Investigating the Role of Translationally Control Tumor Protein in Growth, Development and Differentiation of *Dictyostelium discoideum*

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Translationally controlled tumor protein (TCTP) is a multifunctional protein implicated in various types of cellular processes involving growth and development of an organism. Here, we identified *tctp* gene in *Dictyostelium discoideum* and unraveled its function. The sequence analysis of *D. discoideum* TCTP (*DdTCTP*) showed its conservation among eukaryotes. Transcript of *DdTCTP* was highly expressed at the initial time points of development and protein is localized both in the cytoplasm and nucleus. Disruption of *tctp* was achieved by BSR cassette using double homologous recombination method. Abrogation of *tctp* resulted in reduced cell proliferation but increased cell size. Additionally, development was delayed by 4 h wherein small-sized aggregates and fruiting bodies were produced by *tctp*− cells while larger aggregates and fruiting bodies were produced by *tctpOE* cells concordant with the fact that TCTP regulates prestalk/prespore ratio and cell-type differentiation. *tctp*− cells produced round spores with reduced viability and stalk cells are arranged in septate pattern as compared to polyhedral manner of wild type. Abrogation of *tctp* resulted in aberrant localization of cell type specific markers and show low proclivity toward prespore/spore region, in presence of wild type cells.

**Keywords:** *Dictyostelium*, development, growth, RT-PCR, TCTP, chimera

**INTRODUCTION**

Translationally controlled tumor protein (TCTP; also known as P21, P23, and fortilin) is implicated in various biological activities such as cell growth, cytoskeleton regulation (Gachet et al., 1999; Burgess et al., 2008; Tsarova et al., 2010), cell cycle (Brioudes et al., 2010), protein synthesis (Cans et al., 2003), apoptosis (Li et al., 2001; Liu et al., 2005; Yang et al., 2005), development (Chen et al., 2007; Hsu et al., 2007), and cancer (Tuynder et al., 2002, 2004; Rho et al., 2011). TCTP has a critical role in tumorigenesis, which is now recognized as a therapeutic target for various cancers such as prostate, lungs and breast cancers (Acunzo et al., 2014). TCTP is a ubiquitously expressed highly conserved protein present in eukaryotic system from plant to yeast and protozoans to all invertebrates and vertebrates and involved in various kinds of biotic and abiotic stresses (Rupec et al., 1998; Sturzenbaum et al., 1998; Cao et al., 2010). It is crucial for development of
an organism as the knockout in mice is lethal where the fetus dies in the uterus due to inhibition of cell proliferation and increased cellular apoptosis (Chen et al., 2007; Susini et al., 2008). The embryo suffers reduction in cell number and increased apoptosis at embryonic stage 5.5 and dies around stage E9.5–10.5 (Chen et al., 2007). TCTP mutant mice show defect in neuronal and glial differentiation during CNS development (Chen et al., 2020). Also, knockdown of TCTP in Drosophila results in reduced cell number and organ size causing the larvae to die by first-instar stage (Hsu et al., 2007). High abundance of TCTP transcript is found in retinal ganglion cell and adult axons indicate TCTP regulates the growth of retinal ganglion cell axons (Roque and Holt, 2018). It is associated with the survival machinery of axons and has role in the development of vertebrate neural circuit (Roque et al., 2016). In Drosophila, RNA interference of TCTP causes a reduction in size of eye and wing which are due to decrease of both cell number and size. It also helps in the development of brain, wing and endocycling cells of salivary glands of Drosophila by maintaining the genome stability of mitotic cells (Hong and Choi, 2013). In plant or Arabidopsis, downregulation of TCTP results smaller leaf and flower size, reduced root growth, and leads to delayed development (Berkowitz et al., 2008). TCTP also controls the cytoskeleton, such as microfilament and microtubule by interacting with actin and tubulin proteins (Gachet et al., 1999). Hence, TCTP plays a critical role in cell proliferation in a cell or tissue-type specific manner.

Dictyostelium discoideum encompasses the unicellular and multicellular life forms that makes it tractable for the study of genetic changes occurring at the intersection of uni- and multicellular life (Eichinger et al., 2005). It has been used for the study of chemotaxis, cell biology, and developmental biology (Firtel and Chung, 2000; Cardelli, 2001). In metazoans, multicellularity results by the repeated cell division of the “zygote” but in case of Dictyostelium multicellularity arises due to the aggregation of spatially segregated cells. Dictyostelium feeds on bacteria and divides mitotically to remain as unicellular organism but enters into multicellularity at the onset of starvation (Loomis, 2014). Following starvation, some of the cells start secreting cAMP and in response to this signal other cells start moving toward cAMP gradient (Saran et al., 2002). These cells communicate with each other and form aggregation territories (mounds) comprising of approximately $10^5$ amoebae which undergo various morphogenetic movements to form slugs comprising anterior $\frac{1}{3}$ region of prestalk cells and the remaining posterior region of prespore cells and then fruiting bodies that made up of dead vacuolated stalk cells and viable spores (Raper, 1935; Wittingham and Raper, 1960; Williams et al., 1989).

In this paper, we have identified and characterized Dictyostelium TCTP (DdTCTP) that show conservation to human and yeast TCTP. Disruption of DdTCTP was achieved using Blastidin-S (BSR) cassette to unravel its function in growth and development of this organism. The tctpOE (overexpressor) and tctp− (knockout) cells showed reduced cell proliferation. Disruption of tctp exhibited increased cells mass, cell size and reduced pinocytosis. Both, tctpOE and tctp− cells displayed delay in development resulting in fruiting body formation at 28 h, respectively. Additionally, tctpOE cells formed large-sized aggregates and fruiting bodies with a larger stalk and a small sorus. On the contrary, tctp− cells formed small-sized aggregates and fruiting bodies. tctp− fruiting bodies formed round spores in contrast to elliptical spores of wild type and exhibited only ~25% viability. In chimeras of Ax2 and tctp− cells, mutant cells showed more propensity toward prestalk/stalk region and its derived structures and contributed less in chimeric spore formation.

**MATERIALS AND METHODS**

**Cell Growth, Development and Transformation**

Dictyostelium (Ax2) cells were grown axenically and development was performed as described (Gosain et al., 2012). Log phase cultures were inoculated into fresh media at a density of $5 \times 10^3$ cells/mL and cell proliferation was monitored over 5 days using hemocytometer. For development, log phase cultures were harvested, washed and developed at a density of $5 \times 10^5$ cells/mL, followed by synchronization at 4°C for 4–6 h and further incubated at 22°C to proceed in development. The timings and pattern of development was noted and photographed under Nikon SMZ-1500 stereomicroscope. Analyses were performed using NIS-ELEMENTS AR v. 4.0.

**Preparation of tctp Overexpressing (tctpOE) and tctp Knockout (tctp−) Strains**

Tctp overexpressing (tctpOE) strain was made as described earlier (Kumar et al., 2017). The knockout of tctp gene was created by disrupting the tctp gene with BSR cassette, followed by homologous recombination in the genome of D. discoideum. The 5’ and 3’ homology regions were PCR amplified using gene-specific primer pairs (Supplementary Tables S1, S2) and cloned at the flanking sides of BSR cassette, followed by linearization of the construct (Supplementary Figure S1). The linearized product (5’ homology region + BSR cassette + 3’ homology region) was transformed into Ax2 cells by electroporation and transformants were selected at 10 µg/mL Blastidicin-S (Invitrogen).

**RNA Detection by RT-PCR Analysis**

RNA isolation and cDNA synthesis were performed as described (Gosain et al., 2012). Briefly, Dictyostelium cells were developed and samples were collected at various time points, washed and lysed in TRI reagent (Sigma-Aldrich, United States). cDNA was synthesized and PCR amplification was performed. The gene-specific primer pairs used for RT-PCR reactions were mentioned in Supplementary Table S2.

**Pinocytosis Assay**

Pinocytosis assay was performed as described (Maurya et al., 2020). Briefly, log phase cultures of Ax2 and tctp− cells were harvested, re-suspended in fresh media at a density of $3 \times 10^6$ cells/mL and kept under shaken conditions for 20 min. Furthermore, FITC dextran (2 mg/mL; Mr = 70 kDa;
Cells were developed at a density of $10^7$ of Ax2 and Calcofluor staining of stalk and spores under moist conditions. *Klebsiella aerogenes* and aliquots of 100 spores were mixed with the suspension of agar. Spores from mature fruiting bodies were collected, washed and aliquots of 100 spores were mixed with the suspension of *Klebsiella aerogenes* (bacteria) and allowed to grow for 5 days under moist conditions.

**Cell Dry Weight Analysis**

Cell dry weight measurement was done according to Maurya et al. (2017). Log phase cultures were harvested, washed and pelleted having $5 \times 10^7$ cells were vacuum dried at $55^\circ$C for 1 h and cell weight was scored by subtracting empty tube weight.

**Cell Size Analysis by Flow Cytometer**

Cell size analyses were performed as described (Maurya et al., 2017). Briefly, approximately $1 \times 10^8$ cells were harvested, washed with 1xKK$_2$ buffer, resuspended in 1.5 mL buffer (0.9% NaCl, 2% sucrose, 5 mM EDTA in KK$_2$ buffer) followed by fixation. Just prior to analysis $1 \times 10^6$ cells were washed, treated with RNaseA and incubated at $37^\circ$C for 30 min followed by addition of propidium iodide (Sigma-Aldrich, United States) and incubated at RT. Sample was loaded on BD FACS Calibur flow cytometer and cell size analysis was performed using Cell Quest software.

**Spore Viability Assays**

Spore viability assay was performed according to the procedure as described (Myre et al., 2011). Approximately, $1 \times 10^8$ cells were harvested, washed and developed on 1.5% non-nutrient agar. Spores from mature fruiting bodies were collected, washed and aliquots of 100 spores were mixed with the suspension of *Klebsiella aerogenes* (bacteria) and allowed to grow for 5 days under moist conditions.

**Calcofluor Staining of Stalk and Spores of Ax2 and tctp**

Cells were developed at a density of $5 \times 10^7$ cells/mL on nitrocellulose membrane, already placed on agar till the formation of fruiting bodies, followed by treatment with calcofluor white (Sigma) at 0.01% wt/vol for 10 min. The staining solution was removed and images were captured using fluorescence microscope (Nikon Eclipse TiE).

**Development of Chimeras**

RFP marked Ax2 and GFP marked tctp$^-$ cells were mixed in various proportions (10–75%) and developed as described above. The development of chimeras was monitored under fluorescence microscope (Nikon SMZ-1500) and photographed.

For scoring contribution of Ax2 and tctp$^-$ cells in chimeric spore formation, individual spore-heads were picked on a glass-slide and photographed, both under brightfield and fluorescence using Nikon Eclipse 80i microscope. Red and green fluorescent spores were counted and graph was plotted. A minimum of 10–15 fruiting bodies developed from each ratio mixture per individual experiment were counted.

**β- Galactosidase Staining**

β-galactosidase staining was performed according to the method described by Escalante and Sastre (2006) with minor modifications (Mishra et al., 2017). Log phase cells were developed at a cell density of $5 \times 10^7$ cells/mL on dialysis membrane placed on non-nutrient agar plates. Various developmental structures were collected, fixed with methanol for 30 min followed by washing and permeabilization with NP40 (0.1% in Z buffer) for 1 h at room temperature. Further, structures were washed 2–3 times and staining with X-gal (5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside) was performed. The reaction was stopped when evident staining appeared by washing with Z buffer. Images were captured under the Nikon AZ100 microscope.

**Image Processing and Statistical Analyses**

Image processing and mound size analyses were performed using NIS ELEMENTS ARv.4.0. The statistical analyses were performed with mean, standard deviation, and standard error and values were plotted in graph using GraphPad Prism (GraphPad Software, CA, United States, www.graphpad.com). Student t-test was used where p-values; ***$p < 0.001$, **$p < 0.01$ and *$p < 0.05$ were established as significant.

**RESULTS**

**Identification and Characterization of DdTCTP**

To investigate the similarity between DdTCTP and TCTPs from other species, we performed multiple sequence alignment of protein sequences through MUSCLE program (Edgar, 2004). The protein sequence of DdTCTP is ~171 amino acids (source from dictybase server; Fey et al., 2009). Amino acid sequence alignment of TCTP from social amoeba (*D. discoideum*) and other species (including Arabidopsis thaliana, Drosophila melanogaster, *Schizosaccharomyces pombe*, *Mus musculus*, and *Homo sapiens*) was performed (Figure 1A). We aligned the TCTP protein sequences from seven species that cover lower to higher animals, plant and fungi. Conserved residues showed identity between social amoeba, plants, fungi and lower as well as higher animals. A constructed phylogenetic tree suggested that DdTCTP was closely related to the other members of dictyostelids and protozoa such as *Theileria annulata*, Dictyostelium fasciculatum, and *Plasmodium falciparum* with high bootstrap values (Figure 1B). DdTCTP was also closer to fungi (*Saccharomyces cerevisiae*, *Schizosaccharomyces japonicas*, and *Schizosaccharomyces pombe*). The homology between DdTCTP and *S. pombe* was 46% while with human it showed 42%.

The expression level of tctp mRNA during growth and development of *D. discoideum* was checked using semi-quantitative RT-PCR (Figure 1C). tctp mRNA was expressed throughout development, predominantly expressed at initial time points (0–6 h) then showed reduced expression between 8 and 20 h followed by increased expression till 24 h.
FIGURE 1 | Identification and characterization of DdTCTP. (A) Protein sequence alignment of DdTCTP and other species. Sequences used in this study are: Dictyostelium discoideum (DDB0305046), Arabidopsis thaliana (Uniprot id: P31265), Drosophila melanogaster (Uniprot id: Q9VGS2), Schizosaccharomyces pombe (Uniprot id: Q10344), Mus musculus (Uniprot id: P63028), and Homo sapiens (Uniprot id: P13693). Identical [*], 90% conserved [:], and 50% conserved [.] amino acid residues are indicated in multiple sequence alignment. Residues highlighted (rectangular boxes) in different color involve in different functions which are conserved from lower to higher organisms. Residues crucial for Rheb interaction are shown in yellow color, while residues involved in microtubules and Ca\textsuperscript{2+} are highlighted in green color. Serine residues (Ser46 and Ser64) in mouse and human are known to be phosphorylated by the Plk (Polo like kinase) marked in black dots, are only conserved in mammalians. (B) Phylogenetic analysis of various TCTPs from protozoa and other species. Evolutionary distances between sequences are indicated as given scale in bottom. (C) Semi-quantitative RT-PCR of tctp transcript at different developmental time points. The values represent mean ± standard deviation; n = 3; ***p < 0.001, *p < 0.05.

Generation of tctp Knockout in Dictyostelium discoideum

To explore the functions of TCTP, its deletion mutant was generated by disrupting the tctp gene with BSR cassette through homologous recombination method in the genome of D. discoideum (Figure 2A and Supplementary Figure S1). Approximately, 400 independent clones were screened for positional integration, confirmed by PCR and one out of the two positive clones was used for further experiments (Figures 2B,C).

Dictyostelium TCTP Reduces Cell Proliferation

For measuring cell proliferation, log phase cultures were identifiedly diluted into fresh HL5 media and observed over 5 days (Figure 3A). Wild type and tctp\textsuperscript{OE} cells divides at every 12.3 ± 0.2 and 18 ± 0.6 h and attained stationary phase at ~10.8 × 10\textsuperscript{6} cells/mL and ~7.2 × 10\textsuperscript{6} cell/mL, respectively, while cell proliferation of tctp− cells was very slow with a longer lag phase and reached the stationary phase and at a much lower density of 4.7 × 10\textsuperscript{5} cell/mL (Figure 3A). The doubling time for tctp− cells was 26 ± 2.5 h. Growth of Dictyostelium was accomplished with either increase in cell proliferation or increase in size of the cells. Cell growth and proliferation are regulated independently in the organism. Cell proliferation rate is increase in number while cell growth is increase in size of cell due to the increased molecular mass in the cell. To determine the role of DdTCTP in regulating growth, we estimated the cell mass and observed that Ax2, tctp\textsuperscript{OE}, and tctp− cells have an average cell mass of 9 ± 0.08, 8.1 ± 0.2, and 11.4 ± 0.5 mg, respectively (Figure 3B). Increased cell mass of tctp− cells was attributed to the enhanced cell size revealed by FACS analysis (Figure 3C). The size of the tctp− cells was also observed by microscopy and found significantly more than wild type cells (Figure 3D and Supplementary Figure S2). Furthermore, the reason behind increased cell size was checked via nuclear staining using DAPI and found that there was increase in number of nuclei per cell in case of tctp− cells as compared to wild type cells (Figure 3E). Together, we conclude that tctp− cells display reduced proliferation but enhanced cell growth due to increased number of nuclei. Next, we wanted to check the effect of TCTP knockout on pinocytosis (rate of liquid uptake; Figure 3F). Pinocytosis also known as macropinocytosis is the uptake of liquid media by axenic strain (Ax2) of Dictyostelium cells grown in laboratory. Pinocytosis is a crucial process in the growth of Dictyostelium Ax2 cells, as it
is the only mode of taking the nutrients (Bloomfield and Kay, 2016). Vegetative stage of Ax2 cells grow and divide by actively pinocytosing the external nutrients. It can be measured through the rate of uptake of fluid phase marker, fluorescent dextran (Maurya et al., 2020). The pinocytosis rate of tctp<sup>−</sup> cells was significantly reduced at all time points as measured by decrease in intracellular FITC content in the tctp<sup>−</sup> cells (Figure 3D). Therefore, decreased proliferation could be attributed to defect in nutrient uptake as well.

**Deletion of tctp Cause Developmental Delay and Aorist Fruiting Bodies Formation**

To delineate the role of TCTP in multicellular developmental program of Dictyostelium, wild type, tctp<sup>OE</sup>, and tctp<sup>−</sup> cells were plated at a density of 5 × 10<sup>7</sup> cells/mL on non-nutrient agar and allowed to develop at 22°C following synchronization (Figure 4). Ax2 cells completed its developmental life cycle by 24 h forming aggregation streams by 6 h, aggregates (mounds) by 12 h, slugs by 16 h, and fruiting bodies by 24 h whereas the formation of aggregates was delayed by 4 h in tctp<sup>OE</sup> and tctp<sup>−</sup> strains. The tctp<sup>OE</sup> and tctp<sup>−</sup> cells formed mound at 16 h, slugs at 20 h and fruiting body at 28 h (Figure 4). Additionally, the number of aggregates were significantly reduced in tctp<sup>OE</sup> and tctp<sup>−</sup> strains as compared to Ax2 (Figures 5A,B) but the size of aggregates was 3-fold more in case of tctp<sup>OE</sup> whereas tctp<sup>−</sup> showed comparatively smaller aggregates than Ax2. tctp<sup>OE</sup> cells formed long and slender slugs as compared to wild type whereas tctp<sup>−</sup> slugs was similar to wild type slugs. Then, we analyzed the fruiting body morphology of wild type, tctp<sup>OE</sup> and tctp<sup>−</sup> strains and found that tctp<sup>OE</sup> cells formed fruiting body with long, thin stalk and small sorus while tctp<sup>−</sup> cells developed into fruiting body with a small and thick stalk and a large sorus as compared to wild type fruiting bodies (Figure 5C). Moreover, in case of tctp<sup>OE</sup> and tctp<sup>−</sup> fruiting bodies stalk cells are arranged in a septate pattern as compared to the polyhedral manner in wild type cells.

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**FIGURE 2** Generation of tctp deletion mutant. **A** Schematic representation of tctp disruption using BSR cassette (blasticidin resistance cassette) employing double homologous recombination method. Linearized product (5′ homology region + BSR + 3′ homology region) was transformed into Ax2 cells and transformants were selected at 10 µg/mL blasticidin. **B** Confirmation of tctp<sup>−</sup> clone by PCR using primer pairs for positional integration. **C** Semi-quantitative RT-PCR to check tctp mRNA in Ax2, tctp<sup>OE</sup>, and tctp<sup>−</sup> cells.
FIGURE 3 | Abrogation of TCTP rendered enhanced cell growth but reduced cell proliferation. (A) Cell proliferation rate of Ax2, tctp\textsuperscript{OE}, and tctp\textsuperscript{−} cells was measured every 12 h. (Seeding density- 5 × 10\textsuperscript{5} cells mL\textsuperscript{−1}). (B) Cell dry weight analyses of Ax2, tctp\textsuperscript{OE}, and tctp\textsuperscript{−} cells. Cell dry weight was computed in milligram. (C) Relative cell size analyses of Ax2, tctp\textsuperscript{OE}, and tctp\textsuperscript{−} cells were carried out by FACS. (D) Measurement of cell size of Ax2 and tctp\textsuperscript{−} cells. (E) Quantification of nuclei per cell of Ax2 and tctp\textsuperscript{−} cells. (F) Pinocytosis measurement by FITC-dextran uptake assay in Ax2 and tctp\textsuperscript{−} cells. (The values represent mean ± standard error mean; n = 3; ***p < 0.001, **p < 0.01, *p < 0.05; Student’s t-test; and Scale bar- 10 μm). (Figure 5D). Further, cellulose deposition in fruiting bodies of these strains were checked by calcofluor staining and found less cellulose deposition in the septum of tctp\textsuperscript{−} stalk (Figure 5D and Supplementary Figure S3A). The spores formed by tctp\textsuperscript{OE} cells were elliptical similar to Ax2 but tctp\textsuperscript{−} spores were round in shape (Figure 5D and Supplementary Figure S3B). cotA (spore COaT protein, SP96) and cotB (spore COaT protein, SP70) codes for major spore coat proteins, SP96 and SP70, respectively, that are necessary for the integrity of the spore coat (Srinivasan et al., 2000). Altered levels of any of these coat proteins are responsible for defect in shape and size of spores formed. Since the shape of tctp\textsuperscript{−} spore was largely altered, thus we checked the expression of spore coat genes cotA and cotB at the mRNA level by semi-quantitative RT-PCR in different stages of development (Figure 5F). We observed the expression of cotA remained unaffected, while the transcript of cotB was down regulated in tctp\textsuperscript{−} strain as compared to wild type (Figure 5F). Collectively, the above results suggest that TCTP is required for timely development and appropriate formation of aggregates and fruiting bodies.
Aberrant fruiting body formation by mutant cells led us to ask whether spores formed by mutant cells were viable or not therefore, spore viability assay was performed. Wild type and \( \text{tctp}^{\text{OE}} \) cells produced spores with average spore viability of 85.1 ± 1.7 and 65.3 ± 5.5%, respectively, whereas \( \text{tctp}^{-} \) cells produced 27.7 ± 1.2% viable spores (Figure 5E). Therefore, TCTP is essential for maintaining the viability of \( D. \text{discoideum} \) spores.

**mRNA Expression Analysis of Early Developmentally Regulated Genes**

Initial development of \( D. \text{discoideum} \) is mainly regulated by proteins involved in aggregation such as \( \text{acaA} \) (adenylate cyclase aggregate A), \( \text{carA} \) (cAMP receptor A), cell adhesion molecules \( \text{csA} \) (contact site A), and \( \text{cadA} \) (Calcium-dependent ADhesion). \( \text{acaA} \) and \( \text{carA} \) are major components for cAMP signal relay. Since, we observed delay in initiation of streaming we checked the mRNA expression of \( \text{acaA} \) and \( \text{carA} \) in \( \text{Ax2}, \text{tctp}^{\text{OE}}, \) and \( \text{tctp}^{-} \) strains (Figures 5G,H). The transcript level of \( \text{acaA} \) remain unaffected while \( \text{tctp}^{\text{OE}} \) showed reduced expression at 8 h (Figure 5G). The mRNA expression of \( \text{carA} \) was significantly down regulated in \( \text{tctp}^{\text{OE}} \) at 8 h and \( \text{tctp}^{-} \) at 8, 12, and 16 h. Furthermore, we checked the mRNA expression of cell adhesion molecules \( \text{csA} \) and \( \text{cadA} \) that are required for aggregate formation in early development. \( \text{cadA} \) expression remained unchanged and \( \text{csA} \) level was significantly reduced in \( \text{tctp}^{\text{OE}} \) cells resulting in larger aggregate formation. The \( \text{cadA} \) mRNA was down regulated and \( \text{csA} \) expression was up regulated at 12 h contributing to decrease cell-cell adhesion and stream break up in \( \text{tctp}^{-} \) aggregates (Figure 5H). The above data suggest that reduced \( \text{acaA} \) and/or \( \text{carA} \) may be responsible for the delay in development and altered \( \text{cadA} \) and/or \( \text{csA} \) expression contribute to altered aggregates formation in \( \text{tctp}^{\text{OE}} \) and \( \text{tctp}^{-} \) strains.

**Disruption of tctp Alters Spatial Cell Type Patterning**

Aberrant fruiting body formation by mutant cells incited us to study its role in cell type differentiation and patterning. The spatial distribution of prespore/prespalk cells in multicellular structures developed by wild type and \( \text{tctp}^{-} \) cells was analyzed using \( \text{lacZ} \) reporter fused to promoter of prestalk/prespore specific marker genes such as \( \text{ecmA} \), \( \text{ecmB} \), and \( \text{pspA} \) (d19). The \( \text{ecmA} \)/\( \text{lacZ} \) staining in wild type slugs was observed in the anterior region of slugs and in ALCs (anterior-like cells) of posterior region while in \( \text{tctp}^{-} \) mound it was found in anterior region but absent in ALCs of posterior region of slug. In \( \text{tctp}^{-} \) early culminant and culminant, \( \text{ecmA} \)/\( \text{lacZ} \) staining was absent in the lower cup, basal disk, and the stalk tube (Figure 6A). In wild type, the \( \text{ecmB} \)/\( \text{lacZ} \) staining was observed in the central core region of prestalk cells of migrating slugs and distributed in the basal disk, upper and lower cups of fruiting bodies formed (Figure 6B). In \( \text{tctp}^{-} \) cells, \( \text{ecmB} \)/\( \text{lacZ} \) expression is precocious as found at mound stage when compared with the wild type. The \( \text{ecmB} \)/\( \text{lacZ} \) staining in \( \text{tctp}^{-} \) early and late culminant was found similar to wild type. The \( \text{pspA} \)/\( \text{lacZ} \) staining was found in the prespore/spore region of the multicellular structures developed but the \( \text{pspA} \)/\( \text{lacZ} \) staining region was reduced in \( \text{tctp}^{-} \) slugs, early and late culminant (Figure 6C). Thus, we suggest that TCTP plays an important role in the distribution of cell types in multicellular structures.

**tctp^{-} Cells Exhibit More Tendency Toward Prestalk/Stalk Region**

To trace the fate of \( \text{tctp}^{-} \) cells in a chimera, cell types mixing experiments were performed in which \( \text{Ax2} \) (RFP-tagged) and \( \text{tctp}^{-} \) (GFP-tagged) mixed in varying ratios of 9:1, 3:1, 1:1, and 1:3 and allowed to co-develop (Figure 7 and Supplementary Figures S4–S6). Chimeras are the mixture of cells from two or more genetically different backgrounds.

**FIGURE 4** Overexpression/deletion of \( \text{tctp} \) impaired development of \( D. \text{discoideum} \). Comparative developmental profiles of \( \text{Ax2}, \text{tctp}^{\text{OE}}, \) and \( \text{tctp}^{-} \) strains developed at equal cell density and incubated at 22°C for development. White, red, and yellow arrows denoted mound, slug and fruiting body observed in \( \text{Ax2}, \text{tctp}^{\text{OE}}, \) and \( \text{tctp}^{-} \) strains, respectively (Scale bar=200 μm).
The differential presence of wild type and/or mutant cells in development and differentiation can be clearly observed by chimera study. Here, we wanted to observe the differential preference of mutant cells in presence of wild type cells. During early development, Ax2 and tctp\textsuperscript{−} cells contribute equally in formation of aggregation centers of various chimeras (1:9, 1:3, 1:1, and 3:1 of tctp\textsuperscript{−} and Ax2 cells; Supplementary Figure S4). As the development proceed further, in chimera of 10% tctp\textsuperscript{−} and 90% wild type cells, we observed that tctp\textsuperscript{−} cells occupied the prestalk/stalk region.

FIGURE 5 | TCTP regulates size of aggregates and fruiting bodies. (A) Numbers and (B) size of aggregates formed during development of Ax2, tctp\textsuperscript{OE}, and tctp\textsuperscript{−} strains. (C) Image showing a single fruiting body (n = 5; Scale bar = 50 µm). (D) Stalk and spore anatomy of Ax2, tctp\textsuperscript{OE}, and tctp\textsuperscript{−} strains. (E) Percent spore viability showing less viable spores of tctp\textsuperscript{OE} and tctp\textsuperscript{−} strains as compared to Ax2. (F) Relative transcript levels of spore coat A (cotA) and B (cotB) genes in Ax2 and tctp\textsuperscript{−}. (G) Relative transcript levels of acaA, carA, csA, and cadA early developmentally regulated genes in Ax2 and tctp\textsuperscript{OE} and (H) Ax2 and tctp\textsuperscript{−}. (The values represent mean ± standard deviation; n = 3; ***p < 0.001, **p < 0.01, *p < 0.05; Student’s t-test; Scale bar for stalk = 5 µm; Scale bar for spore= 2.5 µm; V- Vegetative, Str- Streaming, M- Mound, MS- Migratory Slug, EC- Early Culminant, and FB- Fruiting body).
while the Ax2 cells occupied the prespore/spore region. During slug formation, tcp− cells majorly occupied the anterior prestalk region, rear-guard and ALCs (Figure 7 and Supplementary Figure S5). At early culminant stage, tcp− cells occupied the anterior prestalk region, ALCs, and basal disk. At culminant, it majorly contributed to anterior tip region, upper and lower stalk portion and basal disk (Figure 7 and Supplementary Figure S5). This trend was observed as the percentage of tcp− cells increased further as it also occupied entire stalk and basal disk region. Next, the contribution of tcp− cells in spore formation was assessed and it exhibited decreased contribution in the prespore/spore region as compared to wild type cells. The contribution of tcp− cells in chimeras of varying proportion, i.e., 10, 25, 50, 75, and 90% of tcp− cells with Ax2 cells resulted in chimeric spore formation in which tcp− cells contributed only to 1.95 ± 0.35, 11 ± 1, 15 ± 1, 15.5 ± 0.5, and 49 ± 9% (Figure 8). Our results suggested that in chimeras with wild type cells, tcp− cells differentiated into prestalk/stalk cells preferentially, further suggesting that TCTP is a crucial component controlling the equilibrium between prestalk and prespore cell-types.
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**FIGURE 7** | tctp− cells exclusively occupied prestalk/stalk region in chimeras. RFP-tagged Ax2 cells and GFP tagged tctp− cells were mixed in varying proportions (10, 25, and 50% GFP marked tctp− cells and 90, 75, and 50% RFP marked Ax2 cells) were allowed to co-develop on non-nutrient agar until the formation of migratory slug, early and late culminant. Both, DIC and fluorescence images during development were captured using Nikon SMZ1500 fluorescence microscope. (Scale bar- 100 μm; n = 4).

**FIGURE 8** | Percentage (%) spore count of Ax2 and tctp− cells in chimeric fruiting bodies. GFP tagged tctp− cells were mixed in various proportion as given above with RFP-tagged wild type cells and cell suspensions were allowed to co-develop. About 100–200 spores per fruiting body and 10–15 fruiting bodies were counted and scored.

**DISCUSSION**

Translationally controlled tumor protein is ubiquitously expressed, highly conserved protein present in various type of cells and tissues implicating its essential role in normal cells. This conservation is also observed in the protozoans, such as *Plasmodium*, *Toxoplasma*, and *Dictyostelium* (Bhisutthibhan et al., 1998; Zheng et al., 2018). It is involved in regulation of cell cycle, development, cytoskeleton, autophagy, and apoptosis. It plays a vital role in the growth and development of an organism by regulating the main physiological events. However, the biologic function of this protein in social amoeba such as *Dictyostelium* has never been investigated. Sequence analysis suggested that the key residues are conserved throughout the evolution. Previous reports demonstrated that Glutamate12 (Glu12) present in the loop region of *Drosophila* TCTP is an important residue for Rheb (Ras homologue enrichment in brain) function that activates TOR (Target of rapamycin) but it is contradictory that TCTP has role in mTORC1 signaling pathway (Chen et al., 2007; Hsu et al., 2007; Rehmann et al., 2008). Conservation of Glu12 in *DdTCTP* thus open the possibility of Rheb function in lower eukaryotes. Phylogenetic studies suggested that *DdTCTP* is highly conserved protein in eukaryotes. Residues important for microtubules and Ca²⁺ binding are also conserved in different species studied (Bommer and Thiele, 2004; Graidist et al., 2007). In contrary, Ser46 and Ser64 residues which phosphorylated by the Plk are not conserved in lower eukaryotes and found only in mammalian TCTP (Yarm, 2002). Higher expression of *DdTCTP* mRNA was observed in vegetative or initial stages, suggested that *DdTCTP* may have a role in cell proliferation and cell division. As the complete knockout of tctp gene was lethal it limited the understanding of its functions. Here, we have studied
the functions of TCTP during growth and development of *D. discoideum* with the help of making overexpressor and knockout strains. Overexpression and disruption of *tcp* in *Dictyostelium* displayed reduced cell proliferation. Previous studies also suggested that overexpression of TCTP resulted in inhibition of cell proliferation in cell lines (Gachet et al., 1999). It has tubulin binding properties and found in association with tubulin or microtubules in cell cycle phases, its overexpression causes rearrangement of microtubules and stability. Disruption of *tcp* exhibited increased cell-size, which could be due to multi-nucleation. The size of organs or organism depends on the number and size of the cells that constitute them. Overexpression and disruption of *tcp* resulted in delayed aggregation stream formation and decelerated *acaA* and *carA* expression implying that an optimum level of TCTP is essential for normal early gene expression and early development. There are several genes whose timely expression and accurate level is important for regulation of early and late development such as overexpression or disruption of *srsA* exhibited delayed aggregation and reduced expression of *acaA* and *carA* along with aberrant fruiting body formation suggesting its role in the regulation of development (Sasaki et al., 2008). The genes *acaA* (adenyl cyclase A) and *car* (cAMP receptor) require to mediate the cAMP signaling during early development such as aggregation. During aggregation, *Dictyostelium* cells secrete and sense cAMP. ACA (Adenylate cyclase A) is a membrane protein which convert ATP to cAMP and cAMP get transported outside of the cells. Then cAMP bind with surrounding cells through cAR (cAMP receptor). Lacking of *acaA* and *carA* genes result abolish aggregation and delay in early development (Sun et al., 1990; Pitt et al., 1993). *tcp*OE cells form less but large-sized aggregates while *tcp*− cells form less but small-sized aggregates suggesting its involvement in regulation of aggregate size. Previously, various genes of *Dictyostelium* were reported in regulating aggregate size such as *ctnA, srsA,* and *ampk* (Okuwa et al., 2001; Sasaki et al., 2008; Maurya et al., 2017). *tcp*OE cells form long slugs and fruiting bodies with long stalks and small sori that has elliptical spores whereas *tcp*− cells developed into fruiting bodies with small stalks and large sori having round spores as compared to wild type suggesting its role in terminal differentiation. The round spores formed by *tcp*− fruiting bodies could be due to decreased expression of spore coat protein, *cotB* that has been previously known to maintain integrity of spore. Overexpression of *tcp* resulted in long fruiting bodies while deletion of it resulted in small fruiting bodies suggesting the function of TCTP in cell type differentiation and patterning for maintenance and establishment of proper proportioning and patterning. Interestingly, spatial distribution of prespore/prestalk cell type specific markers were aberrant and mis-localized in the multicellular structures formed by *tcp*− cells as compared to Ax2. The aberrant localization of *ecmAO*/*lacZ* in *tcp*− alluding its role in cell type patterning. Our results showed that TCTP is involved in cell type patterning and regulates the ratio of prespore/prespore cells in multicellular structures formed. A number of genes have been shown to play a role in cell type proportioning and spatial patterning in *Dictyostelium*. In Mekka null cells, there was a two-fold increase in pstO compartment and concomitant decrease in prespore region (Chung et al., 1998). Disruption of homeobox containing gene, *warai* led to doubling of pstO compartment (Han and Firtel, 1998). Overexpression of zZIP led to reduce prespore and enhanced prestalk expression (Balint-Kurti et al., 1997; Ginsburg and Kimmel, 1997). Furthermore, cell lineage study with the help of chimeric mixtures of Ax2 and *tcp*− in different proportions showed occupancy of both Ax2 and *tcp*− cells in different regions of multicellular structures developed. Chimeras are mixture of cells from two or more genetically different backgrounds. Our results clearly show the tendency of *tcp* null cells to form prespore cells in presence of wild type. It occupied the prespore regions like rear-guard cells, the anterior-like cells, lower cup, basal disk, and stalk regions. When the percentage of the null cells was increased in the chimeras, *tcp*? cells could contribute to spore cells though the spore forming tendency by the null cells was very low. The spore count from various chimeric mixtures substantiated our results that *tcp*− cells has less proclivity in spore formation in presence of wild type cells. The deletion of *tcp* allowed the cells to occupy the dying cell population suggesting that it may be required to regulate the spore differentiation. Thus, TCTP may contribute toward both spore and stalk cell differentiation but the question arises, how TCTP regulate spore/stalk differentiation in this organism? Whether TCTP is directly involved in regulating differentiation and patterning or it regulates expression of other genes involved in this process so, further studies are needed to be explored to clarify its exact role in cell type differentiation and patterning.

**CONCLUSION**

Our study suggests that TCTP regulates the cell size and cell proliferation in *Dictyostelium*. It is required for timely development and correct aggregation formation. Further, it is involved in maintaining prespore/prestalk or spore/stalk ratio and cell type differentiation and patterning.

**DATA AVAILABILITY STATEMENT**

All datasets presented in this study are included in the article/Supplementary Material.

**AUTHOR CONTRIBUTIONS**

RK, RM, and SS conceived and designed the study, analyzed the data, and wrote the manuscript. RK and RM performed the experiments. All authors read and approved the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcell.2020.00742/full#supplementary-material

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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