An introduction to the methodology of expansion microscopy

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Abstract
Expansion microscopy is a novel, fluorescence imaging technique, which allows three-dimensional nanoscale imaging of specimens on a conventional fluorescence microscope. This is achieved through an innovative sample treatment, which culminates in approximately 4.5-fold expansion of specimens in each dimension. This allows 70nm lateral and 200nm axial resolution. To further develop application of the technique, there has been considerable focus on improving the methodology by i) extending the efficacy of labelling, ii) enabling multi-colour labelling of different biomolecules simultaneously, iii) further improving resolving power through alterations to sample preparation and iv) by combination of expansion microscopy with other well-established super resolution techniques. This review will highlight some of these recent advances and suggest ways that the technique could be developed further in the future.

Key words
Expansion microscopy, super resolution, imaging.

Key facts
1. Expansion Microscopy (ExM) allows three-dimensional nanoscale visualisation of specimens on a conventional fluorescence microscope.
2. By embedding specimens into a polyelectrolyte gel, specimens can be physically expanded 4.5x in each dimension, resulting in decrowding of nanoscale information.
3. Broader application of ExM has been enabled by use of commercially available chemical cross-linkers to anchor labels to the gel meshwork.
4. Use of trifunctional labelling approaches enables retention of fluorescence intensity post-expansion and labelling of a range of biomolecules e.g. lipids, proteins.
5. Higher resolutions can be achieved by increasing expansion factor of gels, or by combining ExM with other super resolution approaches.
1. Introduction

Expansion microscopy (ExM) is a fluorescence imaging technique which enables three-dimensional nanoscale imaging of specimens on a conventional fluorescence microscope. This method relies on embedding samples into a polyelectrolyte gel which can be swollen by the addition of water. This results in a physical increase in specimen size by 4.5x in each dimension (Chen et al., 2015). The net effect of this is decrowding of features within the specimen, meaning super resolution imaging can be performed on a conventional microscope, thus eliminating the need for sophisticated software and expensive hardware as exists with other super resolution microscopy approaches (Gao et al., 2017 & Chozinski et al., 2016). ExM allows for rapid image acquisition, and samples are optically cleared during the process. Clearing eliminates scattering of light through the sample, meaning imaging depth and volumes accessible are orders of magnitude higher than those achievable by other super resolution techniques. As a result, ExM is suited for rapid, scalable super resolution imaging in cells and thicker specimens such as tissue sections.

A schematic summarising the steps required for ExM preparation of samples is given (Figure 1) (Tillberg et al., 2016 & Chozinski et al., 2016). The specimen (e.g. cells, tissue) is labelled through immunofluorescence or genetically encoded fluorescent proteins (Fig 1A & B). Labels are chemically equipped with anchors to allow their incorporation into the network of a polyelectrolyte gel which is formed evenly and densely throughout the specimen (Fig 1C). The polymer-embedded samples are homogenised (proteins are digested) to enable isotropic expansion of the gel (Fig 1D). Addition of water results in the physical expansion of the gel, which now carries anchored labels describing the features of interest in the cell, by up to 4.5x in all three dimensions. This allows visualisation of labelled structures with an effective resolution of ∼70nm laterally, and ∼200nm axially (Gao et al., 2017) (Fig 1E). In this review, we give an introduction to the expansion microscopy technique, and highlight its inherent limitations and recent efforts to address these. We will also give a brief outlook on the scope and future applications of this spectacular new approach for super-resolution imaging.

2. Expansion Microscopy: Techniques and applications

2.1 Ensuring robust, isotropic expansion of specimens

A key consideration when applying ExM is determining that expansion is isotropic on the macro- and nanoscale. Isotropy of expansion is a measure of the uniformity of the expansion at the nanoscale, where anisotropy in the expansion leads to distortion in the expanded image. The expansion isotropy is dictated by two key molecular factors: formation of a dense, homogeneous polymer network through the specimen, and efficient homogenisation of the sample. Expansion isotropy can also be influenced by the mechanical handling of the sample (Vanheusden et al., 2020). In developing ExM, Boyden and co-workers developed a gel composition which provided optimal expansion and mechanical stability (Chen et al., 2015). A key feature of these gels is the meshwork size is estimated to be 1-2nm; well below the size of biomolecules, meaning that biomolecules are pulled apart from each other evenly whilst retaining their relative spatial organisation, down to a precision of 5-10nm for measurements on the micron-scale (Wassie et al., 2019). The gel meshwork is thought to be critical to retention of nanoscale information in ExM prepared specimens (Tillberg & Chen, 2019).

In each variant of ExM, distortions introduced during expansion have been characterised (Chen et al., 2015, Chozinski et al., 2016, Tillberg et al., 2016, Chen et al., 2016, Chang et al., 2017, Pernal et al., 2020). This has been commonly achieved through a correlative imaging approach where images of highly conserved structures (e.g. microtubules) are acquired pre-expansion with a super resolution
technique (e.g. structured illumination microscopy (SIM)) and compared to post-expansion images of the same features in the same cells. Using this approach, and after a single round of expansion, Chen et al. measured 1 – 4% errors in length measurements on the scale of tens to hundreds of microns, a level similar to that seen in other studies listed above. Isotropy of expansion has also been determined by imaging of the nuclear pore complex protein, Nup153, which has a highly conserved molecular configuration meaning that structural comparisons can be made pre- and post-expansion without the need to image the same cells (Pesce et al., 2019). ExM has now been meticulously validated across a range of cell and tissue types and even whole mouse organs (Wassie et al., 2019).

Accurate quantification of the isotropy of expansion is challenging due to the potential for inducing heterogeneities and distortions during the sample preparation (Pesce et al., 2019). Determining expansion factor is hampered by observations of ‘differential expansion’ (Pernal et al., 2020). This is the result of varying ionic conditions in samples, which impact differentially on the expansion factor of those samples. Using particularly challenging samples that had been fixed for pathology, Zhao et al report expansion factors between four- and five-fold, with an average expansion factor of 4.7-fold (Zhao et al., 2017). Pernal et al observe a similar effect across different samples with some wholesale variation in expansion factor, where the fixative (paraformaldehyde) concentration used prior to expansion is varied (Pernal et al., 2020). This renders absolute distance measurements challenging.

Recent work by Martinez et al has shown differential expansion of regions of the hydrogel that either carry or are devoid of cells. However, the same study also showed that nonscale expansion of the gel around the cell is isotropic and, indeed, quantitative measurements in expanded samples can be made using a protein standard (microtubule) to calibrate the expansion factor (Martinez et al., 2020). Works from Pesce et al additionally identified Nup153 as an intrinsic reporter of expansion factor, with an accuracy of 5-10nm. This was determined by combining ExM with STED for nanoscale precision.

Users of ExM, should therefore be conscious of the sources of anisotropy in their experimental setups and, where measurements of distance are being recorded, that protein standards, or comparison to other techniques are used to corroborate them.

2.2 Optimising labelling and anchoring in ExM

Since its introduction, numerous variations of the technique have been reported which broadly simplify the process, and extend the scope of the approach. Successful application of ExM requires a labelling strategy which allows labels to retain their relative spatial organisation through the subsequent digestion and expansion steps (Chen et al, 2015). Approaches to labelling and anchoring in ExM are summarised (Table 1). The choice of approach is dependent on sample type and biomolecules of interest, requiring careful design by users. Chen et al originally achieved retention of labels by using a trifunctional fluorescent probe comprised of an antibody-conjugated DNA oligomer bearing an anchoring moiety for incorporation into the gel and a fluorophore for visualisation. Several trifunctional probes bearing different fluorophores and oligomers were synthesised which allowed multi-colour imaging in cells and tissue sections. However, the requirement for a custom-made labelling probe limited wider application of ExM. This hurdle was overcome by use of commercially available cross-linking monomers as the anchoring moiety (e.g. glutaraldehyde) (Chozinski et al, 2016). The hypothesis was that by applying commercially available anchors following staining with conventional immunofluorescence, sufficient linkages would be formed between the labels and the polyelectrolyte gel to enable retention of fluorescence signal for
detection after expansion. The authors demonstrated good retention of fluorescence signal following digestion and expansion when applying glutaraldehyde or methacrylic acid N- hydroxysuccinimidy ester (MA-NHS) in cultured cells and tissues. Interestingly, fluorescence intensity following expansion was observed to be higher in cells treated with DNA-labelled antibodies than in those treated with glutaraldehyde or MA-NHS (90% compared to 70%). However, the authors conclude the antibodies can conjugate more fluorophore than the DNA oligonucleotide based probes, which mitigates this effect. The retention of antibody labels by simple crosslinking moieties was similarly demonstrated in the development of protein retention ExM (ProExM) using 6-((acryloyl)amino)hexanoic acid (AcX) (Tillberg et al., 2016). These approaches were also shown to be compatible with retention of genetically encoded fluorescent proteins following ExM. GFP and GFP-like proteins have a high stability to proteases (e.g. proteinase K), meaning their fluorescence is retained through digestion. A range of fluorescent proteins were characterised and most tested proteins retained >50% of their fluorescence intensity following ExM preparation. Retention of fluorescent signal from both antibodies and fluorescent proteins was demonstrated in cells and tissues in a reproducible manner using these approaches.

A general limitation of ExM is reduced contrast in images due to a number of compounding factors; 100-fold volumetric dilution of fluorescence signal during expansion (Wassie et al., 2019), loss of fluorescently labelled antibody fragments due to digestion (Wen et al., 2020 & Shi et al., 2019), and destruction of fluorophores due to free radical generation during gel polymerisation (Min et al., 2020) (Figure 2). Whilst the dilution of fluorescence signal due to expansion may seem unavoidable, several approaches have been developed to mitigate this, as well as the loss of labels due to digestion and damage of fluorophores during polymerisation.

An approach to overcome volumetric dilution of fluorescence and destruction of fluorophores during polymerisation is to enable post-expansion labelling of specimens. This was attempted in the development of ProExM in which the non-specific proteolytic digestion was replaced with milder digestion conditions (e.g. Endoproteinase LysC) to retain epitopes for post-expansion labelling (Tillberg et al., 2016). However, epitope preservation was variable and incomplete homogenisation resulted in anisotropic expansion of specimens. Post-expansion labelling has been demonstrated in magnified analysis of the proteome (MAP) and ultra-ExM (U-ExM) techniques (Ku et al., 2016 & Gambarotto et al., 2019). These approaches are based on combining tissue clearing methods and expansion, to retain endogenous proteins and allow for nanoscale imaging of specimens. Applying post-expansion labels following MAP and U-ExM preparation improves signal and contrast in images, but efficacy of labelling depends on antibody identity.

Whilst a portion of the antibody fragments are bound to the hydrogel through chemical anchors, the non-specific proteolytic digestion used for sample homogenisation results in fragmentation and subsequent loss of up to 50% of these fragments during the final expansion step. This issue has been addressed by direct grafting strategies (Wen et al., 2020), and label retention ExM (Shi et al., 2019). These approaches rely on use of trifunctional probes which are comprised of a targeting moiety, a reporter moiety and a polymerisation group. By conjugating the fluorophore directly to the anchoring moiety, any fluorescence loss during digestion and expansion is mitigated. Additionally, this simplifies the process as labelling and anchoring can be performed simultaneously. For example, Shi et al developed trifunctional probes to mitigate loss of fluorescence through antibody fragmentation and these probes were modified to enable labelling using enzymatic tags (e.g. SNAP-tag) in addition to conventional immunofluorescence labelling (Shi et al., 2019). The direct grafting
strategy has been exploited to use small molecules that target actin (e.g. phalloidin conjugates) or membrane lipids (e.g. trivalent fluorescent lipids carrying 1,2-Distearoyl-sn-glycero-3-phosphoethanolamine (DSPE)) (Wen et al., 2020). During ExM, lipid membranes are lost, but by applying a covalently-tethered lipid probe, the signal derived from the membrane is preserved post-expansion.

Many of the labelling approaches described focus on imaging of proteins post-expansion. It has also been demonstrated that ExM approaches are compatible with imaging nucleic acids (Chen et al., 2016). This was demonstrated in the development of expansion fluorescence in situ hybridisation (ExFISH) in which they developed a small molecule linker, LabelX, to allow retention of RNA transcripts through ExM sample preparation. These transcripts were subsequently detected using FISH probes. It was also shown this small molecule linker allowed for expansion of the nuclear compartment. Therefore, by combining ExM methodologies, imaging of DNA, RNA and proteins within the same specimen can be achieved.

The polymerisation reaction is thought to lead to chemical damage of some fluorophores rendering them non-fluorescent, meaning fluorophore choice is crucial. Retention of fluorescence depends on the identity of the fluorophore with some (e.g. Alexa Fluor (AF) 647) being completely destroyed following gelation, whilst others (e.g. AF488) retain approximately 50% of their brightness post-expansion (Min et al., 2020). The best-performing fluorophores for ExM were identified as AF488, Tamra, Atto565 and Atto647N (Chen et al., 2015). This list has since been further extended to include AF405, AF546, AF568 and GFP (Chozinski et al., 2018). In addition, Min et al. methodically assessed brightness of cyanine-based (CF) and Alexa Fluor dyes to achieve four colour imaging with maximal signal-to-noise ratio during standard ExM preparation. Fluorophores and fluorescent proteins characterised for use in ExM are summarised (Table 2). The values in this table are based on the ability of fluorescent labels to survive the polymerisation step required for ExM preparation. Labels may also be applied post-gelation allowing for staining with labels which would not survive the gelation process (Chozinski et al., 2016).

These approaches to ExM have extended the range of labelling strategies and fluorophores compatible with the method, and have largely overcome limitations surrounding fluorescence intensity following expansion.

2.3 Improving resolution of ExM

ExM protocols result in a 4.5-fold expansion of specimens in each dimension, offering a resolution of 4.5-fold smaller than the diffraction limit (250-300nm/4.5 = 55-66nm lateral resolution). Two approaches have been developed which further increase expansion factor and therefore the effective resolution: iterative ExM (iExM) (Chang et al., 2017) and X10 ExM (Truckenbrodt et al., 2018). iExM enables 25nm resolution, facilitated by performing a preliminary gelation and expansion, followed by formation of a second polymer mesh in the spaces generated by the first expansion. All information is transferred from the first to the second gel by chemical means (Wassie et al., 2019). Expansion is then performed on the second gel resulting in a 4.5 x 4.5, or 20x physical expansion. X10 ExM similarly achieves 25-30nm resolution and this is achieved by modifying the gel composition to incorporate N,N-dimethylacrylamide (DMAA) and sodium acrylate. DMAA is a self-crosslinking monomer meaning the gel can be formed without additional multifunctional monomers. These gels have excellent mechanical stability and are superabsorbent, absorbing up to 3000x their weight in water (Cipriano et al., 2014). More uniform distribution of crosslinks in the gel networks reduced local stress and distortions during the expansion process.
ExM can be successfully combined with other super resolution techniques including structured illumination microscopy (SIM) (Halpern et al., 2019), stimulated emission depletion (STED) microscopy (Gao et al., 2018) and stochastic optical reconstruction microscopy (STORM) (Xu et al., 2019). These combined approaches result in lateral resolutions of ~30nm, ~10nm and ~10-20nm, respectively. Such resolutions match or exceed the resolving power of existing imaging techniques when applied individually. When combining ExM with these other techniques, care must be taken to minimise trade-offs in the imaging experiment. In all cases, imaging depth is increased 4.5-fold, necessitating use of water-based instead of oil-based immersion objectives to prevent spherical aberrations. Sample preparation and data acquisition parameters may need to be altered to mitigate lower fluorescence intensity and labelling density caused by ExM and for each combined imaging experiment, specific parameters may need optimisation. For example, ExM gels must be completely immobilised to prevent severe reconstruction artefacts derived from gel movement when combining ExM with SIM. When applying STED to ExM prepared samples, high fidelity dense labelling of epitopes is critical to achieving the best resolution.

3. Conclusions and future directions

Expansion microscopy is a unique approach to super resolution imaging, enabling rapid and easy nanoscale imaging of specimens in 3D. The nature of the preparation offers several technical advantages. Samples are optically clear which reduces the effects of diffraction and scatter (Gao et al., 2017). This enables greater imaging depth with minimal introduction of optical aberrations. Multi-colour applications of ExM are possible with minimal constraints on fluorophore choice (Min et al., 2020), and recent innovation of this technique has extended the range of biomolecules and labelling approaches compatible with it (Wen et al., 2020 & Shi et al., 2020). However, with all of these advances, the key caveat of ExM i.e. its incompatibility with live cell imaging, remains.

At present, ExM allows resolutions comparable to the best-performing super resolution techniques. It has been suggested that higher expansion factors and therefore resolution may be achieved and would supersede other super resolution techniques (Chang et al., 2017 & Truckenbrodt et al., 2018). It has also been demonstrated that ExM approaches may be combined (e.g. ExFISH and ProExM) (Chen et al., 2016). If a unified protocol could be developed, imaging of DNA, RNA, proteins and lipids may be combined to reveal organisation of heterogeneous complexes. The aqueous nature of specimens and the decrowing effect of ExM has been postulated to allow multiplexed readout of molecular information with nanoscale precision. This has been demonstrated using fluorescence in situ hybridisation approaches, and may be extended to being compatible with DNA-PAINT style probes (Wassie et al., 2019). In theory, any biomolecule could be labelled with an oligonucleotide barcode which can be identified post-expansion meaning that nanoscale mapping of biomolecules in a highly multiplexed fashion may be possible.

References


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Figure Legends

Figure 1: Schematic overview of the expansion microscopy method.

A) Cells are fixed and permeabilised and (B) immunolabelled using antibodies, tethered to a DNA oligonucleotide functionalised with a fluorophore (red star) and an anchoring acroloyl group (anchor). C) The labelled sample is infused with acrylamide monomers, sodium acrylate and bis-acrylamide crosslinking reagent and the polyelectrolyte gel is formed in situ, around the fixed sample. The anchoring acryloyl groups are bound into the polymer network. D) The sample is treated with a non-specific protease, which removes the cell and antibodies, leaving the tethered fluorophores in place. E) The sample is immersed in water and expands by approximately 4.5-fold in one dimension, resulting in an enlarged copy of the labelled cell.

Figure 2: Schematic showing possible mechanisms for loss of fluorescent signal from a sample in ExM.

A) Heterogeneity inherent in the labelling of antibodies means only a fraction of labels are retained for imaging. B) Digestion can lead to cleavage of an antibody between the fluorophore and the anchoring group. C) The polymerisation reaction can lead to chemical damage of fluorophores.
Figures

Figure 1

A

B

C

D

E
Figure 2

A

Well-labelled  Unlabelled  Unanchored

B

Digestion

C

Polymerization
Table 1. Table summarising approaches to labelling and anchoring in ExM approaches. Advantages, limitations and validated uses of each approach are provided.

<table>
<thead>
<tr>
<th>Cross-linking/ labelling strategy</th>
<th>Advantages</th>
<th>Limitations</th>
<th>Uses</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glutaraldehyde / MA-NHS approach</td>
<td>Commercially available reagents which are cheap and no complex preparation required. Good retention of fluorescence following digestion and expansion. MA-NHS incorporation into gels highly efficient as structurally similar to the methacryloyl group used previous.</td>
<td>Glutaraldehyde preferred for cells and not tissues as induces higher background than MA-NHS. Incorporation of glutaraldehyde-anchored probes into the polyelectrolyte gel may be affected by glutaraldehyde becoming topologically entangled in the polymer network. Relies on sufficient linkages being formed between proteins and gel so could be variable.</td>
<td>Validated for use in tissues and cells for a range of fluorescent antibodies and fluorescent proteins.</td>
<td>Chozinski et al, 2016.</td>
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Table 2. A summary of fluorophores and fluorescent proteins assessed for use in ExM preparations is provided.

<table>
<thead>
<tr>
<th>Tested Fluorophores and fluorescent proteins</th>
<th>Retention of fluorescence following polymerisation</th>
<th>Method of determining fluorescence retention</th>
<th>Reference</th>
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<tbody>
<tr>
<td>AF488&lt;br&gt;Atto565&lt;br&gt;Atto 647N&lt;br&gt;AF647</td>
<td>57.2%&lt;br&gt;76.2%&lt;br&gt;58.5%&lt;br&gt;-</td>
<td>Fluorescence intensity compared pre-expansion and post-gelation with no expansion.</td>
<td>Chen et al, 2015.</td>
</tr>
<tr>
<td>AF405&lt;br&gt;Atto488&lt;br&gt;AF532&lt;br&gt;AF546&lt;br&gt;AF568&lt;br&gt;GFP&lt;br&gt;YFP&lt;br&gt;DsRed&lt;br&gt;Hoechst 33342&lt;br&gt;SYBR gold.</td>
<td>All fluorophores and fluorescent proteins listed reported to survive the gelation process.</td>
<td>Quantification is not reported for each individual fluorophore or fluorescent protein. Fluorescence retention assessed for antibody (Atto 488) in presence of glutaraldehyde and MA-NHS (&gt;60% retention). GFP assessed and reported &gt;80% retention.</td>
<td>Chozinski et al, 2016.</td>
</tr>
<tr>
<td>Dylight 405&lt;br&gt;CF405M&lt;br&gt;AF488&lt;br&gt;AF546&lt;br&gt;AF594&lt;br&gt;CF633&lt;br&gt;AF647&lt;br&gt;Atto647N&lt;br&gt;EBFP2&lt;br&gt;mTagBFP2&lt;br&gt;mTurquoise2&lt;br&gt;mCerulean3&lt;br&gt;ECFP&lt;br&gt;mTRP1&lt;br&gt;mEmerald&lt;br&gt;EGFP&lt;br&gt;mClover&lt;br&gt;YFP&lt;br&gt;mVenus&lt;br&gt;mCitrine&lt;br&gt;mOrange2&lt;br&gt;LSSmOrange&lt;br&gt;tdTomato&lt;br&gt;mRuby2&lt;br&gt;mCherry&lt;br&gt;mKate2&lt;br&gt;mCardinal</td>
<td>28±5%&lt;br&gt;51±4%&lt;br&gt;48±2%&lt;br&gt;68±3%&lt;br&gt;46±2%&lt;br&gt;51±10%&lt;br&gt;7±3%&lt;br&gt;55±2%&lt;br&gt;62±4%&lt;br&gt;65±9%&lt;br&gt;68±8%&lt;br&gt;69±4%&lt;br&gt;51±2%&lt;br&gt;70±7%&lt;br&gt;53±4%&lt;br&gt;65±5%&lt;br&gt;61±4%&lt;br&gt;64±7%&lt;br&gt;44±5%&lt;br&gt;54±7%&lt;br&gt;32±2%&lt;br&gt;42±3%&lt;br&gt;67±4%&lt;br&gt;90±7%&lt;br&gt;72±3%&lt;br&gt;37±3%&lt;br&gt;36±3%</td>
<td>% fluorescence intensity compared between live cells and those which have undergone gelation and digestion steps of ExM protocol.</td>
<td>Tillberg et al, 2016.</td>
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<tr>
<td><strong>IRFP</strong></td>
<td><strong>14±1%</strong></td>
<td><strong>CF405S</strong> yielded highest fluorescence*</td>
<td><strong>CF405S</strong>**</td>
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<tr>
<td>CF405S, CF405M, CF430, CF440</td>
<td>CF488A and CF514 yielded highest fluorescence*. CF488 higher than CF514.</td>
<td>CF488A and CF514**. CF488A yielded highest fluorescence retention.</td>
<td>CF488A and CF514**. CF488A yielded highest fluorescence retention.</td>
</tr>
<tr>
<td>CF450, CF488A, CF514, CF532</td>
<td>CF568 and CF594 yielded highest fluorescence*. CF568 higher than CF594.</td>
<td>CF568 and CF594**. CF568 yielded highest fluorescence retention.</td>
<td>CF568 and CF594**. CF568 yielded highest fluorescence retention.</td>
</tr>
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</table>

1. **AF = Alexa Fluor**
2. * Fluorescence intensity quantified pre-expansion
3. ** Relative fluorescence retention calculated by comparing post-expansion specimens with pre-expansion