OPTIMAL METHODS OF ANTIGEN RETRIEVAL FOR ORGANIC ANION TRANSPORTERS IN CRYOSECTIONS OF THE RAT KIDNEY

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To localise antigens by immunocytochemistry (IC), the samples of tissues or cells are usually denatured by fixation, and either frozen and cryosectioned, or embedded in paraffin before sectioning. p-Formaldehyde (PFA; formalin) is a common fixative, which preserves antigenicity of proteins, but damages the tissue/cell morphology and “masks” the antibody binding sites (epitopes). In order to “unmask” epitopes, some kind of antigen retrieval (AR) is used. The aim of this study was: a) to find an optimal AR method in cryosections of in vivo PFA-fixed kidneys for organic anion transporters (Oat) that reside in the basolateral (Oat1, Oat3) and brush-border membrane (Oat2, Oat5) of the rat renal proximal tubules, and b) using optimal method, to compare IC staining of Oats in kidneys that had been PFA-fixed in vivo or in vitro. IC staining in untreated cryosections was compared with that following detergent treatment or microwave heating in citrate buffer of pH 3, pH 6, or pH 8, with or without alcohol pre-treatment. The preferred AR method for Oat1, Oat2, and Oat5 was heating of cryosections at pH 6, and for Oat3 heating at pH 3, without alcohol pre-treatment. Compared with tissue fixed in vivo, tissue fixed in vitro exhibited damaged tubule morphology, similar staining intensity of Oat1 and Oat3, and higher staining intensity of Oat2 and Oat5. We conclude that for optimal IC presentation, each Oat in the rat kidney has to be treated individually, with different fixation and AR approach.

KEY WORDS: cell membrane, cryosections, immunocytochemistry, immunofluorescence, membrane transporters, proximal tubule

Immunocytochemistry is a method for determining the exact site of individual proteins in tissues and cells. Tissues/cells for immunocytochemistry are often denatured by fixation with (2 to 10) % p-formaldehyde (PFA, formalin). However, fixation with PFA has some drawbacks; it “masks” the antibody binding sites (epitopes) on some antigens, and alters cellular morphology by damaging the cell structure (1). It is not clear how PFA reacts with proteins, except that it cross-links reactive sites within the same proteins and between adjacent proteins via methylene bridges (2). However, not all proteins are masked by PFA at the same rate and intensity. For instance, monoclonal antibodies fail to label neuro-filaments after one day, lymphocyte antigens LN1, LN2, and LN3 have shown reduced staining after three days, while carcino-embryonic antigen stains even after 14 days of tissue fixation in formalin (3). This problem is more evident for monoclonal then polyclonal antibodies, because polyclonal antibodies can bind to more reactive places on proteins (2). In order to “unmask” cryptic epitopes, one has to apply a series of antigen retrieval techniques for each set of antigen/antibody separately in order to find optimal conditions for the immunocytochemical presentation of an antigen. Antigen retrieval is defined as “a high-temperature heating method to recover antigenicity of tissue sections that have been masked by formalin fixation” (4). Most of the previous
studies of antigen retrieval methods were performed with sections of paraffin-embedded tissues/cells. Depending on the antibodies used, these methods include treatment with various alcohols (to remove paraffin), different heating temperatures, pressures, osmolality, and chemical composition of the reagents, pH, and different timing of treatments (5-10). The pH of reagents seems to play a crucial role in antigen retrieval; pH 8 to pH 9 has been proposed as optimal for starting with a new antigen (9). However, this is not true for all antigens; the optimal condition for labelling thrombospondin in paraffin-embedded sections was antigen retrieval in an acidic buffer (pH 1) (11).

The revealing methods have been less used with cryosections of PFA-fixed tissues, organs, or cells. While some antibodies weakly recognised, or failed to recognise their epitopes in PFA-fixed tissue cryosections, they did it after treatment with a detergent such as sodium dodecyl sulfate (SDS), which is more commonly used in gel electrophoresis. Thus, SDS treatment of cryosections enhanced the immunostaining of Na/K-ATPase and vacuolar H^+-ATPase (56 kDa subunit) in various tubules, but this was the only way to stain caveolin in endothelial cells, as well as the anion exchanger AE2 in the principal cells and Na/K-ATPase in intercalated cells of the collecting duct in rat kidney (1, 12-14). However, the same treatment of cryosections did not affect the staining of megalin and water channels AQP1 and AQP2 (12). These findings have indicated a need for an individual approach to each antigen and its antibody, and for defining an optimal antigen retrieval method and conditions for each protein before its detailed characterisation by immunocytochemistry in tested tissues/cells.

Using cryosections of rat kidneys PFA-fixed in vivo, we (15, 16) and others (17-19) have recently established a detailed distribution of several organic anion transporters (Oats) localized to the basolateral (BLM) (Oat1, Oat3) and brush-border membrane (BBM) (Oat2) of mainly proximal tubules (PT) in the rat nephron. The highest expression of the Oat1 protein was observed in the PT S2 segments in the cortex, where it exhibited male (M)-dominant sex differences, whereas the Oat3 protein was localised basolaterally in various nephron segments, being the strongest and M-dominant in the cortical tubules (15). The Oat2 protein was strongly expressed in the BBM of PT S3 segments, even more so in females (16). However, in these studies the conditions for optimal immunocytochemical presentation of Oats have been only superficially tested, and a detailed study of antigen retrieval has not yet been performed. In addition, we have included another transporter in the BBM of PT S3 segments with F-dominant expression, that is, Oat5 (20), whose distribution along the rat nephron has yet to be described in detail. With these experiments we aim to find an optimal method and conditions for retrieving cryptic antigens, possible non-specific staining, and artifacts. After establishing optimal antigen retrieval conditions, we aim to compare the pattern of localisation and staining intensity of all 4 Oats in cryosections of rat kidneys fixed with PFA in vivo and in vitro in order to test these two modes of fixation for the integrity of tubule morphology and immunostaining efficiency.

MATERIAL AND METHODS

Animals and treatment

The experiments included adult, three-month-old male and female Wistar rats from the breeding colony of the Institute for Medical Research and Occupational Health, Zagreb. Due to significant sex differences in the expression of individual Oats in the rat kidney (15, 16, 20), male rats were used to study Oat1 and Oat3, whereas female rats were used to study Oat2 and Oat5. The animals were bred and maintained according to the Guide for Care and Use of Laboratory Animals (National Institute of Health, Bethesda, USA, 1996). The experiments were approved by the Institute’s Ethics Committee.

Antibodies and chemicals

Polyclonal antibodies against Oat1 (Slc22a6), Oat2 (Slc22a7), Oat3 (Slc22a8), and Oat5 (Slc22a19) were raised in rabbits; their use was described in our previous reports (15, 16, 20). The secondary antibody was CY3-labelled goat anti-rabbit IgG (GARCY3), which was purchased from Jackson ImmunoResearch (West Grove, PA, USA).

Anaesthetics (Narketan and Xylapan) were purchased from Chassot AG (Bern, Switzerland). Other chemicals used were of the highest purity available and purchased from Sigma (St. Louis, MO, USA) or Fisher Scientific (New Jersey, NY, USA).

Tissue fixation and immunocytochemistry

Rats were sacrificed either by decapitation or were anaesthetised (Narketan, 80 mg kg^-1 b.m. + Xylapan,
12 mg kg⁻¹ b.m., *i.p.*), and their circulatory system was perfused *via* the left ventricle of the heart *in vivo* using the Masterflex pump (Cole-Parmer, Chicago, IL, USA), first with aerated (95 % O₂ + 5 % CO₂) and temperature-equilibrated (37 °C) phosphate-buffered saline (PBS; 137 mmol L⁻¹ NaCl, 2.7 mmol L⁻¹ KCl, 8 mmol L⁻¹ Na₂HPO₄, 2 mmol L⁻¹ K₂PO₄, pH 7.4) for ~2 min (to remove blood *via* the incisioned abdominal vena cava) and then with a 100 mL fixative (4 % PFA in PBS) for ~5 min. The kidneys from non-perfused and *in vivo* perfused animals were removed, decapsulated, cut in ~1 mm thick sagittal slices, and the slices were further kept in the same fixative for 24 h at 4 °C, followed by extensive washing in PBS, and stored in PBS containing 0.02 % NaN₃ at 4 °C until use.

Before cryosectioning, tissue slices were infiltrated with 30 % sucrose (in PBS) overnight, embedded in OCT-medium (Tissue-Tek, Sakura, Japan), frozen at -25 °C, and sectioned in a Leica CM 1850 cryostat (Leica instruments GmbH, Nussloch, Germany). Four-micrometer thick cryosections were collected on Superfrost-plus microscope slides, dried at room temperature for 3 h, and stored at 4 °C until use.

The optimal conditions for immunostaining of individual transporters were tested in cryosections of the kidneys fixed *in vivo*; the staining of untreated cryosections was compared with that of cryosections processed with the following three antigen retrieval methods before applying the primary antibody: a) SDS treatment, b) microwave heating in citrate buffers of pH 3, pH 6, and pH 8, and c) treatment with different alcohols (steps used for deparaffinization/delipidation of paraffin-embedded sections), followed by microwave heating in citrate buffers of pH 3, pH 6, and pH 8. Having established the optimal method and conditions, we compared the staining pattern and intensity for each Oat in the kidneys fixed by perfusion *in vivo* with that in the non-perfused kidneys, fixed *in vitro*.

The untreated cryosections were (steps): rehydrated in PBS for 15 min, incubated in BSA (1 % bovine serum albumin in PBS) for 30 min in order to prevent non-specific binding of the antibody, incubated in primary antibody (optimally diluted in PBS) at 4 °C overnight, rinsed in high-salt PBS (PBS + 1.8 % NaCl; 2x5 min) and normal PBS (2x5 min), incubated in secondary antibody (1.6 µg mL⁻¹ in PBS) at room temperature for 1 h, rinsed in high-salt PBS (2x5 min) and PBS (2x5 min), overlayed with a fluorescence fading retardant (Vectashield; Vector Laboratories Inc., Burlingame, CA, USA), covered with the cover glass, and sealed with nail-polish.

The SDS-treated cryosections were (steps): rehydrated in PBS for 15 min, incubated in SDS solution (1 % SDS in PBS) for 5 minutes, rinsed in PBS (4x5 min), blocked with BSA for 30 min, and further processed as described above for untreated cryosections.

The microwave-treated cryosections were (steps): rehydrated in PBS for 15 min, immersed in 10 mmol L⁻¹ citrate buffer (pH 3, pH 6, or pH 8), and heated in a microwave oven at 800 W in 4 cycles (5 min each), then cooled down to room temperature in the same buffer for 20 min, rinsed in PBS (3x5 min), incubated in 0.5 % Triton X-100 in PBS (15 min) and in 2 % Triton X-100 in PBS (30 min), rinsed in PBS (2x5 min), blocked in BSA (30 min), incubated in primary antibody at 4 °C overnight, rinsed in 0.1 % Triton X-100 in PBS (10 min) and PBS (2x5 min), incubated in secondary antibody at room temperature for 1 h, rinsed in 0.1 % Triton X-100 (10 min) and PBS (2x 5 min), overlayed with a fluorescence fading retardant (Vectashield; Vector Laboratories Inc., Burlingame, CA, USA), and further processed as described above for untreated cryosections.

The alcohol-treated cryosections underwent the following consecutive steps: delipidation in a series of alcohols, including 100 % xylol (30 min), 100 % propanol (5 min), 96 % ethanol (5 min), 75 % ethanol (5 min), and 60 % ethanol (5 min), then rinsing in distilled water (5 min), rehydration in PBS (15 min), followed by the steps listed for microwave-treated cryosections.

Immunostaining was inspected with the Opton III RS fluorescence microscope (Opton Feintechnik, Oberkochen, Germany) using a Spot RT Slider camera and software (Diagnostic Instruments, Sterling Heights, MI, USA). The Oat-related immunostaining, imaged in red fluorescence of CY3, and the tubule morphology, imaged in green background fluorescence of the tissue, were taken separately using specific cut-off filters with no bleed-through effects. Due to specific sex and zone differences in the tubular expression of various Oats in rat kidneys (15, 16, 20), the immunostaining of Oat1 and Oat3 was studied in the cortex of male rats, whereas the immunostaining of Oat2 and Oat5 was studied in the outer stripe of female rats. Each set of images, related to the individual Oat, was taken using the same fluorescence-recording parameters. The colour images were then imported into Adobe Photoshop 6.0, assembled in panels, converted into
black and white mode, and labelled. The data shown represent the findings in cryosections from three rats in each experimental group.

RESULTS

Optimal antigen retrieval conditions for basolateral transporters Oat1 and Oat3

Oat1

As shown in Figure 1 for the kidneys fixed in vivo, the untreated cryosections exhibited weak staining of Oat1 in the BLM of individual PT in the cortex (A). The SDS-treatment increased the number of weakly positive tubules (B). In the microwave-treated cryosections, the staining intensity depended on buffer pH, and followed the pattern pH 3<pH 6<pH 8. However, the use of cryosections at pH 8 was strongly compromised by numerous tubules partially (E, arrows) or completely (data not shown) detached from the microscope slides. In cryosections pre-treated with alcohols and then heated in a microwave (F-H), Oat1 staining in the PT BLM followed the same pattern as in the microwave-treated cryosections, but at much lower level of intensity.

PT morphology was compared with Oat1 staining in the same tubules between tissue cryosections of kidneys fixed in vitro and in vivo that had been microwave-treated at pH 6 (Fig. 2). Whereas the tubule profiles in cryosections of the kidneys fixed in vitro showed collapsed lumina, filled with cellular debris (A), the tubules in cryosections of the kidneys fixed in vivo exhibited regular appearance, with open lumina and sharp contours (C). The staining intensity of Oat1 in the PT BLM was similar in both cases (compare B and D), except that some tubules in the kidneys fixed in vitro exhibited shorter basolateral invaginations (B, arrows).

Figure 1 Antigen retrieval conditions for Oat1 in untreated (A), SDS-treated (B), microwave-treated (C-E), and alcohol-treated (F-H) cryosections of the kidneys fixed in vivo. (A) Untreated cryosections show a weak staining of the BLM in individual proximal tubules. (B) SDS-treated cryosections have a higher number of weakly stained tubules. (C-E) Microwave-treated cryosections in citrate buffers of pH 3 (C), pH 6 (D), and pH 8 (E) exhibit strong BLM staining in the cortical proximal tubules, the strongest being at pH 8 (E). (F-H) In cryosections pre-treated with alcohols, followed by microwave heating in citrate buffer of pH 3 (F), pH 6 (G), and pH 8 (H), the staining intensity of the BLM in the tubules is significantly weaker than in the microwave-treated cryosections. At pH 8, in both microwave-treated (E) and alcohol-treated cryosections (H), many tubules are partially (arrowheads) or completely (not shown) detached from the microscope slide (arrowheads). Scale bar=20 µm
Figure 3 shows the data on antigen retrieval for Oat3 in cryosections of the kidneys fixed in vivo. In untreated sections, Oat3 staining in the BLM of various cortical tubules was weak (A). In the SDS-treated cryosections, the specific staining was even weaker, and a high non-specific background staining in all tubule profiles was observed (B). Heating in a microwave showed that Oat3 was best stained in a citrate buffer of pH 3 (C), and less of pH 6 (D) and pH 8 (E). The same pattern, but with lower staining intensity, was observed in cryosections pre-treated with alcohols (F-H).

Figure 4 compares the morphology of cortical tubules and Oat3 staining in the same tubules between kidneys fixed in vitro and in vivo. In the kidneys fixed in vitro, most cortical tubules had irregular and/or collapsed lumina, filled with cellular debris (A), whereas (B), Oat1 in the BLM of the same tubules is strongly stained, but some tubules lost the BLM invaginations (arrows). (C) Proximal tubules in cryosections of the kidneys fixed in vivo are open, with sharp borders and regular appearance, and (D), the intensity of Oat1 staining in the BLM of these tubules is strong. Scale bar=20 µm.

Figure 2 Comparison of proximal tubule morphology (A, C) and Oat1-immunostaining (B, D) in the same tubules between cryosections of the kidneys fixed in vitro (A, B) and in vivo (C, D) that were microwave-treated at pH 6. (A) Proximal tubules in kidneys fixed in vitro show collapsed lumina, filled with cellular debris, whereas (B), Oat1 in the BLM of the same tubules is strongly stained, but some tubules lost the BLM invaginations (arrows). (C) Proximal tubules in cryosections of the kidneys fixed in vivo are open, with sharp borders and regular appearance, and (D), the intensity of Oat1 staining in the BLM of these tubules is strong. Scale bar=20 µm.

Figure 3 Antigen retrieval conditions for Oat3 in untreated (A), SDS-treated (B), microwave-treated (C-E), and alcohol-treated (F-H) cryosections of the cortex of kidneys fixed in vivo. (A) Untreated cryosections show a weak staining of the BLM in various cortical tubules. (B) SDS-treated cryosections exhibit a significant non-specific background staining, and a hardly visible specific staining of the BLM. (C-E) Microwave-treated cryosections in citrate buffers of pH 3 (C), pH 6 (D), and pH 8 (E) exhibit significant BLM staining in various cortical tubules, the strongest being at pH 3 (C). (F-H) In cryosections pre-treated with alcohols, followed by microwave heating in citrate buffer of pH 3 (F), pH 6 (G), and pH 8 (H), BLM staining in the cortical tubules exhibits a similar pattern, but weaker intensity than in the microwave-treated cryosections. At pH 8, in both microwave-treated and alcohol-treated cryosections many tubules exhibit signs of detachment from the microscope slide (not shown). Scale bar=20 µm.
basolateral Oat3 staining in various cortical tubules, however, was similarly strong between the kidneys fixed in vitro (B) and in vivo (D).

**Optimal antigen retrieval conditions for brush-border transporters Oat2 and Oat5**

**Oat2**

Figure 5 shows antigen retrieval for immunostaining Oat2 in cryosections of the kidneys fixed in vivo. In the untreated cryosections (A), no significant staining was observed, whereas in the SDS-treated cryosections (B), Oat2 was weakly stained in the BBM of PT S3 segments in the outer stripe. In the microwave-treated cryosections in citrate buffer of pH 3 (C), we observed a significant non-specific background staining in all S3 segments. In addition, strongly stained were the BBM and non-identified intracellular organelles, mainly localised at the basal cell domain. However, at pH 6 (D), non-specific background staining was much lower, and only the BBM was positive, whereas
at pH 8 (E) only a weak staining of the BBM was observed. Cryosections pre-treated with alcohols, and microwave heated in citrate buffer, showed much weaker Oat2 staining intensity in the BBM at all pH values than the microwave only-treated cryosections.

Cryosections treated with microwave at pH 6 were used to compare morphology of the PT S3 segments and Oat2 staining in the same segments between kidneys fixed in vitro and in vivo (Figure 6). In the kidneys fixed in vitro, various tubule profiles had collapsed lumina, filled with cellular debris (A), whereas in the kidneys fixed in vivo, the tubules had regular appearance, with open lumina and sharp borders (C). The intensity of Oat2 staining in the BBM was stronger in the tubules of the kidneys fixed in vitro (compare B and D).

Oat5

Figure 7 shows antigen retrieval for immunostaining Oat5 in the tubules of the outer stripe in cryosections of

Figure 6 Comparison of proximal tubule S3 segment morphology (A, C) and Oat2 immunostaining (B, D) in the same tubules in the outer stripe between cryosections of the kidneys fixed in vitro (A, B) and in vivo (C, D) that were microwave-treated at pH 6. (A) All S3 segments in the kidneys fixed in vitro have collapsed lumina, filled with cellular debris, whereas (B), Oat2 in the BBM of the same tubules is strongly stained. (C) The proximal tubule S3 segments in cryosections of the kidneys fixed in vivo are open, with sharp borders, and regular appearance, and (D), the intensity of Oat2 staining in their BBM is much weaker than in cryosections of the kidneys fixed in vitro. Scale bar=20 μm

Figure 7 Antigen retrieval conditions for Oat5 in untreated (A), SDS-treated (B), microwave-treated (C-E), and alcohol-treated (F-H) cryosections of the outer stripe of kidneys fixed in vivo. (A) Untreated cryosections show no significant staining in the tubules of the outer stripe. (B) SDS-treated cryosections exhibit a weak staining of the BBM in the proximal tubule S3 segments. (C-E) Microwave-treated cryosections exhibit a significant background in all the tubules at pH 3 (C), and strong staining of the BBM in the proximal tubule S3 segments, whereas the individual cells (arrows) are strongly stained. At pH 6 (D), there is a strong staining of BBM, without a significant background. At pH 8 (E), the staining of BBM in the S3 is weak. (F-H) In cryosections pre-treated with alcohols, followed by microwave heating in citrate buffer of pH 3 (F), pH 6 (G), and pH 8 (H), the staining intensity of BBM in the proximal tubule S3 segments at pH 3 and pH 6 is much weaker than at pH 8, and similar to the staining in the respective microwave-treated cryosections. Scale bar=20 μm
the kidneys fixed in vivo. In the untreated cryosections, the tubules showed no significant staining (A), whereas in the SDS-treated cryosections, a weak staining of the BBM in PT S3 segments was observed (B). In the microwave-treated cryosections, overall staining intensity in the PT S3 segments was pH-dependent, exhibiting the pattern pH 3>pH 6>pH 8 (C, D, and E, respectively). The same pattern was observed for the intensity of non-specific background staining. In addition, at pH 3, individual cells were heavily stained in toto (C, arrows). In cryosections pre-treated with alcohols, followed by microwave heating in citrate buffer, background and BBM staining was much less intense, and exhibited an inverse pattern (pH 3<pH 6<pH 8).

**DISCUSSION**

Harsh antigen retrieval methods, such as delipidation with alcohols, microwave heating in buffers of different pH, and autoclaving, are usually used to recover cryptic epitopes in sections of formalin-fixed, paraffin-embedded tissues (1-10, 12). Such techniques have not regularly been used with cryosections. More is known about the use of SDS in cryosections of the PFA-fixed rat tissues and cultured cells. As shown in previous publications (12-14), SDS effects in cryosections are antigen-dependent; some invisible antigens can be revealed, the immunostaining of some antigens can be enhanced, whereas the staining intensity of others remains unchanged.

In order to define optimal conditions for immunostaining various Oats in cryosections of the PFA-fixed rat kidneys, we applied two stronger methods in this study in addition to SDS, that is, microwave heating and delipidation in alcohols plus microwave heating in citrate buffers of different pH. After applying these experimental conditions on cryosections of rat kidneys fixed in vivo, the data showed the following overall pattern of staining intensity of Oats: untreated < SDS-treated ≤ alcohol-treated < microwave-treated. Specifically, in untreated cryosections all Oats were barely stained, and SDS treatment slightly increased the staining of Oat1, Oat2, and Oat5, but not of Oat3. Alcohol pre-treatment and microwave heating enhanced the staining effect in all tested Oats. These data indicate that Oat epitopes may interact with lipids, and/or are deeply embedded (masked) in the cell membrane matrix, which protects from antibody binding, and are unmasked either by detergent treatment (weakly) or after denaturation by alcohols and/or heating in a microwave (stronger).

Due to lower background and strong BBM immunostaining in the BBM of PT S3 segments was higher in cryosections of the kidneys fixed in vitro.

**Figure 8** Comparison of proximal tubule S3 segment morphology (A, C) and Oat5 immunostaining (B, D) in the same tubules in the outer stripe between cryosections of the kidneys fixed in vitro (A, B) and in vivo (C, D) that were microwave-treated at pH 6. (A) All S3 segments in the kidneys fixed in vitro have collapsed lumina, filled with cellular debris, whereas (B) Oat5 in the BBM of the same tubules is strongly stained. (C) Proximal tubule S3 segments in cryosections of the kidneys fixed in vivo are open, with sharp borders and regular appearance, and (D) the intensity of Oat5 staining in their BBM is weaker than in cryosections of the kidneys fixed in vivo. Scale bar=20 µm

Compared to microwave treatment alone, pre-
treatment of cryosections with alcohols, followed by microwave heating, resulted in weaker staining intensity of all tested Oats, possibly because alcohols damaged the epitopes.

Although our results showed that antigen retrieval for rOat1 was best with microwave heating in citrate buffer of pH 8, this pH also caused a partial or complete detachment of cryosections from the glass slides. Therefore, it is not favourable for routine work.

We conclude that microwave treatment alone, in citrate buffer of pH 6 best reveals Oat1, Oat2, and Oat5 epitopes, whereas pH 3 is best for Oat3 epitopes. With this method and conditions, specific staining was strong, non-specific background staining was weak, and no staining artefacts were observed with the Oat-specific antibodies. At this point, we have to admit that our previous studies of Oat1, performed with SDS-specific antibodies. At this point, we have to admit that our previous studies of Oat1, performed with SDS-specific antibodies. At this point, we have to admit that our previous studies of Oat1, performed with SDS-specific antibodies. At this point, we have to admit that our previous studies of Oat1, performed with SDS-specific antibodies. At this point, we have to admit that our previous studies of Oat1, performed with SDS-specific antibodies.

As expected, tubules in cryosections from the kidneys fixed in vitro had collapsed lumina full of cell debris, possibly due to a hypoxic damage caused by prolonged fixation time in vitro. Judging by the open lumina and regular morphology, tubules fixed in vivo did not sustain such damage. However, under optimal immunocytochemical conditions, the staining intensity of basolateral transporters Oat1 and Oat3 was similar between cryosections of kidneys perfused in vitro and in vivo, whereas the staining intensity of brush-border transporters Oat2 and Oat5 was higher in cryosections of the kidneys fixed in vitro. These data suggest that PFA-fixation by perfusion in vivo is the method of choice for immunostaining basolateral transporters, whereas PFA-fixation in vitro seems to be better for presenting brush-border transporters. The reason for these differences is not clear, but several explanations come to mind: a) during fixation in vivo, the BBM may be overfixed, and the epitopes need a stronger “unmasking” treatment, b) the hypoxic damage of the BBM during slower fixation in vitro may have a kind of “digestive” effect and expose more epitopes, and c) the well known differences in the fluidity of these two membrane domains, determined by different content of phospholipids and cholesterol (21, and references in there), may be critical for epitope sensitivity to PFA fixation, heating, and pH.

In conclusion, our studies teach us that: a) some harsh antigen retrieval techniques, previously used with sections of formalin-fixed, paraffin-embedded tissues, can be efficiently used with cryosections of the PFA-fixed tissues, and b) specific Oats, and probably other renal transporters in the rat kidney and other mammalian organs have to be treated individually, with different fixation and antigen retrieval methods in order to define the optimal method and conditions for their immunocytochemical presentation.

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Sažetak

OPTIMALNE METODE OTKRIVANJA ANTIGENA ZA PRIJENOSNIKE ORGANSKIH ANIONA U KRIOSTATSKIM NARESCIMA ŠTAKORSKOG BUBREGA

Za imunocitokemijsku (IC) lokalizaciju antigena uzorci tkiva ili stanica obično se denaturiraju fiksacijom, a potom se odmah smrznu i naresci režu kriostatom ili se uklope u parafin prije rezanja narezaka. Parafin se kasnije odstrani alkoholima. p-Formaldehid (PFA; formalin) čest je fiksativ, koji čuva antigeničnost proteina, ali oštećuje morfologiju stanica i “maskira” vezna mjesta za protutijela (epitope). Za “demaskiranje” epitopa potrebno je primijeniti neku od metoda otkrivanja (regeneracije) antigena (RA). Namjera ove studije jest: a) odrediti optimalnu metodu RA u kriostatskim narescima PFA-fiksiranih bubrega in vivo za prijenosnike organskih aniona (Oat) smještene u bazolateralnoj (Oat1, Oat3) i četkastoj (Oat2, Oat5) membrani proksimalnih kanalića u bubrezima štakora i b) rabeći optimalnu metodu, usporediti IC bojenje Oat-a u bubrezima koji su bili PFA-fiksirani in vivo ili in vitro. IC bojenje u neobrađenim kriostatskim narescima uspoređeno je s bojenjem nakon obrade detergentom ili mikrovalnim kuhanjem u citratičnom puferu pH 3, pH 6 ili pH 8, s prethodnom obradom alkoholima ili bez nje. Optimalni RA-uvjet za Oat1, Oat2 i Oat5 bio je kuhanje narezaka pri pH 6, a za Oat3 pri pH 3, bez prethodne obrade alkoholima. U usporedbi s tkivom fiksiranim in vivo, tkivo fiksirano in vitro imalo je: oštećenu morfologiju kanalića, sličnu jačinu bojenja Oat1 i Oat3, a jače bojenje Oat2 i Oat5. Zaključujemo da svaki Oat u bubregu štakora treba obraditi pojedinačno, s različitim fiksacijom i metodom RA, kako bi se našli optimalni uvjeti za njegovo IC prikazivanje.

KLJUČNE RIJEČI: imunocitokemija, imunofluorescencija, kriostatski naresci, membranski prijenosnici, proksimalni kanalići, stanična membrana

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