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Elimination of Autofluorescence in Immunofluorescence Microscopy with Digital Image Processing¹

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Introduction

Autofluorescence is a very common phenomenon and can interfere in microscopic studies using fluorescence labeling. It is caused by certain biomolecules, such as elastin, fibronectin, and lipofuscin, and by aldehyde fixation (1). Autofluorescence can be very intense and often results in masking of the specific fluorescence signals. Only a few reports have dealt with the elimination of autofluorescence, most of them related to flow cytometry. Addition of crystal violet, for example, has been used to quench autofluorescence of macrophages in flow cytometry (2). Dual-wavelength flow cytometry has also been applied to correct for autofluorescence (3,4). In fluorescence microscopy, several parameters, such as excitation wavelengths, excitation power, microscope apertures, and buffers, have been optimized to minimize autofluorescence (5). However, autofluorescence cannot be fully eliminated in this way. Probably the most sophisticated method to eliminate autofluorescence is time-resolved fluorometry. This procedure is based on fluorescent lanthanide chelates with long decay times, which can be used to suppress the fast-decaying autofluorescence (6,7). This technique, however, needs a sophisticated fluorescence microscope to visualize the time-resolved emission (7).

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Immunostaining. For indirect immunofluorescence, tissue samples were frozen in isopentane cooled with liquid nitrogen and stored at −70°C. Cryosections (2-12 μm) were stored at −70°C. Cryosections were rehydrated for 10 min in PBS, pH 7.4, and primary and secondary antibodies were applied for 90 min in PBS containing 1% BSA. After each incubation, sections were washed in PBS (three times for 5 min). Anti-heparan sulfate antibody and FITC-conjugated secondary antibodies were used at a dilution of 1:100. Anti-desmin and 3H6 antibodies were used at a dilution of 1:50. In sections of quadriceps muscle, autofluorescence was enhanced by incubation with 3% formaldehyde in 150 mM phosphate buffer (pH 7.2) for 1 hr (1).

Comparison of Sections on the Basis of Protein Content. Because the micrometer setting of a microtome is not an accurate way to obtain sections of a certain thickness, we measured section thickness also by determining the surface and the protein content of a section. Sections were checked for homogeneity, i.e., for the presence of the same structural elements in the sections. This was done to exclude the possibility that sections differed from each other in structural composition, which may give rise to differences in measurable protein per unit of volume. Only sections consisting almost completely of muscle cells were taken. The surface of a cryosection was marked with a marker pen and copied by a photocopier onto a piece of paper. The image of the sections was cut out and weighed. The surface was calculated by using the weight of 1 cm² of paper as a standard. After surface analysis, the tissue section was taken up in demineralized water and the protein content was determined according to Lowry et al. (9). The section thickness was expressed in μg protein/mm².

Digital Subtraction of Autofluorescence. Microscopic (fluorescence) images were digitized using the MagiCal hardware and TARDIS software of Joyce–Loebl (Gateshead, UK). Images were captured with a resolution of 256 x 256 datapoints and 256 gray levels each. The images were processed with ImageCalc, an MS-Windows application designed for this purpose (10).

To substract the autofluorescence signal, two images were digitized. For an FITC-immunostained tissue section, one image, representing the total fluorescence image (i.e., the specific FITC image plus the autofluorescence image), was obtained by excitation at 480 nm and emission at 515 nm. The second image, representing the autofluorescence image, was obtained by excitation at 380 nm and emission at 515 nm. For optimal results, two additional background images were digitized using the same parameters and representing images of an object slide without tissue. These images were used for noise and shading correction (11). Camera gain and sensitivity were adjusted to obtain optimal signal-to-noise ratios for both excitation wavelengths. Both images were captured immediately after each other, each image representing the average of eight individual images (to reduce noise). Total image capture of both images was achieved within 1 sec.

Because autofluorescence intensity depends on the excitation and emission wavelength utilized, a correction factor for the use of two wavelengths was applied by excitation of a non-immunostained cryosection at 480 and 380 nm. After determining the ratio of emission at 515 nm for these wavelengths (R_{480/380}) the specific fluorescence can be calculated using the equation:

\[ SF = TF - (R_{480/380} \times AF). \]

where SF is the specific fluorescence, TF is the total fluorescence, and AF is the autofluorescence.

Results

Elimination of Autofluorescence

Human Lung Parenchyma. Human lung parenchyma sections display a very strong autofluorescence, mainly caused by elastic fibers (12). This autofluorescence masked the linear heparan sulfate basement membrane staining with antibody JM403 (Figure 1A), which was clearly visible on glomerular and tubule basement membranes (8). Digital subtraction of the autofluorescence image (Figure 1B) from the total fluorescence image revealed an autofluorescence-free image (Figure 1C). In this image, the linear basement membrane staining was clearly visible. The autofluorescence image lacked any specific basement membrane staining.

Human Heart Muscle. Human heart muscle, especially from individuals of older age, contains the strongly autofluorescent pigment lipofuscin (13). Immunostaining of heart tissue sections with monoclonal antibody 3H6 showed staining of the intercalated disks. This staining, however, was locally completely masked by the autofluorescent lipofuscin particles (Figure 2A). Subtraction of the autofluorescence image (Figure 2B) unmasked the specific staining of the intercalated disks (Figure 2C).

Figure 1. Elimination of autofluorescence (mainly caused by elastin fibers) in human lung parenchyma sections. Sections were stained with mouse anti-heparan sulfate antibodies, which were detected with FITC-labeled rabbit anti-mouse Ig antibodies. The autofluorescence image (B) was subtracted from the total fluorescence image (A), resulting in an autofluorescence-free image (C). Arrows indicate areas where the basement membrane location of heparan sulfate becomes visible only after subtraction of autofluorescence. The correction factor R_{480/380} was 0.7. Bar = 25 μm.
DIGITAL ELIMINATION OF AUTOFLUORESCENCE

Because the autofluorescence signal can be obtained separately, we investigated if the intensity of this signal could be used as a measure of the thickness of a section. Since this requires an equal distribution of autofluorescence over the section, we used sections of quadriceps muscle in which the autofluorescence was enhanced by pre-incubation with formaldehyde. To obtain precise measurements, the section thickness was accurately measured by analysis of surface and protein content. We found a positive correlation between the section thickness (measured as µg protein/mm²) and the autofluorescence intensity (Figure 3). We evaluated this observation further on sections of human quadriceps muscle stained with anti-desmin antibodies. Whereas the specific fluorescence and the autofluorescence showed a clear correlation with the apparent thickness of the sections (indicated by the microtome), the ratio between these fluorescence parameters was independent of the apparent section thickness (Figure 4).

Figure 2. Elimination of autofluorescence caused by lipofuscin in human heart sections. Tissue sections were immunostained with the mouse monoclonal antibody 3H6, which is reactive with intercalated disks. Antibodies were detected with FITC-labeled rabbit anti-mouse Ig antibodies. The total fluorescence image (A) minus the autofluorescence image (B) results in an autofluorescence-free image (C) completely lacking the fluorescence of lipofuscin particles. Note that the specific fluorescence at some intercalated disks becomes visible only after elimination of autofluorescence (arrow). The correction factor R480/560 was 1.0. Bar = 25 µm.

Use of Autofluorescence as a Measure of Tissue Thickness

Because the autofluorescence signal can be obtained separately, we investigated if the intensity of this signal could be used as a measure of the thickness of a section. Since this requires an equal distribution of autofluorescence over the section, we used sections of quadriceps muscle in which the autofluorescence was enhanced by pre-incubation with formaldehyde. To obtain precise measurements, the section thickness was accurately measured by analysis of surface and protein content. We found a positive correlation between the section thickness (measured as µg protein/mm²) and the autofluorescence intensity (Figure 3). We evaluated this observation further on sections of human quadriceps muscle stained with anti-desmin antibodies. Whereas the specific fluorescence and the autofluorescence showed a clear correlation with the apparent thickness of the sections (indicated by the microtome), the ratio between these fluorescence parameters was independent of the apparent section thickness (Figure 4).

Figure 3. Autofluorescence as a measure of section thickness. Autofluorescence was established in sections of rat quadriceps muscle with various thickness (expressed as µg protein/mm²). Autofluorescence was enhanced by pre-incubation of the sections with formaldehyde. Values are mean ± SEM of four individual observations. Note that there is a positive correlation (r = 0.93; p<0.0001) between autofluorescence and section thickness.

Figure 4. The specific fluorescence (●), the autofluorescence (■), and the ratio of specific fluorescence and autofluorescence (□) as a function of the apparent thickness of sections of human quadriceps muscle stained with rabbit anti-chicken desmin antibodies and FITC-labeled goat anti-rabbit IgG antibodies. Section thickness was derived from the microtome setting. A positive correlation is present between both the specific fluorescence and the autofluorescence and the apparent section thickness (r = 0.69, p<0.001; r = 0.51, p<0.01, respectively). The ratio of specific fluorescence (SF) and autofluorescence (AF) is independent of the thickness of the sections (r = 0.21; p<0.7). Autofluorescence was enhanced by pre-incubation of the sections with formaldehyde. Values are mean ± SEM of four individual observations.
Discussion

In this report we demonstrate that autofluorescence in immunofluorescence microscopy can be efficiently eliminated by subtraction of the autofluorescence image, captured using an excitation wavelength outside the excitation spectrum of the fluoroprobe. This principle has been used for elimination of autofluorescence in flow cytometry (3,4), and we show here that it is also applicable in immunofluorescence microscopy of tissue sections. Using this technique, we were able to eliminate the strong autofluorescence of elastin and lipofuscin without affecting the specific fluorescence. An alternative way to eliminate autofluorescence is time-delayed fluorometry (6). This procedure, however, needs a sophisticated and expensive fluorescence microscope (7), whereas our technique requires only a digitization apparatus (e.g., a CCD camera) connected to a standard fluorescence microscope.

Elimination of autofluorescence can simplify the interpretation of immunostained sections, especially in tissues with an abundance of autofluorescent molecules, such as heart and brain (lipofuscin) and lung (elastin). The method is also useful in cases where the specific staining is weak (e.g., in tissues with a small amount of antigen or in situ hybridization). Because the digitization process is usually performed within a single second, fading of the fluorescence probe is minimized, which is essential for quantitative analysis (14). In addition, specific fluorescence can be corrected for section thickness using the autofluorescence signal (Figure 4).

In conclusion, autofluorescence in tissue sections treated for immunofluorescence microscopy can be effectively eliminated by digitalization of the fluorescence signal at two different wavelengths. Correction for section thickness and minimal fading of the fluoroprobe make this method suitable for (semi)quantitative analysis of stained sections.

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Literature Cited