Cyto-architecture constrains the spread of photoactivated tubulin in the syncytial Drosophila embryo

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ABSTRACT Drosophila embryogenesis begins with nuclear division in a common cytoplasm forming a syncytial cell. Morphogen gradient molecules spread across nucleo-cytoplasmic domains to pattern the body axis of the syncytial embryo. The diffusion of molecules across the syncytial nucleo-cytoplasmic domains is potentially constrained by association with the components of cellular architecture. However, the extent of restriction has not been examined. Here we use photoactivation (PA) to generate a source of cytoplasmic or cytoskeletal molecules in order to monitor the kinetics of their spread in the syncytial Drosophila embryo. Photoactivated PA-GFP and PA-GFP-Tubulin generated within a fixed anterior area diffused along the antero-posterior axis. These molecules were enriched in the cortical cytoplasm above the yolk-filled center, suggesting that the cortical cytoplasm is phase separated from the yolk-filled center. The length scales of diffusion were extracted using exponential fits under steady state assumptions. PA-GFP spread a greater distance as compared to PA-GFP-Tubulin. Both molecules were more restricted when generated in the center of the embryo. The length scale of spread for PA-GFP-Tubulin increased in mutant embryos containing short plasma membrane furrows and a disrupted tubulin cytoskeleton. PA-GFP spread was unaffected by cyto-architecture perturbation. Taken together, these data show that PA-GFP-Tubulin spread is restricted by its incorporation in the microtubule network and intact plasma membrane furrows. This photoactivation based analysis of protein spread allows for interpretation of the dependence of gradient formation on syncytial cyto-architecture.

KEY WORDS: syncytium, photoactivation, Drosophila, embryogenesis, morphogen gradient, Bicoid

Introduction

Insect embryos initiate their development in a large syncytial cell where multiple nuclei undergo nuclear divisions in a common cytoplasm without forming complete cells. The cytoplasm is thought to mix uniformly in the syncytial cells. However, syncytial Drosophila embryos have distinct domains of gene expression patterns in nuclei despite being in this common cytoplasm (Shvartsman et al., 2008). Several tissues in different organisms, for example, plant endosperm cells, animal muscle cells and fungal hyphae, also contain syncytial cells. Syncytial nuclei in fungi maintain distinct cell cycle stages (Anderson et al., 2013; Dundon et al., 2016). The spatially separated daughter nuclei in these fungi continue to proceed through the cell cycle synchronously by maintaining a similar concentration of cell cycle components (Lee et al., 2013). Syncytial nuclei in muscle cells have a differential expression of mRNAs as compared to their neighbors (Pavlath et al., 1989). These studies indicate that several components of the cytoplasm have local function and are likely to be generated and sequestered in the vicinity of the syncytial nuclei. It is of interest to understand the cellular mechanisms that regulate compartmentalized distribution of molecules despite being in a common cytoplasm.

The syncytial embryos of Drosophila provide a tractable system to decipher the extent to which different cellular components are shared across nucleo-cytoplasmic domains. Drosophila embryogen-
esis begins with 9 nuclear division cycles deep within the embryo during the preblastoderm stage. Nuclei along with centrosomes migrate to the cortex in nuclear cycle 10 and the nuclear division cycles 11-14 occur beneath the cortex in the syncytial blastoderm embryo (Foe and Alberts, 1983; Foe, Odell and Edgar, 1993; Karr, 1986; Sullivan and Theurkauf, 1995; Warn, 1986). Each interphase nucleus of the syncytial blastoderm embryo is surrounded by apical centrosomes and a microtubule array in an inverted basket configuration. Astral microtubules reach out from the centrosomes towards the cortex and overlap with the astral microtubules originating from neighbouring nuclei (Cao et al., 2010). F-actin is present in caps above the nuclei and centrosomes. Lipid droplets and yolk are enriched at the bottom of the basket (Kuhn et al., 2015; Mavrakis et al., 2009a; Schmidt and Grosshans, 2018; Welte, 2015). Each interphase cycle begins with 9 nuclear division cycles deep within the embryo (Foe and Alberts, 1983; Foe, Odell and Edgar, 1993; Karr, 1986; Sullivan and Theurkauf, 1995; Warn, 1986). Each interphase cycles 11-14 occur beneath the cortex in the syncytial blastoderm embryo (Chowdhary et al., 2009a; Schmidt and Grosshans, 2018; Welte, 2015). Each interphase nucleo-cytoplasmic domain in the blastoderm embryo is associated with organelles such as the endoplasmic reticulum, Golgi complex and mitochondria (Chowdhary et al., 2017; Frescas et al., 2006; Mavrakis et al., 2009b). The microtubule and the actin cytoskeleton remodel during prophase and metaphase of the syncytial division cycle. The centrosomes move laterally during prophase and give rise to spindles during metaphase. Actin is enriched along the cortex and at the extending plasma membrane furrows (Foe, Odell and Edgar, 1993). The short furrows present in interphase between adjacent nuclei extend deeper between spindles in metaphase. Molecules in the plasma membrane, ER, Golgi complex and mitochondria have limited exchange between adjacent nucleo-cytoplasmic domains in the syncytial *Drosophila* embryo (Chowdhary et al., 2017; Frescas et al., 2006; Mavrakis et al., 2009b).

Analysis of exchange of molecules in the cytoplasm or cytoskeleton across the syncytial nucleo-cytoplasmic domains remains to be documented in a systematic manner, though several studies have probed various cytoplasmic properties. Fluorescent dextran of various sizes when injected in the cytoplasm of the syncytial blastoderm embryo has been used to estimate the rate of cytoplasmic diffusion in the embryo (Gregor et al., 2005). Micro-rheology based measurements of cytoplasmic viscosity have found that cytoplasmic viscosity is three times higher than that of water in the region between nuclei and yolk of the syncytial *Drosophila* embryo. In addition, microtubules, but not actin contribute to the observed viscosity (Wessel et al., 2015).

Morphogen gradient formation in the syncytial *Drosophila* can be used as a paradigm to estimate properties of the embryo cytoplasm. Bicoid forms a gradient in the antero-posterior axis, patterning the head of the embryo (Gregor et al., 2007). The Bicoid

**Fig. 1.** Cytoplasmic green fluorescent protein (GFP) and mCherry-Tubulin are enriched cortically in the syncytial division cycles. (A) Schematic showing sagittal imaging of the embryo. (B-E) Characterization of cortical distribution of GFP and mCherry-Tubulin in the syncytial division cycles. Images are shown from different cycles (NC11, 12, 13, 14) of embryos ubiquitously expressing GFP (B) or maternally expressing mCherry-Tubulin (similar trends were observed for n=3 movies) (D). Kymographs show cortical enrichment of fluorescent signal for GFP (C) and mCherry-Tubulin (E) over time. Scale bar, 5 µm, 600 s. (F,G) Quantification of cortical enrichment of fluorescent signal in GFP and mCherry-Tubulin for nuclear cycles 11th to 14th interphase. Graph shows relative intensity profile for GFP(F) and mCherry-Tubulin (G) obtained from a line drawn from the cortical region towards the center of the embryo. The dashed line shows a point at which the intensity drops to 30% of the maximum intensity. Note that the signal remains above the region containing the dark yolk filled vesicles. The images are shown in a 16 color intensity rainbow where blue represents the lowest intensity and red represents the highest intensity. Scale bar, 10 µm.
gradient is present primarily in the cortical region of the embryo (Cai et al., 2017). The dorso-ventral gradient formed by Dorsal is compartmentalized to each nucleo-cytoplasmic domain (DeLotto et al., 2007) and modelling studies show that plasma membrane furrows could restrict Dorsal gradient spread (Daniels et al., 2012). The Dorsal gradient formation on the ventral side depends on specific binding partners on the ventral side (Carrell et al., 2017). These studies together imply that the syncytial blastoderm cortex has gradients whose properties depend upon sequestration due to interaction with other cytoplasmic components or the syncytial cyto-architecture.

In this study, we attempt to elucidate the extent of spread of molecules across nucleo-cytoplasmic domains of the syncytial Drosophila embryo using a comparison between cytoplasmic PA-GFP and PA-GFP-Tubulin. Fluorescently labelled tubulin incorporates well in the microtubule network and is also present in the cytoplasm. We use photoactivation to generate a fixed population of PA-GFP or PA-GFP-Tubulin and find that both spread in the cortical region as compared to the yolk filled central region of the syncytial blastoderm embryo. The spread of PA-GFP-Tubulin is more restricted as compared to PA-GFP in the antero-posterior axis. PA-GFP and PA-GFP-Tubulin molecules have a decreased spread when generated in nucleo-cytoplasmic domains in the middle of the embryo as compared to the anterior. The PA-GFP-Tubulin spreads to a greater distance in mutants showing a loss of plasma membrane furrows and disruption of the microtubule network. The PA-GFP spread is not affected in these mutants. Our study provides a framework for assessing the regulation of gradient formation by its interaction with the syncytial cytoarchitecture components and has implications on the spread of morphogen gradients.

Results

**Cytoplasmic GFP and mCherry-Tubulin are enriched cortically in the syncytial division cycles in the Drosophila embryo**

The syncytial Drosophila blastoderm embryo has a characteristic arrangement of microtubules around each nucleus. Microtubules emanate from the apical centrioles and spread vertically covering the nuclei in an inverted basket like arrangement (Karr, 1986; Sullivan and Theurkauf, 1995). In order to test the extent of spread of molecules in the cytoplasm we imaged embryos expressing GFP ubiquitously under the control of the ubiquitin promoter. GFP is expected to be present primarily in the cytoplasm and is not known to interact with any cytoplasmic components (Verkman, 1999). We compared the expression of cytoplasmic GFP to fluorescently labelled tubulin as it would partition into the cytoplasm and also incorporate into the microtubule cytoskeleton. For this we imaged live embryos expressing fluorescently tagged alpha-Tubulin (UASP-mCherry-Tubulin) (Rusan and Peifer, 2007) with mat-Gal4-vp16 (mat-Gal4) in the sagittal view (Fig. 1A). Fluorescence intensities were displayed with respect to time as kymographs. Relative intensities in each syncytial cycle were computed to estimate the depth of spread in the embryo. We found that cytoplasmic GFP was enriched cortically and accumulated inside the cortical nuclei (Fig. 1B). Accumulation of GFP occurs passively inside the nucleus as a result of its small size which allows it to pass through the nuclear pore complex (Ruiwen Wang, 2007). The fluorescence intensity of cytoplasmic GFP progressively increased near the cortex as syncytial division cycles progressed but remained above the yolk filled region (Fig. 1C, Movie S1). We noticed GFP fluorescence dropped to approximately 30% between 32 to 36 μm in syncytial cycle 14 (Fig. 1F).

mCherry-Tubulin was enriched on apical centrioles, in microtubules spreading vertically from the cortex and in the cytoplasm in the syncytial division cycles (Fig. 1D, Movie S2). Progressive accumulation of fluorescence signal was seen near the cortex as the syncytial cycles progressed (Fig. 1E). mCherry-Tubulin fluorescence dropped to 30% between 25 to 27 μm beneath the cortex in syncytial cycle 14 (Fig. 1G). Cytoplasmic GFP spread to a greater depth as compared to mCherry-Tubulin from the cortex in the embryo most likely because of not being incorporated in the cytoskeleton. In summary cytoplasmic GFP and mCherry-Tubulin were concentrated near the cortex and further enriched during the progression of the nuclear cycles. In addition, they were present in a separate cortical layer of cytoplasm on top of and distinct from the inner yolk-filled region of the embryo.

**Photoactivation generates a source of PA-GFP and PA-GFP-Tubulin at the anterior that spreads cortically along the antero-posterior axis**

Labelled tubulin had a cytoplasmic and a microtubule bound fraction, in contrast to GFP, which had a cytoplasmic fraction in the Drosophila syncytial blastoderm embryo. This gave us an opportunity to assess the diffusion of these two proteins in the cytoplasm across nucleo-cytoplasmic domains. Computational simulations have predicted that binding to microtubule network and movement on motors is sufficient for partitioning the cytoplasm, in the absence of membrane boundaries in the syncytial blastoderm embryo (Chen et al., 2012). We therefore asked whether tubulin which partitions partially into microtubules could be more restricted as compared to GFP in the syncytial blastoderm embryo.

Photoactivation of cytoplasmic and cytoskeletal proteins have been used to generate a local source of protein molecules for monitoring their directional spread in axons (Gauthier-Kemper et al., 2012; Gura Sadovsky et al., 2017). In order to test the differential spread of cytoplasmic and cytoskeletal proteins in the syncytial blastoderm embryo, we used continuous photoactivation to create a local source of fluorescent PA-GFP or PA-GFP-Tubulin at different locations of the embryo (Fig. 2A). Unlike morphogens such as Dorsal and Bicoid, GFP and tubulin are not differentially distributed in the syncytial embryo. PA-GFP and PA-GFP-alpha-Tubulin84B (PA-GFP-Tubulin) were expressed individually in embryos by crossing the transgenic flies to mat-Gal4. Expression of cytoplasmic GFP, mCherry-Tubulin, PA-GFP and PA-GFP-Tubulin did not hamper development of the syncytial cycles. A fixed area was continuously photoactivated to form fluorescent PA-GFP/PA-GFP-Tubulin, thus creating a local source of PA-GFP/PA-GFP-Tubulin at the anterior pole of the embryo (Fig. 2B,D, Movie S3,4). PA-GFP and PA-GFP-Tubulin spread from the source of generation forming a gradient of visible fluorescent molecules and could be used to quantitatively assess the extent of spread in the syncytial embryo. Photoactivation in each embryo allowed generation of 2 estimates of spread from the anterior to posterior pole via dorsal or ventral surface thus allowing two length scale estimates. The movies of PA-GFP photoactivation also showed the presence of a strong autofluorescent signal at the base of the cortex comprising of yolk (Movie S3). The movies of PA-GFP-Tubulin showed an increase in fluorescence in the cytoplasm and PA-GFP-Tubulin was also incorporated in
microtubules in interphase and in metaphase spindles (Movie S4). Both PA-GFP and PA-GFP-Tubulin increased in concentration away from the source across the syncytial division cycles. A kymograph obtained at the source of photoactivation showed a distinct increase in amount of photoactivated molecules over time (Fig. 2 C,E). The kymograph also showed that the fluorescent signal was enriched near the cortex and did not enter the central yolk filled region of the embryo. A 30% drop in the signal was obtained between 40 to 45 µm for PA-GFP and 20 to 25 µm for PA-GFP-Tubulin beneath the cortex. An analysis of the directionality of spread showed that both molecules spread to a greater distance cortically along the antero-posterior axis (XY) as compared to the depth within the embryo (XZ) (Fig. 2 F,G). The cytoplasm of syncytial Drosophila blastoderm embryo has a biphasic distribution with cortical nucleo-cytoplasmic domains present above a barrier comprising of yolk and other unknown components (Foe and Alberts, 1983; Wessel et al., 2015). This organization possibly allows for greater spread along the cortex in the antero-posterior axis as compared to the center.

**Anteriorly photoactivated PA-GFP and PA-GFP-Tubulin shows an exponential spread that is steeper for PA-GFP-Tubulin.**

We next attempted to quantify the spread of photoactivated molecules in the antero-posterior axis when generated anteriorly. We found that the photoactivated probes spread further as the syncytial cycles progress (Fig. 3 A,B). The fluorescence intensity of PA-GFP and PA-GFP-Tubulin increased with time at different locations in the embryo (Fig. 3 C,D). The concentration of PA-GFP and PA-GFP-Tubulin when measured at 11 µm from the photoactivation source, increased with time and reached saturation. The time taken to reach steady state increased as we moved away from the photoactivation source. Temporal evolution of fluorescence at x=38 µm approached a steady state at later time points. In contrast, for x=165 µm, the concentration did not reach a steady value (Fig. 3 C,D). This is also apparent from the temporal evolution of the rate of change of the concentration at these different locations (Fig. 3 E,F). We used the

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**Fig. 2. Anteriorly photoactivated PA-GFP and PA-GFP-Tubulin produces a cortical gradient.**

(A) The photoactivation method to create an ectopic source of PA-GFP and PA-GFP-Tubulin. Photoactivation was carried out, continuously with an interval ~2 seconds, in a fixed area (373 µm²) in the anterior or the center of the syncytial embryo. A kymograph monitoring the increase in signal was drawn at the source (green bar). A cortical region was drawn to estimate the change in intensity in the antero-posterior axis (orange). The exponential function was fit to estimate the length scale of spread for the gradients. (B-E) Anteriorly photoactivated PA-GFP and PA-GFP-Tubulin forms a gradient. Images for NC11,12,13,14 of embryos from expressing PA-GFP (B) or PA-GFP-Tubulin (D) are shown after photoactivation at the anterior pole. Kymograph shows increase in cortical fluorescence over time in PA-GFP (C) and PA-GFP-Tubulin (E) expressing embryo. Scale bar, 50 µm, 60 s. (FG) PA-GFP and PA-GFP-Tubulin spreads preferentially at the cortex. Graph quantifying the spread of photoactivated protein fluorescence in the antero-posterior XY axis vs depth or XZ direction for PA-GFP (F) and PA-GFP-Tubulin (G) with a line drawn across either XY or XZ direction from the activated region. The raw data is in a lighter color and the averaged data is in a darker color, error bars represent standard error on means (n=6 from 3 embryos for PA-GFP-Tubulin and PA-GFP each, the PA experiment in 1 embryo gave 2 estimates of spread in the AP direction from the dorsal or ventral side). The images are shown in a 16 color intensity rainbow where blue represents the lowest intensity and red represents the highest intensity. Scale bar, 50 µm.
steady state concentration profile and extracted the characteristic length scales by fitting it to an exponential decay equation (Fig. 3 G,H). PA-GFP and PA-GFP-Tubulin formed gradients of distinct length scales after activation at the anterior pole (Fig. 3I). We found that the length scale for PA-GFP was significantly higher than PA-GFP-Tubulin (Fig. 3J). The estimated diffusion coefficient for PA-GFP was 44.25 \( \mu m^2/s \) and PA-GFP-Tubulin was 20.87 \( \mu m^2/s \) (refer to Materials and Methods). This is likely to be because PA-GFP-Tubulin, in addition to being present in the cytoplasm is also engaged in forming the microtubule cytoskeleton and this turnover makes it less available to diffuse as compared to PA-GFP alone.

**Photoactivated PA-GFP and PA-GFP-Tubulin when generated in the middle of the Drosophila embryo spreads less as compared to the anterior activation**

The syncytial Drosophila embryo has three domains containing distinct patterns of density of nuclei and packing (Blankenship and Wieschaus, 2001; Rupperecht et al., 2017). The domains show different speeds of furrow extension during cellularization. The anterior domain contains nuclei at a lower density as compared to the middle domain and the cells formed have a shorter plasma membrane furrows as compared to the middle domain in cellularization. This difference in architecture across the antero-posterior axis is regulated by the patterning molecules Bicoid, Nanos and Torso (Blankenship and Wieschaus, 2001). Further, the protein dynamics

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**Fig. 3.** Anteriorly photoactivated PA-GFP and PA-GFP-Tubulin shows an exponential spread with PA-GFP-Tubulin being more restricted as compared to PA-GFP (A,B). Quantification of the photoactivated signal across nuclear cycles. Graph shows intensity for PA-GFP (A) and PA-GFP-Tubulin (B) for one embryo with a line drawn across the cortical region in the syncytial nuclear cycles. Similar profiles were observed in multiple embryos (n=3 for each). (C,D) PA-GFP and PA-GFP-Tubulin increases in concentration over time. The graph depicts an increase in PA-GFP (C) or PA-GFP-Tubulin fluorescence intensity over time as measured at different locations (11, 38, 165 \( \mu m \) from the source of photoactivation at the anterior). (E,F) Graph shows the rate of change in concentration of photoactivated PA-GFP (E) and PA-GFP-Tubulin (F) to assess if the steady state has reached. Each plot is a derivative of the corresponding plot in (C,D). (G,H) Anteriorly photoactivated PA-GFP and PA-GFP-Tubulin shows an exponential spread. Raw experimental values (red) were fit to an exponential function (blue) for each probe. (I) Quantification of intensity profile of photoactivated probe measured at the end of the experiment for PA-GFP and PA-GFP-Tubulin. Graph shows raw data in a lighter color and averaged data in a darker color, error bars represent standard error on means (n=3 embryos each for PA-GFP and PA-GFP-Tubulin). (J) Scatter plot of length scales extracted after fitting an exponential decay function to the intensity profiles seen in (I) (n=6, 3 i.e 6 length scales extracted from 3 embryos in the AP direction on the dorsal or ventral side for PA-GFP, PA-GFP-Tubulin, Two tailed Mann-Whitney non-parametric test with p value=0.0007).
Fig. 4. Photoactivation of the cytoplasmic PA-GFP and PA-GFP-Tubulin in the middle of the Drosophila embryo. (A-D) Monitoring spread of center photoactivated PA-GFP and PA-GFP-Tubulin. Images from NC11, 12, 13, 14 expressing PA-GFP (A) or PA-GFP-Tubulin (C) and photoactivated at the center of the embryo are shown. Kymograph shows increase in cortical fluorescence with time for PA-GFP (B) and PA-GFP-Tubulin (D) embryo. Scale bar, 50 μm, 60 s. (E,F) Quantification of evolution of photoactivated signal in syncytial nuclear cycles. The graph depicts the fluorescence intensity for PA-GFP (E) and PA-GFP-Tubulin (F) from one embryo for a line drawn from the source along the antero-posterior axis. Similar profiles were observed in multiple embryos (n=6,3 embryos for each). (G,H) Quantification of photoactivated protein in XY vs XZ direction for PA-GFP (G) and PA-GFP-Tubulin (H). Graph shows intensity profile of a line drawn in the XY or XZ direction from the activated region. The raw data is shown in a lighter color and the averaged data is shown in a darker color, error bars represent standard error on means (n=6,3 for PA-GFP-Tubulin and PA-GFP each).
would encounter difference in geometric shape at the conical anterior region as compared with the flatter central region of the embryo (He et al., 2010). This difference in the density of nucleo-cytoplasmic domains and geometry prompted a comparison of the extent of photoactivated molecule spread, when it originates in the middle of the embryo versus when it originates in the anterior domain (Fig. 2A).

We tested if there was a difference in the spread of photoactivated molecules when generated in the middle of the embryo (Fig. 4) as compared to the anterior (Fig. 2,3). For this we continuously photoactivated PA-GFP and PA-GFP-Tubulin containing embryos in a fixed region in the cortical region in the middle of the embryo (Fig. 2A, 4 A,C). Photoactivation produced a cortical spread with a progressive increase in concentration spread across the syncytial division cycles (Fig. 4 B,D-F, Movie S5,6). Each experiment performed allowed generation of two estimates of spread from the center to anterior and center to posterior pole leading to two length scales. PA-GFP and PA-GFP-Tubulin spread to a greater extent in the antero-posterior axis (XY) as compared to the depth of the embryo (XZ), away from the region of photoactivation (Fig. 4 G,H).

Length scale values were extracted by fitting an exponential equation and it was found that the extent of spread for both the probes was lower than that observed when photoactivation was performed anteriorly (Fig. 4 I-K). We further analysed the difference in the extent of spread formation from the center towards the anterior versus center towards the posterior pole (Fig. 4L,M). We found that the length scales of the spread did not differ in either direction (Fig. 4N). These analyses show that the molecules form gradients uniformly across the syncytial nucleo-cytoplasmic domains towards the anterior pole and the posterior pole of the Drosophila embryo, negating the presence of any cytoplasmic flows or currents. In summary, photoactivated molecules generated in the middle spread to a smaller distance as compared to when they were generated at the anterior pole.

**Anteriorly photoactivated PA-GFP-Tubulin spread increases in embryos containing an overexpression of RhoGEF2 on loss of pseudocleavage furrows**

The gradients produced by PA-GFP and PA-GFP-Tubulin provided a framework to test the role of syncytial cytoarchitecture in regulating their spread. Each cortical nucleus in the syncytial blastoderm embryo of Drosophila contains a small ingression of the plasma membrane around it. Astral microtubules support ectopic furrows (Barmchi et al., 2005; Cao et al., 2008; Crest et al., 2012). The plasma membrane furrows ingress deeper in metaphase to form pseudocleavage furrows (Schmidt and Grosshans, 2018). To test the role of furrows in regulation of extent of spread across to which the fluorescence is confined. Scale bar, 50 μm, 60 s. (F,G) Quantification of intensity profile of photoactivated probe as measured at the end of the experiment for PA-GFP (F) and PA-GFP-Tubulin (G) in embryos overexpressing RhoGEF2. The raw data is shown in a lighter color and the averaged data is shown in a darker color, error bars represent standard error on means (n=3 embryos for PA-GFP-Tubulin and PA-GFP each). The graph for photoactivation of PA-GFP and PA-GFP-Tubulin in control embryos is the same as that shown in Fig. 3I and is repeated here for comparison. (H) Scatter plot of the length scales extracted after fitting an exponential decay function to the intensity profiles seen in (F,G). The values of length scales for PA-GFP and PA-GFP-Tubulin for anterior photoactivation in control embryos are repeated from Fig. 3I (n=6,3 for PA-GFP in RhoGEF2-OE, 6,3 for PA-GFP-Tubulin in RhoGEF2-OE. Kruskal-Wallis nonparametric test of variance p value=0.0064, followed by Dunn’s posttest at 95 percent confidence interval).
the syncytial nucleo-cytoplasmic domains, we performed photoactivation experiments in embryos defective in furrow formation. RhoGEF2 is a Rho-GTP exchange factor specifically needed for the formation of furrows in the syncytial embryo (Barmchi et al., 2005; Cao et al., 2008; Crest et al., 2012). Depletion of RhoGEF2 leads to shortened furrows but they are not lost (Sherlekar and Rikhy, 2016; Zhang et al., 2018). An overexpression of RhoGEF2 causes increase in active Myosin II leading to increased contractility and loss of furrow formation (Dey and Rikhy, 2020). We overexpressed RhoGEF2 by crossing flies containing mat-Gal4 and UASp-RhoGEF2. Embryos overexpressing RhoGEF2 showed missing or short furrows in 100% of the embryos in metaphase (Fig. 5A) and 87% (n=200) of the embryos were lethal at 24 hours. The metaphase spindles were comparable to controls in RhoGEF2 overexpressing embryos (Fig. 5A).

Next, we generated embryos expressing PA-GFP or PA-GFP-Tubulin along with RhoGEF2 overexpression. We performed continuous photoactivation at the anterior pole in a fixed area and followed the resultant spread of photoactivated molecules across time (Fig. 5 B,D, Movie S7,8). We found that similar to the control embryos (Fig. 2), the gradients evolved over time (Fig. S1 A,B). Further, in spite of major contractions in the embryo yolk (Movie S7,8), the activated fluorescent molecules remained near the cortex and did not mix with the underlying inner yolk region of the embryo (Fig. S1 C,D). This was also evident from the kymographs which showed undulations in the fluorescence of the cortical layer, yet maintaining a

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**Fig. 6.** PA-GFP-Tubulin spreads to a greater extent in *eb1* mutant embryos. (A) *eb1* RNAi expressing embryos show perturbed tubulin architecture: Surface and sagittal views of fixed control or *eb1* RNAi embryos, stained with Tubulin (Red), Scribbled (Green) and DNA (Grey), show perturbed spindles in metaphase (100%, n=25 embryos). Zoomed inset of spindles show loose organization in *eb1* RNAi expressing embryos. Scale bar, 10 µm, inset scale bar, 5 µm. (B-E) Images from syncytial cycles of an *eb1* RNAi expressing embryo co-expressing PA-GFP (B) or PA-GFP-Tubulin (D) with photoactivation at the anterior pole. Kymograph for PA-GFP (B) and PA-GFP-Tubulin (E) shows increase in cortical fluorescence across time while sometimes changing the extent to which the fluorescence is confined. Scale bar, 50 µm, 60 s. (F,G) Quantification of intensity profile of photoactivated probe as measured at the end of the experiment for PA-GFP (F) and PA-GFP-Tubulin (G) in *eb1* RNAi embryos. The raw data is shown in a lighter color and the averaged data is shown in a darker color, error bars represent standard error on means (n=3 embryos for PA-GFP-Tubulin and PA-GFP each). The graph for photoactivation of PA-GFP and PA-GFP-Tubulin in control embryos is the same as that shown in Fig. 3I and is repeated here for comparison. (H) Scatter plot of length scales extracted after fitting an exponential decay function to the intensity profiles seen in (F,G). The values of length scales for PA-GFP and PA-GFP-Tubulin for anterior photoactivation in control embryos are repeated from Fig. 3J. (n=6,3 embryos for PA-GFP in *eb1* RNAi, 6,3 for PA-GFP-Tubulin in *eb1* RNAi. Kruskal-Wallis nonparametric test of variance p-value=0.0008, followed by Dunn’s post test at 95 percent confidence interval).
separation from the embryo’s inner yolk region (Fig. 5 C,E). The PA-GFP spread did not change in the embryos over-expressing RhoGEF2 (Fig. 5F). The PA-GFP-Tubulin spread however changed significantly (Fig. 5G) when compared to the respective spread in control embryos (Fig. 2). The length scales were extracted on fitting an exponential function to the concentration profile obtained. It was found that PA-GFP-Tubulin spread to a greater extent in embryos over expressing RhoGEF2 (Fig. 5H). It is interesting to note that the length scale of PA-GFP-Tubulin in mutant embryos increased from its control value and became similar to PA-GFP. RhoGEF2 overexpression led to loss of plasma membrane furrows and loss of restriction of PA-GFP-Tubulin spread in the syncytial Drosophila embryo.

Anteriorly photoactivated PA-GFP-Tubulin spread increases in EB1 mutant embryos showing aberrant tubulin organization

Microtubules emanate from the centrosome at the apical side and spread vertically downwards in the syncytial blastoderm embryo (Kellogg et al., 1988; Sullivan and Theurkauf, 1995). EB1 is present at the growing end of microtubules and its depletion is likely to disrupt the microtubule organization (Rogers et al., 2002). We depleted embryos of EB1 by combining eb1 RNAi to mat-Gal4. 60% (n=300) of the embryos expressing eb1 RNAi were lethal at 24 hours. The microtubule staining in embryos expressing eb1 RNAi was reduced in interphase (Fig. S2A). The spindles appeared loosely organized in metaphase of the syncytial cycle (Fig. 6A, Movie S9,10). Also Scribbled levels at the plasma membrane levels were lowered indicating a loss of furrows (Fig. 6A and S2A). Live imaging of eb1 RNAi expressing embryos showed an aberrant spacing of nuclei and an increase in spindle defects (Figure S1B, Movie S10). Thus the tubulin cytoskeletal organization showed increased defects in embryos expressing eb1 RNAi.

We combined the eb1 RNAi with flies expressing PA-GFP or PA-GFP-Tubulin and performed anterior photoactivation experiments in a fixed area (Fig. 6B,D, Movie S11,12). We found that similar to the control embryos (Fig. 2), the gradient evolved over time (Fig. 6 B,D,S2 C-F). Similar to RhoGEF2 overexpression embryos, in spite of major contractions in the embryo yolk, the activated fluorescent molecules remained near the cortex (Fig. 6 C,E) and did not mix with the underlying yolk region of the embryo (Fig. S2 G,H). Length scales were extracted by fitting these gradients (Fig. 6 F,G) to an exponential function. We saw that the PA-GFP spread did not change, while PA-GFP-Tubulin spread increased significantly in mutant embryos (Fig. 6H). The length scale analysis showed that PA-GFP-Tubulin in eb1 mutant embryos was similar to PA-GFP. In summary, eb1 mutant embryos had a disrupted microtubule architecture and showed a loss of restriction of PA-GFP-Tubulin spread in the syncytial Drosophila embryo.

Discussion

In this study, we have examined the distribution and spread of cytoplasmic components of the Drosophila syncytial blastoderm embryo. We have used photoactivation of cytoplasmic PA-GFP to analyze its distribution and diffusion across nucleo-cytoplasmic domains of the syncytial Drosophila embryo and further compared it to PA-GFP-Tubulin, which is present in the cytoplasm and is also incorporated in microtubules. We find that the cytoplasmic components have an increased concentration at the cortex near the nucleo-cytoplasmic domains. Photoactivated molecules show diffusion to a greater distance in the antero-posterior axis in the cortex as compared to the depth of the embryo. Also photoactivated molecules spread less when generated at the center of the embryo as compared to the anterior. The spread of photoactivated tubulin is constrained by interaction with the cyto-architecture components of the syncytial blastoderm embryo (Fig. 7).

Photoactivation as a method to study regional differences in kinetics of gradient formation in the syncytial Drosophila embryo

The use of photoactivatable GFP molecules allows for the creation of localized ectopic gradients and enables us to follow their evolution in real time across the syncytial nuclear cycles. Photoactivation has been used previously to analyze the spread of morphogens in similar contexts. Photoactivation of Dorsal-PA-GFP allowed an analysis of the extent of its spread in the dorsal versus ventral side of the syncytial blastoderm embryo. Sequestration of Dorsal by signaling components and nuclear capture on the ventral side gave a more constrained spread for Dorsal-PA-GFP as compared to the dorsal side of the embryo (Carrell et al., 2017). In our study we used two photoactivatable proteins which
are incorporated in all nucleo-cytoplasmic domains. This allows us to quantify the differences in their spread due to inherent differences in association with cyto-architecture of the embryo. We found that PA-GFP and PA-GFP-Tubulin had a smaller length scale when activated at the center as compared to the anterior of the syncytial blastoderm embryo. The restricted spread at the center of the embryo could be a result of a difference in relative crowding of nucleo-cytoplasmic domains in these two regions (Blankenship and Wieschaus, 2001; Rupprecht et al., 2017). An increase in the density of nucleocytoplasmic domains in the center could lead to greater sequestration of cytoplasmic components in general, leading to a smaller length scale. Alternatively this could also come about due to differences in protein degradation or sequestration machinery or even geometry of the embryo between these two regions. Whether the difference in density of nucleo-cytoplasmic domains also leads to change in viscosity in the two regions remains to be examined.

**Organization of cytoplasm in cells**

The cytoplasm of majority of living cells can be described as an inhomogeneous, multi-phasic medium. Images of different components when drawn to scale (Goodsell, 2013) clearly convey the fact that the cytoplasm is quite contrary to the earlier picture of a freely flowing medium. The cytoplasm can be likened to a complex medium comprising of physical constraints and constraints due to binding and crowding. Fluorescent dextran of various sizes when injected into cells partitions based on their size (Luby-Phelps, 2000). This further corroborates the fact that the space available for various cytoplasmic components is constrained depending on their size. The metabolic state can also change cytoplasmic properties in a bacterial cell into either a glass-like or fluid-like state (Parry et al., 2014). Cytoplasmic distribution can change depending on the ability and strength of a cytoplasmic molecule to bind to other components. A modelling based study showed that binding to negative end directed dynein motors on the mitotic spindle was sufficient to partition the cytoplasm into two halves even without the presence of any membrane bound compartments (Chen et al., 2012).

Our finding that PA-GFP-Tubulin spreads less as compared to PA-GFP suggests that cytoplasmic components having multiple interactors are more confined in their diffusion. For the syncytium, this property is beneficial, as components produced from a syncytial nucleus tend to remain near their parent nucleus, with no clear boundaries being present in the shared cytoplasm. This observation suggests that different components in a cell could be restricted by distinct mechanisms, some binding to microtubules, some to actin or some being sequestered in the nuclei or other organelles ultimately resulting in restricting their action in space and time. The restricted spread of PA-GFP-Tubulin in our study was lost on abrogation of the metaphase furrows and microtubule cytoskeleton in embryos over-expressing RhGEF2 and eb1 RNAi, which highlights how binding and sequestration were responsible for PA-GFP-Tubulin restriction. Loss of plasma membrane furrows could also lead to disorganization of astral microtubules (Cao et al., 2010; Crest et al., 2012) in the periphery thereby increasing the effective diffusion of PA-GFP-Tubulin.

Further, the observation that cytoplasmic components are cortically enriched corroborates previously reported data about Bicoid movement in the cortex and its dependence on the actin and the microtubule cytoskeleton of syncytial blastoderm embryos (Cai et al., 2017). SEM images from cross sectioned Drosophila embryos show the presence of biphasic compartments (Figard et al., 2013; Turner and Mahowald, 1976). Filamentous actin and non-muscle myosin are concentrated in the 3-4 μm and 1-2 μm region of the “yolk-free” cytoplasm just beneath the plasma membrane of the preblastoderm embryo, respectively (Foe, Odell and Edgar, 1993). The cortical yolk-free cytoplasm increases in its depth as the syncytial cycles progress (Foe, Odell and Edgar, 1993). Our study is a characterization of protein mobility in these phases, and we show that the cortical cytoplasm and yolk beneath it seem to form two separate phases, and do not mix in spite of major contractions in mutant embryos. The size of the cortical cytoplasmic region as determined by the concentration of cytoplasmic GFP in our study, is approximately 40 μm. This observation raises further questions about how these two phases are separate and the mechanisms that contribute to maintaining their integrity.

**Implications on morphogen spread**

The observation of the presence of two separate phases of cortical cytoplasm and embryo yolk provides an interesting perspective to our current understanding of morphogen gradients in the early embryo, namely, Bicoid, Dorsal and Torso. The Bicoid gradient has been extensively studied using the framework of the synthesis, diffusion and degradation (SSD) (Durrieu et al., 2018; Gregor et al., 2007; Grimm et al., 2010) and related models. Our finding implicates a restriction of the effective volume in which Bicoid gradient develops and matures. It also raises the possibility that various cytoarchitectural components could impinge on its formation. For example, perturbations in furrows or cytoskeletal structures can change the effective concentration of morphogens in the cortical cytoplasm, leading to changes in the morphogen profiles, specifically for Bicoid.

There have also been various studies, implicating the size and shape of the mRNA source in Bicoid gradient formation (Fahmy et al., 2014; Little et al., 2011; Spirov et al., 2009). Photoactivation allows creation of different sized sources which can produce PA-GFP/PA-GFP-Tubulin or morphogen gradients at different rates and provides an opportunity to study the effect of the source on the gradient shape and dynamics.

The observation of distinct length scales of PA-GFP-Tubulin versus PA-GFP points to another facet of morphogen gradient formation, namely decrease in the diffusivity of morphogens based on their interactions. FGF gradient is known to interact with Heparan sulfate proteoglycans which changes the effective diffusivity of the morphogen. The removal of these proteoglycans leads to an increase in the morphogen spread (Balasubramanian and Zhang, 2016). We can interpret the difference between the PA-GFP and PA-GFP-Tubulin profiles as being a consequence of increased binding of tubulin to the microtubule architecture. This leads to increase in its residence time by sequestration and thus a lower effective diffusion and consequently, a smaller length scale. It would be interesting to analyse the effect of removal of binding interactions for well-studied morphogen like Bicoid. It is notable that Dorsal gradient is known to be modulated depending on the presence or absence of a dimerizing GFP (Carrell et al., 2017). In conclusion, these and our studies recommend a systematic analysis of the impact of local architectural properties in the formation and maintenance of morphogen gradients.
Materials and Methods

Drosophila stocks and crosses

Drosophila stocks were maintained in standard corn meal agar at 25°C. All crosses were setup at 25°C, except eb1 RNAi (29°C). mat-gal4-VP16; mat-gal4-VP16 (Girish Rnaparkhi, IISER, Pune, India) was used to drive m-Cherry-alpha-TubulinA1B (m-Cherry-Tubulin) (#25774), PA-GFP (gift from Prof. Gerald M. Rubin, Janelia Research Campus, VA, USA), PA-GFP-alpha-TubulinB84B (PA-GFP-Tubulin) (#32076), UASp-RhoGFP2 (#9386) and eb1 RNAi (#36599). GFP expressed under ubiquitin promoter (ubi-GFP, #1681) was imaged directly.

Microscopy

1.5 hour old embryos were collected on sucrose agar plates, washed, dechorionated using 100% bleach, mounted on coverglass chambers (LabTek, Germany) in PBS (Mavrikas et al., 2008) and imaged on Plan-Neofluor 40x/1.30 oil objective on Zeiss LSM780(fixed imaging) or Plan-Apochomat 25x/0.8 Oil Immersion on LSM710 systems(photostimulation experiments). PA-GFP and PA-GFP-Tubulin were photoactivated using the 405 nm diode laser using the bleach module on the LSM software. PA-GFP and PA-GFP-Tubulin thus produced was imaged using the 488 nm laser. ROI size was kept constant at 373μm². Photostimulation iterations were kept constant at 8 iterations per frame with activation being performed continuously after every frame. The photoactivation was carried out for 0.36 s (10 iterations). 512 pixel X 512 pixel images were acquired after that with a scan speed of 1.97 seconds per frame. Mid sagittal sections were imaged. 8 bit images were acquired with mean line averaging of 2. The gain and laser power were adjusted to be cover the dynamic range of each fluorescent tag and care was taken to not reach 255 on the 8 bit scale. Pinhole was kept open at 180μm.

Immunostaining

F1 flies were selected from Gal4 and mutant crosses were transferred to embryo collection cages (Genesee Scientific, CA, USA) with 2.5% sucrose agar supplemented with yeast paste. Embryos were washed, dechorionated using 100% bleach for 1 min, washed and fixed in heptane: 4% paraformaldehyde (1:1) in PBS (1.8 mM KH2PO4, 137 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4) for 20 mins at room temperature. Embryos were devitellicized by vigorously shaking in heptane:methanol (1:1) for anti-Tubulin and Scribbled immunostaining. 2% Bovine Serum Albumin (BSA) in PBS with 0.3% Triton X-100 (PBST) was used for blocking. Following primary antibodies were diluted in the block solution: anti-Tubulin (Anti-mouse, Sigma-Aldrich, Bangalore, India,1:1000), anti-Scrib (Anti-Rabbit, Kind gift by Prof. Kenneth Prehoda, University of Oregon,OR,USA,1:1000). Fluorescently coupled secondary antibodies (Alexa Fluor 488, 568, 647 coupled anti-rabbit and anti-mouse, Molecular Probes, Bangalore, India) were used at 1:1000 dilution in PBST. Embryos were imaged using LD LCI Plan-Apochomat 25x/0.8 Imm Korr DIC M27 objective on the Zeiss LSM710/780.

Image analysis

Sagittal sections were imaged. The script rescales the embryo length from 0 to 1 in the anterior/posterior direction, subtracts the minimum intensity value, rescales it with the maximum and smoothens the intensity values using sliding window averaging.

Sampling and Statistics

3 or more embryos indicated in the corresponding figure legends were imaged and quantified for each experiment. Graphpad Prism 5.0 was used for Statistical analysis and plotting.

Theory

Estimation of length scales from concentration profiles

We analyse the time evolution of the concentrations of the photo-activated molecules within the framework of the standard one-dimensional Synthesis-Diffusion-Degradation (SDD) model (Driever and Nüsslein-Volhard, 1988; Gregor et al., 2007) in a domain of length L. Before the initiation of photoactivation, the system consists of unlabelled molecules which are assumed to be present at a uniform concentration across the embryo. When photoactivation is initiated, these unlabelled molecules are converted to fluorescent molecules at the position of photoactivation. Thus photoactivation generates a localised source for fluorescent molecules. Unlabelled molecules enter the photoactivation region to get activated in subsequent photoactivation events. Photoactivated molecules spread outwards from this source point with a diffusion coefficient D, and are degraded at a rate k. This degradation rate can, in general, be a combination of the conversion from fluorescent to unlabelled molecules due to loss of fluorescence, or due to degradation of the protein itself. The time evolution of the concentration of the fluorescent molecules is then described by,

\[ \frac{\partial C(x,t)}{\partial t} = D \frac{\partial^2 C(x,t)}{\partial x^2} - \kappa C(x,t) \]  

where, \(c(x,t)\) represents the concentration of the photoactivated species as position x at time t. The mean lifetime of the molecule \(\tau\) is the inverse of the degradation rate, \(\tau = 1/k\). This equation is to be solved subject to the appropriate boundary conditions, accounting for the presence of a localised source of fluorescent molecules at the anterior pole of the embryo, and reflecting boundary conditions at the posterior pole,

\[-D \frac{\partial C}{\partial x} |_{x=0} = Q \quad \text{and} \quad D \frac{\partial C}{\partial x} |_{x=L} = 0 \]  

and the appropriate initial condition reflecting the absence of any photo-activated molecules for \(t \leq 0\), \(c(x, t \leq 0) = 0\).

At long enough times, the concentration profile evolves to a steady state (Fig. 3C,D). The steady state solution of the SDD model for a semi-infinite domain is given by,

\[ c_{ss}(x) = \frac{Q}{2D} \exp \left(-x/\lambda\right) \]  

where the characteristic length-scale lambda is defined as, \(\lambda = \sqrt{D/\kappa}\). The semi-infinite assumption holds if the characteristic length-scale is much smaller than the size of the domain, \(\lambda \ll L\).

If the length scale is comparable to the system size, then the steady state solution depends on the length of the domain (size of embryo) and is given by,

\[ c_{ss}(x) = \frac{Q}{2D} \left[ \exp \left( e^{\mu/\lambda} \right) - \exp \left( e^{\nu/\lambda} \right) \right] \]  

where the characteristic length scales \(\mu, \nu\) are much smaller than the system size, the length scales obtained from Eq.3 and Eq.4 will be comparable.

Hence, the length scales reported in this manuscript were obtained using the single exponential solution for the infinite boundary case. Since the magnitude of the source strength is unknown, we use a fitting parameter, A, to model the source strength. Additionally the embryo has a background fluorescence intensity, B, which arises due to baseline fluorescence activ-
ity of the embryo, and is not described within the SDD framework. We therefore fit the experimentally obtained fluorescence intensity profiles using the following expression for the steady state, \( C_s(x) = A \exp \left( -\frac{x}{\lambda} \right) + B \) (Houchmandzadeh et al., 2002).

In order to ensure that the concentration profiles have reached a steady state, we plot the concentration versus time plots and the rate of change of concentration for both PA-GFP and PA-GFP-Tubulin. The time taken to reach the steady state depends on the position along the AP axis, and is smaller for locations closer to the anterior pole. We first show the results for PA-GFP-Tubulin (Fig. 3F). As can be seen from the figures, the tubulin concentration reaches a steady state quickly, justifying the assumption of the steady state for fitting the concentration profile. The time taken to reach the steady state can be determined by the time at which the derivative \( dc/dt \) approaches zero (see Fig. 3E and 3F).

A similar analysis can be performed for PA-GFP (Fig. 3E). The situation in this case is more complex, with the locations closer to the anterior pole having reached a steady state, while locations further away still evolving at the final time point of the experiments. The larger time taken to reach the steady state for PA-GFP can be understood from the fact that the length-scale for PA-GFP is much larger than PA-GFP-Tubulin and hence it takes a correspondingly larger time for the concentration profile as a whole to reach steady state. In this case, since the locations closer to the anterior pole have reached a steady state, we can fit the concentration profile in a localised region closer to the anterior pole.

The concentration profiles at the last time point are fitted by this steady state formula to obtain the characteristic length-scale \( \lambda \). The fits are shown for PA-GFP (Fig. 3G) and PA-GF-Tubulin (Fig. 3H). This yields

\[ \lambda_{GFP} = 145 \pm 24.6 \text{ \mu m} \quad \text{and} \quad \lambda_{Crest} = 86 \pm 11.28 \text{ \mu m}. \]

The PA-GFP spreads to a much larger distance from the anterior pole than Tubulin-PA-GFP.

**Estimation of diffusion constant from concentration profiles**

For the SDD model, the time taken to reach the steady state can be estimated theoretically (Berezkhkovskii et al., 2010), and is given by

\[ \tau_{dd}(x) = \frac{2}{\lambda} \left( 1 + \frac{2}{\lambda} \right) x \]

where \( \tau_{dd}(x) \) is the time taken to reach steady state at location \( x \), and the mean lifetime of the molecule is denoted by \( \tau \), as before. The above formula also supports the notion that locations further away from the source at the anterior pole, take longer time to reach steady state. For distances much smaller than the characteristic length-scale, \( x < \lambda \), the above equation reduces to \( \tau_{dd}(x) = \sqrt{2x} \), and hence the mean lifetime can be read off from the concentration plots (Fig. 3C,D) and its derivative plots (Fig. 3 E,F) by noting the time taken to reach steady state for both PA-GFP and PA-GFP-Tubulin for \( x = 1 \times (x < \lambda) \). This yields

\[ \tau_{GFP} \approx 600 \text{ s} \quad \text{and} \quad \tau_{Crest} \approx 200 \text{ s}. \]

Combining the estimates of the length scale \( \lambda \) and the lifetime \( \tau \), we can then independently obtain an estimate of the diffusion constant, \( D = \lambda^2/\tau \).

This yields

\[ D_{GFP} \approx 44.25 \text{ \mu m}^2/\text{s} \quad \text{and} \quad D_{Crest} \approx 20.87 \text{ \mu m}^2/\text{s}. \]

The estimation of the time taken to reach steady state makes certain assumptions. Firstly, the fluctuations in the concentration can be significantly high in certain embryos, which results in a large variation of the time estimate. Secondly, a characteristic feature of the time evolution of concentration profiles is that there is a sharp initial increase followed by a slow increase in the concentration. This suggests that there may be other biological processes beyond those described by the SDD model that affect the evolution of the concentration to the steady state. While estimating the diffusion coefficient, we neglect the slower variation and have chosen the onset of this slow increase as the steady state time.
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