Simultaneous ultrastructural analysis of fluorochrome-photoconverted diaminobenzidine and gold immunolabelling in cultured cells

M. Malatesta, C. Zancanaro, M. Costanzo, B. Cisterna, C. Pelliccieri

Abstract

Diaminobenzidine photoconversion is a technique by which a fluorescent dye is transformed into a stably insoluble, brown, electron-dense signal, thus enabling examination at both bright field light microscopy and transmission electron microscopy. In this work, a procedure is proposed for combining photoconversion and immunoelectron microscopy: in vitro cell cultures have been first submitted to photoconversion to analyse the intracellular fate of either fluorescent nanoparticles or photosensitizing molecules, then processed for transmission electron microscopy; different fixative solutions and embedding media have been used, and the ultrathin sections were finally submitted to post-embedding immunogold cytochemistry. Under all conditions the fluorescently labelled structures get antigen were properly detected in the same photoconversion reaction product and the targets of the fluorochrome (or the fluorochrome-conjugated probes) for a molecular target is the prerequisite for its accurate localization at the ultrastructural level by means of the photoconversion version procedures. However, investigating the dynamic interaction of cellular organelles with different molecular factors involved in cell mechanisms often requires the simultaneous detection in situ of more than one molecule.

Introduction

Photoconversion is a procedure by which a fluorescent dye is transformed into a stably stained product, thus enabling a long-lasting analysis of fluorescently-labelled cellular structures at both bright field light microscopy and transmission electron microscopy. It exploits the property of fluorescent dyes to generate reactive oxygen species under intense illumination, and is accomplished by illuminating a fluorescently labelled sample at the excitation wavelength of the fluorochrome in the presence of O₂ and of the chromogen 3,3′ diaminobenzidine (DAB). Under these conditions, the dye molecules generate reactive oxygen species which oxidize DAB, thus inducing the formation of insoluble, brown, electron-dense precipitates. As DAB is only oxidized in the close proximity of the fluorochrome molecules (due to the very short half-life of the reactive oxygen species), this technique enables to precisely localize at high resolution the fluorescently labelled structures.

This procedure was first used by Maranto to convert the intracellularly injected fluorochrome Lucifer Yellow into a stable signal for light and electron microscopic examination. Subsequently, this technique has been extended to a variety of fluorochromes used either as cell tracers or antibody markers, thus allowing fine investigation of e.g. neuronal networks, synaptic vesicle turnover, endosomal enzyme activity. The conventional photoconversion method has been also associated with techniques of fluorescence recovery after photobleaching (FRAP) or of 3D electron tomography or adapted for confocal laser scanning microscopy. The specificity of the fluorochrome (or the fluorochrome-conjugated probes) for a molecular target is the prerequisite for its accurate localization at the ultrastructural level by means of the photoconversion version procedures. However, investigating the dynamic interaction of cellular organelles with different molecular factors involved in cell mechanisms often requires the simultaneous detection in situ of more than one molecule.

To this aim, in the present work we combined DAB photoconversion and post-embedding immunocytochemistry to simultaneously assess the distribution of photoconverted nanoparticles or photosensitizing molecules and gold-labelled antigens allowing a specific identification of subcellular organelles at transmission electron microscopy.

Materials and Methods

Cell cultures and treatments

Human HeLa cells and rat neuronal B50 were grown in 25 cm² flasks in Dulbecco Modified Eagles Medium (DMEM) supplemented with 10% foetal bovine serum, 1% glutamine, 100U of penicillin and streptomycin (Celbio s.r.l., Milan, Italy), in a humidified air atmosphere containing 5% CO₂.

HeLa cells were seeded onto glass coverslips in six-multiwell plates (1x10⁵ per well) 24 h before being incubated for 60 min in the dark with 10⁻⁴ M Hypocrellin B acetate (HypB-Ac) (as reported by Croce et al.). HypB was purchased from Axxora.com (Vinci-Biochem, Florence, Italy) and HypB-Ac was prepared according to Croce et al. The addition of two acetate groups to the native HypB facilitates cell internalization compared to the native molecule, and makes HypB-Ac to act as a fluorogenic substrate: the modified compound cannot serve, per se, as a photosensitizer but once inside a cell the bound groups are enzymatically removed by esterases, and the photophysical properties of the native molecules are restored. HypB-Ac was diluted at the final concentration in serum-free culture medium to prevent the hydrolysis of acetate groups by serum esterases. At the end of the incubation times, slides were removed from the medium, rinsed with PBS, and processed for transmission electron microscopy (see below). To avoid unwanted photobleaching and/or phototoxicity

Correspondence: Manuela Malatesta, Dipartimento di Scienze Neurologiche e del Movimento, Sezione di Anatomia e Istologia, Università di Verona, Strada Le Grazie 8, 37134 Verona, Italy. E-mail: manuela.malatesta@univr.it

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effects, these operations were performed under low light illumination.

Rat neuronal B50 were trypsinized when subconfluent, and seeded on glass coverslips in twelve-multiwell dishes (1×10³ per well). Two days after seeding, the initial medium was removed and the cells were incubated for 24 h with 450 µL of fresh medium plus 50 µL of suspension of fluorescein 5(6)-isothiocyanate (FITC)-containing chitosan nanoparticles (as reported by Malatesta et al.39).

DAB photoconversion and sample processing for transmission electron microscopy

HypB-Ac-loaded HeLa cells and nanoparticle-loaded B50 cells on coverslips were fixed with 2.5% (v/v) glutaraldehyde and 2% (w/v) paraformaldehyde in 0.1 M phosphate buffer, pH 7.4, at 4°C for 1 h, washed and incubated with DAB (20 mg/10 mL in Tris HCl 0.05 M, pH 7.6) under simultaneous irradiation with two 8W Osram Blacklite 350 lamps for 2 h at room temperature (these lamps have emission peaks of high intensity at 430-450, 550 and 580 nm, thus being suitable for FITC and HypB excitation). In order to avoid cell damage due to heating effect of irradiation, the multiwell dishes were maintained on ice.

After irradiation, the cells were post-fixed with 1% OsO₄ and 1.5% potassium ferrocyanide at room temperature for 1 h, dehydrated with aceton and embedded in Epon resin. Some B50 cell samples were fixed with 4% (w/v) paraformaldehyde in 0.1 M phosphate buffer, pH 7.4, at 4°C for 1 h, incubated with DAB under light irradiation, treated with 0.5 M NH₄Cl in PBS at 4°C for 30 min to block free aldehydes, dehydrated with ethanol and embedded in LRWhite resin polymerized for 24 h at 60°C. As controls, some samples were processed as described above but omitting both DAB incubation and exposure to light.

Post-embedding immunocytochemistry

Ultrathin (70-90 nm thick) sections of HeLa or B50 cell samples were collected on 200 mesh nickel grids coated with a formvar-carbon layer. Before starting the immunocytochemical procedure, Epon-embedded samples were pretreated with a 0.2 M aqueous solution of sodium metaperiodate for 60 min at room temperature in order to improve antibody binding.44

Both LRWhite- and Epon-embedded samples were then processed for immunocytochemistry by using human autoimmune sera recognizing either the lysosomal/endosomal proteins corresponding to the autophagy marker LC3B35 or the mitochondrial 70 kDa E2 subunit of the pyruvate dehydrogenase complex.36 Sections were floated on 0.1% ovalbumin (Fluka, St. Louis, MO, USA) in PBS for 3 min, and then incubated for 17 h at 4°C with the primary antibody diluted in PBS containing 0.1% bovine serum albumin (Fluka) and 0.05% Tween 20 (the anti-lysosome/endosome serum was diluted 1:10, the anti-mitochondria serum 1:50). After rinsing in PBS-Tween and in PBS, the sections were floated on ovalbumin, and then reacted for 45 min at room temperature with 12 nm-gold-conjugated protein A (British Biocell International, Cardiff, UK) diluted 1:20 in PBS. Finally, the sections were extensively rinsed in PBS and water, and air-dried. As controls, the primary antibodies were omitted from test sections.

Ultrathin sections were weakly stained with 4.7% aqueous solution of uranyl acetate for 2 min for better visualization of the photoconversion reaction product. All samples were observed in a Philips Morgagni transmission electron microscope (FEI Company Italia S.r.l., Milan, Italy) operating at 80kV and equipped with an Olympus Megaview II camera for digital image acquisition.

Results and Discussion

In all samples, DAB photoconversion gave rise to a finely granular electron dense product; this confirmed the presence of photoactive HypB molecules inside endosomes, multivesicular bodies, late lysosomes and mitochondria (Figure 1 a-c) as well as in the cytosol of HeLa cells (as already reported in a previous paper45), and the occurrence of FITC-containing nanoparticles in the cytosol and inside endosomes and late lysosomes of B50 cells30,31 (Figure 1 d,e).

The electron dense granular precipitate was evident in cells submitted to post-fixation with osmium tetroxide and potassium ferrocyanide, and embedded in Epon resin (Figure 1 a-d) as well as in cells embedded in the LRWhite resin without post-fixation (Figure 1 e). The control samples (where both DAB incubation and light exposure were omitted) did not show any reaction product (not shown). In photoconverted samples embedded either in Epon or in LRWhite resin, the post-embedding immuno-
cytochemical procedure gave positive results, thus demonstrating that the cell molecules preserve the ability to react with specific antibodies after the photoconversion procedure applied in our experiments: the serum recognizing lysosomal/endosomal proteins specifically labelled endosomes, multivesicular bodies and late lysosomes (Figure 1 a,c,e), while the probe recognizing the pyruvate dehydrogenase complex specifically labelled mitochondria (Figure 1 b). Notably, the immunogold signal also occurred on organelles devoid of the photoconversion product, thus excluding interference of the DAB precipitates with the antibody binding.

It is known that the fixation and embedding procedures we applied to the Epon-embedded cells are optimal for good morphological preservation, but may cause denaturation of many molecules which could become barely detectable by antibodies. In the Epon-embedded samples, the pre-treatment with sodium metaperiodate (which is able to unmask the antigenic sites on osmicated thin sections without altering their ultrastructural preservation)30 successfully improved antibody binding both increasing the specific signal and decreasing background, although the gold labelling was always weaker than in LRWhite-embedded sections. However, it is worth noting that the treatment with oxidizing agents which solubilise osmium molecules leading to their removal from the embedded tissue,24 markedly decreased the electrodensit of the photoconversion product, sometimes making the finest granular deposits hardly visible, especially when photosensitizing molecules occurred in the cytosol or in mitochondria. This limitation should be taken into consideration in selecting the fixation/embedding procedure for photoconverted samples to be processed for immunocytochemistry, particularly when the photoconverting molecules are supposed to be scanty and/or scattered. LRWhite-embedded samples proved to be optimal for the simultaneous detection of photoconversion products and immunolabelled molecules. In LRWhite-embedded cells the photoconversion deposits were well visible: this demonstrates that the morphology and distribution of the DAB precipitates are not altered by the mild fixation with paraformaldehyde (in the absence of osmication) and the embedding in an acryl hydrophilic resin which is known to improve antibody-antigen interactions while preserving a sufficiently good ultrastructure.

In order to optimise the observation of both photoconversion and immunogold signals, the Epon-embedded and LRWhite-embedded cells were weakly stained with uranyl acetate to increase sample contrast without masking the sometimes minutely dispersed photoconversion precipitates. Uranyl acetate was preferred to lead citrate, which often gives rise to a tingly granular staining31 that could closely resemble the photoconversion product. The interference of uranyl acetate staining in the visualization of the reaction product was excluded by observing unstained samples after DAB photoconversion and immunogold labeling (Figure 1c). Photoconversion and immunohistochemistry have seldom been combined in previous studies, but pre-embedding techniques were only used to label photoconverted neurons.32,33 The simple and reliable procedure of the present work allows to exploit a photoconverted sample to detect a variety of antigens in different ultrathin sections; it will also be possible to simultaneously localise with high precision and sensitivity several antigens on a single section of a photoconverted sample by using probes conjugated with gold grains of different sizes.

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