



# Direct Visualization of Actin Filaments and Actin-Binding Proteins in Neuronal Cells

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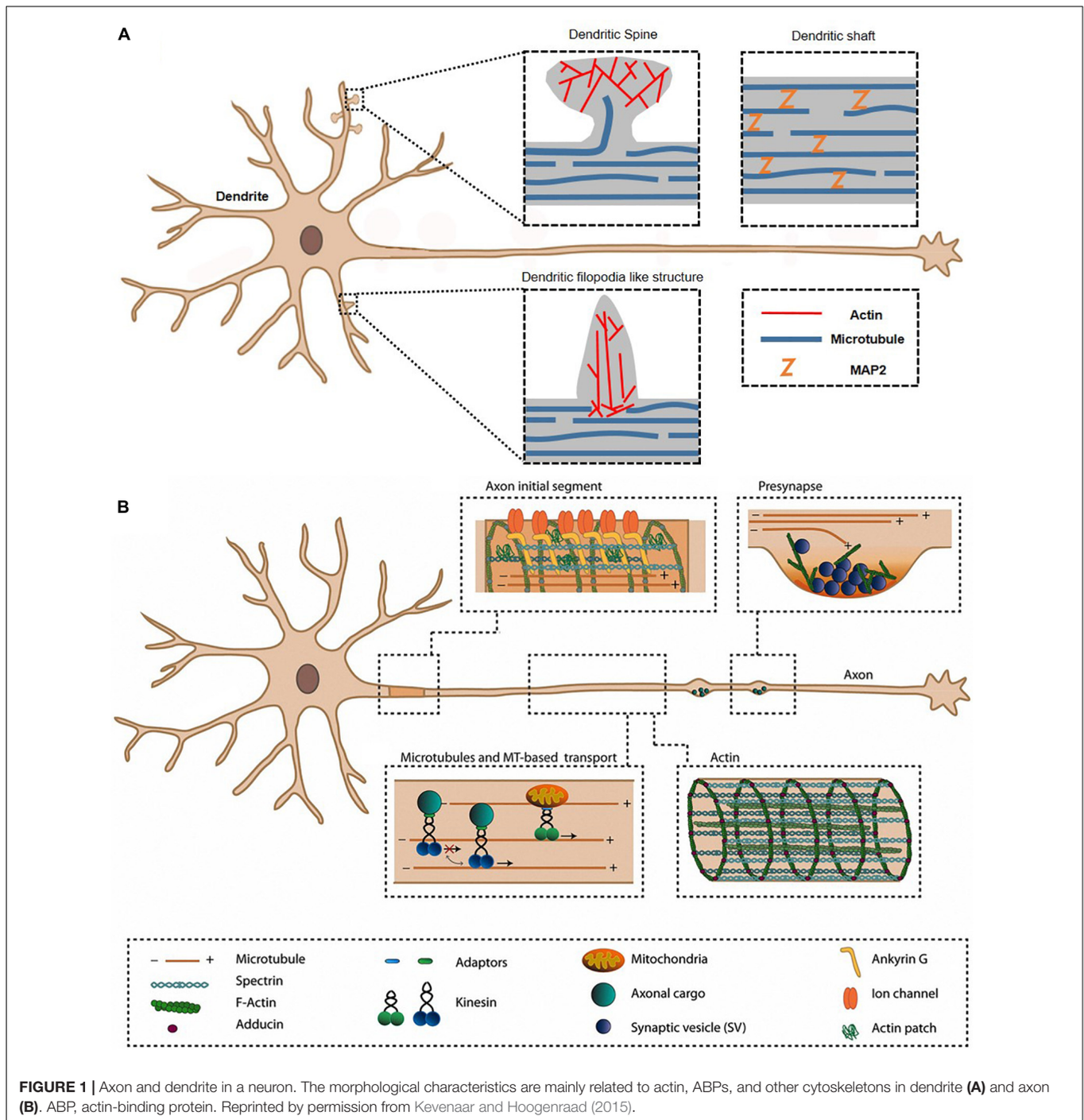
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Actin networks and actin-binding proteins (ABPs) are most abundant in the cytoskeleton of neurons. The function of ABPs in neurons is nucleation of actin polymerization, polymerization or depolymerization regulation, bundling of actin through crosslinking or stabilization, cargo movement along actin filaments, and anchoring of actin to other cellular components. In axons, ABP–actin interaction forms a dynamic, deep actin network, which regulates axon extension, guidance, axon branches, and synaptic structures. In dendrites, actin and ABPs are related to filopodia attenuation, spine formation, and synapse plasticity. ABP phosphorylation or mutation changes ABP–actin binding, which regulates axon or dendritic plasticity. In addition, hyperactive ABPs might also be expressed as aggregates of abnormal proteins in neurodegeneration. Those changes cause many neurological disorders. Here, we will review direct visualization of ABP and actin using various electron microscopy (EM) techniques, super resolution microscopy (SRM), and correlative light and electron microscopy (CLEM) with discussion of important ABPs in neuron.

**Keywords:** electron microscopy, super-resolution microscopy, correlative light and electron microscopy, actin binding protein, actin, neuronal cell

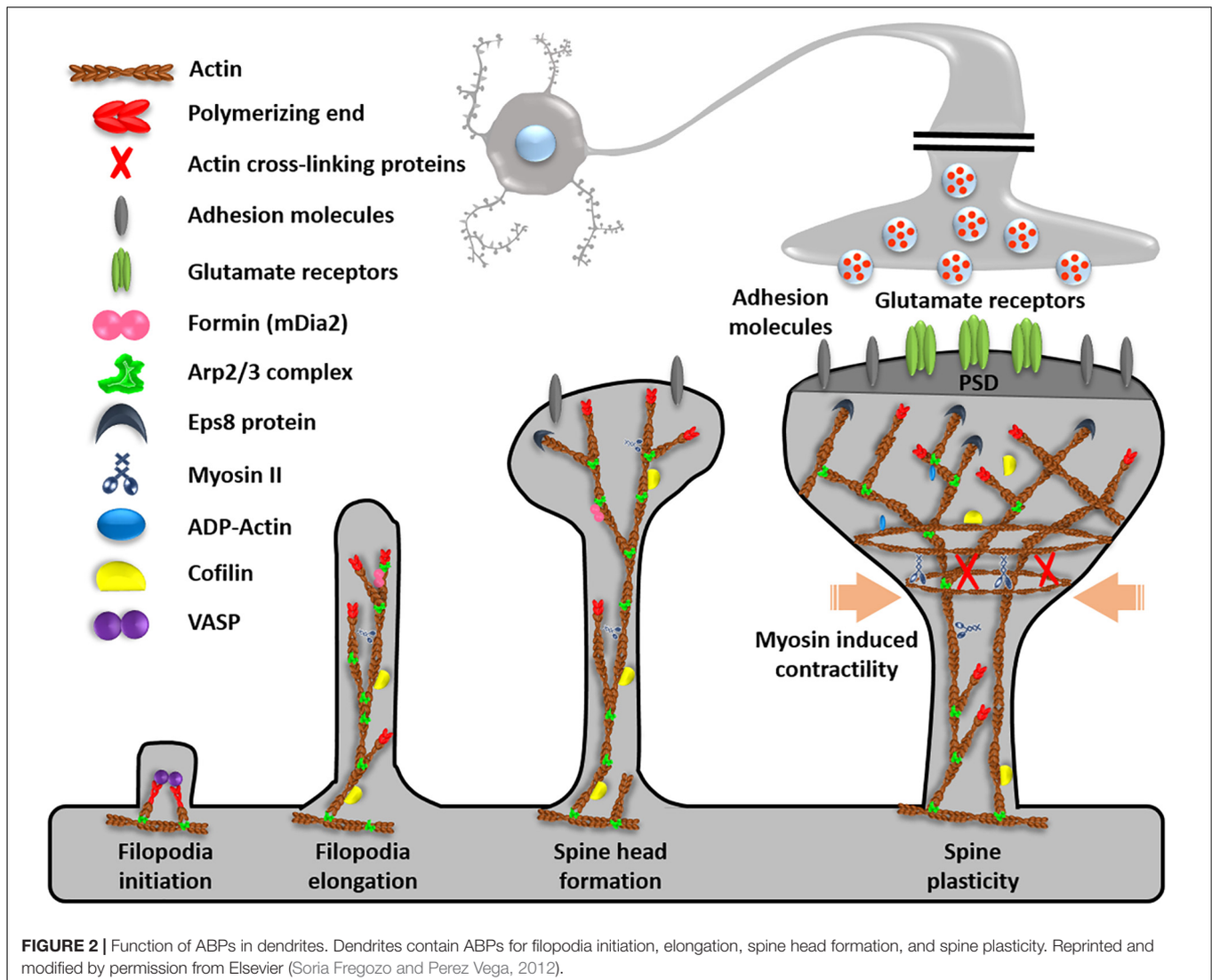
## INTRODUCTION

Neurons are specialized cells with long processes or connections in the nervous system. Each neuron has two types of cytoplasmic protrusions from the neuronal cell body, an axon and a dendrite (**Figure 1**). The length of an axon is the distance between one neuron and a target neuron, which varies from tens of micrometers to 1 m. The function of axon for transmission of information to different neurons is essential for brain function, therefore axons normally form a connection with the correct target and maintain their structure. Because each neuron has a single axon, axon dysfunction is both the result and the main cause of many neurological disorders, such as loss of cognitive ability, general paralysis, paraplegia, and loss of sensory function. Another protrusion from neuron is dendrites, and the main function is receiving signals from other neurons. In dendrite, spines receiving signals can be classified into mature spines, mushroom-shaped and immature spines (filopodia), or thin spines with a hairpin-like fine head <3 μm in length (**Figure 2**). Actin filaments are a part of the cytoskeleton and form the specific axon and dendrite morphology (Dogterom and Koenderink, 2019). In developing neurons, dynamic regulation of actin polymerization and organization mediates axon morphogenesis and path finding to synaptic targets. Changes in the axon shape such as branching, branch retraction, axonal arbor morphology depend on actin filament dynamics. Actin filaments do not function in a naked state, and actin binding proteins (ABP) s regulate all aspects of actin, that is, actin filament dynamics and



organization are regulated by various ABPs. In mature axons, stable F-actin and ABPs play a scaffolding role and maintain axon integrity through actin ring and help transport organelles. In addition, F-actin and ABPs mediate vesicular trafficking and regulate neurotransmitter release in mature axon terminals. Actin remodeling through ABPs in synaptic boutons also plays a crucial role in postsynaptic terminal plasticity. In postsynaptic terminal, the dendritic spine length, branching, spine density, shape of spine, and spine distribution (actin-rich protrusions

from the dendritic shaft), and motility respond morphologically to various physiological stimuli depending on actin and ABPs. Dysfunction of synapse plasticity causes neurological and psychiatric disorder, which includes Alzheimer's disease (AD), Parkinson's disease (PD), Huntington's disease (HD), schizophrenia, autism, and depression (Ma et al., 2012; Borovac et al., 2018). Therefore, researchers are trying to understand the mechanism of interaction of actin and ABPs in axon and dendrite in order to find ways to prevent neuronal disorders.



As ABPs, the Arp2/3 complex binds with F-actin and nucleates a new filament from an existing actin filament. Arp2/3 complex deficiency induces excessive growth, exclusive axon branching, and loss of dendritic spine maturation and enlargement (Strasser et al., 2004; Pinyol et al., 2007). Formins are unbranched actin nucleators (Kuhn and Geyer, 2014), and involved in proper axon development (Matusek et al., 2008) through deep actin network formation. A high formin amount causes long F-actin bundles to push out filopodia and promotes long actin filament polymerization (Hotulainen et al., 2009). Filament-severing proteins, such as actin-depolymerizing factor (ADF) and cofilin-1 accelerated actin turnover (Sarmiere and Bamberg, 2004), because cofilin is a major actin depolymerization factor for regulating actin length (Cichon et al., 2012). Cofilin-1 activity is necessary for neurite growth and growth cone turning, receptor trafficking, dendritic spinogenesis, and dynamic synaptic plasticity (Noguchi et al., 2016). As a linker to form actin networks, there are reports about ABPs tropomodulin (Gray et al., 2017), filamin

(Cho et al., 2015), neurofascin (Winkelman et al., 2014), spectrin (Unsain et al., 2018), adducin (Leite et al., 2016), and ERM proteins (ezrin, radixin, moesin) (Marsick et al., 2012). Among them, we will show details of main proteins such as Arp2/3, spectrin, neurofascin, and adducin through direct visualization techniques.

Electron Microscopy (EM) is one of the primary methods of choice for visualizing actin and ABP, as it is capable of achieving a nanometer resolution. However, the application of conventional EM for live or wet biological samples is limited due to the need for high-vacuum conditions (Kim et al., 2015). Moreover, the molecular specificity is limited in EM imaging as electron-dense heavy metals used for increasing the electron contrast of biological samples often stain lipids or proteins non-specifically. Although several molecular structures can be identified by their characteristic shapes from advanced EM images, such identifiable molecular structures are relatively few, and the characteristic structures of macromolecules are not known in most cases (Kim

et al., 2015). Immuno-EM using antibodies conjugated with gold nanoparticles can be used to localize specific targets in EM, but it suffers from a low labeling efficiency and limited multi-target imaging (Griffiths and Hoppeler, 1986; Robinson et al., 2000). These limitations can be overcome using advanced EM techniques such as cryo-techniques, light microscopy (LM) with super resolution, and correlative light and electron microscopy (CLEM). Although LM exhibits diffraction-limited spatial resolution ( $\sim 300$  nm), it enables the simultaneous visualization of multiple targets with high molecular specificity. Live and wet biological samples can also be imaged using light microscopy. The recent development of super-resolution fluorescence microscopy (SRM) has enabled substantially increased spatial resolution of light microscopy ( $\sim 10$  nm) by overcoming the diffraction limit (Hell and Wichmann, 1994; Betzig et al., 2006; Rust et al., 2006; Huang et al., 2009).

In this review, we focused on the application of (1) EM such as single particle analysis, electron tomography, unroofing and etching with cryo-techniques, (2) SRM such as STORM, PALM, and STED, (3) CLEM to the study of the molecular architecture of actin and ABP, since these are the most widely used techniques for such molecular ultrastructural studies.

## VISUALIZATION TECHNIQUES IN THE NANOSCALE TO STUDY ACTIN AND ACTIN-BINDING PROTEINS

### Electron Microscopy

Electron microscopy is good tool for study actin and ABPs because of its nano scale resolution. Traditionally, actin and actin ABPs were observed by conventional EM such as negative stained EM and etching techniques. However, they have some limitations as we mentioned in introduction. Recently, advanced techniques including cryo-TEM, cryo-electron tomography, and CLEM have been used for the study. We will discuss three methods including single particle analysis, electron tomography with freeze-etching and unroof, and immuno gold staining among various EM techniques with their findings. As a single particle analysis with negative stained EM or cryo-EM showed proteins' interaction in angstrom resolution level. It can reveal of ABPs' binding mechanism to actin and their competition. Electron tomography with unroof and etching showed intervals the protein's interaction to actin with higher resolution. Immuno-gold labeling showed specific ABP's detailed locations in cells.

### Single Particle Analysis Using Negative-Stained EM and Cryo-EM

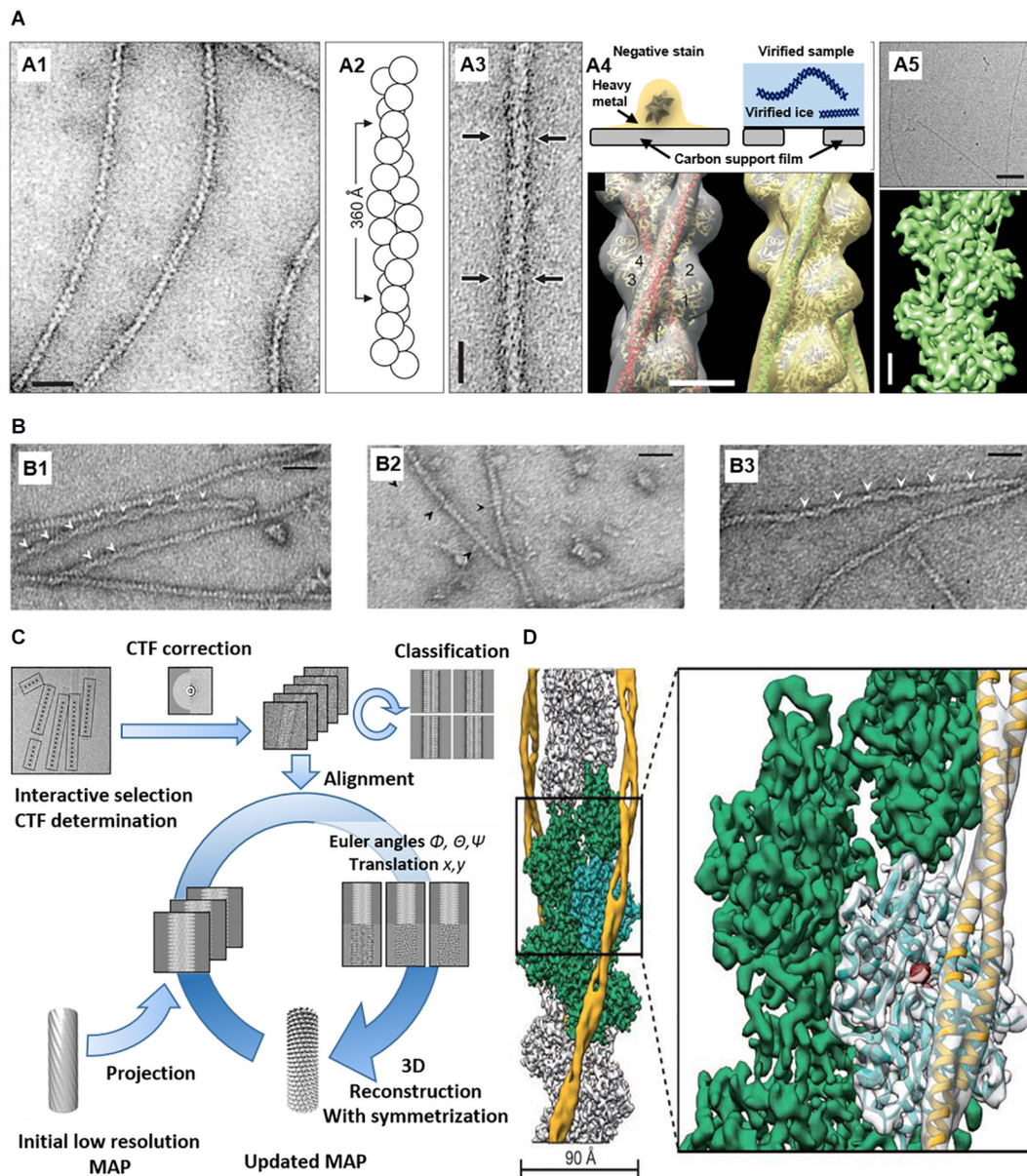
Simple and reliable EM technique combined by image processing and 3D reconstruction (Figure 3) can be used to visualize ABPs. EM and 3D reconstruction of purified ABPs and actin show ABP-binding sites for actin filaments *in vitro* (Narita and Maeda, 2007; Egelman, 2010; Behrmann

et al., 2012) are used. Negative-stained EM of F-actin itself and F-actin with ABPs has been conventionally used for structural studies of actin with specific ABPs. Negative staining is a simple sample preparation method in which protein samples are embedded in a thin layer of dried heavy metal salt to increase specimen contrast. After EM imaging in low-dose mode, the straightened single filaments from the images are used for helical reconstruction or iterative helical real space reconstruction (IHRSR). Averaging the reconstructions shows the 3D structure density, and the known X-ray crystal structure of each protein can be fitted to the 3D structure. The technique shows the binding sites of cortactin (Pant et al., 2006), ADF/cofilin (Galkin et al., 2011), vinculin (Thompson et al., 2014), drebrin (Grintsevich et al., 2010) to actin.

Cryo-EM is applied to reveal further details because of its high resolution without staining and dehydration. Compared to negative-stained EM with  $\sim 20$  Å resolution, cryo-EM can reach 1–2 Å resolution (Bartesaghi et al., 2015; Merk et al., 2020). For decades, structural biologists have conventionally used X-ray crystallography involving crystallizing proteins. X-ray crystallography gives high-quality images of structures, but it is difficult to make crystals. Cryo-EM does not use crystallized proteins, and uses vitrified pure water including a protein solution in a hydrophilic carbon film, which is plunged into a cryogen, such as liquid ethane. Frozen, hydrated protein samples are embedded in a thin layer of vitreous ice, which is imaged by cryo-EM using direct electron detection with complementary metal oxide semiconductor (CMOS)-based sensors or a charge-coupled device (CCD). The first cryo-EM of actin was performed by Trinick et al. (1986) at 40 Å resolution, and recent data of phalloidin-bound F-actin was reconstructed at 3.3 Å resolution (Das et al., 2020). Using this technique, Galkin et al. (2008) investigated actin–fimbrin/plastin interaction, and Chou et al. (Chou and Pollard, 2019) investigated ADP–actin binding. Cofilin (Tanaka et al., 2018), spectrin (Avery et al., 2017), and Arp2/3-binding sites (Zimmet et al., 2020) was also investigated using cryo-EM.

3D reconstruction techniques use Interactive Helical Real Space Reconstruction (IHRSR). The 3D reconstruction software using IHRSR has a heterogeneous structure compared to traditional Fourier–Bessel approaches. However, there is still limitation, because when the protein has large heterogeneity, a large proportion of densities become smeared out in the 3D structure after averaging. Therefore, researchers often use fixation to get tight ABP–actin binding or better alignment accuracy by checking the assigned azimuthal rotation angle of segments (Yang et al., 2016).

Structural studies of actin and ABPs using negative-stained EM and cryo-EM provide new insights into the mechanism underlying ABP–actin binding to regulate actin's function. In addition, the structures show the competition between ABPs to regulate actin filaments' interaction. Cofilin–actin and cofilin–actin bundles were also investigated using negative-stained EM



**FIGURE 3 |** EM imaging and 3D reconstruction of actin with ABPs *in vitro*. **(A)** **(A1)** Negatively stained F-actin. **(A2)** F-actin model based on EM. **(A3)** Negatively stained actin and ABP (arrows). **(A4)** (Top) Comparison between negative-stained EM and cryo-EM. (Bottom) 3D reconstruction of actin and ABPs based on negative staining data. **(A5)** Cryo-EM of F-actin (top) and 3D reconstruction at  $\sim 5.5$  Å resolution (bottom) Reprinted by permission from Korean Society of Microscopy (Craig, 2017). Scale bars = 200 Å (**A1,A3**), 50 Å (**A4**), 500 Å (**A5**), top), 20 Å (**A5**), bottom). **(B)** **(B1)** Actin filaments bound with cofilin-rod. Arrowheads show the intersection of the cofilin-rod clusters. **(B2)** Cofilin-rod molecules are rarely bound to actin filaments and identified by a rod-shaped structure (black arrowheads). **(B3)** Actin filaments with bound cofilin molecules (without rod fusion). Scale bars = 25 nm. Reprinted by permission from Ngo et al. (2015). **(C)** 3D reconstruction workflow. Reprinted by permission from Elsevier (Fromm and Sachse, 2016). **(D)** 3D reconstruction of actin with ABP (tropomyosin) from cryo-EM. PDB fitted to 3D EM volume. Scale bars = 90 Å. Reprinted by permission from Springer Nature (Von Der Ecken et al., 2015).

(Minamide et al., 2010; Ngo et al., 2015). Hyperactive cofilin after dephosphorylation shows conformational changes forming rod-shaped cofilin-actin bundles, which can affect binding of other ABPs (**Figure 3B**; Ngo et al., 2015). The changes induce vesicle transport blocking, an increase in secreted A $\beta$ , phosphorylated tau accumulation, neurofibrillary tangle (NFT) formation, and other pathological hallmarks of AD (Bamburg et al., 2010).

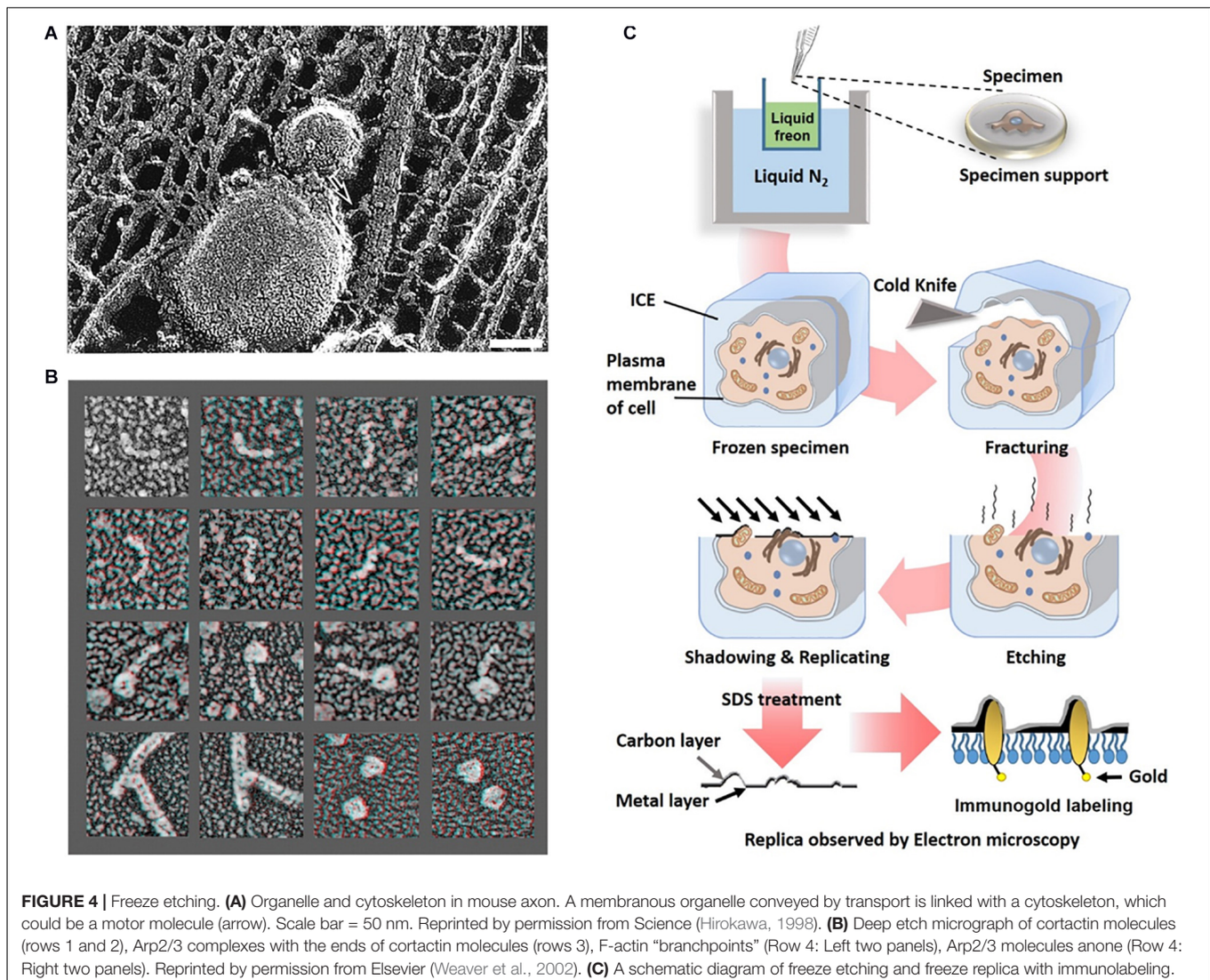
### Electron Tomography Using Quick Freeze and Deep Etching

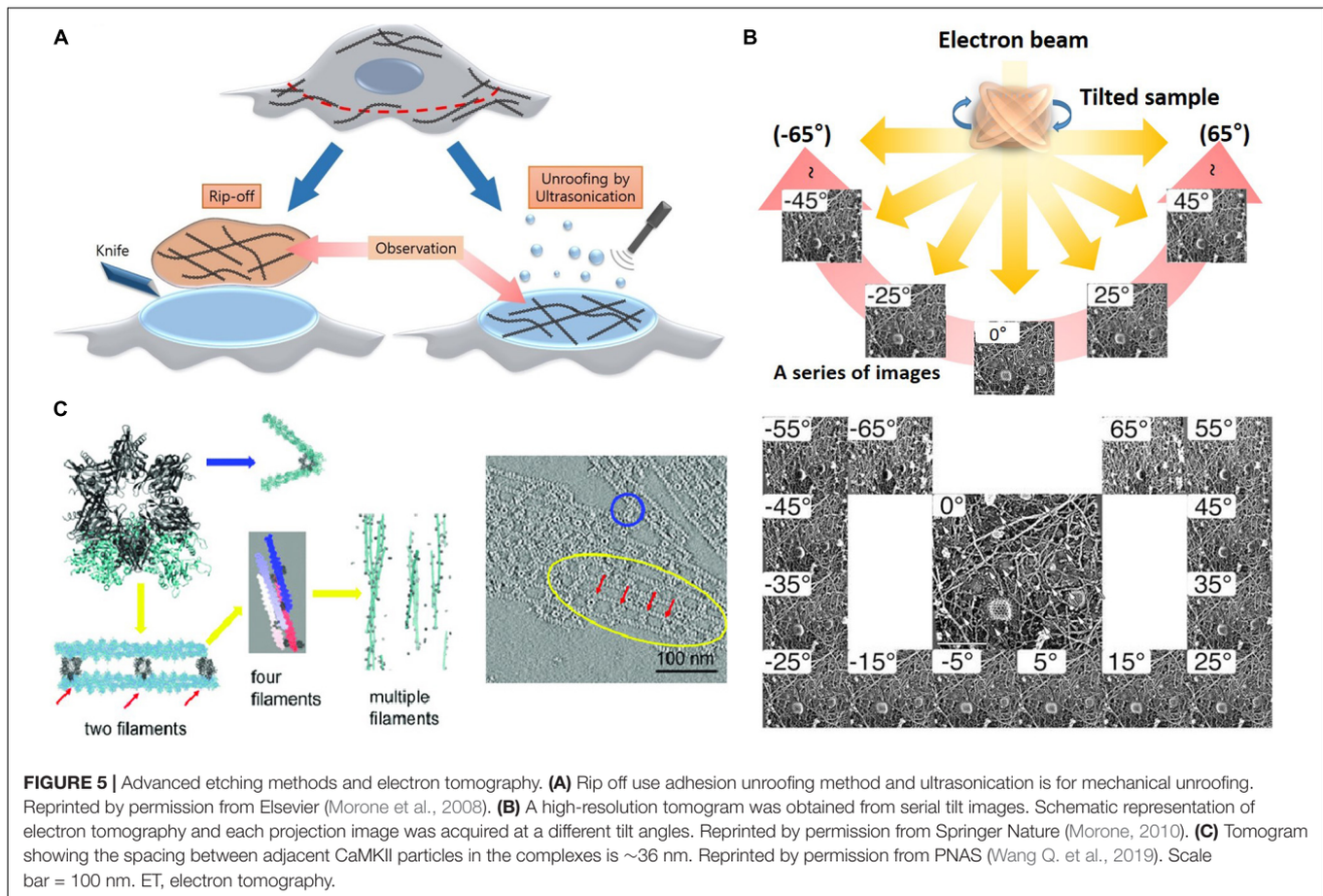
Determining the organization of axonal actin filaments in neurons was challenging. Hirokawa used quick-freeze and deep-etch (QFDE) methods to prepare frog spinal nerve axons for EM (Hirokawa, 1982). While conventional transmission electron microscopy (cTEM) requires chemical fixation, dehydration, and

plastic embedding and results in artifactual changes, QFDE EM provides a pseudo 3D appearance detailed structures. Freeze fracture involves breaking a frozen sample to reveal its intercellular structures. Freeze etching is the sublimation of surface ice under vacuum to reveal originally hidden details of the fractured face (**Figure 4C**). With platinum replication using platinum, freeze-etched samples can be viewed with high dimensional stability and preservation quality capturing subtle structural changes, compared to cTEM (Hirokawa et al., 1988; Svitkina, 2009). QFDE EM can show a central region of microtubules, NFs, interconnected membranous organelles, and a dense actin filament and ABP network (**Figure 4A**). Therefore, QFDE EM has been used for decades to investigate cytoskeletal proteins (Weaver et al., 2002). Weaver et al. used deep-etch EM of cortactin, N-WASP (Neural Wiskott Aldrich Syndrome Protein), and the Arp2/3 complex and showed that cortactin and N-WASP can bind simultaneously to the Arp2/3 complex for actin assembly activation (**Figure 4B**). In addition, immunogold or fluorescent marker-labeled platinum-replica

electron microscopy (PREM) used to study the specific ABPs. Labeling technique is a powerful tool to investigate the function of specific proteins (Hirokawa et al., 1988; Robenek and Severs, 2008; Vassilopoulos et al., 2019) in actin binding, and they will be each explained immuno EM and CLEM section.

Advanced methods (**Figure 5**) for visualizing cytoplasmic surface of the neuronal cell membrane (Morone et al., 2008) include mechanical shearing and adhesion unroofing. The mechanical shearing of a neuronal cell to expose the cytoplasmic surface of the neuronal cell membrane, and it enables observation of membrane cytoskeletons by cryo-EM and freeze-etch EM. In sonication unroofing, fine bubbles are generated, which adhere electrostatically to apical neuronal cell surfaces and remove the apical neuronal cell membrane (**Figure 5A**). In adhesion unroofing, positively charged grids that tightly adsorb neuronal cells are peeled from the cells. These techniques allowed visualization of the cytoplasmic surface and successfully shows many details of the cytoskeleton (Morone, 2010). In addition, it clearly shows the distribution of actin,



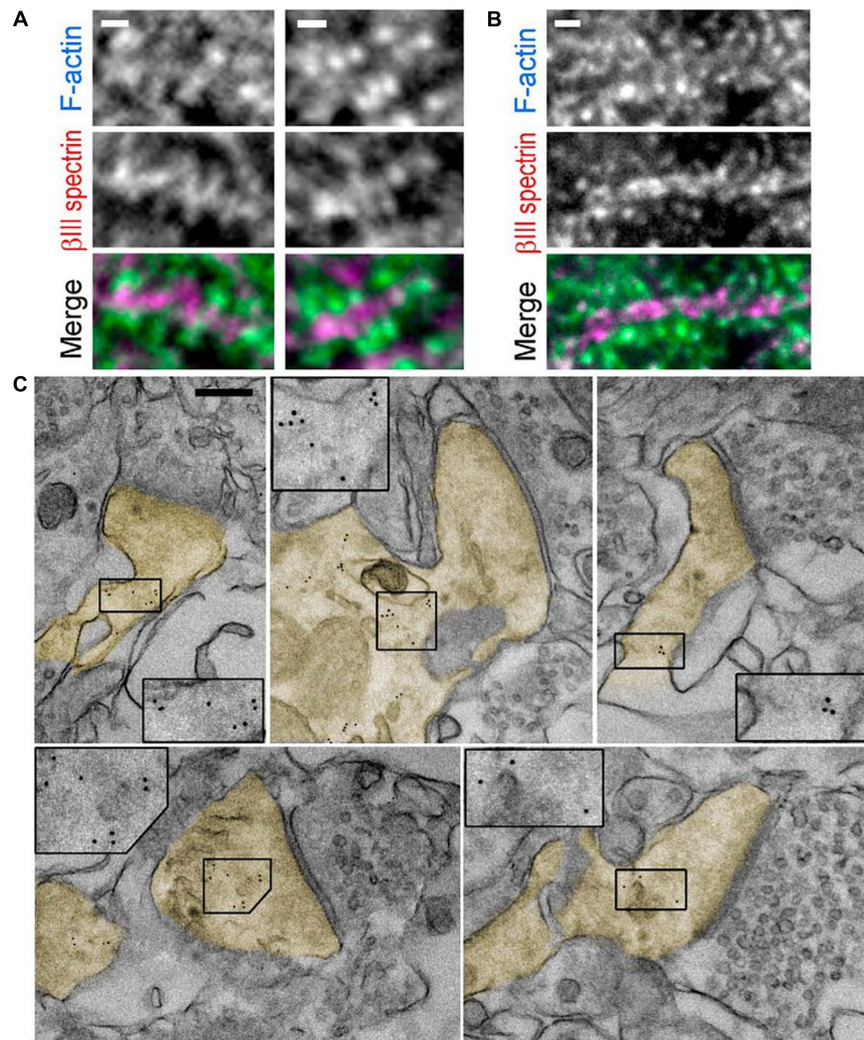


ABPs, transmembrane proteins, membrane lipids, and some organelles. Advanced techniques using freeze-etch combined with electron tomography (ET) can visualize the 3D molecular architecture of membrane-associated structures (e.g., membrane skeleton, clathrin-coated pits, and caveolae) at high resolution (Morone, 2010). The nanometer resolution of the tomogram in **Figure 5** shows that the spacing between adjacent proteins in the complexes is  $\sim 36$  nm. Freeze-etch electron tomography (ET) and ET using thin sectioning are good tools for high-resolution structural analysis. In ET, detailed 3D structures of subcellular macromolecular objects are obtained, because ET uses TEM images at incremental degrees of rotation around the target sample center. Images series of projections at different tilt angles (tilt series) (**Figure 5B**) are automatically aligned by cross-correlation or semiautomatic alignment with or without fiducial markers, such as gold particles. After the tilt-axis direction is determined by alignment, a tomogram is reconstructed using weighted back-projection (WBP), the algebraic reconstruction technique (ART), or the simultaneous iterative reconstruction technique (SIRT). Visualization of the computed tomogram is difficult because of neuronal cell complexity, so various segmentation techniques, such as artificial intelligence with manual segmentation, have been developed. A tomogram image showed actin filaments had in parallel or branching arrangements by ABPs, such as CaMKII (**Figure 5C**),

in a neuron at a nanoscale resolution. ET showed improved resolution after sample preparation using quick freezing and cryo-EM (**Figure 5C**).

### Immuno-Electron Microscopy

Immuno-EM is used to label proteins of interest using the antibody-antigen reaction like in indirect immunofluorescence. The first antibody of the antigen of interest and a second antibody-tagged colloidal gold particle are used to locate specific proteins for actin binding. Gold is used because of its high electron density, and the ABP location shows black dots in EM (**Figure 6**). Different sizes of gold particles can be used for multiple staining. Antibodies and gold particles cannot penetrate the resin used to embed samples, so before staining, thin sectioning using ultramicrotomy is necessary. For better antigenicity, cryofixation, freeze-substitution, low-temperature embedding, or cryo-sectioning is used in immuno-EM (Skepper and Powell, 2008). In addition, there are two different immunogold-labeling procedures, which can be divided depending on the embedding procedure. In pre-embedding, micrometer-thick tissue is immuno-stained prior to plastic embedding for ultrathin sections to be observed by TEM. In contrast, in post-embedding, tissue is embedded in a plastic resin and ultrathin sections are observed and then immunolabeled. Pre-embedding is better to get more immunolabeling, while



**FIGURE 6 |** Immunogold labeling. (A,B). Fluorescence staining with  $\beta$ III spectrin antibody (magenta) and phalloidin (cyan) of tissue sections from mouse hippocampus. (C) Immunogold EM of  $\beta$ III spectrin staining in thin sections of the mouse brain. Dendritic spines are pseudocolored in yellow. Inset: Boxed regions are enlarged to show 5 nm gold particles. Scale bars = (A,B), 1  $\mu$ m; (C), 200 nm. EM, electron microscopy. Reprinted by permission from Society for Neuroscience (Efimova et al., 2017).

post-embedding gives better ultrastructure (Webster et al., 2008). Immunogold labeling shows the detailed location of spectrin in the synapse (Figure 6C; Efimova et al., 2017) to study the function of  $\beta$ III spectrin in neurons.

## Super-Resolution Fluorescence Microscopy

Super resolution microscopy can be divided into two categories; single-molecule localization microscopy (SMLM) and illumination pattern engineering methods. The SMLM use photo-switchable fluorophores to locate single molecules with high precision, and include stochastic optical reconstruction microscopy (STORM) (Rust et al., 2006), photo-activated localization microscopy (PALM) (Betzig et al., 2006), and fluorescence photoactivation localization microscopy (FPALM)

(Hess et al., 2006). The illumination pattern engineering methods exploit a non-linear response to excitation fluorophores to overcome the diffraction limit. Such methods include stimulated emission depletion (STED) microscopy (Hell and Wichmann, 1994), [saturated] structured-illumination microscopy ([S]SIM) (Gustafsson et al., 1995), and reversible saturable optical fluorescence transitions (RESOLFT) microscopy (Hofmann et al., 2005). These SRM methods have also been applied to the study of actin and ABP by providing molecular-specific images at high resolution.

## Stochastic Optical Reconstruction Microscopy (STORM)

The diffraction limit arises from the diffraction of light passed through optical systems, ultimately resulting in a blurred image from the spatial overlapping of point spread functions (PSF) of

single molecules. To distinguish these molecules individually, STORM takes advantage of photo-switchable fluorescence dyes to temporally separate them (Figure 7A; Rust et al., 2006; Huang et al., 2009). The temporally separated individual molecules are localized with high precision, and a super-resolution image can be constructed from the collections of localizations from multiple fluorophores. This method enables the visualization of the ultrastructure of actin and ABP that was previously inaccessible with conventional optical methods and EM (Xu et al., 2013; Zhong et al., 2014; Sidenstein et al., 2016; Han et al., 2017; Pan et al., 2018; Wang G. et al., 2019). For example, Xu et al. (2013) first examined the periodic ring-like structure of actin, spectrin, adducin, and sodium channels in axons with a periodicity of ~180–190 nm using STORM, which had not been resolved by conventional light microscopy (Figures 7B–D). They also performed two-color STORM imaging for spectrin and actin (or adducing), which showed alternating periodic structures. Such a quasi-1D, periodic, actin-spectrin cytoskeleton was not observed in dendrites; thus, it allowed the authors to speculate that this structure may play an essential role in providing stable and elastic mechanical support for axons, which need to withstand mechanical strains on their long and thin structures.

In a later study by the same research group, Zhong et al. (2014) used STORM to elucidate the developmental mechanism of this membrane-associated periodic skeleton (MPS), such as how periodic cytoskeletons form and why they primarily form in axons (Figure 7E). The STORM images revealed that this periodic structure is formed early on in the axon development, followed by propagation from the proximal to distal ends of axons. They also demonstrated that the local concentration of spectrin regulates this structure, and overexpression of spectrin induces this periodic structure in dendrites, which contain very low concentrations of spectrin under normal conditions. The authors also found that knocking out ankyrin B triggered the periodic structure in dendrites; thus, finding that ankyrin B is critical for the polarized distribution of  $\beta$ II spectrin in neurites.

Spectrin organization was also examined in a variety of neuronal and glial cell types by He et al. using the same methods (Sidenstein et al., 2016). The authors showed that spectrin has a long-range and periodic distribution throughout all axons, whereas small patches of periodic spectrin structures are present in the sub-regions of dendrites and in four types of glial cells. These findings allow the authors to conclude that the periodic organization of spectrin is conserved across a wide range of invertebrate and vertebrate animal species.

The spatial organization of spectrin, actin, and adducin in the dendrites and soma at different developmental stages of cultured hippocampal neurons was also investigated by STORM (Figure 7F; Han et al., 2017). Han et al. observed that one-dimensional MPS slowly develops in the dendrites of mature neurons at a much slower rate than that in axons. Interestingly, the authors also found that they form a two-dimensional polygonal lattice structure in the somatodendritic compartment at a much slower rate than that in the 1D MPS in axons. This result suggests that

actin-based membrane skeletons organized with spectrin and ABP are differentially regulated across different sub-regions of neurons. A two-dimensional polygonal lattice structure was also observed from a native ultrastructure of the cytoskeleton in erythrocytes (Pan et al., 2018). Pan et al. used STORM imaging to resolve the ultrastructure of the cytoskeleton, including  $\beta$ -spectrin, F-actin, protein 4.1, tropomodulin, and adducin. They revealed a junction-to-junction distance of ~80 nm, which is in agreement with the relaxed spectrin tetramers; actin and its capping proteins occupy subsets of junctional complexes.

Recently, Wang G. et al. (2019) investigated the behavior of MPS in sensory axon degeneration using the same super-resolution microscopy. From the STORM images of  $\beta$ II spectrin, they found that trophic deprivation (TD) caused the rapid disassembly of MPS, which can be initiated by actin destabilization. In contrast, knockout of  $\beta$ II spectrin prevented TD-induced retrograde signaling to protect axons from degeneration by inhibiting MPS formation.

Collectively, substantial progress has recently been made in identifying the developmental mechanism of the periodic actin-spectrin-based membrane skeleton in different sub-compartments of neurons by taking advantage of the high molecular specificity and spatial resolution of STORM.

### Photo-Activated Localization Microscopy (PALM)

Another widely used SMLM method is PALM, which shares many similarities with STORM. PALM also exploits the temporal image separation of individual fluorophores based on the on/off behavior of fluorescent proteins (Betzig et al., 2006; Huang et al., 2009). Although the only difference between these two methods is the type of fluorescent label (i.e., photo-switchable fluorescent proteins for PALM and photo-switchable organic dyes for STORM), they are essentially similar in terms of configuration and differences are blurred as they use both fluorescent organic dyes and fluorescent proteins. PALM is also used in the study of actin, and most investigations have focused on the visualization of actin dynamics using single-molecule tracking (Tatavarty et al., 2009; Frost et al., 2010; Izeddin et al., 2011).

Tatavarty et al. (2009) for instance, utilized PALM to observe the kinematic (physical motion of actin filaments) and kinetic dynamics of F-actin in the dendritic spines of hippocampal neurons, which had been hindered by the small size of the dendritic spines in diffraction-limited conventional light microscopy (Figures 8A–C). Single-molecule tracking by PALM showed highly heterogeneous kinematic dynamics of F-actin in dendritic spines at a single-filament level, in which simultaneous actin movements were observed in both retrograde and anterograde directions. In contrast, the movements of filaments integrate into a net retrograde flow at the ensemble level, suggesting short actin filaments in dendritic spines.

Frost et al. (2010) also investigated the movement of individual actin molecules within living spines (Figures 8D–F). In a similar manner to Tatavarty et al., they used PALM-based single-molecule tracking to measure the velocity of single actin molecules along filaments as an index of the

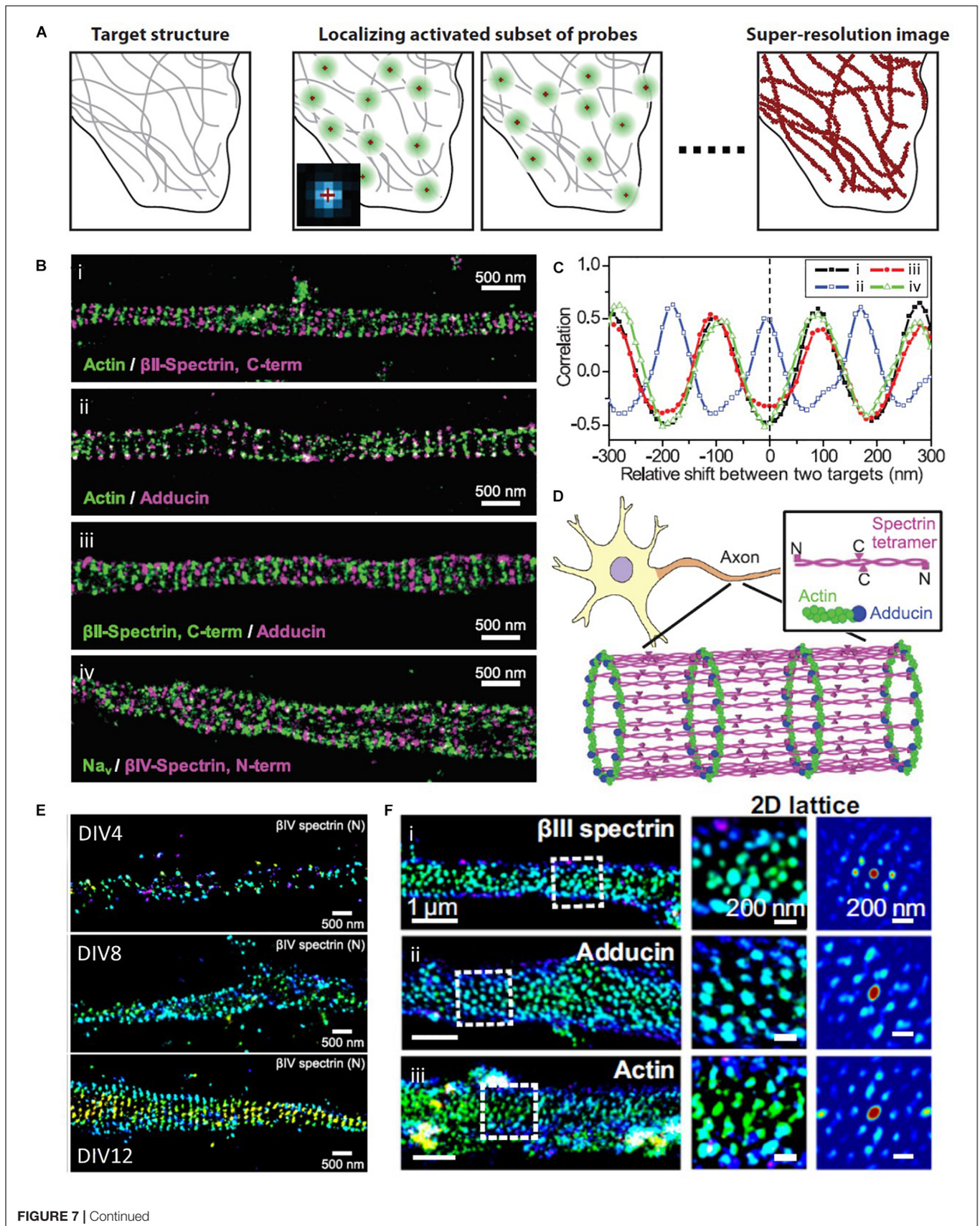
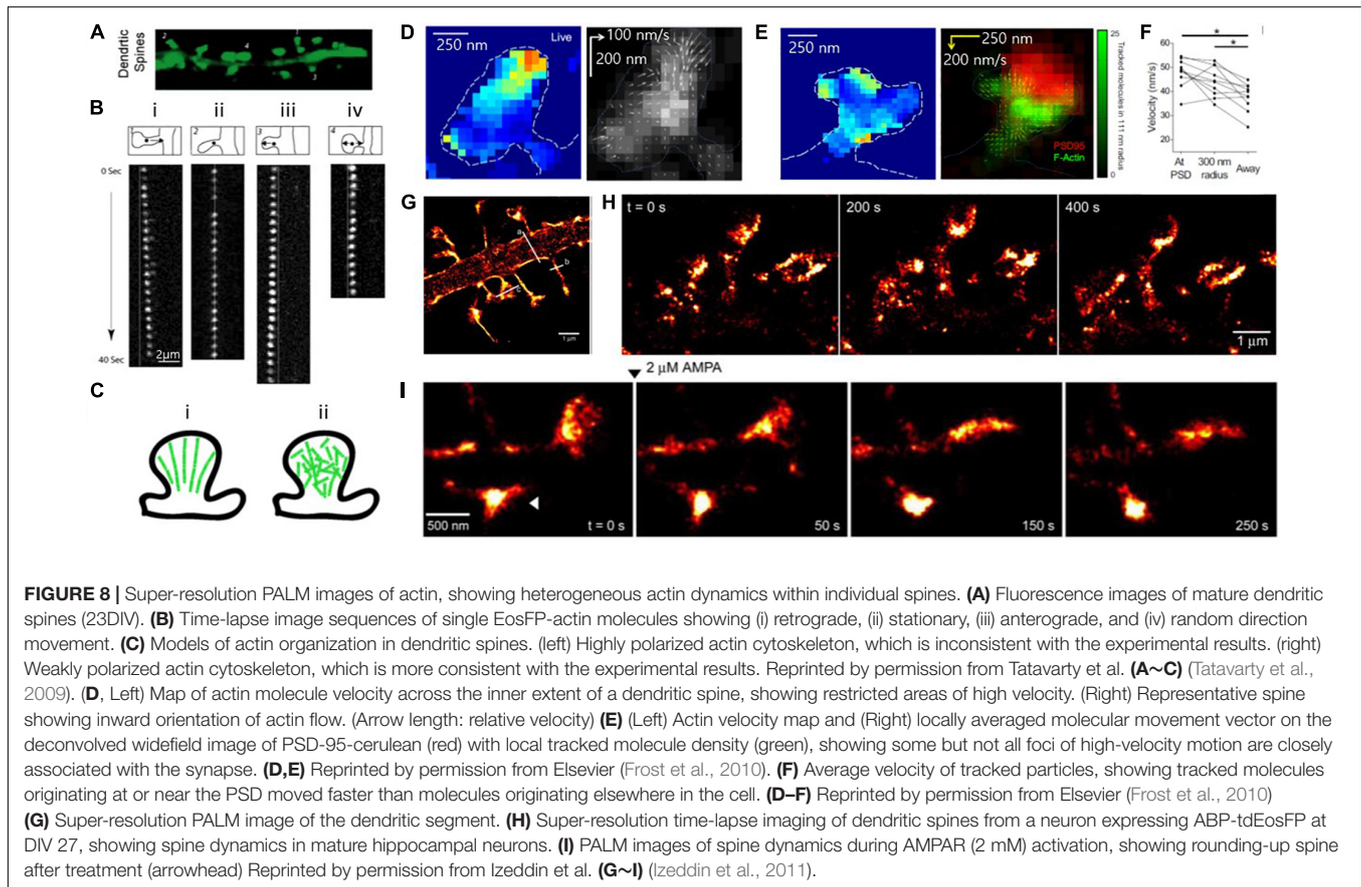


FIGURE 7 | Continued

**FIGURE 7 |** Super-resolution STORM images of actin and ABPs. **(A)** The principle of SMLM, such as STORM, PALM, and FPALM. Reprinted by permission from Annual Reviews (Huang et al., 2009). **(B)** Two-color STORM image of (i) actin (green) and  $\beta$ II-spectrin C terminus (magenta), (ii) actin (green) and adducin (magenta), (iii)  $\beta$ II-spectrin (green) and adducin (magenta), and (iv) sodium channels ( $Na_v$ , green) and  $\beta$ IV-spectrin N terminus (magenta). **(C)** Spatial correlations between actin and the  $\beta$ II-spectrin C terminus [(i), black], between actin and adducin [(ii), blue], between adducin and the  $\beta$ II-spectrin C terminus [(iii), red], and between sodium channels and the  $\beta$ IV-spectrin N terminus [(iv), green], which are calculated for varying relative shifts between the two color channels along the axons. **(D)** A model for the cortical cytoskeleton in axons, forming ring-like structures wrapping around the circumference of the axon with a periodicity of  $\sim 180$  to  $190$  nm. Reprinted by permission from the American Association for the Advancement of Science (B~D) (Xu et al., 2013). **(E)** Representative STORM images at different developmental stages, showing assembly of AIS components into the periodic lattice structure during late developmental stages. Reprinted by permission from Zhong et al. (2014). **(F)** MPS components form 2D polygonal lattice structures in some somatodendritic regions of neurons. (Left) 3D STORM image of a dendritic region from a DIV 28 mouse neuron stained for (i)  $\beta$ III-spectrin, (ii) adducin, and (iii)  $\beta$ III-spectrin. (Middle) Zoom-in image of the region indicated by the white dashed box in the Left. (Right) 2D autocorrelation function of this boxed region, which shows a 2D periodic lattice pattern. Reprinted by permission from PNAS (E,F) (Han et al., 2017).



**FIGURE 8 |** Super-resolution PALM images of actin, showing heterogeneous actin dynamics within individual spines. **(A)** Fluorescence images of mature dendritic spines (23DIV). **(B)** Time-lapse image sequences of single EosFP-actin molecules showing (i) retrograde, (ii) stationary, (iii) anterograde, and (iv) random direction movement. **(C)** Models of actin organization in dendritic spines. (left) Highly polarized actin cytoskeleton, which is inconsistent with the experimental results. (right) Weakly polarized actin cytoskeleton, which is more consistent with the experimental results. Reprinted by permission from Tatavarty et al. (A~C) (Tatavarty et al., 2009). **(D)** (Left) Map of actin molecule velocity across the inner extent of a dendritic spine, showing restricted areas of high velocity. (Right) Representative spine showing inward orientation of actin flow. (Arrow length: relative velocity) **(E)** (Left) Actin velocity map and (Right) locally averaged molecular movement vector on the deconvolved widefield image of PSD-95-cerulean (red) with local tracked molecule density (green), showing some but not all foci of high-velocity motion are closely associated with the synapse. **(D,E)** Reprinted by permission from Elsevier (Frost et al., 2010). **(F)** Average velocity of tracked particles, showing tracked molecules originating at or near the PSD moved faster than molecules originating elsewhere in the cell. **(D~F)** Reprinted by permission from Elsevier (Frost et al., 2010). **(G)** Super-resolution PALM image of the dendritic segment. **(H)** Super-resolution time-lapse imaging of dendritic spines from a neuron expressing ABP-tdEosFP at DIV 27, showing spine dynamics in mature hippocampal neurons. **(I)** PALM images of spine dynamics during AMPAR (2 mM) activation, showing rounding-up spine after treatment (arrowhead) Reprinted by permission from Izeddin et al. (G~I) (Izeddin et al., 2011).

filament polymerization rate. They found that the actin velocity on filaments was substantially elevated at the synapses. In contrast, no enhanced polymerization activity was observed in the endocytic zone, implying a highly heterogeneous rate of actin molecules.

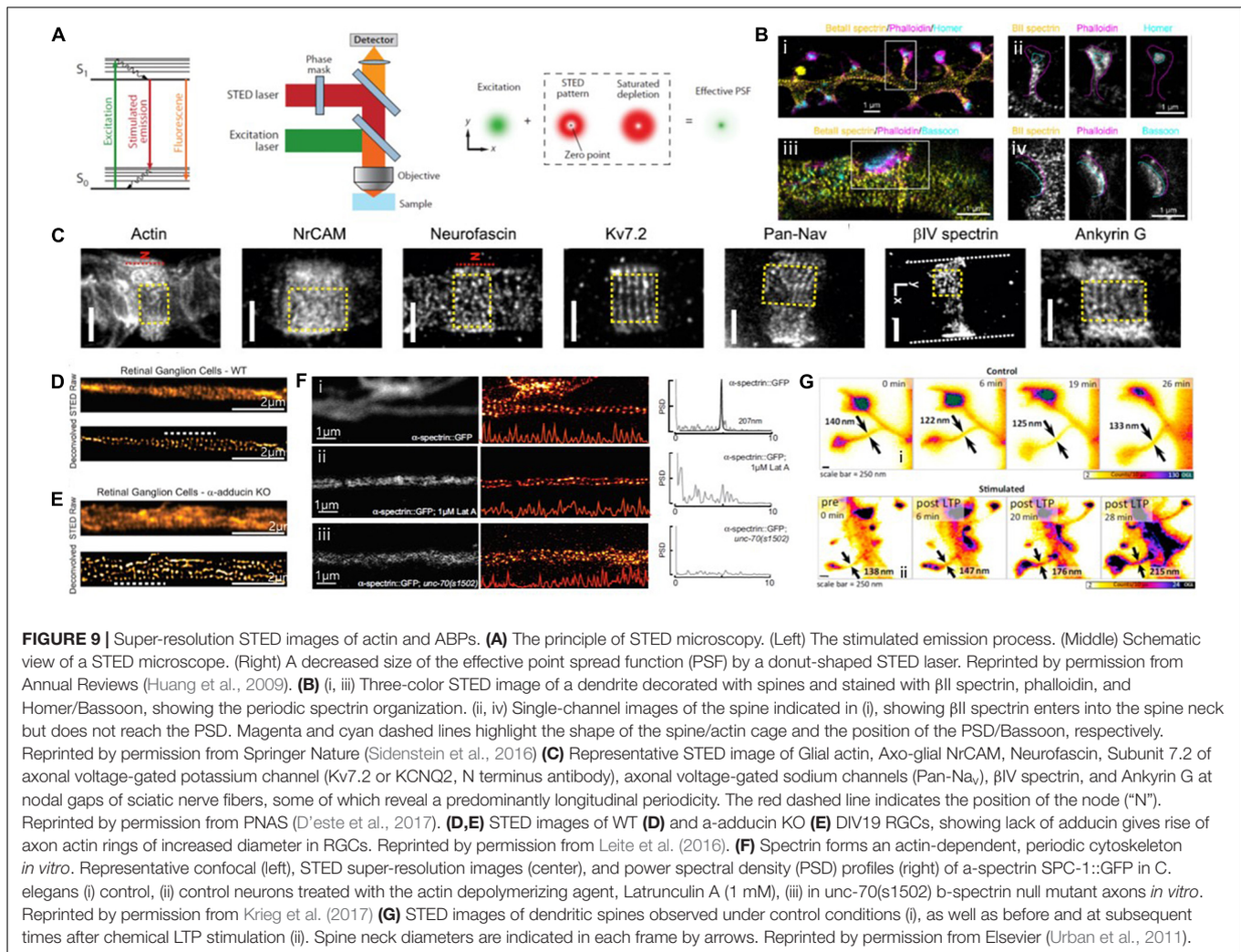
The morphological changes of the spine cytoskeleton were also investigated at super-resolution by Izeddin et al. (2011) (Figures 8G–I). By combining PALM imaging with quantum dot tracking for long-term imaging, they simultaneously observed the cytoskeleton and spine membranes, and their evolution during pharmacologically induced synaptic activity.

Although all studies discussed in this section visualize actin but not ABP in live cells, they demonstrate the possibility of future studies regarding ABP dynamics in relation to the actin in live neurons as a further application of PALM imaging.

## Stimulated Emission Depletion (STED)

Another widely used SRM is STED, which was developed by Hell et al. In STED microscopy, a donut-shaped depletion light is used to suppress the fluorescence emission from the fluorophores as a pattern surrounding the focal spot of the excitation laser (Figure 9A; Hell and Wichmann, 1994; Huang et al., 2009). Such a negative pattern reduces the size of the fluorescent region and scanning of this sharpened focal spot generates a super-resolution image. Through the use of STED, actin and ABP have been thoroughly investigated (Urban et al., 2011; Lukinavičius et al., 2014; Leite et al., 2016; Sidenstein et al., 2016; Barabas et al., 2017; D'este et al., 2017; Krieg et al., 2017; Martínez et al., 2020).

For instance, the subcortical cytoskeleton organization at synaptic sites was previously investigated using three-color multilevel STED nanoscopy (Figure 9B; Sidenstein et al., 2016).

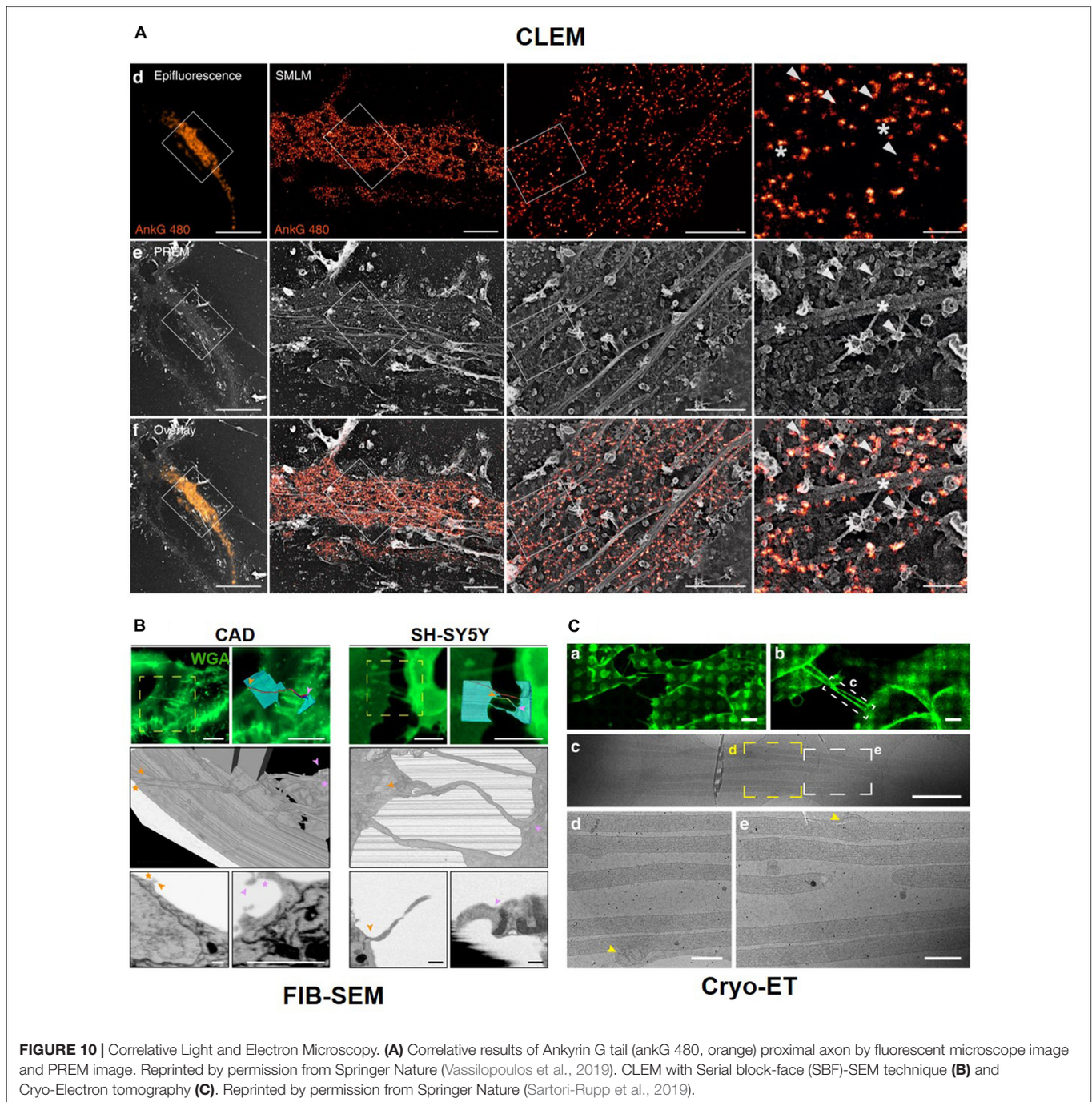


This technique was developed based on an intrinsically co-aligned multicolor imaging scheme to overcome the limitations of multi-color imaging by STED, such as the requirement for complicated setups and analysis algorithms. This technique was successfully demonstrated by imaging neurofascin/spectrin and actin in cultured hippocampal neurons. The multicolor STED images revealed the  $\sim 190$  nm periodic actin/ $\beta$ II spectrin lattices along dendrites and spines, consistent with previously reported STORM images, whereas this periodic pattern was absent at presynaptic and postsynaptic sites. Such protein periodic structures in super-resolution images can now be automatically quantified in terms of the abundance and regularity of the MPS. For example, Barabas et al. (2017) presented a method for the automated quantification of the quality and abundance of periodic protein structures in super-resolution images as an open-source image analysis tool that provides the distributions of correlation coefficients for the spectrin membrane-associated periodic skeleton as an indication of the dynamic assembly and disassembly of the MPS.

D'este et al. (2017) also used STED to map the nanoscale molecular organization of 12 proteins at the nodes of Ranvier

of sciatic nerve fibers, including cytoskeletal proteins of the axon, axonal, and glial nodal adhesion molecules, sodium and potassium channels, axonal proteins, and glial proteins (Figure 9C). The STED images showed that  $\beta$ IV spectrin and ankyrin G revealed a known one-dimensional longitudinal order at nodal gaps, whereas neurofascin-186 and neuron glial-related cell adhesion molecule (NrCAM) revealed a two-dimensional hexagonal-like lattice with a  $\sim 190$  nm periodicity, which has not yet been explored. The authors also observed that both glial proteins (neurofascin-155 and ankyrin B) and axonal proteins ( $\beta$ II spectrin, Caspr) form quasi-1D periodic arrangements at paranodes. These results suggest the importance of the lateral organization of proteins at the nodes of Ranvier.

The defects in ABP have also been investigated by STED imaging to better understand their roles. For example, Leite et al. (2016) observed an increased actin ring without changing the inter-ring periodicity in neurons in mice lacking adducin (Figures 9D,E). This result suggests a model in which adducin is responsible for controlling the diameter of actin rings and axons, but it is not necessary for actin ring assembly and periodicity. Krieg et al. (2017) reported that periodic spectrin structures



were absent due to the loss of UNC-70  $\beta$ -spectrin and actin depolymerization from STED images (Figure 9F). These results allowed the authors to conclude that mechanical neuroprotection depends on spectrin-dependent longitudinal tension, and the defects in  $\beta$ -spectrin may sensitize neurons to damage.

Live cell imaging for actin was also performed using time-lapse STED imaging. To investigate synapses in brain slices or brains under realistic conditions, Urban et al. (2011) used aberration-reducing optics for STED imaging and demonstrated this for deep inside biological tissue imaging

(Figure 9G). This method enabled the visualization of distinct actin distributions inside dendritic spines in living organotypic brain slices at depths of up to 120  $\mu$ m with 60–80 nm spatial resolution, revealing that structurally heterogeneous and highly dynamic actin in spine necks were modulated by neuronal activity. However, long-term STED imaging of live biological samples is often limited due to its high laser power requirement for high spatial resolution. To overcome this limit, Lukinavičius et al. (2014) introduced fluorogenic probes for actin and tubulin (SiR-actin, SiR-tubulin) that exhibited

minimal cytotoxicity, excellent brightness, and photostability. They successfully revealed the one-dimensional periodic actin organization in the axons and the nine-fold symmetry of the centrosome of cultured rat neurons. These advances are expected to allow further investigations into ABP dynamics *in vivo* in the near future.

## Correlative Light and Electron Microscopy (CLEM)

As previously stated, fluorescence light microscopy (LM) and EM have been extensively used for biological studies to probe cellular structures and obtain a better understanding of biological functions. Each of these techniques has its distinct strengths and weaknesses that complement each other, as previously discussed. Fluorescence LM provides protein-specific information with high molecular specificity and sensitivity. Multicolor fluorescence LM offers a promising approach to monitor potential molecular interactions between spectrally different fluorescent labels, and live cell LM imaging can reveal the spatiotemporal dynamics of a protein. Moreover, recently developed SRM techniques allow images to be taken with a higher resolution than that with the diffraction limit, as described in the previous section.

Although such advances in SRM have made remarkable headway in improving spatial resolution by order of magnitude or moreover, the diffraction limit, EM generally achieves a higher spatial resolution in protein localization compared to that with SRM, revealing the subcellular anatomy at resolutions of nanometers. However, conventional fixation and staining methods of traditional EM do not usually provide protein-specific information because of the limited molecular specificity. Moreover, EM is limited to live cell imaging, since the electrons destroy the samples, and the sample needs to be processed and fixed for EM imaging.

As the strengths and weaknesses of LM and EM are complementary, these two methods can be effective if used together for the same sample. This combination is the so-called correlative light and EM (CLEM). It provides different and complementary information by merging the specific protein localization advantages of LM with the ultrastructural information of EM.

Correlative light and electron microscopy is often compared to immuno-EM because this has been the method of choice for protein localization at an ultrastructural level. However, the application of immuno-EM is significantly limited by several technical difficulties, including the destruction of antigens by strong fixation, inaccessibility of antigens in the plastic, poor survival of epitopes, limited preservation of morphology, incomplete labeling, non-specific binding of antibodies, and non-specific background. Therefore, only a subset of target molecules are successfully detected by immuno-EM. These drawbacks have prompted the development of various CLEMs for protein localization at an ultrastructural level. Compared with immuno-EM, CLEM significantly improves the detection efficiency and sensitivity with sufficient reliability and specificity. Moreover, when EM is combined with SRM, such correlative SRM-EM approaches provide a more meaningful correlation between two

images due to their closer resolution match compared to that by CLEM (Kim et al., 2015).

Such CLEM methods have also allowed for the identification of neuronal cell structures and processes of interest in whole-cell images with high molecular specificity and high spatial resolution. This dual examination provides valuable unique and complementary information for actin and ABP. For instance, Vassilopoulos et al. (2019) utilized PREM to visualize the dynamic actin network and combined this with STORM to specifically localize ankyrin G (**Figure 10A**). This correlation imaging method revealed the molecular organization of the actin rings and MPS components along the axonal plasma membrane. In particular, the association of MPS with microtubules via ankyrin G was visualized along the AIS by correlative STORM and PREM.

Sochacki et al. also showed the technique of CLEM with PREM to observe a clathrin lattice and actin (Sochacki et al., 2017). CLEM is also used to obtain high-resolution images using serial sectioning. For example, tunneling nanotubes (TNTs), which are actin-rich membranous protrusions, play a role in the intercellular transport of various cargoes between neuronal cells and can be observed by CLEM, which reveals their structural identity using LM with focused ion beam (FIB)-scanning electron microscopy (SEM) in neuronal cells (**Figure 10B**). FIB-SEM is a serial block-face (SBF)-SEM technique using serial sectioning. Another SBF-SEM technique uses an ultramicrotome embedded inside a SEM. The advantage of these techniques is that we can easily obtain serial ultrathin sections, but the sections can be observed at one time. CLEM can be combined with cryo-ET after cryo-FIB sectioning. CLEM can also be combined with confocal or fluorescent microscopy and super-resolution microscopy, such as stochastic optical reconstruction microscopy (STORM) (Zhanghao et al., 2019) and stimulated emission depletion microscopy (STED) (Urban et al., 2011). Cryo-EM and cryo-FIB-SEM with LM show open-ended TNTs labeled with anti-*N*-cadherin antibodies and 3D structures of TNTs. Labeled TNTs are filled with parallel actin bundles and membrane-bound mitochondria (**Figure 10C**; Sartori-Rupp et al., 2019).

## CONCLUSION AND FUTURE PERSPECTIVES

In conclusion, various microscopy techniques have been successfully demonstrated to visualize actin filaments and ABPs in neuronal cells. First of all, EM has been the primary tool to study actin and ABPs because of its nanometer resolution, for example, single particle analysis, electron tomography with freeze etching and unroof, and immuno-gold staining. Using these EM techniques, ABP binding sites for actin filaments and the organization of axonal actin filaments in neurons have been investigated. However, molecular specific imaging and live-cell imaging have been limited to EM techniques. To overcome these limitations, the recently developed SRM has been applied to the study of actin and ABP, including STORM, PALM, and PALM. This method enables the visualization of the ultrastructure of actin and ABP that was previously inaccessible with conventional optical methods and EM, for example, the periodic ring-like

structure of actin and ABPs even in live neurons with high molecular specificity at high resolution. As the strengths and weaknesses of LM and EM are complementary, the CLEM has also been applied to the study of actin and ABP. Combining these two methods has allowed for the identification of neuronal cell structures and processes of interest in whole cells with high molecular specificity and high spatial resolution.

In the future, we foresee that *in situ* or wet-environment EM would be a goal in this field to study actin and ABPs. Studies have reported progress in observing proteins' structural changes (Fitzpatrick et al., 2013) and the ultrastructure of living cells in the Four-dimension (4D), time (De Jonge and Ross, 2011; Ross, 2015). Although the resolution is poor, live cells or proteins *in situ* have been observed so that actin and ABPs would be the next challenge for this 4D EM-combined CLEM in the near future.

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## AUTHOR CONTRIBUTIONS

MJ, DK, and JM wrote the manuscript. All authors contributed to the article and approved the submitted version.

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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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