Revealing t-tubules in striated muscle with new optical super-resolution microscopy techniques

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Abstract

The t-tubular system plays a central role in the synchronisation of calcium signalling and excitation-contraction coupling in most striated muscle cells. Light microscopy has been used for imaging t-tubules for well over 100 years and together with electron microscopy (EM), has revealed the three-dimensional complexities of the t-system topology within cardiomyocytes and skeletal muscle fibres from a range of species. The emerging super-resolution single molecule localisation microscopy (SMLM) techniques are offering a near 10-fold improvement over the resolution of conventional fluorescence light microscopy methods, with the ability to spectrally resolve nanometre scale distributions of multiple molecular targets. In conjunction with the next generation of electron microscopy, SMLM has allowed the visualisation and quantification of intricate t-tubule morphologies within large areas of muscle cells at an unprecedented level of detail. In this paper, we review recent advancements in the t-tubule structural biology with the utility of various microscopy techniques. We outline the technical considerations in adapting SMLM to study t-tubules and its potential to further our understanding of the molecular processes that underlie the sub-micron scale structural alterations observed in a range of muscle pathologies.

Key Words: cardiac muscle, skeletal muscle, t-tubules, excitation-contraction coupling, super-resolution microscopy, human, rat, mouse, rabbit, horse

Cardiac myocytes and skeletal muscle fibres are among the largest cell types in vertebrate species. Ventricular myocytes can be ~20 µm in width and >100 µm long while skeletal muscle fibres are found in diameters in excess of 100 µm, extending over millimetres to centimetres from tendon to tendon. Forceful contraction of such striated muscle relies on the rapid and synchronised opening of the primary calcium (Ca^{2+}) release channels of the sarcoplasmic reticulum (SR) - the ryanodine receptors (RyRs).\(^1\) RyRs are clustered into dense arrays adjacent to t-tubules in both cardiac\(^2\) and skeletal muscle.\(^3\) A series of tubular membrane invaginations of the surface sarcolemma, collectively known as the transverse tubular system (t-system or t-tubules), is responsible for the synchronisation of excitation-contraction (E-C) coupling. They achieve this by the delivery of the excitation (action potential) to the Ca^{2+} release sites in junctions with terminal SR either at the z-line (in mammalian myocardium) or at the A-band boundary (in mammalian skeletal muscle).\(^4\) Junctions between t-tubules and nuclear envelope/ER also facilitate their putative role in excitation-transcription (E-T) coupling\(^5\) via pathways like store-operated Ca^{2+} entry dependent NFAT activation,\(^6,7\) IP3-receptor/Calmodulin Kinase-II mediated nuclear signalling\(^8\) and β-adrenergic stimulation.\(^9\) The structure of the t-tubules can greatly affect their roles in the above signalling mechanisms; for example, the widespread network structure of the t-system determines how well it synchronises the Ca^{2+} release in the muscle cell interior. Similarly, the tortuosity and local tubule diameters can also affect tubular luminal ion concentrations and effectively regulate t-system excitability and conduction properties.\(^10\) Models that simulate such ion/protein dynamics within a geometrically accurate environment are particularly useful for predicting the structural constraints on EC coupling.\(^11\) This approach has been used recently with models of tubular ionic movements that have been constructed either solely on microscopy data of t-tubules\(^12\) or a combination of experimentally-traced and simulated t-tubule features.\(^13\) Therefore we see such models as a primary motivator for the microscopy analysis of t-tubules.
Microscopy techniques that allow the visualisation of t-tubules either for accurate structural measurements (useful for modelling its function) or for functional studies have therefore been an integral part of muscle biology for well over 100 years (see reviews of history). Some of the pioneering work carried out since the 1950s, used high-contrast staining methods combined with electron microscopy (EM). In more recent decades, fluorescence techniques that are compatible with live cell imaging have become popular (see discussion below). The main appeal in fluorescence microscopes includes high-contrast images, high specificity of fluorescent labelling methods and 3D imaging applications with optical sectioning. However, the resolution achievable with these methods is fundamentally limited to approximately half the wavelength of the light (e.g. 250 nm) due to the wave properties of light (i.e. diffraction). In practical terms, we observe a considerable loss of the sub-micron scale features of muscle structure in conventional fluorescence micrographs, especially of t-tubules. Several recent optical microscopy methods that promise EM-like resolution, high specificity in mapping molecular targets while circumventing this “diffraction-limit” are optical super-resolution imaging (see recent reviews,). In the following we discuss the application of one of these approaches, single molecule localisation microscopy (SMLM) for visualising multi-scale morphologies of cardiac and skeletal muscle t-tubules, a technique which we have adopted in our own work.

**Localisation microscopy: breaking the diffraction limit**

SMLM utilises the stochastic photo-switching or modulation of individual fluorophore emission to localise individual fluorophores within the sample. This is typically achieved with a combination of light-induced photoactivation, photobleaching and/or supplementing the sample immersion medium with a ‘switching buffer’ that may consist of oxygen scavenging species and/or reducing agents (e.g. thiols). Photoswitching individual fluorophores typically yields a diffraction-limited spot of fluorescence with an intensity profile that reflects the point spread function (PSF) of the microscope (often approximated by a 2D-Gaussian), lasting up to a few frames in a fast and sensitive camera. In-focus fluorophore positions can be localised at nanometre-accuracy by fitting these 2D spots (“events”) with a model that approximates the PSF but one must be careful that the distance between adjacent events is larger than the minimum resolvable distance of the microscope (i.e. > half the wavelength) to avoid distortions by overlap. The resultant fluorophore coordinates can be reconstructed into greyscale images that, similar to standard fluorescence micrographs, report intensity in proportion to the local event densities but at a resolution of 10-30 nm. Multicolour SMLM has also been possible by spectral unmixing of fluorophore emissions. Chromatically separating the emissions of multiple fluorophores and examining the ratios of their respective emissions in two channels provides an emission ratio that is typically unique to each fluorophore – which can then be used for assigning the spectral identity to a given event. The primary appeal of this method to the muscle biology community is that it is compatible with a range of commercially available organic fluorophores (e.g. AlexaFluor and cyanine dyes), membrane dyes (e.g. Dil), genetically expressed reporters such as GFP, mEos (see a partial list of common SMLM markers: Table 2 of ). Depending on the mechanism of photoswitching and/or labelling the SMLM technique may be known by one of many acronyms such as dSTORM, PALM, GSDM (see Table 1 of for a summary).

The simplicity of this concept meant that an appropriate sample could be imaged with a basic fluorescence microscope and a camera sufficiently sensitive to record single molecule events. Total internal reflection fluorescence (TIRF) microscopes, but used with a sub-critical beam angle are suitable for imaging muscle tissue that are often thicker than the cultured cell samples that are typically used in the majority of SMLM studies to date. This configuration allows an effective optical sectioning of the sample enabling the imaging of structures further away from the cover slip with an axial resolution similar to confocal microscopy. Electron multiplying charged coupled devices (EMCCD) or scientific CMOS cameras are currently the most widely used detectors for these experiments because of their high sensitivity, comparatively low read-noise and the high frame rates that can be obtained. Mapping the full complement of fluorophores within an image region may require acquisition of 10-50,000 frames over 5-30 minutes depending on the photoswitching rates. The computationally demanding features of fluorophore detection, fitting (localisation), spectral unmixing and reconstruction algorithms often require a powerful computer (readily available and very affordable as compared to microscope and camera hardware) with large storage capacity that can hold the large image sequences. However, increasingly, open-source or free analysis software packages are becoming available (e.g. rapid STORM), which should help introduce non-experts to this approach and commercial systems are also available.

The use of SMLM has grown exponentially over the 8 years since its initial proof of principle (see recent reviews discussing the expansion of SMLM). Additional to the obvious improvement in the resolution, images are often “background free” because fluorophore localisation above a standing non-uniform
background implicitly exclude any sarcomeric autofluorescence patterns (e.g. mitochondria). The stochastic nature of the photoswitching means that the fluorophore coordinates can be used for quantitative analysis of the underlying protein/fluorophore densities or distances to the surrounding structures, although fully quantitative approaches need to carefully avoid artefacts, such as overcounting and failing to reject multi-emitter events. Overestimation of co-localising fractions between two proteins is avoided (to a larger extent) with the superior resolution while image binarisation tends to be more robust. Combined with lipophilic membrane dyes such as DiI, volumetric dyes (e.g. wheat germ agglutinin), SMLM provides potentially a very convenient method of resolving the structure or tracking any remodelling of t-tubules of living muscle preparations at nanometre resolution. Further advancements in the field include 3D SMLM methods that, with fairly straightforward optical modifications, can produce 3D super-resolution images with >10 fold finer axial resolution compared to confocal images that can be particularly useful for imaging a 3D structure like the t-system.

Achieving the super-resolution often promised by SMLM, however, requires consideration of several aspects with respect to sample preparation and imaging protocols. The effective resolution in the image is ultimately dependent on the localisation error and the effective labelling density of the underlying structure. The localisation error is approximately proportional to \( \frac{\sigma}{\sqrt{N}} \) where \( \sigma \) is the standard deviation of the best fit Gaussian (i.e. the diameter of the PSF) and \( N \) the number of photons collected from the event. Maximising \( N \) is typically achieved by increasing the excitation, integrating events lasting multiple frames, chemically increasing the lifetime of the fluorophore events and generally using an optical detection path that aims to minimise light losses. However, in a sample that has substantial background fluorescence (e.g. very high labelling densities or out of focus fluorescence, typically in ‘thick’ samples) the localisation error rises with the background intensity. Therefore, samples that are ultra-clean or ‘thin’ are ideal for SMLM performed with TIRF or Light-sheet microscopy. However, we have demonstrated that SMLM is compatible with isolated cardiomyocytes, single skeletal muscle fibres and ‘thin’ (by confocal standards, i.e. 5-10 \( \mu \)m) tissue sections of ventricular and skeletal muscle to achieve ~30 nm resolution.

A perhaps more serious and frequently encountered issue is “under-labelling” which reduces the detectability of nanoscale features of t-tubules. This becomes a particularly concerning issue during 3D SMLM where sufficient fluorophore densities in the axial dimension are also vital towards a super-resolution 3D image. Fig. 1 illustrates a simulation that summarises the effect of labelling density and background fluorescence on the reconstructed super-resolution image. It is noteworthy that in an equivalent

**Fig 1.** Simulating the effect of labelling densities and background fluorescence on diffraction-limited and SMLM images. (A) A phantom t-tubule with a varying diameter (a = 130 nm and b = 280 nm) and a 20 nm-thick layer of fluorescent markers (similar to combined thickness of a layer of primary and secondary antibodies bound to a t-tubular protein; e.g. caveolin-3) is simulated in (i) confocal micrographs and (ii) typical 2D dSTORM. Images were simulated for labelling densities of (B) 0.12/nm², typical of very high fluorophore densities, (C) 0.008/nm², typically achieved in experimental samples and (D) 0.0005/nm² to simulate a 15-fold poorer labelling than typical. Notice the gradual loss of detail in the dSTORM images with the diminishing labelling density and little observable difference between the confocal images (B-i & C-i). To simulate the effect of higher background intensity (E), the background intensity was set to 50% of the foreground intensity and a foreground labelling density comparable to the simulation in panel-C. Notice the lack of detail on the dSTORM image (E-ii compared to C-ii) while little change is seen in the morphologies in the confocal images (E-i compared to C-i). Scale bar: 500 nm.
confocal micrograph under each condition, the structural features appear often unaffected and mainly result in a dimmer image (which will, however, affect the signal-to-noise ratio, but may be overcome to some extent by “turning up the laser” or increasing integration time). This underscores the greater emphasis that SMLM places on the quality of the samples for achieving super-resolution. Table 1 collates a summary of (our view on) advantages, technical considerations and limitations of SMLM with a particular view towards imaging in muscle.

**Progress in sub-micron-scale microscopy of cardiac and skeletal muscle t-tubules**

Historically, electron microscopy (EM) data have been instrumental to understanding of the fine ultrastructure of muscle, especially t-tubules. Optical microscopy techniques on the other hand have offered the muscle biologist a means for high contrast, 3D images over comparatively large regions of fibres (or tissue), ideal for high spatial sampling (many sarcomeres or large portions of the muscle cell width) and 3D visualisation of the structures of the t-system in the sub-micron to sub-millimetre scales, however, with the resolution limitations as discussed above. Therefore, EM, which is generally not compatible with live-cell imaging and often has limited molecular specificity (but excels at imaging membranes with suitable staining protocols), has remained the primary method to resolve its detailed features.

Soeller and Cannell used two-photon (diffraction-limited) micrographs (similar to Fig 2A &B), digital deconvolution and the entry of a volumetric dye (membrane impermeable fluorescein-dextran) into the rat cardiac t-system (previously used by Endo) to overcome the lack of spatial detail in fluorescence micrographs to some degree. Whole-cell images like these (Fig 2A) were instrumental in observing t-tubules that were predominantly transverse in their orientation but can also run in longitudinal and oblique directions (see magnified image; Fig 2B). By calibrating the fluorescence intensity to the local tubule volume and geometry, they were able to characterise the full distribution of the diameters of t-tubules, many of which were well below the diffraction-limit (blue bar graph in Fig 2C). Direct measurements from antibody-labelled super-resolution images (unpublished data) have since validated these measurements (green bar graph, Fig 2C). A similar method has been used by Savio-Galimberti et al, to characterise the t-tubules of rabbit ventricular myocytes that were typically twice as wide as rat t-tubules and mostly resolvable with confocal microscopy. Their data were the first optical images indicating localised dilatations on t-tubules and flattened topologies that may act as reservoirs to compensate for potential ionic accumulations and/or depletion.

Similar structures can be better resolved by imaging the myocytes in an “end-on” orientation (i.e. in transverse optical sections) which limits artefacts that may be introduced into the images by the asymmetry of the confocal point spread function (PSF). The transverse view of the t-tubule network across the cell widths in these approaches, in our experience, provides

**Fig 2. Confocal fluorescence microscopy of t-tubules of isolated living myocytes.** (A) A whole cell longitudinal confocal image of a rat ventricular myocyte immersed in a dextran-linked fluorescein solution similar to the method used by Soeller and Cannell. (B) Magnified view illustrates the membrane impermeable dextran-fluorescein entry into the t-tubules reporting local tubule geometry and volume against the dark background of the myoplasm. Note the oblique (asterisk) and longitudinal tubules (arrowheads) that are clearly visible with this approach. (C) The percentage distribution of local tubule diameter estimates from such diffraction-limited data (blue bars; Soeller & Cannell) has a similar mean to that based on dSTORM data. Note the limited measurements (poor detection) of tubule diameters that are <70 nm in the diffraction-limited analysis (arrow), while the dSTORM analysis is able to detect >2-times the fraction of tubules narrower than 70 nm. Scale bars: A: 20 µm, B: 5 µm.
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a clearer view of the tubule geometries. The main benefits of this approach are the superior resolution and the greatly minimised geometry dependent intensity variations in the transverse tubules (see discussion in²); therefore we suggest it should be used in preference over the longitudinal (“side-on”) method of imaging t-tubules and dyads in every muscle type that has a predominantly transverse layout for these structures. We used this method to quantify the local variations in the Na+/Ca²⁺ exchanger density in the t-

Table 1. Features of localisation microscopy techniques and their advantages and limitations towards t-tubule imaging in muscles

| Features | Advantages | Limitations |
|----------|------------|-------------|
| Nanometre resolution across sub-millimetre sized areas | >10 fold improvement over diffraction-limited fluorescence techniques Direct measurement of subcellular structures Minimises over-estimation of protein co-localisation due to optical blurring | Resolution is only limited by detectable light yield from a single event, size of the fluorescent label/complex and the local density of the fluorescent probes. |
| Can be performed in aqueous and non-aqueous sample environments | Compatible with correlative imaging with EM and other light microscopies Minimise shrinkage artefacts Customisable mounting medium to optimise photoswitching | Abbreviating the fluorophore ON time further than a few milliseconds reduces photon yield, compromising the localisation accuracy. Prolonging the ON and/or OFF time prolongs the image acquisition time and the image file size. The latter may limit the speed of real-time image reconstruction. Potentially many parameters and variables to chose from; may be confusing to non-specialist users |
| Fluorophore photoswitching kinetics are adjustable | Can be modulated by excitation intensity, chemical constituents in the mounting buffer and/or a secondary activation laser. Optimal photoswitching rate can be achievable depending on the densities of the underlying fluorophores in the samples. Ability to improve localisation accuracy by avoiding simultaneous photoswitching of overlapping fluorophores or suppressing background/out of focus molecules | |
| Image is generated based on localised map of fluorophore position and localisation error | Images are (in principle) quantifiable for in situ biochemical analysis Minimises geometry-dependent non-linearities in fluorescence intensity in diffraction-limited imaging Robust clustering or image binerisation methods based on local event densities | Quantifiability of image data hinges on accurate localisation of overlapping events and correcting for local fluorophore density dependence of repeated photoswitching or limited yields of the molecules. |
| Single molecule events consist of a unique spectral signature | Spectral filtering can ‘unmix’ between multiple fluorophores or autofluorescence to achieve greater specificity in visualising the target structure. | |
| Reconstructed greyscale images are “background free” | More robust binarisation of images Local intensity is less dependent on gradients/patterns in the background compared to standard fluorescence microscopies | Any background fluorescence limits localisation accuracy (hence, resolution) Non-specific binding of fluorescent probes or antibodies are indistinguishable from specific binding; hence requiring appropriate control experiments (an issue common to all marker binding methods) |
| Ability to resolve structures in “thick” samples | Compatible with immunocytochemistry and tissue samples sectioned with standard microtomes (typically 10-20 microns in thickness) | Light scattering, spherical aberration and out of focus fluorescence typically seen in thick samples diminish the localisation accuracy. Therefore, thin samples are ideal (e.g. cultured cells). The native t-tubule structure is poorly preserved in cultured myocytes or myoblasts. |
| 3D localisation | Suitable for resolving the three-dimensional complexities of t-tubules | Thin samples (<10 microns) with dense t-tubule labelling with low background fluorescence work best. |
| Photoswitchability of endogenous, genetically encoded or introduced fluorophores in aqueous physiological saline environments | Compatible with real-time imaging t-tubules in living/functional preparations Conducive to photo-tracking of labelled receptor/protein/lipid mobility in t-tubules | Photodamage to the cells/fibres from prolonged intense light exposure and oxidants generated from the photoswitching (but yet to be quantified if any worse than other imaging approaches) Spatial distortions from drift or local contractures of the cells are difficult to correct for. Visualisation of fast events (protein trafficking, t-tubule membrane remodelling) may be limited by fluorophore photobleaching and photoswitching rates. |
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Fig 3  Comparison of micron- and nanometre-scale morphologies in adult mammalian cardiac t-systems. Transverse (i) Confocal and (ii) dSTORM images blurred with a 2D Gaussian PSF equivalent to a confocal PSF (left panels) and super-resolution image (right). Shown myocytes of (A) C57BL/6 mouse and (B) Wistar rat were stained with a combination of NCX1 and CAV3. Fixed ventricular tissue sections from (C) New Zealand White rabbit, (D) Human (54-year old donor with normal echocardiogram) and (E) Horse (12-year old female New Forest pony) were stained with fluorescent wheat germ agglutinin to visualise the t-tubules. Note the nanometre- and micron-scale t-tubule dilatations in mouse and rabbit myocytes respectively (arrowheads). Cell surface in each example is indicated by asterisks. Scale bars: i-panels: 2 µm; ii: 1 µm.

tubules of rat myocytes and identified a small fraction of non-dyadic RyR clusters.43 This protocol also allowed us to trace the zigzag branchings of t-tubules in regions of misregistrations between adjacent myofibrils and measure the near-4-fold greater t-tubule density and branching in the rat t-system, which, similar to mouse, is highly tortuous and reticular (Fig 3 A&B) compared to the spoke-like t-tubules that project towards the cell centre in ventricular myocytes of larger mammals including human44 (Fig 3C-E). Intriguingly, extensive t-tubular networks have also been observed in atrial myocytes of larger mammals such as horse, sheep, cow and human.45 Analysis of transverse fluorescence images of t-tubules in these atrial samples has revealed a weak correlation between tubule density and cell size; however it is yet to be understood why a near-complete lack of tubules is seen in the atria of smaller animals (e.g. mouse, rat).45 Here it is noteworthy that one of the prime limitations in fluorescence microscopy of t-tubules in fixed samples (typically resorting to immunocytochemistry) is the lack of a gold-standard stain that reports the full extent of the t-system. Poor retention of fixable membrane stains in cardiac t-tubules fixed with formaldehyde-based fixatives and the incomplete staining patterns with common labels (e.g. caveolin-3 or wheat germ agglutinin) have led us to propose that cardiac t-tubules in small animals (e.g. rat and mouse) are often better visualised with the combined fluorescence of two or more independent stains.44

More recent advancements in t-tubule imaging have been achieved both with the newer optical super-resolution microscopy techniques and more recent EM volume imaging approaches. An example is tomographic EM that has enabled 3D reconstructions which have been instrumental in resolving the finer features of t-tubules. These include nanometre-scale reconstructions of the t-tubule membranes in dyad regions to visualise the contact areas of RyRs with transverse46,47 and longitudinal tubules.48 Serial block face scanning EM (SBF-SEM) more recently has produced 3D reconstructions of the t-system in cardiomyocytes from rat and sheep ventricles at nanometre resolution.49 These data volumes demonstrate the relative differences in the tubule widths between sheep and rat t-tubules and, for the first time, resolve local variations in tubule diameters that were previously not directly visualised in fluorescence micrographs due to the diffraction limit (although see our previous discussion of rat t-tubules,39 Sache et al.41 of rabbit; see arrowheads in Fig 3C-ii for example dSTORM image). Similar nanometre-scale dilatations were observed in the junctional segments of mouse ventricular t-tubules using tomographic EM and dSTORM data50 (arrowheads in Fig 3A-ii). More recent tomographic reconstructions suggest that these may be folds of the t-tubules that putatively improves the contact between RyR and the trigger.52 While no data are shown to clarify how consistently these folds are observed among all of the dyads or their role in EC-coupliong, these structures are apparently lost or
not formed in the absence of Bin-1, resulting increased arrhythmogenesis.\textsuperscript{51} Alternative methods for imaging the nanoscale ultrastructure of t-tubules have been based on other optical super-resolution methods. Wagner et al.\textsuperscript{52} used STED microscopy to resolve the fine details of t-tubules stained in live mouse ventricular myocytes with the lipophilic membrane dye di-8-ANEPPS. \textless 60 nm in-plane resolution provided them with high-contrast images of the tubule membranes (and hollow cores) that were relatively convenient to analyse with automated algorithms.

The skeletal muscle t-system, in comparison has classically been visualised in great detail in pioneering EM studies that used high-contrast membrane stains (e.g. Golgi stain).\textsuperscript{53} Characteristic of the mammalian skeletal muscle t-system are the doublets of tubules within each sarcomere (located at each A-band boundary; see Fig 4A), each with a flattened cross-sectional shape and flanked by two SR terminal cisterns in a structure classically known as a triad. In a recent study, we reconstructed the 3D structure of the rat fast-twitch fibre t-system using the calibrated confocal fluorescence of a membrane impermeable dye trapped within sealed t-tubules (Fig 4B).\textsuperscript{54} These reconstructions demonstrated a dense network of t-tubules lining the nuclear envelope, that were previously shown to bear junctions\textsuperscript{55} that may be important in excitation-transcription coupling. These 3D data volumes of mammalian muscle were also useful for estimating a mean tubule diameter of \textasciitilde 86 nm, which predicted a \textasciitilde 38 nm longitudinal width and a 132 nm transverse (flattened) on the assumption that these tubules had a cross-sectional aspect ratio of 3.5. These values were comparable with transverse widths of triads measured using dSTORM images of RyR1 labelling in rat EDL fibres.\textsuperscript{3} However, one major technical difficulty using the confocal imaging approach are the 3-4 times narrower diameters of the skeletal muscle t-tubules in comparison to the cardiac equivalent. As a result, the small tubular volumes and associated noise in confocal micrographs appeared to increase the uncertainty of detection (and correct calibration) of a small fraction of tubules that were on average narrower than 40 nm (in mean width; see Fig 4C). This limitation would be circumvented by using a super-resolution microscopy method as detailed above, which should overcome this size-dependent detection bias. We subsequently used dSTORM for the first time in a recent study to characterise the nano-architecture of the previously detected subsarcolemmal components\textsuperscript{48} of the mammalian skeletal muscle t-tubules that had been filled with fixable fluorescent dextrans.\textsuperscript{37} Facilitating the fast propagation of action potentials,\textsuperscript{57} buffering ion exchange between the external space and deeper t-tubules and/or communication with the tubular extensions of subsarcolemmal mitochondria (recently visualised with deconvolved confocal micrographs and focused ion beam SEM reconstructions\textsuperscript{58}) are all possible functional roles of these tubules. dSTORM images of deeper tubules in rat skeletal muscle have since emerged,\textsuperscript{59} although quantitative measurements are yet to be described.

Altered t-tubule structure in health and disease: motivating renewed interest in t-tubules

One reason why there has been a resurgent interest in the structural imaging of t-tubules, using most of the imaging modalities mentioned above, are observations of changes in the tubule structure in a range of

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**Fig 4.** Visualisation of t-tubules in mammalian skeletal muscle. (A) A longitudinal confocal micrograph of a “mechanically skinned” rat extensor digitorum longus (EDL) fibre with membrane impermeable dye trapped in the t-system. (B) A dSTORM image of t-tubule organisation in the flanks of a single z-line of a similar fibre. T-tubules were filled with fixable dextran and the fibre was skinned to trap the dye prior to fixation. (C) Shown is a percentage histogram of the mean diameter of t-tubules in adult rat EDL fibres calculated by modelling the tubules as flattened cylinders. Notice the left-skewed shape of the histogram, which is a result of the poor detection efficiency of tubules with a mean diameter narrower than \textasciitilde 40 nm. More tubules are expected to be detected with smaller diameters (expected fractions approximated by dashed lines) with the super-resolution method. Scale bars: A: 5 \(\mu\)m; B: 0.5 \(\mu\)m.
pathophysiologica conditions. Loss of t-tubule density and concurrently diminished Ca\(^{2+}\) signalling have been observed in a wide range of human \(^{6,62}\) and animal \(^{63-65}\) cases of heart failure. Fluorescence (confocal) microscopy studies have commonly reported abnormalities that include focal loss of tubules (detubulation), loss of transverse elements of the t-system, a concurrent proliferation of longitudinal tubules and/or a general loss in spatial regularity of the t-system in a range of cardiac pathologies. These include myocardial hypertrophy,\(^{56,67}\) hypertensive cardiomyopathy,\(^{53}\) myocardial infarction,\(^{52,64,66}\) tachycardia-induced dilated cardiomyopathies,\(^{59}\) idiopathic cardiomyopathies \(^{61,62}\) and types of long-QT syndrome\(^{70}\) (see more comprehensive reviews).\(^{71,72}\) Commonly-described t-tubule abnormalities in skeletal muscle include widening of triads \(^{73,74}\) and micron-scale dilatation (vacuolation).\(^{75,76}\) Such abnormalities have been observed in connection to pathologies such as human myotubular myopathies\(^ {73}\), myotonic muscular dystrophy,\(^{77}\) rippling muscle disease\(^{78}\) and a range of genetic myopathies (see review\(^ {79}\)). Vacuolation is often seen in non-pathological conditions as fatigue \(^ {80}\) or in muscle damage due to eccentric contractions.\(^ {80}\) Several recent studies have focused on the molecular mechanisms of t-tubule alterations; commonly implicated proteins include junctional proteins (JPH), bin-\(^ {1}\) l and teletholin. Such proteins either display altered expression or cleavage in correlation with the pathology. In the case of junctional protein, overexpression provided some resistance against t-tubule changes and knock-down led to alterations similar to those seen in failure.\(^ {65,66,81-83}\) While the altered role(s) of such proteins is becoming clearer, questions relating to cause and effect are less well resolved.

### A case for improving imaging methods

Study of the molecular machineries listed above and their dysfunction in muscle pathologies have indicated a tight relationship between the cellular scale topology of the t-system (as characterised, e.g. by the t-power\(^ {56}\) and other tubule orientation indices, e.g. as provided by AutoTT,\(^ {84}\) and other approaches\(^ {51,62}\)) and the nanoscale features of the t-system and associated structures (e.g. junctions). For example, overexpression of JPH could counter t-tubule alterations in failing myocytes but altered the nanoscale structure of dyads.\(^ {85}\) Mechanistically this relationship between the micron-scale and nanometre-scale features of t-tubules is still poorly understood. This has probably been aggravated by the fact that widely used confocal fluorescence methods are excellent at detecting micron scale alterations but nanoscale differences generally require higher resolution. Table 1 of the review by Guo \etal\(^ {71}\) provides a good summary of the wide use of diffraction-limited microscopy modalities to characterise the t-system alterations that occur in heart failure. We argue that a more complete understanding of the changed ultrastructure is only possible with an imaging approach that appropriately resolves also smaller scale changes. While this can be remedied to some extent by using EM techniques an approach that includes both scales (micron-scale and nano-scale) would be beneficial. Optical super-resolution methods and whole-cell tomographic EM techniques have already demonstrated their utility in fulfilling this need for a multi-scale analysis of t-tubule structure in such studies. One such example is a recent Stimulated Emission-Depletion (STED) microscopy study that demonstrated dilation of t-tubules in addition to the aforementioned re-arrangement in failing mouse ventricular myocytes.\(^ {53}\) While dilated t-tubules had been visualised in larger species with naturally larger t-tubules (e.g. human, canine\(^ {62,85,86}\), this was one of the first clear demonstrations of nanometre-scale “remodelling” of t-tubules in a small (murine) species and could be performed in intact myocytes. Whether it is approached with a single microscopy technique or a combination of microscopies, we propose that future studies examining t-tubule changes should generally address both these spatial scales.

With the benefits outlined in Table 1, SMLM is a modality that can be used for imaging the same fluorescent samples for observing the t-tubule nanoarchitecture and the distributions of accessory proteins that are implicated in the cardiac and skeletal muscle pathologies. Less demanding hardware\(^ {39}\) in contrast to STED (e.g.\(^ {55}\)), compatibility with generic fluorescence labelling protocols (e.g. standard immunofluorescence\(^ {51}\) and freely available image analysis and reconstruction software (e.g. rapidSTORM\(^ {26}\)) are key features of this technology that encourage muscle biologists to examine these samples with SMLM. It must be emphasised however that SMLM is not a replacement to other microscopy techniques such as EM or tomographic EM. Correlative or supplemented imaging SMLM samples with EM, whenever possible, promises additional information such as 3D nanostructure and relationship with other organelles as well as a means of validating the nano-scale measurements, e.g.\(^ {20}\) We also underscore the potential utility of SMLM and other optical super-resolution modes in supplementing (diffraction-limited) confocal fluorescence microscopy of calcium micro- and nanodomains (e.g. imaging Ca\(^{2+}\) sparks with intra-dyadic Ca\(^{2+}\) sensors\(^ {35}\)) to correlate functional measurements with the local ultrastructure.

### Conclusion

SMLM is emerging as a fluorescence microscopy technique capable of resolving nanometre-scale features of cardiac and skeletal muscle t-tubules that were previously restricted to EM. In a short period of application of this technique (<5 years) many new observations have been made to advance our
understanding of the membrane and protein organisation of t-tubules. Sample preparation (ensuring high labelling densities, thin samples and low background fluorescence) and accurate localisation of single fluorophore locations are pivotal for obtaining high quality super-resolution images from SMLM data. However, its compatibility with hydrated samples is ideal for examining the nanometre- to micron-scale alterations in the t-tubule structure in a range of pathologies. The potentially quantitative nature of the data also means that image-based measurements of tubule structure and protein densities are likely to be particularly useful for computational simulations of t-tubule related structure-function relationships.

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