Combining FISH with localisation microscopy: Super-resolution imaging of nuclear genome nanostructures
Combining FISH with localisation microscopy: Super-resolution imaging of nuclear genome nanostructures

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Abstract The optical resolution of conventional far field fluorescence light microscopy is restricted to about 200 nm laterally and 600 nm axially and has been thought for many decades to be an insurmountable barrier for the quantitative spatial analysis of cellular and hence also nuclear constituents. Novel approaches in light microscopy have now overcome this barrier. Here, we report on a special method of localisation microscopy, spectral precision distance/position determination microscopy and its combination with fluorescence in situ hybridization to analyse the spatial distribution of specific DNA sequences in human cell nuclei at the macromolecular optical resolution level. As an example, repetitive DNA sequence DYZ2 located within the heterochromatin region on human chromosome Yq12 was labelled with clone pHY2.1. Between 300 and 700 single-probe molecules were resolved in individual chromatin domains, corresponding to a detected molecule density around 500/μm², i.e., many times higher than resolvable by conventional fluorescence microscopy. A mean localisation accuracy of about 20 nm indicated a mean optical resolution in the 50 nm range. Beyond new perspectives for light microscopic studies of specific chromatin nanostructures, this may open a new avenue towards the general analysis of copy number of specific DNA sequences in small regions of individual interphase nuclei.

Keywords FISH · Chromatin nanostructure · Optical resolution · Super-resolution · Localisation microscopy · Spectral precision distance microscopy (SPDM)

Abbreviations

DAPI 4′,6-Diamidino-2-phenylindol
DNA Deoxyribonucleic acid
FISH Fluorescence in situ hybridisation
FCS Foetal calf serum
FWHM Full width at half maximum
NA Numerical aperture
PBS Phosphate-buffered saline
PSF Point spread function
SPDM Spectral precision distance/position determination microscopy
STED Stimulated emission depletion
Introduction

In recent years, nuclear genome structure has emerged as a central theme of cell biology (for reviews, see Lamond and Earnshaw 1998; Cremer and Cremer 1998; Egner et al. 2002; Betzig et al. 1998; Brien et al. 2003; Misteli 2005, 2007; Rouquette et al. 2010). To study chromatin nanostructure, methods of ultrastructure analysis using ionising radiation, in particular electron microscopy, have proven to be a valuable tool for imaging beyond the conventional optical resolution of light microscopy, until recently thought to pose an absolute limit of about half a wavelength or 200 nm in the object plane and around 600 nm in the direction of the optical axis (Abbe 1873; Rayleigh 1896). In the recent past, however, a variety of laser-optical far-field microscopy techniques based on fluorescence excitation has been developed to overcome the ‘Abbe-limit’ of 200 nm.

Some well-known methods are confocal 4Pi-Laser Scanning Microscopy (Cremer and Cremer 1978; Hell et al. 1994, 2003; Hänninen et al. 1995; Egner et al. 2002; Bewersdorf et al. 2006; Baddeley et al. 2006; Lang et al. 2010), structured/patterned illumination microscopy (Heintzmann and Cremer 1999; Gustafsson 2000; Baddeley et al. 2007; Schemmelleh et al. 2008), STED microscopy (Hell and Wichmann 1994; Schrader et al. 1995; Hell 2007; Schmidt et al. 2008) or localisation microscopy techniques using far-field fluorescence microscopy (Cremer et al. 1996, 2010; Bornfleth et al. 1998; van Oijen et al. 1998; Edelmann et al. 1999, Edelmann and Cremer 2000; Esa et al. 2000, 2001). Using these techniques, an effective optical resolution in the 10–20 nm regime has been obtained. For the first time, this wavelength used) and ca. 50 nm axial effective resolution (Esa et al. 2000): Sites with a 3D distance of about 50 nm were still discriminated; their (xyz) positions were determined independently from each other with an error of a few tens of nanometers, including the correction for optical aberrations.

Compared with related ideas (Burns et al. 1985; Betzig 1995), in the SPDM approach, the focus of application was (1) to perform super-resolution analysis by the direct evaluation of the positions of the spectrally separated objects; (2) to adapt the method to far-field fluorescence microscopy at temperatures in the 300 K range; (3) to specify spectral signatures to include all kinds of fluorescent emission parameters suitable, from absorption/emission spectra to fluorescence life times (Heilemann et al. 2002) or any other method allowing ‘optical isolation’ (Cremer et al. 2001).

In the last few years, approaches based on principles of SPDM and related methods have been considerably improved by several groups, especially by using photoswitchable fluorochromes. By imaging fluorescent bursts of single molecules after light activation with a first wavelength, using a second excitation wavelength the position of the molecules was determined with a precision significantly better than the full width at half maximum (FWHM, a parameter describing the width of a peak in a curve, in this case the microscopic Point Spread Function (PSF). The PSF is the normalised intensity distribution of a fluorescent point source; the FWHM of the...
PSF is a measure for the optical resolution at a given spectral signature. These microscopic techniques were termed photo-activated localisation microscopy (PALM) (Betzig et al. 2006), fluorescence PALM (Hess et al. 2006), stochastic optical reconstruction microscopy (Rust et al. 2006) or PALM with independently running acquisition (Geisler et al. 2007, Andresen et al. 2008). All these approaches use special fluorochromes which can be photo-switched between two different spectral states. Recently, we applied SPDM in the nanometer resolution scale to conventional fluorochromes that show reversible photobleaching, which is a general behaviour of several fluorescent proteins, e.g., CFP, Citrine or eYFP (Patterson and Lippincott-Schwartz 2002; Sinnecker et al. 2005; Hendrix et al. 2008). In such cases, the fluorescence emission of certain types can be described by assuming three different states of the molecule: A fluorescent state $M_{fl}$, a reversibly bleached state $M_{rbl}$ and an irreversibly bleached state $M_{ibl}$. By illumination, some of the molecules are switched to $M_{rbl}$ or to $M_{ibl}$. The stochastical transfer of the molecules from $M_{rbl}$ to $M_{fl}$ and then to $M_{ibl}$ can be used for optical isolation of the detected molecules and thus their positions can be determined. With velocity constants $k_1$, $k_2$ and $k_3$, the transition scheme can be assumed as follows

\[ M_{rbl} \xrightarrow{k_1/k_i} M_{fl} \xrightarrow{k_2} M_{ibl} \]

We demonstrated recently that reversible photo-bleaching can be used for super-resolution imaging (SPDM) of cellular nanostructures labelled with conventional fluorochromes such as Alexa dyes (Reymann et al. 2008; Baddeley et al. 2009) or fluorescent proteins (Lemmer et al. 2008, 2009; Gunkel et al. 2009). This effect may be obtained by appropriate physical conditions, such as an exciting illumination intensity in the 10 kW/cm² to several 100 kW/cm² range; only one exciting wavelength is necessary both for activation and read out of a given molecule type. Under these conditions, a large number of images or frames is taken rapidly one after the other (typical frame rates around 20 fr/s or more, up to a total of 1,000 to 4,000 frames); in each frame, under appropriate conditions the distance between the diffraction images of individual fluorescent molecules is typically larger than the ‘Abbe limit’, i.e., larger than the FWHM of the PSF of the optical system used. Recently, a related technique (Fölling et al. 2008) has been described for several other dyes like Atto532, Citrine or PhiYFP called ground state depletion microscopy followed by individual molecule return.

In the present study, we combined SPDM with fluorescence in situ hybridization (FISH)-labelling to quantitatively analyse the spatial distribution of specifically FISH-labelled DNA sequences in small regions of human cell nuclei. Heterochromatin region Yq12 was labelled with a cloned DYZ2 fragment (pHY2.1, Cooke et al. 1982) in human fibroblast nuclei. Repetitive DNA sequence DYZ2 is a major component of the heterochromatic region on human Yq12 and comprises approximately 20% of the Y chromosomal DNA (Cooke 1976). Arranged in tandem arrays, the 2.47 kb repeat is distributed in about 2,000 copies over the entire length of Yq12 (Schmid et al. 1990).

**Materials and methods**

**Cell culture and specimen preparation**

Human diploid fibroblast cells VH7 (kindly provided by Prof. Dr. Beauchamp from DKFZ, Heidelberg) were routinely cultivated in Dulbecco’s Modified Eagle medium supplemented with 10% FCS and penicillin/streptomycin at 37°C in a humidified CO₂ incubator. For specimen preparation, cells were seeded onto cover slips, grown until 70–80% confluency and subsequently fixed with 4% formaldehyde in PBS. Permeabilisation steps included treatment in 0.5% Triton-X in PBS, 20% glycerol in PBS, repeated freezing–thawing in liquid nitrogen and incubation in 0.1 N HCl, according to standard protocols for the preparation of 3D preserved nuclei (Bridger and Lichter 1999; Solovei et al. 2002a). Cover slips were stored in 50% formaldehyde in 2×SSC (pH 7) at 4°C until usage.

**Probe preparation**

Clone pHY2.1 (Cooke et al. 1982), composed of the 2.1 kb HaeIII fragment inserted into plasmid pBR328 (a kind gift from Dr. G. Rappold, Institute of Human Genetics, University of Heidelberg) was labelled by standard nick translation using AlexaFluor®568
(Invitrogen, Germany) as fluorescent dye. Labelled DNA was co-precipitated with salmon sperm DNA serving as carrier DNA and resuspended in hybridisation buffer consisting of 50% deionised formamide, 10% dextran sulphate in 4×SSC.

**FISH and immunocytochemistry**

Probe DNA was denatured at 78°C for 7 min. Cellular DNA was denatured in 70% formamide in 2×SSC (pH 7) at 72°C for 2 min. Subsequently, the denatured probe was placed onto a cleaned slide and the cover slips carrying the cells were lifted onto it. To prevent drying, cover slips were sealed with rubber cement. Hybridisation was performed in a humidified chamber at 37°C for 2 days. Post-hybridisation washings were done in 4×SSC/Tween20 (0.2%) at 42°C and 1×SSC at 60°C. For the analysis of the cell cycle, a mouse monoclonal antibody against nuclear protein pKi67 (Dako, Denmark) was applied. pKi67 displays differential patterns according to cell cycle stage (Gerdes et al. 1983). Detection was done with an Alexa Fluor®488-conjugated goat anti-mouse antibody (Invitrogen, Germany). Cells were mounted in Pro-Long® Gold antifade reagent (Invitrogen, Germany) and sealed with nail polish.

**Data registration**

For localisation microscopy, a microscope system was used allowing spatially modulated illumination with three laser lines (Baddeley et al. 2007). For the measurements reported here, this instrument was used in the single objective lens mode. A schematic of the microscopy setup is shown in Fig. 2.

The setup is equipped with three laser sources providing laser light at \( \lambda = 488 \text{ nm}, \lambda = 568 \text{ nm} \) and \( \lambda = 647 \text{ nm} \) (Lexel Laser, USA). Lenses L2 and L3 (Linos Photonics, Germany) expand the laser beam to a diameter of approximately 20 mm. A 50:50 beam splitter (Edmund Optics, Germany) evenly distributes the beams onto the two interferometer arms. The left optical path provides wide-field illumination: The beam is directed to the objective (O1), passing through an achromatic lens (L4) and a mirror (S2), resulting in collimated illumination of the sample, positioned between the two objectives. Currently in use are 100× oil objectives with an NA of 1.4 (Leica, Germany). Movement of the sample is provided by a 3-axis stepper motor stage and a 2-axis piezo stage (Physik Instrumente, Germany). The right optical path is utilised for localisation microscopy: In addition to an achromatic lens (L5) focusing the beam into the back focal plane of the objective (O2), a second lens (L3) provides the higher laser illumination intensities needed for this type of localisation microscopy. For the illumination of dyes, intensities were in the order of 10 kW/cm². The emission light from the sample is collected by the right hand objective lens (O2), passes through the dichroic mirror D4 (AHF, Germany) reflecting the laser lines, and is focused by a ×1 tube lens (Leica, Germany) onto the CCD chip of a highly sensitive camera (PCO Imaging, Germany). Remaining laser light and out of band fluorescence are blocked by blocking filter BF (AHF, Germany). A typical time series encompassing 1,000 2D images (frames) was acquired with a repetition rate between 10 and 16 Hz.

**Data evaluation**

In localisation microscopy, the essential concept to surpass the resolution limit of conventional light microscopy is the optical isolation of single fluorescent molecules by using appropriate spectral signatures, which may include photo-physical or photo-chemical properties of the fluorescent dye molecules (e.g., photoswitching or photoconversion). This means that at any given time, the distance between two molecules of the same spectral signature has to be larger than the conventional optical resolution (FWHM of the PSF). In conventional microscopy, on the contrary, the image of the fluorescence-labelled object emerges from the simultaneous detection of all fluorescent molecules, i.e., also from molecules with a distance smaller than the conventional resolution: in this case, due to diffraction, the object appears blurred and the resulting loss of structural information cannot be reversed. In localisation microscopy, however, a time series of only a few single molecules per image (i.e., with distances larger than the conventional resolution) is acquired. The result is a large number of single images or frames of sparsely distributed molecules with non-overlapping diffraction patterns. This allows to determine the positions of the individual fluorophores with a precision mainly depending on the detected photon numbers (Bornfleth et al. 1998; Edelmann and Cremer 2000; Thompson et al. 2002). All positions obtained from the
individual frames are assigned to the same ‘localisation map’. Thus, an image of the measured object (positions of all molecules detected) can be reconstructed that is not limited by diffraction.

The individual steps for the position determination of a single fluorophore are as follows:

1. Segmentation of the signal within the wide-field data
2. Fitting of an appropriate model function to the data to determine the individual molecule positions.

For the detection and segmentation of the signals, established algorithms such as CLEAN (Högbom 1974) can be used.

As a model function \( f(x, y) \) for the PSF, a 2D gaussian is approximated using a linear estimate of the background noise. Thus, the model function is

\[
f(x, y) = p_1 \exp \left( -\frac{(x - p_2)^2 + (y - p_3)^2}{2p_4^2} \right) + p_5 + p_6(x - p_2) + p_7(x - p_3)
\]

where \( p_1 \) to \( p_7 \) are, respectively, the signal amplitude, the \( x \) and \( y \) coordinates of the barycenter (gravity center of fluorescence intensity), the standard deviation as a measure of the spread of the signal, the constant contribution of background noise and the

![Fig. 1 Principle of SPDM. a, d Three point-like objects are located in the xy plane within 50 nm distance to the middle one. They are labelled with the same spectral signature in (a), or with three different, unique spectral signatures in (d). The computed “virtual microscopy” responses of the objects in (a) and (d) (parameters for computation are NA=1.4 and system magnification of ×63) are shown in (b), (e) and (f). In (f), the different spectral components are imaged together whereas in (e), the same signals are recorded independently from each other. Normalized linescans through the objects in (b) and (e) are shown in (c) and (g). Figures from (Kaufmann et al. 2009) adapted from (Cremer et al. 1999), based on the SPDM concept of (Cremer et al. 1996)]
linear approximation of the background signal in the $x$ and $y$ directions. By using this model function with appropriate parameters, signals above and below the object plane are excluded, i.e., an ‘optical section’ of about 600 nm thickness is obtained. For further details on the localisation algorithm, please refer to Kaufmann et al. (2009) and Lemmer et al. (2009). The algorithms used for data evaluation of the localisation microscopy images were implemented in MATLAB (7.0.1, The MathWorks, Natick, USA). In addition, measurements on the segmented data were performed using the DipImage toolkit (http://www.ph.tn.tudelft.nl/DIPlib/).

The simulation in Fig. 3 illustrates the basic concept of localisation microscopy with emphasis on the relationship between the localisation accuracy and the number of detected photons.

**Fig. 2** Experimental setup used for localisation microscopy. $D1$–$D4$ dichroic elements, $L$ lasers, $L1$–$L6$ lenses or lens systems, $B1$–$B4$ field diaphragm, $W$ beam splitter, $O1$–$O2$ objective lenses, $PS$ piezo translation stage, $PO$ piezoelectrical stage, $S2$ mirror, $BF$ blocking filter, $P$ periscope, $K$ CCD camera. The appropriate laser intensities required are gained by adjusting lens L3. For a detailed description, see text

**Fig. 3** Virtual localisation microscopy (computer simulation). *Upper panel:* a simulated structure, b simulated wide-field image, c interpolated image of (b). *Middle panel:* Simulated wide-field images of a small number of individual single molecule signals differing in detected photon number; the level of background noise remains constant: In (d), the average number of detected photons/molecule assumed was $N_{\text{phot}}=15$, in (e) $N_{\text{phot}}=100$ and in (f) $N_{\text{phot}}=1,500$. *Lower panel:* Virtual localisation microscopy of the simulated structure given in (a) with varying localisation accuracies (standard deviations): In (g), it was $\sigma=50$ nm, in (h) $\sigma=20$ nm and in (i) $\sigma=10$ nm. The size of the scale bars is 250 nm. Note that the effective pixel size in the individual images differs. For the simulated wide-field images in (b) and (d–f) it is 100 nm, whereas in (a, c) and (g–i), it is 5 nm.
In Fig. 3(a), the simulated structure and the positions of the individual fluorophores are shown. The total number of fluorescent molecules amounts to 9,289, and the area covered by the object is 0.77 \( \mu \text{m}^2 \), resulting in a fluorophore density of \( \frac{12,060}{\mu \text{m}^2} \). Figure 3(b) shows the blurred diffraction limited image as simulated for wide-field microscopy. The accompanying interpolated image is depicted in Fig. 3(c). The wide-field images were simulated assuming a wavelength of 500 nm, an NA
of 1.46 and 150 detected photons on average per fluorophore. This choice of parameters almost resembles ideal conditions and admittedly is rather optimistic. For the convolution of the simulated structure (a), the Airy disc was utilised as intensity PSF. The Airy disc is the image of an infinitely small fluorescent object in the focal plane of a perfect optical system (Rayleigh 1896). Its rotational symmetric intensity distribution follows from the equation

\[ I(r, \lambda) = I_0 \left( \frac{2J_1(2\pi NAr/\lambda)}{2\pi NAr/\lambda} \right)^2 \]

where \( J_1 \) is the first order Bessel function and \( r \) is the distance to the centre in nm.

The middle panel of Fig. 3 shows the effect of detected photon numbers per signal on localisation accuracy, assuming equal background levels in each case. For the PSF, the function described above was used. In (d), the photon number \( N_{\text{phot}} \) of the fluorescence signal of active molecules averages to 15, in (e) to 100 and in (f) to 1,500 photons. For each molecule, a variation in detected photon number, expressed as standard deviation of a normal distribution, of 30% of the mean was assumed. In (d), it is not possible to differentiate between background and signal. In (e), signals become recognisable but the precise number of molecules cannot be determined. In (f), the number and the position of the individual signals can be readily identified.

The effect of localisation accuracy on the attainable resolution is illustrated in the lower panel. For clarity, the corresponding images of the middle and lower row are simulated with consistent average photon numbers, i.e., in (g), it averages to 15, in (i) to 100 and in (h) to 1,500 photons. In (g), a localisation accuracy of 50 nm along with the detection of a few tens of photons provided no additional structural information about the object, when compared to the conventional microscope image shown in (b). By the
detection of around 100 photons/molecule and a resulting localisation accuracy of 20 nm, substructures become distinguishable (h) which are certainly not observable in the conventional wide-field image. The detection of more than 1,000 photons (i) is sufficient for a localisation accuracy better than 10 nm. In combination with an adequate number of detected molecules, this allows a good reconstruction of the given nanostructure.

One of the fundamental quantitative parameters of a fluorescent chromatin nanostructure is its overall size (area). This can be done in different ways, e.g. by adapting appropriate model functions (Baddeley et al. 2010) or by interpolation. Figure 4 illustrates the impact of an interpolation procedure on the result of the segmentation process. Both raw data of an experimental wide-field image of a FISH signal (4a) and the accompanying interpolated image data (4c), obtained by a standard procedure of image processing termed “lossless interpolation” were segmented with a conventional threshold based method. The same threshold value was used in both cases. As a result of the interpolation procedure, the number of pixel within the images increases and the effective pixel size decreases accordingly. For Fig. 4(a), the pixel size is 65 nm, and 13 nm for Fig. 4(c). Provided the image is Nyquist sampled (i.e., the pixel size is chosen in such a way, that the smallest possible structural component of the object is sampled at least 2-fold), the information of all intermediate values is available and can be used as additional landmarks in the segmentation procedure. The spaces between pixels can thus be filled while all morphological information is retained.

The area of the labelled DNA region (b, d) is extracted from the wide-field images (a, c) by an operation from the field of mathematical morphology termed morphological closing (Matheron 1975; Serra 1982). As structuring element, a circular disc in the plane was utilised. Basically, this results in the elimination of small holes and the filling of gaps in contours. In addition, a certain amount of smoothing on an object contour is effected.

Without interpolation, the area of the segmented region is calculated to $0.942 \mu m^2$ and with interpolation to $1.049 \mu m^2$. This means the area is 11.3% larger in the interpolated image. It is easily observable that the non-interpolated images (Fig.4 a, b) appear rather pixelated, whereas the images subjected to interpolation are smoother to some extent, and the edges are shown in far more detail, thus allowing the extraction of considerably more structural information. For this,

**Fig. 7** Representation of the data processing procedure for a typical experimentally obtained FISH signal of DNA probe pHY2.1. a Wide-field image at low magnification; b corresponding interpolated magnification; c reconstruction of the localisation microscopy (SPDM) data, blurred with a Gaussian corresponding to the mean localisation accuracy; d segmented image of the SPDM data in (e); e binary image of the area obtained by morphological closing of the SPDM image in (d); f overlay of the image of the segmented region as seen in (d), displayed as a “heat map” encoding the molecule density, and the wide-field image (gray scale) shown in (b).
However, a reliable discrimination of object and background and thus a good signal-to-noise ratio is important.

In Fig. 5, the line profiles, denoted in Fig. 4(c) are presented, indicating the excellent signal-to-noise ratio of the FISH-labelled chromatin structures.

Localisation microscopy, as every other fluorescence microscopical technique, is affected by non-specific labelling. For a correct quantitative evaluation of the chromatin nanostructure images obtained by SPDM, it is necessary to eliminate such non-specific signals from further consideration. For the investigation of the FISH-labelled regions, differences in the local fluorophore density were used to distinguish between specifically and non-specifically bound dye molecules. The employed automatic segmentation algorithm determined the number of neighbours in a defined vicinity for the individual molecules and used these numbers subsequently as “density threshold” values ($s$): The higher $s$, the lower the number ($n$) of remaining molecules. Figure 6 shows the principal approach: On the assumption that the fluorophore density is significantly higher within the labelled structure than in the surrounding area, the total number ($n$) of remaining molecules will decrease rapidly with increasing density threshold ($s$); this decrease will slow down considerably when all non-specifically bound molecules are eliminated (Fig. 6a). To obtain a quantitative, objective measure for the density threshold ($s_0$), derivative functions were applied. Figure 6(b) shows the first derivative, i.e., the alteration of the number of remaining molecules depending on the applied threshold value ($f'(n) = \frac{dn}{ds}$, where $n$ is the number of remaining molecules and $s$ the density threshold value). This curve first falls down rapidly to a minimum and then rises versus zero values (i.e. there $n$ changes only slightly with increasing density threshold values $s$). To determine the density threshold $s_0$ where the decline of ($n$) starts to slow down, the first derivative (Fig. 6b) was differentiated again mathematically to obtain the second derivative of the curve in Fig. 6(a). Accordingly, $f''(n) = \frac{d^2n}{ds^2}$ was calculated and plotted in Fig. 6(c). The density threshold value at the second zero crossing of the second derivative was then used as a measure for $s_0$. The entire process was established as an automatic algorithm and subsequently used for the segmentation of the localization microscopy images of the FISH-labelled chromatin regions.

Fig. 8 Selection of typical experimentally obtained FISH signals for DNA probe pHY2.1 and the corresponding processed data. Columns from I to IV: I wide-field images of the FISH signal; II segmented SPDM images; III binary images of the areas in (II); IV wide-field images of pKi67 indicating cell cycle phase and the nuclear boundary

Figure 7 outlines the entire data processing procedure as applied in this study using an experimentally obtained FISH signal. Prior to the localisation measurements a conventional wide-field image is obtained (Fig. 7a), providing an overview of the imaged region. The corresponding interpolated magnification is shown in Fig. 7(b). The reconstructed localisation image shown in Fig. 7(c) is subjected to the automatic segmentation procedure outlined above (Fig. 6); the result is presented in Fig. 7d. Morphological information of the labelled region is extracted from the binary image presented in Fig. 7e. Figure 7(f) depicts the good agreement of the image of the segmented region and the wide-field image. In this case, the segmented region is represented as a “heat map”: The colour encodes the density of the detected molecules; the wide-field image is displayed in gray scale.

Results

Localisation microscopy (SPDM) was performed on the heterochromatic region on Yq12 labelled with Alexa568 dye in 25 human diploid fibroblast cell nuclei and evaluated using appropriate algorithms as described above. As an indication of cell cycle phase, nuclear protein pKi67 was detected by Alexa488 coupled antibodies. In Fig. 8, two galleries of typical data resulting from a localisation microscopy experiment are shown. In each case, the wide-field image of the chromatin region, the segmented SPDM image, the binary image of the segmented SPDM image and the wide-field image of nuclear protein pKi67, indicating the cell cycle phase are shown. Both wide-field images were acquired at low illumination intensities before the localisation measurements. From the distribution pattern of pKi67 (Gerdes et al. 1983) nuclei in Fig. 8(a–f) and Fig. 8(g–i) were assigned to G1/S phase, whereas nuclei in Fig. 8(j–l) were assigned to G2.
The quantitative, automated evaluation procedures applied allowed the extraction of various structural parameters such as variations in size, shape, and signal number. For a description of the shape, we calculated for every FISH signal the area, extent and anisotropy. The anisotropy can be defined as the ratio between the spread along the major and minor axes. In Fig. 9, the histograms of the segmented areas of the chromatin region (a) and the number of single molecule signals within these areas are shown (b).

Both histograms display several maxima at various positions. The analysis of the area yielded values ranging from 0.50 to 1.65 μm² and a mean value of 0.86±0.28 μm². The number of signals was found to range from 281 to 713 with an average of 443±114. The comparison of FISH labelled area and cell cycle stage (as determined from the pKi67 pattern) indicates that all nuclei with areas smaller than 1.1 μm² are in G1/S phase.

The scatter diagrams in Fig. 10 depict the relationship between different structural parameters.

In Fig. 10(a), the number of detected signals was plotted against the area of the chromatin regions. The basic trend of the graph shows that the number of signals increases linearly with the area. Figure 10(b) displaying the anisotropy of the chromatin region reveals a strong linear relationship with a slope of 0.972. This suggests that on average the DNA regions do not show a shape deviating considerably from a sphere.

Another possibility to obtain structural information from the SPDM data is to utilise the position data directly. This eliminates the necessity to determine intensity thresholds as in conventional epifluorescence microscopy segmentation. As a measure for the extension of the FISH-labelled region, the spread of the molecule positions along each of the image axes is used. To quantify the spread, the standard deviation of the individual signal coordinates in the x and y directions was used. First, the position of the barycenter \( P(x, y) \) of the individual molecule positions of the segmented SPDM images was determined by

\[
\bar{x} = \frac{\sum x_i}{N} \quad \text{and} \quad \bar{y} = \frac{\sum y_i}{N}
\]

where \( N \) is the total number of signals. Then the spread of the molecule distances \((X,Y)\) to the respective barycenters was determined. The basic principle was adapted from the Multi point model for the evaluation of conventional confocal data (Baddeley et al. 2010).

As an example, Figure 11 (a, b) shows the histograms of the \( X \) and \( Y \) positions of the combined single molecule signals obtained for the FISH signal depicted in Fig. 4. As expected from a structure with many molecules positioned at different distances to the barycenter, a broad distribution was obtained.

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**Fig. 9** Distributions of **a** the segmented areas of the FISH signals and **b** the number of detected molecules obtained per FISH signal.
Whether the frequency variations indicate a specific substructure remains to be elucidated. The same data as in Fig. 11(a, b) are presented in Fig. 11(c, d) as quantile plots (i.e., accumulated frequency histograms). On the ordinate the distances \((X,Y)\) of the signal to the respective barycenter is given as a function of the part of the distribution; i.e., every value of the distribution is plotted against the proportion of all smaller values. From this, in the \(x\) direction, a medium value of 495 nm was obtained while in the \(y\) direction, the medium value was 405 nm. This suggests an anisotropy of 495/405 = 1.22.

In Fig. 12 (a, b), the distributions of the standard deviations in the \(x\) and \(y\) directions measured for the positions of all the molecules detected in the FISH-labelled regions for all 25 nuclei analysed are presented. In order to obtain a value that can be compared to other size measurements, the individual values can be converted to the FWHM of a Gaussian distribution by multiplication with a factor of 2.35.

In contrast to Fig. 9, displaying the distribution of segmented areas for all 25 nuclei, the distributions of the standard deviations in Fig. 12 appeared to be more homogeneous, suggesting a bimodal distribution of nuclei with standard deviations smaller than 300 nm and of nuclei with standard deviations larger than 300 nm. It may be noted that the nuclei with a FISH-labelled Yq12 area larger than 1.1 \(\mu m^2\) (Fig. 9) are represented by standard deviations larger than 325 nm in both directions.

**Discussion**

In a previous study, we have shown that localisation microscopy (SPDM) can be accomplished with conventional fluorochromes such as Alexa dyes or fluorescein derivatives (Reymann et al. 2008; Lemmer et al. 2009; Cremer et al. 2010). Besides their application in immunocytochemical staining methods, these dyes are widely used to specifically label chromatin structures by fluorescence in situ hybridisation (for review, see Cremer and Cremer 2001). Many of these structures, however, have a dimension too small for conventional light microscopical techniques. Examples for such structures in nuclear cell biology are small gene domains which may change their conformation due to differentiation or regulatory mechanisms. In this study, we have assessed and demonstrated the capability of the combination of fluorescence in situ hybridisation with localisation microscopy. Heterochromatin region Yq12 was labelled with a probe for DYZ2, a tandemly repeated sequence with a length of 2.47 kb, which is distributed in about 2,000 copies over the entire length of band Yq12. In addition, we described a data processing procedure that minimises the loss of structural information present in the original data and hence produces an accurate representation of the measured structure. Several structural parameters were extracted, describing heterochromatin region Yq12 in far more detail than can be gained by conventional microscopical techniques. In particular, the extent of this region was...
determined directly from the positions of individual hybridised molecules; the number of the molecules hybridised and detected was determined for each individual FISH-labelled region, despite a mean molecule distance much smaller than the conventional optical resolution $d_{\text{conv}}$ of approximately 200 nm: a density of 500 molecules/μm$^2$ (Fig. 10a) corresponds to a mean distance of about 45 nm, i.e. about five times smaller than $d_{\text{conv}}$. From the mean localisation accuracy $\sigma$ of about 20 nm, an average optical resolution of about 2.35 $\sigma$ corresponding to about 50 nm can be estimated. This means that so far the data do not allow drawing detailed conclusions on the substructure of the Yq12 domain. It is anticipated that this problem can be overcome by technical improvements of the SPDM microscopy technology presently in development (Pertsinidis et al. 2010). Another current limitation is the number of detected molecules: From a FISH labelled Yq12 region containing a total number of 2000 repeats, only about 300–700 signals are obtained. Some reasons for this may be the labelling efficiency of the probes, the inherent optical sectioning capacity of the technique: a ‘slice’ of about 600 nm thickness was analysed while the entire Yq12 domain had a diameter of about 1 μm; or the efficiency of detection of bound probe molecules. Despite these limitations, the method allows currently a large variety of applications where the determination of relative molecule numbers is sufficient.

This paper presents the first detailed description of the application of localisation microscopy (SPDM) for the nanostructural analysis of specific FISH-

**Fig. 11** Distribution of the $X$ and $Y$ positions of all single molecule signals obtained from the labelled region shown in Fig. 4. Figure 11 (a, b) present the frequency distribution while

**Fig. 11** (c, d) show the corresponding accumulated frequencies (quantiles). The distance is given in nm

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labelled chromatin domains. This approach may be combined with other SPDM techniques where fluorescent proteins were applied to study general features of nuclear nanostructure (Kaufmann et al. 2009; Gunkel et al. 2009; Bohn et al. 2010). For example, one could analyse specific domains of active/silenced genes or of gene deserts (Shopland et al. 2006) in the context of the histone distribution. Another interesting field of application is the analysis of break point regions induced e.g. by ionising radiation. In contrast to early attempts to use the SPDM methods (Esa et al. 2000, 2001) where only a very few spectral signatures were available, the present FISH-SPDM technique described here would allow positioning multiple molecules in a nuclear area as small as 200 nm diameter. Another attractive field of application is to study condensation variabilities on the macromolecular resolution level, by artificially changing the degree of compaction of the chromatin. This can be done with chemicals such as Trichostatin A, an inhibitor of histone deactetylase, inducing decondensation of heterochromatin regions (Görisch et al. 2005), or Berenil, an AT-specific DNA ligand, inhibiting the condensation of the heterochromatin block on Yq12 (Haaf et al. 1989). The quantitative data obtained this way may be used to test numerical models of chromatin nanostructure (Odenheimer et al. 2009; Bohn et al. 2010) A further appealing perspective is to count copy numbers of specific DNA sequences on a single cell nucleus basis. So far, copy numbers are mainly assessed by PCR and array-based techniques, which have the drawback that they typically show an average over a large number of cells. Changes in copy numbers attract increasing interest, as it becomes more and more apparent that these contribute to several diseases, including cancer (Dear 2009).

The FISH localisation microscopy approach presented here is particularly suited for the analysis of small gene regions in intact cell nuclei. Many of these domains are thought to change their conformation due to activation but so far incontrovertible evidence is still elusive. Furthermore, extending the SPDM approach to two or more colours, each allowing the localization of a large number of molecules, will make possible to study on the nanoscale the interaction of specific chromatin structures such as the widely discussed problem of intermingling of chromosome territories (Branco and Pombo 2006; Rouquette et al. 2010).

The major concerns in the application of FISH-based SPDM to study nuclear nanostructure are changes induced by the fixation and denaturation process (Solovei et al. 2002b). This problem might be overcome in various ways, e.g., by using COMBO or PNA FISH with non-denaturating conditions (Hausmann et al. 2003; Schmitt et al. 2010) or by using thin cryosections (Branco and Pombo 2006).

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