Localization of Nuclear Subunits of Cyclic AMP–dependent Protein Kinase by the Immunocolloidal Gold Method

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ABSTRACT An immunocolloidal gold electron microscopy method is described allowing the ultrastructural localization and quantitation of the regulatory subunits RI and RII and the catalytic subunit C of cAMP-dependent protein kinase. Using a postembedding indirect immunogold labeling procedure that employs specific antisera, the catalytic and regulatory subunits were localized in electron-dense regions of the nucleus and in cytoplasmic areas with a minimum of nonspecific staining. Antigenic domains were localized in regions of the heterochromatin, nucleolus, interchromatin granules, and in the endoplasmic reticulum of different cell types, such as rat hepatocytes, ovarian granulosa cells, and spermatogonia, as well as cultured H411E hepatoma cells.

Morphometric quantitation of the relative staining density of nuclear antigens indicated a marked modulation of the number of subunits per unit area under various physiologic conditions. For instance, following partial hepatectomy in rats, the staining density of the nuclear RI and C subunits was markedly increased 16 h after surgery. Glucagon treatment of rats increased the staining density of only the nuclear catalytic subunit. Dibutyryl cAMP treatment of H411E hepatoma cells led to a marked increase in the nuclear staining density of all three subunits of cAMP-dependent protein kinase. These studies demonstrate that specific antisera against cAMP-dependent protein kinase subunits may be used in combination with immunogold electron microscopy to identify the ultrastructural location of the subunits and to provide a semi-quantitative estimate of their relative cellular density.

Cyclic AMP (cAMP) is the intracellular mediator of the regulatory signal of several polypeptide hormones and catecholamines. It plays a key role in the modulation of nuclear events such as the specific regulation of transcription of a number of cAMP-induced enzymes and proteins (1–4), phosphorylation/dephosphorylation of histones and nonhistone chromosomal proteins (5, 6), and nuclear mechanisms that may be involved in cAMP-regulated control of cellular proliferation (7). However, the precise sequence of molecular steps initiated by cAMP in these nuclear events remains to be elucidated. From our present knowledge of cAMP action it appears that cAMP-dependent protein kinase, the only identified mediator of cAMP action in eukaryotic cells, forms an important link in these nuclear events. Two isoenzyme forms of cAMP-dependent protein kinase (type I and type II) are found in mammalian tissues in various ratios depending on the cell type (8). These tetrameric holoenzymes contain a regulatory subunit dimer (R2) and two catalytic subunits (C) (9, 10). The type I and type II isoenzymes differ only in their regulatory subunits (RI and RII) which both are the major cAMP-binding proteins in eukaryotic cells identified so far. The potential relevance of the cAMP-dependent protein kinases to the understanding of differentiation and cellular growth is suggested by findings that the relative amounts of type I and type II isoenzymes change at times of physiological transitions (11–16) implying selective regulatory roles for the RI and RII subunits.

Although the cAMP-mediated nuclear events imply a functional association of protein kinase subunits with nuclear components, it is difficult to unambiguously identify a nuclear location of cAMP-dependent protein kinase subunits by clasp-
tical biochemical techniques involving cell homogenization and differential centrifugation. Possible perturbation of cellular structure during homogenization, the common use of nonphysiological homogenization media, and the susceptibility of the cytosol to protein kinase subunit redistribution during subcellular fractionation make an unequivocal demonstration of the presence or interaction of protein kinase subunits with the nucleus technically difficult. A valuable complementary approach to the problem of interaction and localization of protein kinase subunits within the cell is the application of histochemical or immunocytochemical techniques. With the introduction of immunogold-linked colloidal gold solutions (17) the localization and quantitation of antigen-antibody complexes has been greatly facilitated (18-22). Sufficiently electron-dense to be readily visible in the nucleus (23-25), immunogold reagents may be used in nuclear antigen labeling studies. In this paper we report the nuclear localization of the catalytic and regulatory subunits of cAMP-dependent protein kinase and demonstrate the applicability of immunogold electron microscopy in the identification and localization of the subunits at discrete nuclear substructures and to provide a semi-quantitative estimate of their cellular density.

MATERIALS AND METHODS

Cell Culture: A cloned cell line (H4IIE) derived from Reuber H35 hepatoma was kindly provided by Dr. Daryl Granner (University of Iowa). The cells were grown as monolayer cultures in Swimm's 77 medium supplemented with 2.5% fetal calf serum, 2.5% newborn calf serum, 50 U/ml of penicillin, 50 μg/ml of streptomycin, 2.4 mM CaCl2, 60 mM cystine, and 60 mM N-acetyl-L-cysteine (26). Cells were dehydrated with ethanol and embedded in a low viscosity resin according to the method of Spurr (26).

Preparation of Tissues: Liver, ovaries, and testes were taken from adult Sprague-Dawley rats (150-200 g) that had been killed by cervical dislocation. Small tissue sections (~1 mm3) were fixed according to the protocols given in the text (see Tables I and II). After fixation, the tissue sections were dehydrated in ethanol, and embedded in a low viscosity resin (26). The embedding plastic had no effect on the distribution of immunogold staining using all three antisera (Tables I and II).

Preparation of Tissue Sections and Staining Protocol: Thin sections (70-80-nm thick) were cut on an ultramicrotome and mounted on 200-nm nickel grids. To localize antigens, we processed mounted tissue sections by an indirect immunogold staining method similar to ones previously described (18, 27-29). All grids were floated with the tissue section face down on 50-μl drops at room temperature. Mounted tissue sections were first etched with 10% H3O2 for 10 min to enhance immunostaining (30-31). After a rinse in distilled water, nonspecific sites of protein absorption were blocked with a 1:10 dilution of normal goat serum in PBS for 5 min. After the blocking step, the grids were incubated for 60 min with the respective antiserum or preimmune serum diluted in PBS as indicated in the text. After a rinse in PBS, the grids were floated for 30 min on a 1:2 dilution of goat anti-rabbit immunoglobulin linked to 20-nm colloidal gold particles (Janssen Pharmaceuticals, Beerse, Belgium). After a rinse with phosphate-buffered saline (PBS), nuclear morphology was enhanced by staining with 3% aqueous uranyl acetate for 10 min followed by lead citrate for 5 min (32). The sections were examined using a JEOL JEM-100 CX electron microscope operated at 80 keV and photographed with Kodak 4489 EM film.

Quantitative Evaluation by Morphometric Analysis: Morphometric quantitation of immunogold staining was carried out in at least 10 different cells from each of three different immunogold preparations. Each micrograph was photographed at an initial magnification of 7,200 and analyzed at a final magnification of 72,000 by placing each micrograph negative in an optical enlarging bench equipped with a Numonics digitizer operated with a Hewlett-Packard 9820A computer programmed to record the number of gold particles per unit area. Immunogold labeling is expressed as the number of colloidal gold particles per cellular component unit area.

Purification of Antigen: The regulatory subunits RI and RII were prepared and purified from rat liver according to Dills et al. (33). Bovine catalytic subunit was purified as described previously (34). Anti-catalytic subunit RI and RII were prepared in rabbits using highly purified RI and RII. Non-immune control

Table I. Staining Intensity in Rat Hepatocytes Obtained for the Catalytic Subunit of cAMP-dependent Protein Kinase Using Different Fixation Conditions

<table>
<thead>
<tr>
<th>Fixation</th>
<th>Postfixation</th>
<th>Media</th>
<th>Intensity of labeling (gold particles/μm² ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.5% Glutaraldehyde</td>
<td>1% OsO4</td>
<td>Spurr</td>
<td>16.3 ± 1.4, 15.9 ± 1.3, 2.5 ± 0.2</td>
</tr>
<tr>
<td>2.5% Glutaraldehyde</td>
<td>-</td>
<td>Spurr</td>
<td>39.8 ± 1.2, 23.0 ± 1.0, 2.5 ± 0.2</td>
</tr>
<tr>
<td>2.5% Glutaraldehyde</td>
<td>1% OsO4</td>
<td>PEG-GMA</td>
<td>ND</td>
</tr>
<tr>
<td>1.0% Glutaraldehyde</td>
<td>-</td>
<td>Spurr</td>
<td>20.5 ± 2.2, 15.4 ± 1.2, 1.1 ± 0.2</td>
</tr>
<tr>
<td>1.0% Glutaraldehyde</td>
<td>-</td>
<td>PEG-GMA</td>
<td>ND</td>
</tr>
<tr>
<td>4.0% Paraglutaraldehyde + 0.5% Glutaraldehyde</td>
<td>-</td>
<td>Spurr</td>
<td>9.1 ± 1.0, 9.7 ± 1.1, 1.3 ± 0.1</td>
</tr>
<tr>
<td>Dimethylsuberimidate (20 mg/ml)</td>
<td>1% OsO4</td>
<td>Spurr</td>
<td>11.3 ± 0.8, 12.5 ± 0.2, 0.8 ± 0.1</td>
</tr>
<tr>
<td>Dimethylsuberimidate (20 mg/ml)</td>
<td>-</td>
<td>Spurr</td>
<td>9.0 ± 0.7, 6.9 ± 1.3, 1.1 ± 0.1</td>
</tr>
<tr>
<td>Embedding medium alone</td>
<td>-</td>
<td>Spurr</td>
<td>0.3 ± 0.1</td>
</tr>
</tbody>
</table>

NO, not determined because of high nonspecific binding over resin.

Table II. Staining Intensity Obtained for the Regulatory Subunits RI and RII of cAMP-dependent Protein Kinase in Rat Hepatocytes

<table>
<thead>
<tr>
<th>Subunit</th>
<th>Fixation</th>
<th>Intensity of labeling (gold particles/μm² ± SEM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RI</td>
<td>2.5% Glutaraldehyde</td>
<td>39.8 ± 1.4, 49.1 ± 3.3, 2.9 ± 0.2</td>
</tr>
<tr>
<td>RII</td>
<td>2.5% Glutaraldehyde</td>
<td>27.2 ± 1.7, 31.6 ± 1.7, 2.4 ± 0.3</td>
</tr>
</tbody>
</table>

Embedding medium: 0.2 ± 0.1, 0.3 ± 0.1
immunoglobulins were prepared by the same method from sera of uninjected rabbits. Affinity purification of the antisera was carried out as described by Miles et al. (36). The specificity of the antisera was tested extensively by Ouchterlony double-diffusion analysis (37), double-antibody precipitation, indirect enzyme-linked immunosorbent assay (ELISA) (33), and immunoblot analysis (38).

**Immunoblot Analysis:** Purified antigens and either whole cell protein or nuclear protein extracts were subjected to electrophoresis on SDS 10% polyacrylamide gels (39). The proteins were transferred to nitrocellulose paper (38) using an Electroblot apparatus (E-C Apparatus Corp., St. Petersburg, FL). Experiments were carried out to determine the efficiency of protein transfer using [35S]methionine-labeled H4IIE cytosol. After 3 h of electroblotting, we observed complete transfer of the soluble [35S]-labeled proteins from the gel to the nitrocellulose membrane (see Fig. 2). After electroblotting, the nitrocellulose membrane was stored overnight at 4°C in transfer buffer (0.025 M Tris, pH 8.3, 0.192 M glycine, and 20% methanol [vol/vol]). Remaining protein binding sites on the membrane were blocked with 3% bovine serum albumin in 0.05 M Tris, pH 7.5, and 0.2 M NaCl (TBS) for 1 h at room temperature using gentle agitation. After blocking, the membrane was placed into a primary antibody solution (anti-RI [1:100], anti-RII [1:100], or anti-C [1:250] in TBS) for 24 h at 4°C with gentle agitation. The membrane was then washed three times in TBS containing 0.1% Tween 20 (15 min/wash). Identification of antigen was carried out using the BRL Immunodetection Kit (Bethesda Research Laboratories Inc., Rockville, MD). Accordingly, secondary antibody (affinity-purified biotinylated goat anti-rabbit IgG) was diluted to 3 μg/ml in TBS. The membrane was incubated in secondary antibody solution for 1 h at room temperature using gentle agitation. After incubation, the membrane was washed three times in TBS and placed in a solution of streptavidin-HRP enzyme conjugate (diluted 1:500 in TBS) for 30 min at room temperature. The membrane was again washed three times in TBS and then immersed into substrate solution (0.5 mg/ml of 4-chloro-1-napthol in TBS). The membrane was incubated in substrate solution without agitation at room temperature for 30 to 60 min or until a blue-purple color was observed indicating antibody-antigen conjugation.

**RESULTS**

**Analysis of Antisera**

The preparation and characterization of antiserum against bovine heart catalytic subunit and its cross-reactivity with rat liver catalytic subunit have been described in detail (34, 35). Antisera against rat liver regulatory subunits RI and RII were raised in rabbits by injecting them with purified antigen using an identical procedure. The specificities of the antisera were determined in several ways. Double immunodiffusion Ouchterlony analysis of the antisera and purified rat liver RI, RII, or C and crude rat liver 105,000 g supernatant fraction showed in each case a single continuous immunoprecipitation band (results not shown). This suggested the presence of a single antibody-antigen system. For the competitive ELISA, the RI and RII antisera were preincubated with either RI or RII and then added to the antigen-coated microtiter wells. As shown in Fig. 1, only the complementary antigen eliminated the reactivity of the respective antisera toward RI or RII. The noncomplementary antigen did not compete even at 100-fold the concentration of complementary antigen with which competition was first detected. Western blot analysis of the affinity-purified RI, RII, and C antisera showed that all three specifically reacted with their complementary antigen in a whole cell protein extract from rat liver, ovary, or H4IIE cells (Fig. 2, lower panels, A–C). Preimmune serum showed no reactivity (Fig. 2, lower panels, D). Similarly, all three antisera reacted with their complementary antigen in a rat liver nuclear protein extract (Fig. 3) and only minor apparent nonspecific bands are detected (the peak with the shoulder in scan E represents the phospho- and dephosphoforms of RII). It is important to note that Western blot analysis was done using either a 1:100 dilution (anti-RI or anti-RII) or a 1:250 dilution (anti-C) of antiserum, whereas immunogold antigen detection was done with a substantially greater dilution of antiserum, further diluting out any potential cross-reacting contaminating antibodies.

**Effect of Tissue Fixation**

The labeling intensity on thin sections fixed under various conditions is presented in Tables I and II. Best results were obtained with a fixation protocol including 2.5% glutaraldehyde and elimination of osmium tetroxide. Intense, specific staining was obtained with 2.5% glutaraldehyde as fixative in nuclear and cytoplasmic areas, whereas only negligible labeling was found in sinusoidal areas or the embedding medium alone. The optimal dilutions of the antisera and corresponding preimmune sera for a 60-min incubation time with the tissue sections were 1:500 for the catalytic subunit antisera and 1:2,000 for the RI and RII regulatory subunit antisera. Higher concentrations usually led to an increase of nonspecific background labeling. Counterstaining of thin sections with uranyl acetate and lead citrate did not affect the staining intensity (data not shown). For polyethylene glycol–glycol methacry-
FIGURE 2 Analysis of antibody specificity by immunoblotting. (Upper panels) Protein transfer to nitrocellulose. H4IIIE cells were incubated with 50 μCi of [35S]methionine for 24 h. Cytosol protein was prepared, resolved on SDS 10% polyacrylamide gels, and transferred to nitrocellulose by electroblotting. The blotted gel and nitrocellulose membrane were dried and placed on a Kodak X-omat XRP-5 film for 3 d to obtain autoradiographs. The transferred 35S-labeled proteins are shown in lane 1. Labeled proteins remaining in the gel after blotting are shown in lane 2. (Lower panels) Immunodetection of protein kinase subunits. Whole cell extracts from rat liver, rat ovary, or cultured H4IIIE cells were separated by electrophoresis on SDS 10% polyacrylamide gels and transferred to nitrocellulose by electroblotting. Soluble proteins resolved by electrophoresis from rat liver (lane 1), H4IIIE cells (lane 2), and rat ovary (lane 3) were incubated with a 1:100 dilution of either anti-RII (A), anti R-I (B), a 1:250 dilution of anti C (C), or preimmune serum (D). Immunocomplexes were visualized with the BRL immunodetection kit as described in Materials and Methods. The molecular weights (× 10^3) of RI, RII, and C are indicated and were determined using standard protein markers.

FIGURE 3 Laser beam densitometric scans of nuclear protein immunoblot autoradiograms. Rat liver nuclear 0.35 M NaCl extracts, late (PEG-GMA)–embedded thin sections, staining of all cellular structures was always accompanied with high background staining over the resin which was not the case when Spurr’s embedding medium was used (Table I).

Controls for Specificity of Immunogold Staining

Three different types of controls were performed to test for the specificity of immunogold labeling. (a) After preincubation of thin sections with IgG from the respective preimmune sera followed by colloidal gold–linked goat anti-rabbit immunoglobulin only minimal immunogold staining was observed (Fig. 4c). (b) Almost complete abolition of staining was observed after preincubation of thin sections with the respective antigen or direct application of the colloidal gold–linked goat anti-rabbit immunoglobulin (micrographs not shown). (c) Finally, preabsorption of antiserum with excess prepared as described previously (7), were separated by electrophoresis on 10% polyacrylamide gels and transferred to nitrocellulose by electroblotting. Scan A was stained for protein with Amido black. The additional scans were incubated with a 1:100 dilution of either preimmune serum (B), anti-C (C), anti-RI (D), or anti-RII (E). Immunocomplexes were visualized with the BRL immunodetection kit as described in Materials and Methods. The molecular weights (× 10^3) of standard protein markers are indicated at the top of the figure.
Liver tissue thin sections were prepared from male Sprague-Dawley rats (150–200 g) that had received an intraperitoneal injection of either 0.1 ml of saline or 0.4 mg of glucagon (Eli Lilly & Co., Indianapolis, IN) for 2 h or had been partially hepatectomized for 16 h as described (7). Fixation of tissue was carried out with 2.5% glutaraldehyde and embedding was in low-viscosity resin according to the method of Spurr (26). (a) Micrograph of a hepatocyte from a saline-injected rat stained with anti-C serum. Nu, nuclear area; Cy, cytoplasmic area. (Inset) Low magnification showing preferential localization of the catalytic subunit over the peripheral area of the nucleolus. Arrows in the inset denote the region of higher magnification. Bar, 0.5 μm; × 10,750. (b) Micrograph of a hepatocyte from a glucagon-treated rat. (Inset) Low magnification showing relatively dense catalytic subunit staining. Arrows in inset denote the region of higher magnification. Bar, 0.5 μm; × 51,600; (inset) bar, 1.0 μm; × 12,000. (c) Micrograph of a hepatocyte from a saline-injected rat. The tissue section was treated with pre-immune serum diluted 1:500. Note absence of immunostaining. × 41,000. (d) Micrograph of a hepatocyte from a saline-injected rat. The tissue section was stained with anti-C serum that had been incubated with excess purified catalytic subunit. Note absence of immunostaining. Bar, 1.0 μm. × 41,000. (e) Micrograph of nuclear area in a hepatocyte from a partially hepatectomized rat showing staining over nucleolus and in peripheral areas of the nucleolus. Bar, 0.5 μm; × 50,000.
of rat tissues with gold are shown in Figs. 4-7. The well-described biochemical characteristics of the cAMP-dependent protein kinase in the tissues studied (7, 12, 16) made them an ideal choice for the immunogold labeling studies. The ultrastructural localization of the subunits was accomplished through the use of the specific antisera in combination with a 20-nm colloidal gold-linked goat anti-rabbit antibody. In Fig. 4 cytoplasmic and nuclear areas of a hepatocyte from either saline-treated (Fig. 4a) or glucagon-treated (Fig. 4b) rats are shown stained with anti-C. Immunogold staining was associated with the endoplasmic reticulum (see especially Fig. 4b), with heterochromatin and with nucleolar regions (see arrows in Fig. 4a). After glucagon treatment, a higher labeling density of catalytic subunit was observed within the nucleus (compare inset of Fig. 4a with inset of Fig. 4b), and to a lesser extent also in cytoplasmic regions. Association of catalytic subunit with the nucleolus and heterochromatin areas in hepatic nuclei from partially hepatectomized rats is shown in Fig. 4c. Immunostaining of catalytic subunit in euchromatin appeared at low levels in all hepatocyte nuclei examined.

The staining and distribution of the regulatory subunits RI and RII in hepatocytes from regenerating rat liver are shown in Fig. 5. Both regulatory subunits are located over the endoplasmic reticulum (Figs. 5, a-c) and over nuclear areas particularly in heterochromatin and nucleolar areas (Figs. 5, a and b) and perichromatin regions (Fig. 5c, arrows and higher magnification of this region shown in Fig. 5d).

Fig. 6 shows the localization of the catalytic and RII sub-
units in H4IIE hepatoma cells. These cells contain relatively low levels of type I cAMP-dependent protein kinase (Squinto, S. P., and Jungmann, R. A., unpublished observation) and RI staining in the nucleus was generally much lower than RII and C staining (see also Table IV). Regulatory subunit RII was distributed over cytoplasmic and nuclear areas (Fig. 6d, inset and higher magnification). The catalytic subunit showed a preferential association with nuclear areas, particularly with heterochromatin (Fig. 6a, top of micrograph) and with interchromatin granules (Fig. 6a, arrows, and higher magnification of that area shown in Fig. 6b). After dibutyryl cAMP treatment of H4IIE hepatoma cells, the catalytic subunit was distributed rather densely over heterochromatin (lower left corner of Fig. 6c) and over interchromatin granules (Fig. 6c, arrow and inset).

The subunit distribution was also examined in rat ovarian granulosa cells and rat spermatogonia. Fig. 7c shows a rather dense distribution of catalytic subunit in the heterochromatin and perichromatin region of the nucleus of a spermatogonium. The staining density in the cytoplasm, in contrast to nuclear areas, is considerably less. Rat granulosa cells show staining for the catalytic subunit (Fig. 7a), the regulatory
subunit RII (Fig. 7b), and RI (Fig. 7d). All three subunits appear to be primarily associated with heterochromatin and perichromatin regions (see, for instance, Fig. 7b, arrow) and in areas of the endoplasmic reticulum (Figs. 7, a and b).

Morphometric Analysis of Subunit Density

Table III lists the subunit densities of spermatogonia from two-wk-old rats and of thecal and granulosa cells from a small antral follicle isolated from the ovary of 33-d-old rats. While thecal cells show a slightly higher RI subunit density, the regulatory subunit RII density is markedly higher in granulosa cells. This is in apparent agreement with biochemical studies in which relatively higher levels of RII than RI were identified in rat granulosa cells (40).

The quantitative evaluation of immunogold labeling over rat hepatocytes from glucagon-treated and partially hepatectomized rats has previously been published (41, 42). A modulation of subunit labeling density was observed in hepatocytes after glucagon administration as well as after partial hepatectomy. After glucagon administration, the labeling density of the catalytic subunit in the nucleus increased about 2.5 to 3-fold and to a lesser degree (1.5- to 1.9-fold) in extranuclear areas. RI and RII labeling densities were, however, not affected by glucagon. Partial hepatectomy elicited a marked increase of staining density of RI and C but not of RII over
the nuclear area. Cytoplasmic staining density was not markedly altered after partial hepatectomy.

Quantitation of the subunit density in dibutyryl cAMP-treated H4IIE hepatoma cells is shown in Table IV. These data allow a comparison of the relative staining densities of each subunit in untreated versus dibutyryl cAMP-treated cells. The staining density of all three subunits changed to varying degrees after dibutyryl cAMP treatment. The nuclear catalytic subunit staining density increased significantly \((p < 0.001) \) 10 min (1.7-fold increase) after stimulation. Additionally, the nuclear RI subunit showed an increased labeling at 15 min (2.6-fold increase, \( p < 0.01 \)). RI labeling peaked 10 min (2.3-fold increase, \( p < 0.001 \)) and 30 min (1.5-fold increase, \( p < 0.05 \)) after cyclic nucleotide treatment. There was a slight decrease of cytoplasmic RI staining in the cytoplasm, whereas cytoplasmic RII and C staining did not change significantly.

**DISCUSSION**

Through the use of a newly developed post-embedding immunogold technique, we have localized all three subunits of cAMP-dependent protein kinase within the nucleus and in extranuclear areas of hepatocytes, ovarian granulosa and thecal cells, spermatogonia, and cultured H4IIE hepatoma cells. Evaluation of the immunogold staining patterns showed that the three subunits are selectively localized over several different nuclear substructures: nucleolus, condensed (hetero) chromatin, interchromatin granules, and perichromatin regions. The specificity of antigen–antibody interaction was carefully evaluated and demonstrated by the ELISA and immunoblot analysis. Immunoblotting is limited to the transfer of soluble protein antigens. Therefore, it is conceivable that insoluble potential cross-reacting antigens are not transferred from the gel to the nitrocellulose membrane. However, we find no indication of this from the autoradiographs (Fig. 2). Controlled studies using preimmune sera or staining of tissue with antisera that had been preadsorbed with purified antigen revealed insignificant nonspecific adsorption of the colloidal gold reagent to any intracellular compartment. In addition, staining of sinusoidal areas of rat liver sections was at background level.

The ultrastructural localization of all three subunits in the nucleus of various cell types was achieved using a colloidal gold–linked second antibody reagent. The immunogold technique has been used previously to investigate the localization of antigen in the nucleus of euakaryotic cells (23–25, 41, 42, 43). Immunogold reagents are ideal probes for localizing antigen–antibody complexes and exhibit distinct advantages over immunofluorescent techniques or conventional stains such as horseradish peroxidase and ferritin. The discrete size of the colloidal gold reagent and the high electron density of gold allows for easy visualization of antibody binding sites over electron-dense regions of the nucleus such as heterochromatin and the nucleolus. Additionally, in contrast to immunofluorescence, immunogold complexes offer a high degree of resolution with a low degree of nonspecific staining and allow a semi-quantitative evaluation of staining density. The availability of differing sizes of colloidal gold–linked secondary antibody has allowed for the design of experiments using more than one antisera label. Thus, we are presently defining the exact spatial relationships of the RI, RII, and C

<table>
<thead>
<tr>
<th>Cell type</th>
<th>Nucleus</th>
<th>Cytoplasm</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>RI</td>
<td>RII</td>
</tr>
<tr>
<td>Granulosa cells</td>
<td>66.2 ± 2.5</td>
<td>83.6 ± 2.4</td>
</tr>
<tr>
<td>Thecal cells</td>
<td>81.0 ± 2.7</td>
<td>69.1 ± 5.3</td>
</tr>
<tr>
<td>Spermatogonia</td>
<td>15.2 ± 0.6</td>
<td>12.7 ± 1.1</td>
</tr>
</tbody>
</table>

Tissue sections were fixed in 2.5% glutaraldehyde and processed using the immunogold procedure as described in Materials and Methods. The spermatogonia examined were from the testicular tissues of a 2-wk-old rat. Granulosa and thecal cells were from small antral follicles isolated from 33-d-old rats.

<table>
<thead>
<tr>
<th>Time after dibutyryl cAMP treatment</th>
<th>Staining Intensity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Nucleus (Gold particles/μm² ± SEM)</td>
</tr>
<tr>
<td></td>
<td>RI</td>
</tr>
<tr>
<td>0</td>
<td>13.4 ± 1.6 (6)</td>
</tr>
<tr>
<td>2</td>
<td>22.1 ± 1.9 (6)</td>
</tr>
<tr>
<td>5</td>
<td>25.6 ± 2.3 (6)</td>
</tr>
<tr>
<td>10</td>
<td>20.9 ± 3.3 (5)</td>
</tr>
<tr>
<td>15</td>
<td>34.9 ± 2.5 (5)</td>
</tr>
<tr>
<td>30</td>
<td>14.1 ± 2.2 (5)</td>
</tr>
<tr>
<td>60</td>
<td>25.5 ± 2.6 (6)</td>
</tr>
<tr>
<td>120</td>
<td>41.3 ± 5.3 (5)</td>
</tr>
</tbody>
</table>

H4IIE hepatoma cells were stimulated with 0.5 mM dibutyryl cAMP for the time periods indicated. Cells were fixed in 2.5% glutaraldehyde and processed as described in Materials and Methods. The numbers in parentheses represents number of cells analyzed.
subunits within the nucleus.

Optimal nuclear immunolabeling with an acceptably preserved nuclear ultrastructure was obtained with 2.5% glutaraldehyde as fixative and a low viscosity embedding resin. Under these experimental conditions the three nuclear subunits are relatively resistant to the destructive potential of fixation, dehydration, and embedding. Cytoplasmic labeling is similarly optimal under these experimental conditions. However, direct quantitative comparisons between nuclear and cytoplasmic subunit staining density should be done with caution. Possible differences of the binding affinities of the protein kinase subunits to nuclear substructures and to extranuclear regions, and a possible preferential loss of cytoplasmic antibody during tissue processing, may make these comparisons imprecise. Also, because of differing antibody–antigen affinities, a direct comparison of the labeling densities of the individual subunits should be done with caution.

Quantitative evaluation of the immunogold staining patterns by morphometric analysis demonstrated several notable changes of nuclear antigen labeling densities at times of cellular stimulation by effector agents. As we have previously shown, glucagon stimulation elicited an increased nuclear staining of only the catalytic subunit (41), whereas the staining density of both the nuclear catalytic and regulatory subunit RI was increased 16 h after partial hepatectomy (42). In the present study, we additionally demonstrated that dibutyryl cAMP treatment of H4IIIE hepatoma cells leads to a modulation of nuclear RI, RII, and C subunit densities (see Table IV).

The altered staining densities of the nuclear cAMP-dependent protein kinase subunits that occurred in response to partial hepatectomy correlate well with the observed subunit modulation during the prereplicative phase of rat liver regeneration identified by biochemical and immunochromatographic measurements (7). Similarly, glucagon treatment of rats has been shown to increase the nuclear activity level of catalytic subunit (44, 45).

Several factors may be responsible for the changing antigen densities. Firstly, the increased nuclear staining density is most likely the result of a net increase of antigen molecules in the nucleus as the consequence of translocation of antigen from extranuclear sites into the nucleus (46). The localization of antigen over nuclear substructures and cytoplasm and changing antigen densities in the nucleus may indicate that the antigens shuttle between nuclear and cytoplasmic structures and that this may play an important functional role in nuclear–cytoplasmic communication. Secondly, although the RI, RII, and C-antiseras used in our studies have identical affinities for their complementary antigen in either the nonassociating holoenzyme form or in the dissociated form as identified by the ELISA (34, 35), they could conceivably interact only with the dissociated subunits in tissue sections. Should this be the case, increased staining density would occur after cellular stimulation reflecting the formation of free subunits as the result of cAMP-mediated dissociation of the cAMP-dependent protein kinase holoenzymes. However, we did not observe a concomitant modulation of cytoplasmic staining density which, therefore, makes this latter possibility unlikely.

In conclusion, the aim of the present study has been directed toward the immunocytochemical identification of the unique nuclear sites of RI, RII, and C subunit binding using specific antisera and immunogold electron microscopy. The results reported here demonstrate the usefulness of immunogold reagents in the cytochemical study of cAMP-dependent protein kinase subunits and extend earlier work on nuclear cAMP-dependent protein kinase subunit localization using immunofluorescence at the light microscopy level (47–49). The technique appears sensitive and specific and the resolution allows a precise correlation between antigen and antigenic binding sites. Potential applications of this technique are numerous, particularly for the identification of the nuclear sites of action of the cAMP-dependent protein kinase subunits under a variety of physiological conditions.

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REFERENCES


