1.1–1.6. The length-wise abundance of major tncRNA classes was studied in different plants (Fig. 2B). The bulk of the identified tncRNAs belonging to the tRF-5 and tRF-3 series, fall in a smaller length range up to 25 nt and 21 nt respectively. The high frequency of specific-sized tncRNAs indicates cleavage preference by the endoribonuclease machinery at a certain position in the loop/stem region. Also, more than 92% of the total identified tRF-3 sequences contained terminal end CCA in all plants. The high abundance of CCA ending fragments indicates that the terminal CCA might also play a role in the structural and functional integrity of fragments generated from the 3’end. We also observed a remarkable fraction of fragments generated from internal portions of mature tRNA. These fragments constitute nearly 45% of the total tncRNA sequences identified in each plant (Supplementary Fig. 1A–F). Longer tRHs occupy less than 10% of the tncRNAs. Compared to 3’tRHs, the 5’tRHs were more abundant. The remaining fraction (<1%) consisted of tncRNAs generated from pre-tRNAs. The length diversity of tncRNAs indicates their prospective roles besides canonical miRNA-like gene expression
Summary of identified groups of transfer-RNA-derived non-coding RNAs (tncRNAs) abundance in six plants.

| Plant species     | Total Samples | tncRNAs Total/Unique | tRF-5 Total/Unique | tRF-3(CCA) Total/Unique | tRF-1 Total/Unique | 5’tRH Total/Unique | 3’tRH(CCA) Total/Unique | Leader tRF Total/Unique | Other tRFs Total/Unique |
|-------------------|---------------|----------------------|--------------------|-------------------------|--------------------|-------------------|------------------------|------------------------|-------------------------|
| A. thaliana       | 1676          | 3,362,128/50619       | 760,155/2180       | 782,722/4513            | 22,505/693         | 164,074/662       | 28,835/840             | 2359/238               | 1,601,478/4102          |
| C. arietinum      | 21            | 54,390/10627         | 8837/1012          | 11,763/1792             | 77/35              | 2125/306          | 778/175                | 26/12                  | 30,774/7310             |
| M. truncatula     | 127           | 306,816/11971        | 63,157/1182        | 82,262/1967             | 77/92              | 15,232/321        | 423/139                | 137/32                 | 140,857/8270            |
| O. sativa         | 243           | 406,496/23117        | 100,828/1812       | 98,089/2787             | 153/189            | 19,070/447        | 2595/294                | 53/26                  | 184,308/17585           |
| S. lycopersicum   | 160           | 265,435/23312        | 62,838/1753        | 59,833/3323             | 1878/688           | 19,298/560        | 2429/264                | 130/78                 | 119,029/16664           |
| Z. mays           | 221           | 428,193/18304        | 95,748/1819        | 99,510/2651             | 1959/265           | 37,768/618        | 7457/354                | 164/24                 | 185,587/12584           |

2.3. Cleavage of tncRNAs is highly specific and conserved amongst plants

Nucleotide composition and the ratio of the constituent monomer units is an important characteristic feature for studying nucleic acids. For studying the tncRNA cleavage pattern, we looked into the nucleotide composition of tncRNA::RNA junction for highly recurring tRF-5, tRF-3(CCA), 5’tRH, and 3’tRH(CCA) sequences (Fig. 3). A common conserved motif was observed in tRF-5::RNA interface among all plants i.e. AG::TGG (first row in Fig. 3) in tRF-5 s ranging between 15 and 19 nt. Also, the last nucleotide in tRF-5 was observed to be A/G rich (first three rows from top in Fig. 3). In tRNA::tRF-3(CCA) junction, CGA[N];(N):TC pattern was commonly observed across all species (fourth to sixth rows in Fig. 3). Similarly, in the majority of the 5’ tRNA halves, the breakpoint region in the middle (fifth and sixth position) is either TC (seventh to ninth rows in Fig. 3), whereas the first nucleotide in 39 nt 3’tRH (CCA) is A rich in all species (sixth nucleotide in the tenth rows in Fig. 3). Also, a common conserved pattern, C1/G3/AA/CC was observed in the majority of 40–41 nt length 3’tRHs (eleventh and twelfth rows in Fig. 3). Despite the variability due to the occurrence of anticodon triplets among various isoacceptors, we have seen common motifs at tRNA cleavage points among different plants indicating that the cleaving enzymes mostly prefer specific isoacceptors for spawning tRHs. Also, the majority of tRF-1 sequences identified in rice, maize, and tomato samples had terminal ends rich in T residues (Supplementary Fig. 3). It has been reported that G at 18th and 19th position in tRF-5 plays a crucial role in the inhibition of protein translation in both human and Arabidopsis [46,47]. The production of tncRNAs from pre-tRNAs has not been well studied in plants. In humans, tRF-1/pre-tRF-3U usually end in a short stretch of T residues due to the release of polymerase III [6,57]. Based on observed motifs and specific nucleotide composition of tncRNAs, it is worth speculating that the enzymatic cleavage might first recognize or prefer a particular motif located on parental tRNA to cleave precise fragments for at least a reasonable number of genuine tncRNAs.

2.4. Specific anticodons, not the whole tRNA repertoire are responsible for tncRNA generation

Out of the 64 triplet codons, 61 are sense codons that code for 20 amino acids. We observed that not all 61 codons generate tncRNAs. Out of these 61 codons, Arabidopsis, chickpea, and Medicago utilized 49, 41, and 44 isoacceptors respectively, while 48 isoacceptors contributed to the production of different tncRNAs in tomato, rice, and maize (Fig. 4). Different isoacceptors contributed to the production of different tncRNA types and varied from one plant to another. Although coding for the same amino acids, there was marked variation among the isoacceptors for contributing to the tncRNA generation, e.g. in Arabidopsis, among five tRNA-Arg isoacceptors viz. ACG, CCG, CCT, TCG, and TCT, tRNA-ArgACC generated the most number of tRF-5 s. Also, many anticodons can generate more than one class of tncRNAs, e.g. GluCTC, and ProCCTC gave rise to tRF-5 and tRF-3, while GluCTC, GlyTCC, and LeuTAA generated 5’tRH and 3’tRH in Arabidopsis. By comparing the number of tncRNAs generated per the number of copies for each tRNA isoacceptor, we observed that the tncRNAs generation showed a slightly positive correlation with the increasing abundance of isoacceptor copies (Fig. 5). Pearson correlation coefficient ranged from 0.37 to 0.66, which implies that the number of
Fig. 2. Abundance of the identified transfer-RNA-derived non-coding RNAs (tncRNAs) in different plants. A Relative distribution of samples, total and unique tncRNA, tRF-5, tRF-3(CCA), tRF-1, 5’tRH, 3’tRH(CCA), leader, and other tRFs analyzed in this study. B Size distribution of major tncRNA categories viz. tRF-5, tRF-3(CCA), tRF-1, 5’tRH, 3’tRH(CCA), and other tRFs in individual plants.
tncRNAs generated is not directly proportional to tRNA frequency. Still, there is a possibility that the tRNA isoacceptors with more copy numbers may contribute more to tncRNAs generation.

The tRNA and tncRNA generating regions were visualized on the genome (Fig. 6). Various classes of tncRNAs are generated from distinct portions of genomic tRNAs. We found many regions on the genome that showed lesser tRNA density but generated more tncRNAs and vice-versa. For example, tRNA-dense regions can be seen particularly on chromosome 1 (chr1) in Arabidopsis but most of the tncRNAs were generated from chr2. Similarly, despite low tRNA gene density, a large number of tncRNAs originated from chr4 in tomato, chr6 and 8 in chickpea, chr3 in Medicago, chr7 and 11 in rice, and chr1, 5, and 8 in maize. It can be observed that the tncRNAs are not necessarily generated from tRNA-rich regions of the genome. Interestingly, some regions generated several classes of tncRNAs, like chr2 in Arabidopsis, chr1 in tomato, chr4 in Medicago and rice, and chr5 in maize, while some regions gave rise to specific tncRNA classes. For example, more 5’ containing tncRNAs, i.e. tRF-5 and 5’tRHs, on chr3 in Arabidopsis, chr6 in tomato, chr4 in Medicago, chr11 in rice, and chr5 in maize. It is quite conceivable that some of these regions may govern rapid turn-over of tRNAs into tncRNAs. These findings give clues about the additional insights into the genomic organization of tRNA into clusters and their implications on tRNA functions. The prevalence of tRNA gene clusters in all kingdoms of life reveals various insights into evolutionary history and tRNA functions. Whether tRNA gene clusters impact the generation of tncRNAs in the living system, including plants, is worth intriguing.

Fig. 3. Web logo representation depicting conserved cleavage sites for four tncRNA subclasses viz. tRF-5, tRF-3(CCA), 5’tRH, and 3’tRH(CCA). Web Logos were made using top-three recurring tncRNAs by taking 5 nt stretches from progenitor tRNA and respective tncRNA fragments viz. tRF-5::tRNA; 5’tRH::tRNA; tRF-3(CCA)::tRNA; 3’tRH(CCA)::tRNA. To study the breakpoint and its flanking nucleotide composition, for tRF-5 and 5’tRH classes, the first five nucleotides (from left to right) represent the last five nucleotides of tRF-5/5’tRH and the remaining stretch of nucleotides is the tRNA (sixth to the eleventh nucleotide). For tRF-3(CCA) and 3’tRH(CCA) classes, the first five nucleotides (from left to right) represent progenitor tRNA sequence, and the remaining stretch represents the first sixth nucleotide of tRF-3(CCA)/3’tRH(CCA) (sixth to the eleventh nucleotide). The red arrow represents the breaking point of each tncRNA fragment from its parental tRNA. All the sequences shown are oriented in the 5’→3’ direction. G/T; T/C; A/A breakpoint was commonly found in all types of tncRNAs. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
2.5. Modification may play a role in regulating the tncRNA generation and function

As tRNAs are heavily modified molecules with over 100 known modifications, we scrutinized the modifications present in our identified tncRNAs. For this, we utilized HAMR [48], a high utility tool for predicting modified nucleosides in high throughput sequencing datasets. Various modifications were detected by HAMR on tncRNAs in different plants, viz. 3-methylcytosine (m3C), pseudouridine (Y/W), dihydrouridine (D), N2-methylguanosine (m2G), N2-dimethylguanosine (m22G), 1-methylguanosine (m1G), 1-methyladenosine (m1A), 1-methylinosine (m1I), 2-methylthio-N6-isopentenyladenosine (ms2i6A), N6-isopentenyl adenosine (i6A), Threonylcarbamoyladenosine (t6A) in various isoacceptors generating tncRNAs. The visualization of various modifications on tncRNAs concerning their position on the respective consensus sequence of mature tRNA in 49 isoacceptors in Arabidopsis has been shown in Supplementary Fig. 4. The plots for modification distribution in the other five angiosperms have been provided in Supplementary Figs. 5–9. We observed that most modifications were predicted from 20th to 40th position and 50th to 60th position, i.e. containing the cleavage sites for tRF-5, tRHs, and tRF-3, indicating a high probability of tncRNA generation to be regulated by tRNA modifications. The recurring tRNA modifications in tncRNAs from Arabidopsis samples have been highlighted in Supplementary Fig. 10. It was observed that some modifications were specific for certain anticodons and were found in abundance at specific positions. For example, in the D region, Guanosine (G) was modified to m1G on GlyGCC and iMetCAT (Supplementary Fig. 10). Also, pseudouridine (Y) modification on GlyGCC tRNAs was highly abundant. In the anticodon loop region, Y and D were abundant in tncRNAs originating from GluTTC tRNAs. In the T region, m1A|m1I|ms2i6A were observed in a large number of tRNAs.

We also looked for modifications found common in all six plants selected for this study (Supplementary Fig. 11 A-U). Adenosine (A) replaced with m1A|m1I|ms2i6A modifications were found...
in 20 isoacceptors viz. tRNA ArgACG|CCG|TCG|TCT (58th position), AspGTC (57th position), GlnTTG (57th position), iMetCAT (57th position), LeuCAA (69th position), Leu-CAG (66th position), Leu-TAG (65th position), LysCTT (58th position), LysTTT (57th position), SerAGA|CGA|TGA (67th position), ThrTGT (57th position), TyrGTA (58th position), and ValAAC|CAC (59th position). G is modified to m1G on the T loop at 58th position in TyrGTA and 59th position in ValAAC|CAC. These modifications were prominent in the region of the T loop, which is the site of cleavage for tRF-3 generation. Also, on the 9th nucleotide, G was replaced with m1G in ArgACG, GlyGCC, and iMetCAT in all plants. Am nucleoside (20-O-methyladenosine) is reported to be induced by salt stress in a variety of land plants, including Arabidopsis, Brachypodium, poplar, and rice [49]. 20-O-methylguanosine (Gm), 5-methyluridine (m5U), and 5-methylcytidine (m5C) are also tightly linked to plant development [50]. Overall, the findings suggest the possibility of a very delicate cross-talk among the genetic machinery involved in tncRNA biogenesis, functioning, as well as tRNA nucleoside modification in response to plant growth, development, as well as environmental changes.

2.6. tncRNAs are conserved amongst plants

In our entire analysis, we observed that various tncRNAs were identical in all six plant species. We found a total of 252 tRF-5, 351 tRF-3, 46 5tRH, and 12 3tRH common sequences among all six plant species included in our analysis. The sequence conservation provides a clue to the conservation in the regulatory function of these tncRNAs. They can act as post-transcriptional regulators by complementary binding to their messenger RNA (mRNA) targets leading to mRNA degradation. Thus, conserved sequences were utilized for target prediction and pathway enrichment analysis to gain some functional insights into gene expression regulation by tncRNAs. The detailed list of conserved tRF-5 s and tRF-3 s along with their respective target genes have been provided in Supplementary sheets 2.1 and 2.2, respectively. We observed that several target genes were associated with the molecular function, biological process, and cellular components for tRF-5 and tRF-3. The tRF-5 target genes were related to 9 distinct molecular functions, 25 types of biological processes, and six cellular components, while tRF-3 target genes were associated with 13 molecular functional categories, 26 biological activities, and seven cellular components (Supplementary Fig. 12). A large number of target genes for the tRF-5 series were involved in catalytic (7 1 1), transferase (3 3 9), kinase (1 3 8) activities (Supplementary sheet 2.3). The target genes for the tRF-3 series were mostly associated with catalytic (9 0 3), and transferase activities (4 1 6) (Supplementary sheet 2.4). For a more organized visualization of our enrichment results, we clustered our enriched genes into enrichment maps, and clusters of similar pathways representing major biological processes were generated. The tRF-5 targets were enriched in growth and development, transport and localization, and cellular metabolic processes (Supplementary Fig. 13), while the tRF-3 targets formed two distinct clusters related to developmental and cellular metabolic processes (Supplementary Fig. 14). It can be predicted that many tRF-5 and tRF-3 may act as potent regulators of the genes involved in plant development, and cellular metabolic machinery in a tissue/condition-specific manner. The top one target analysis reveals that conserved tRF sequences can bind to different transcripts in plants, but the genes targeted by the same tRF don’t need

![Fig. 5. Correlation graph showing the total tncRNAs identified vis per tRNA gene copies in individual plants.](image-url)
to have an identical function in different plants (Supplementary sheets 2.5 and 2.6). It can be speculated that tRFs can interact with multiple genes, and may regulate various activities in the plant system.

We also examined the conserved tRF-5 and tRF-3 against full transposable elements (TEs) from Arabidopsis. Many tRF-5 and tRF-3 sequences were associated with several TE transcripts (Supplementary sheets 2.7 and 2.8). A large number of tRF-5 showed association with various transponson families (Fig. 7). As compared to tRF-5 s, tRF-3 showed a weaker association with TEs. TE belonging to the LTR/Gypsy superfamily were the most frequent targets of the tRF-5 s. With this superfamily, TE from ATHILA2, ATHILA6A, ATHILA4C, ATHILA4A, ATLANTYS1, ATHILA6B, and ATLANTYS2 families were found to be associated with a large number of TEs. Apart from them, TE superfamilies like RC/Helitron (ATREP3 and ATREP15) and DNA/Mudr (VANDAL2 and VANDAL3) also show considerable association with TEs. Smaller length TEs (mostly 19–22 nt) originating from AspGTC/ATC, GlnTTG, ProAGG/CGG/TGG, HisGTG, LeuCAA/TAG, and SerGCT tRNAs were associated with TEs in high numbers. tncRNAs binding with TE can bestow genome stability to plants. Interestingly, two recent reports have shown that TEs target transposable elements (retrotransposons) in both plants and mammals, indicating their potential as epigenetic regulators [51]. For instance, in Arabidopsis, a 19 nt tRF-5 from tRNA-MetCAT cleaves its target LTR retrotransposon transcript, Athila6A [52]. Deciphering the synchronization among the tncRNA generation, TE expression, and their implications in plants under normal and stress conditions can unveil additional regulatory functions of this class of ncRNAs.

### 2.7. tncRNAs profiling

We checked the organization of tncRNA sequences in each tncRNA class for tissue samples using t-Distributed Stochastic Neighbor Embedding (tSNE). The t-SNE plots revealed numerous clusters for individual tncRNA classes in different plants. In Arabidopsis, seedlings and seed clusters were frequently seen in tncRNAs belonging to tRF-5, tRF-3, tRF-1, and 5′tRH (Fig. 8). As compared to the tRNA halves, smaller fragments particularly belonging to tRF-5 and tRF-3 classes showed enriched clustering for more than ten different tissues. It was interesting to observe that vasculature, and epidermal tissue clusters were found to be in close proximity in tRF-5, tRF-3, and 5′tRH (Fig. 8). As compared to the tRNA halves, smaller fragments particularly belonging to tRF-5 and tRF-3 classes showed enriched clustering for more than ten different tissues. It was interesting to observe that vasculature, and epidermal tissue clusters were found to be in close proximity in tRF-5, tRF-3, and 5′tRH. Surprisingly, seedling, seed, shoot, rosette leaf, inflorescence, silique, and flower clusters were also seen in the tRF-1 series. Some clusters belonging to various tissues like leaf, anther, and seedling were also observed in rice, tomato, and maize (Supplementary Figs. 15–17). The distribution pattern of tncRNAs belonging to various tissues indicates that spatiotemporal expression of tncRNAs occurs in a tissue-specific manner at normal physiological conditions.

### 2.8. Differentially expressed (DE) tncRNAs

We checked the expression of tncRNAs under various abiotic and biotic stresses in Arabidopsis, tomato, rice, and maize (Supplementary sheet 3.2–3.5). Significant differences in tncRNA expression were observed in different plants under different conditions. Besides up-regulated fragments, a significant number of tncRNA were down-regulated in various samples. We observed stress,
time, or tissue-dependent tncRNA expression in different plants. For instance, in Arabidopsis, only 3 tncRNAs showed dysregulation after 0.5 h of heat stress but their number rose to 42 after a time duration of 6 h. Likewise, only 25 tncRNAs were expressed during drought stress at 20% field capacity (FC), whereas 182 DE tncRNAs were detected when FC was 30% drought stress. On the other hand, less than 10 DE tncRNAs were detected in Magnaporthe oryzae infected root tissues (9 and 7 DE tncRNAs after 30 and 120 min respectively). However, the numbers were highly elevated in infected leaves as compared to root tissues (87 and 62 tncRNAs after 30 and 120 min respectively). The time-dependent difference in expression was also observed in few tissues in maize under heat stress (2 h vs 48 h). Among all the conditions studied, we found that heat exposure leads to the maximum number of differentially expressed tncRNAs in different maize tissues, both in vegetative and reproductive stages when compared with control. The highest among all with a total of 1748 tncRNAs were differentially expressed in tassel in reproductive stage upon exposure to 48 h of heat stress, among which 35 nt tRF derived from GlyTCC, was significantly downregulated with a log2FC of ~30. The tomato plant when subjected to Tomato Mosaic Virus (TMV) infection showed 757 DE tncRNAs, of which the majority of the fraction were contributed by 15 nt tRFs. In addition, it was noticed that most of the DE tncRNAs were generated from HisGTG in Arabidopsis, GlnCTG in maize, and GluTTC in rice as well as tomato. We also observed the upregulation of 19 nt tRF-5 (ArgCCT) under drought (30% FC) and NaCl stress in Arabidopsis. It also showed elevated expression in maize leaf subjected to heat stress. In earlier studies, it was reported that this tRF was expressed at high levels in Arabidopsis seedlings under drought conditions and cold-treated rice inflorescences [22]. The findings suggest that the accumulation of evolutionarily conserved plant tncRNAs is regulated by biotic and environmental stresses.

3. Discussion

We have developed a workflow/pipeline to accurately identify genuine tncRNAs from sRNA-seq datasets. The workflow has been designed for convenient tncRNA mining and can be tailored as per the requirement. In the past two decades, various research groups have contributed to the tncRNA identification in the animal world, particularly in humans. Some databases e.g. tRex and PtRFdb have been developed in the recent past for the exploration of tncRNAs in plants. Also, an annotation pipeline, SPORTS1.0 was developed for profiling canonical small RNAs like miRNAs, piRNAs, rRNAs, and tRNA-derived sRNAs in various organisms including some plants [53]. In this study, we have reported 43,153 novel tncRNAs (Supplementary Fig. 18) by using our workflow/pipeline (i.e. tncRNA Toolkit), and tested it on previously published datasets [52]; and further compared it with SPORTS 1.0. Both pipelines (i.e. tncRNA...
Toolkit and SPORTS 1.0) detected the previously validated 19 nt tRF-5 (AlaAGC) in all five replicates in Arabidopsis pollen. Comparison of various features, and outputs of the tncRNA Toolkit with already existing databases (i.e. tRex and PtRFdb), and methodologies (SPORTS 1.0) reveals the clear difference in the number of reported tncRNAs because of the differences in mapping strategies, read count, read length, and read classification parameters (Supplementary sheets 4.1–4.3). SPORTS1.0 profiled canonical tncRNAs coming from mature tRNAs and categorized them in four categories based on whether they are derived from 5′ end, 3′ end, 3′ CCA end of tRNAs, or from other regions within the mature tRNA. The various tncRNA subtypes and their respective counts from different regions of mature tRNAs from both pipelines have been provided in Supplementary sheet 4.3. By comparing the overall results, it can be stated that tncRNA Toolkit identify and classify the tncRNAs into various subclasses sourcing from both mature as well as pre-tRNAs, and can detect potential tncRNAs including experimentally validated ones. Also, with tncRNA Toolkit, the read count cut-off, number of allowed mismatches, and threshold of suppressing multi-mapped alignments can be altered as per the user requirements, making this a superior tool. Moreover, in addition to the application of computational approaches for the prediction of tncRNAs, studying and validating the tncRNAs in the future will provide the opportunity for better benchmarking the tools, and pipeline using real positive datasets.

In recent years, many studies associated with eukaryotic tRNA fragments have revealed newer complexities and functions linked with the conserved tRNA molecules beyond their established role in the canonical translation process [54–56]. However, structured searches for tncRNAs have so far been restricted to their biogenesis and functional role as regulatory molecules analogous to miRNAs. The existence of tRNAs with the same anticodon sequence (isodecoders) necessitates further exploration of their role in tncRNAs biogenesis. Our observations confirm the existence of a heterogeneous pool of tncRNAs in plants. Organellar tncRNAs identified in this pool demand further attention for exploring their probable functions in the cellular milieu. In plants, the existence of chloroplastic tRNAs in addition to mitochondrial tRNAs adds another layer of intricacy in terms of their biogenesis, localization, and functions. Conservation of cleavage consensus sites in different plants, and codon-dependent tncRNA generation support that they are the product of non-random ribonucleolytic tRNA cleavage. Also, the DE tRF-1 reported for the first time in our study encourages us to commence the exploration of these fragments in planta. The identification of tncRNA targets, as well as DE tncRNAs, is just a head start to explore innumerable pathways that may be critically regulated by tncRNAs in various cellular processes. The tissue-specific organization of tncRNAs suggests their potential biological role in plants. Further validation of DE tncRNAs and their putative targets will pave the way for a better understanding of the underlying mechanism in regulating stress responses. Differences in tncRNAs expression suggest their multifarious physiological roles particularly in relevance to stress responses. The tncRNA sequences particularly in the length range of 19–25 nt have been previously studied for the Argonaute proteins (AGOs) interaction and Argonaute-immunoprecipitation (AGO-IP) libraries have been observed to be rich in tRNA fragments [21,22]. However, in our study, we have seen the abundance of tncRNAs in other libraries as well. This indicates that AGOs may or may not associate with all tncRNAs. Apart from conventional AGO binding mediated RNA interference, many tncRNAs may be functionally independent of AGO, unlike miRNAs. Also, the expression of tncRNAs in non-AGO

Fig. 8. t-Distributed Stochastic Neighbor Embedding (t-SNE) for the visualization of the tncRNA organization in Arabidopsis tissue samples. Individual t-SNE plots were made for tRF-5, tRF-3, tRF-1, 5′tRH, and 3′tRH classes. The tncRNAs from a total of 525 samples from 13 different tissues were utilized for this analysis. Tissue cluster observed have been shown by similar color-coded dashed circles.
ILPs indicates the existence of some unrevealed tncRNA regulatory mechanisms that might be involved in fine-tuning gene expression. tRNA is post-transcriptionally modified at various nucleotides, their vital role in structural integrity and the translation is well established, but the detailed mechanisms showing how these tRNA modifications affect the regulation of their cleavage remain unclear. Our study provides a future perspective for better understanding the structural and functional impact of tRNA modifications in tRNA biology.

Although the in-silico approach will help in identifying these novel molecules across a wide range of organisms, experimental validation and characterization are vital for distinguishing more and more bona fide tncRNAs. Switching from conventional RNA-seq to tRNA-seq to overcome modification biases will be advantageous for the identification of tncRNAs. To study the molecular mechanisms of tncRNAs, apart from classical molecular biology methods, advanced ribonomics [57] like cross-linking, ligation, and sequencing of hybrids (CLASH) may lay the foundation for studying tncRNA-target hybrids in plants. As tncRNAs are ubiquitous and conserved across different domains of life, our comprehensive study is believed to substantiate research of these novel molecules. Their global identification will facilitate deciphering common conserved pathways in eukaryotes and their mechanism of unexplored regulatory action. Leveraging computational power together with molecular biology techniques will augment the current understanding of the vast tncRNAomic landscape in the living system.

4. Methods

4.1. Data retrieval

Genome assembly for Arabidopsis thaliana (TAIR10.1), Solanum lycopersicum (nuclear: SL4.0; organellar: SL3.0), Cicer arietinum (nuclear: ASM33114v1; organellar: ASM33114v1), Medicago truncatula (nuclear: MedtrA17_4.0; organellar, MtrunA17r5.0-ANR), Oryza sativa (Build 4.0; organellar: IRGSP-1.0), and Zea mays (Zm-B73-REFERENCE-NAM-5.0) were downloaded, and IDs were transformed for starting with ‘chr’, to make them foundation for studying tncRNA-target hybrids in plants. As tncRNAs are ubiquitous and conserved across different domains of life, our comprehensive study is believed to substantiate research of these novel molecules. Their global identification will facilitate deciphering common conserved pathways in eukaryotes and their mechanism of unexplored regulatory action. Leveraging computational power together with molecular biology techniques will augment the current understanding of the vast tncRNAomic landscape in the living system.

4.2. tRNA annotation and genome pre-processing

Nuclear tRNAs were predicted by tRNAscan-SE [44] (v2.0.6) using the eukaryotes model, while organellar tRNAs were detected by using the ‘-O’ option in this tool. The tRNA pseuodogenes and tRNAs with a score less than 50 were eliminated, and filtered tRNA genomic region flank. These ‘stockholm’ format files were used to create postscript files available for the abovementioned plants were downloaded using NCBI SRA Toolkit (v2.10) [58].

4.3. Modification site prediction

The single-end small RNA reads were processed by Trim Galore v0.6.6 (https://github.com/FelixKrueger/TrimGalore) to remove the low-quality bases and trim the adapter sequences. To permit error-tolerant and uniquely mapping reads alignments, processed reads were aligned to the artificial genome index using bowtie [60] with ‘-v2’ and ‘best’ options. The SAM output was sorted and converted to BAM by samtools [61] (v1.10). The modification sites were predicted by HAMR (v1.2) (python hamr.py IN.bam REF.fa models/euk_trna_mods.Rdata OUT ath 30 10 0.05 H4 0.01 0.05 0.05) [48,62]. Further, modification sites only on mature tRNA and leader/trailer regions were counted.

4.4. tncRNAs identification and classification

The mapped reads from previously generated SAM files were fetched to create a FASTA file of unique reads with the count, using SAM flag 0 & 16 for single-end reads. Mapped unique reads were aligned to +ve strand only with no mismatch, and multi-mapped reads were discarded if alignment occurred greater than 50 times, using bowtie1 with ‘-anchor -v 0 -m 50’ arguments. The bowtie combinatorial arguments ‘-best’ and ‘-strata’ were used, which guarantees that reported singleton alignments are best in terms of the stratum. Then, the output was used to identify locus, location, length of read mapped to mature tRNA, and their 50 bp upstream and downstream flank; based on that, reads were classified into different classes of tncRNAs. The tRNA halves were classified by cleavage at the anticodon loop (2 nt + 3 nt of anticodon + 2 nt = 7 nt), and information for modification site was added for tncRNAs. Reads per million (RPM) was calculated for each tncRNA by using the formula:

\[
\text{Per Million Factor (PMF) = Total mapped reads} / 10^6
\]

\[
\text{RPM = Number of reads mapped to progenitor tRNA} / \text{PMF}
\]

A local alignment file of tncRNA to the sequence of origin was also created for visualization using the pairwise2 biopython module [63]. The alignment score is given as per identical = 1, non-identical = −1, gap-open = −1, gap-extend = −0.5

4.5. tRNA model for consensus sequence and structure

The FASTA files were generated for specific isoacceptor tRNAs and provided as an input to the LocARNNA [64] (v1.9.2) software package with ‘--stockholm’ option for consensus sequence study. These ‘stockholm’ format files were used to create postscript files by RNAAlifold [65] (v2.4.11) for the consensus tRNA sequence. The modification sites predicted for each tRNA isoacceptor have been shown on the consensus tRNA structures.

4.6. Target prediction, GO, and pathway enrichment

The psRNATarget [66] software was utilized for tncRNA target prediction (2017 release; default parameters). Common tncRNAs (tRF-5 and tRF-3 series; 18–25 nt long) were used as a query, while Arabidopsis CDNA and transposons sequences were used as probable targets. The protein targets were analyzed for pathway enrichment analysis using g:Profiler [67] and visualized in Cytoscape [68] (EnrichmentMap [69] and AutoAnnotate [70]).

4.7. t-Distributed Stochastic Neighbor Embedding (t-SNE) plot

The psRNATarget software was utilized for tncRNA target prediction (2017 release; default parameters). Common tncRNAs (tRF-5 and tRF-3 series; 18–25 nt long) were used as a query, while Arabidopsis CDNA and transposons sequences were used as probable targets. The protein targets were analyzed for pathway enrichment analysis using g:Profiler [67] and visualized in Cytoscape [68] (EnrichmentMap [69] and AutoAnnotate [70]).

4.7. t-Distributed Stochastic Neighbor Embedding (t-SNE) plot

t-SNE is a non-linear technique for dimensionality reduction that is particularly well suited for the visualization of high-dimensional datasets. We employed t-SNE to study datasets comprising of unique tncRNA sequences belonging to tRF-5, tRF-3, tRF-1, 5’tRH, and 3’tRH in major tissues from the Arabidopsis, rice, tomato, and maize. The tissues with at least 10 samples (Supplementary sheets 5.1–5.4) were utilized for matrix generation. A binary matrix, for presence (1) and absence (0), was created for unique tncRNA sequences from each tissue sample. From this, the plot was generated for each aforementioned class of tncRNAs.
4.8. Differential expression study

DESeq was utilized for the identification of differentially expressed tncRNAs under stress conditions [71]. Only those samples were selected in which at least three biological replicates for each control and treatment were present (Supplementary sheet 3.1). In DESeq2, the p-values attained by the Wald test are corrected for multiple testing using the Benjamini and Hochberg method by default. For our analysis, only those tncRNAs with a p value less than 0.05 were considered to be significant. Only significant tncRNAs were reported in expression analysis. tncRNAs with log2FC value greater than 1 (>1), and less than –1 (<–1) were considered up- and down-regulated, respectively.

5. Code and Data availability

All pipeline scripts, codes, data generated, and analyzed for each of the species are freely available at our website (URL: http://nipgr.ac.in/tncRNA). The codes and usage are also available at https://github.com/skbinfoc/tncRNA-ToolKit.

Author contributions

S.Z. and A.S. designed the pipeline, wrote the codes, and performed all the analysis. N.P. contributed to the data analysis. S.Z. and SK wrote the manuscript. SK conceived the study and coordinated the project.

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Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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