Effect of Colchicine on Internalization of Prolactin in Female Rat Liver: An In Vivo Radioautographic Study

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ABSTRACT Binding and internalization of 125I-ovine prolactin into hepatocytes of female rats was visualized by the in vivo radioautographic method (Bergeron, J. J. M., G. Levine, R. Sikstrom, D. O'Shaughnesssey, B. Kopriwa, N. J. Nadler, and B. I. Posner, 1977, Proc. Natl. Acad. Sci. USA, 74:551-555). Receptor-mediated internalization of label was observed into lipoprotein-filled vesicles in the Golgi/bile canalicular region of the hepatocyte. Colchicine treatment had no effect on the internalization of label into the lipoprotein-filled vesicles. However, the location of the radio-labeled lipoprotein-filled vesicles was altered from the Golgi/bile canalicular region to subsinusoidal. Radioactive content of hepatocytes decreased as a function of time after injection of 125I-prolactin; however, colchicine treatment markedly retarded this loss of label. Subcellular fractionation experiments indicated that colchicine treatment led to decreased levels of 125I-prolactin accumulation in microsomes but augmented the accumulation of label in the L fraction.

It is concluded that in normal female rats prolactin is internalized into lipoprotein-filled vesicles in the Golgi region before degradation of the hormone. Colchicine treatment accumulates labeled lipoprotein-containing vesicles in a subsinusoidal region and retards hormone catabolism. The labeled vesicles observed after colchicine treatment may correspond to the unique vesicles previously observed in the L fraction and found to be enriched in prolactin receptors (Khan, M. N., B. I. Posner, A. K. Verma, R. J. Khan, and J. J. M. Bergeron, 1981, Proc. Natl. Acad. Sci. USA, 78:4980-4981).

Prolactin receptors have been shown to be highly concentrated in microosomal fractions isolated from homogenates of livers from female as opposed to male rats (19, 28). Using subcellular fractionation procedures, we observed that 125I-ovine prolactin (oPRL) administered iv was internalized, via a receptor-mediated process, into rat liver and concentrated in a substantially intact form in Golgi vesicular elements (17, 29). Colchicine treatment was shown to decrease the initial uptake of 125I-oPRL into Golgi vesicles (26). Since other investigators using radioautography have described the uptake of 125I-labeled hormones into secondary lysosomes (1, 10, 15) we examined the uptake and distribution of 125I-oPRL in liver tissue after hormone injection by the in vivo radioautographic technique (5, 8). A major aim was to correlate our observations obtained with subcellular fractionation procedures (17, 26, 29) and those with in vivo radioautography in order to define with increased certainty those structures accumulating internalized hormone.

Our studies demonstrate that 125I-prolactin is bound to female rat hepatocytes to a much greater degree than male, in conformity with previous observations of in vitro binding (19, 28). They further show that 125I-prolactin is rapidly internalized into hepatocytes and concentrated in lipoprotein-containing vesicles in the Golgi region of the cell. Colchicine treatment did not prevent the progressive accumulation of 125I-prolactin into lipoprotein-containing vesicles but altered their cellular location from the Golgi complex to the subsinusoidal region of the hepatocyte.

MATERIALS AND METHODS

Animals: Young adult male and female Sprague-Dawley (Canadian Breeding Farms, St. Constant, Quebec) or Sherman (McGill Anatomy colony) rats (180-280 g) were used. Colchicine was purchased from Sigma Chemical Co. (St. Louis, MO) and a freshly prepared solution of colchicine in saline was administered intraperitoneally s.c. 10 mg/ml per 100 g body weight 3 h before injection of 125I-oPRL.

Hormone: oPRL (NIH-P-S10, 26.4 IU/mg) was generously provided by the Pituitary Hormone Distribution Program of the National Institute of Arthritis, Metabolism and Digestive Diseases and iodinated as described previously (17, 29) to specific activities of 114-152 μCi/μg.

Injection of Radio-labeled Hormone and Tissue Fixation: The protocol has been described previously (5, 8). Briefly, the rats were...
anaesthetized with i.p. injections of 0.06 mg Nembutal per 100 g body weight. The radio-labeled hormone, or radio-labeled mixed with unlabeled hormone, was injected into the external jugular vein, usually the left, following a paramedian supraclavicular exposure. The hormone was allowed to circulate for 2 to 60 min. The heart was exposed by a midline ventral incision, anterior incision of the diaphragm and paramedian severance of ribs. Lactated Ringer's solution was perfused via the left ventricle until the liver visibly blanched (~30 s), at which time the perfusate was changed to 2.5% glutaraldehyde in 0.1 M Sörenson's buffer/0.1% sucrose, pH 7.3. Small samples of the midquadrate lobe were removed and diced into 1- to 2-mm cubes. After 2 h in the same fixative as the perfusate, the samples were washed in 0.1 M cacodylate buffer, pH 7.4, and immersed in 2% OsO4 (0.1 M cacodylate, pH 7.4) or reduced OsO4 (11) for 2 h. After further rinsing in buffer the samples were stained en bloc with uranyl acetate or galloyl glucose (5, 8). Following dehydration the blocks were embedded in Epon 812.

For some experiments organs were dissected out after glutaraldehyde perfusion, blotted, weighed, and radioactive content was evaluated by gamma spectrometry (Packard autogamma counter; 40.5% efficiency).

Light and Electron Microscope Radioautography: For light microscopy, semithin (0.5-µm) sections were pre-stained with iron haematoxylin and coated with Kodak NTB2 emulsion (23) and developed with freshly prepared Kodak D-170 developer (23). For electron microscopy, thin sections (silver-gray) were cut on an LKB-Huxley ultramicrotome (LKB Instruments, Inc., Rockville, MD), and a monolayer of Ilford L-4 emulsion was applied (21). Development was carried out either for large filamentous grains (chemical development) or for fine grains ("solution-physical" development) as described by Kopriwa (22). Sections were poststained with uranyl acetate and/or lead citrate and photographed with either a Siemens 1A, Siemens 101 or Philips 400 electron microscope.

Quantitative analysis of light microscope radioautographs was carried out as described previously (5, 8). Exposed light microscope slides were evaluated at × 1,000 using a calibrated ocular equivalent to a frame size of 11 × 11 µm on the slide. For light microscopy greater than 1,000 grains were counted for each determination.

The electron microscope radioautographs when developed for filamentous grains (2 min time point) were photographed (× 10,000) in such a way as to ensure that an hepatocyte cell membrane was in view whenever possible. The distribution of filamentous grains from the nearest cell membrane was determined by placing a pinhole in the geometric center of the grain and measuring the distance to the nearest hepatocyte plasmalemma (5). Filamentous grains were also analyzed by scoring the structure beneath the grain (5, 8). The micrographs developed for fine grain analysis (2-30 min) were photographed differently, i.e. wherever there were grains, with no consideration as to whether a cell membrane was in view. The results were based on the analysis of 266 filamentous grains and 2,943 fine grains. A total of 22 cellular compartments were analyzed, of which only the major labeled compartments are noted in Results.

Figure 1 Light microscope radioautographs of Epon sections of livers from female (a) and male (b) rats (285 g body weight, 82 d of age) at 2 min after injection of 125I-oPRL (1.8 × 10⁶ dpm; 156 µCi/µg). At 40-d exposure, the silver grain density was 3.54 ± 0.28 grains per frame (11 µm) for the female and 0.81 ± 0.12 grains per frame for the male. Student's t test revealed a significance of P < 0.001. × 1,000.

Figure 2 Light microscope radioautographs of Epon sections from livers of adult female rats at 2 min after the injection of 125I-oPRL (5.61 × 10⁸ dpm; 152 µCi/µg) (a) and injected with the same dose of 125I-oPRL but with an excess (1,000 µg) of unlabeled ovine prolactin (b). At 16-d exposure, silver grain density was 8.01 ± 0.36 grains per frame for the sections from the experimental animals (a) as compared to 1.79 ± 0.12 grains per frame for the control (b). Student's t test revealed this difference as significant (P < 0.001). × 1,000.
Subcellular Fractionation: Liver homogenates from colchicine-treated and control female rats were fractionated into nuclear-mitochondrial pellets, L fraction, microsomes (P), and final supernatant (S) following the protocol described previously (20) based on the method described by Wattiaux et al. (39) and de Duve et al. (12).

RESULTS

Initial Binding In Vivo

Injection of $^{125}$I-oPRL into female rats resulted in silver grains over liver sections at a 4.4-fold higher grain concentration than over liver sections from identically injected males (Fig. 1). In experiments in which excess unlabeled prolactin was coinjected with labeled hormone into female animals (Fig. 2) a 4.5-fold reduction in grain concentration was observed when compared to sections from female animals injected with $^{125}$I-oPRL alone. In contrast no competition by excess unlabeled prolactin was observed over the proximal convoluted tubule of the kidney (Fig. 3).

In a separate experiment carried out to evaluate the proportion of injected $^{125}$I-oPRL in kidney and liver it was found (average from two female rats) that 17.2% of the injected dose ($393 \times 10^6$ dpm [disintegrations per minute]) at 2 min was found in kidney and 15.8% in liver as assessed by gamma spectrometry of the fixative-perfused organs (data not shown).

Initial Peripheral Localization

Electron microscope radioautography with the chemical development method revealed filamentous silver grains close to the hepatocyte plasmalemma (Fig. 4). A plot of the distribution of all grains revealed 93% within 2 $\mu$m of the cell membrane with the first 50% of the these within 1,792 $\AA$ (Fig. 5). Direct scoring indicated 61% of the total grains over the hepatocyte plasmalemma compartment. By directly attributing all grains according to how close they were to either the sinusoidal lateral or bile canalculus surface, 84.9% of the grains at 2 min were...
FIGURE 4  Low power electron microscope radioautograph of liver at 2 min after injection of $^{125}$I-oPRL. Silver grains (arrows) are observed close to the periphery of the hepatocyte ($H$), with few over endothelial cells ($E$). The bile canaliculus is indicated ($bc$). Intracytoplasmic lipid drops ($L$) are retained by the fixative employed (galloyl glucose). An erythrocyte ($RBC$) within the liver sinusoid ($S$) is also indicated. $\times 9,900$.

attributed to the sinudoidal surface with 14% attributed to the lateral surface, and 1.13% attributed to the bile canaliculus.

_Hormone Internalization_

_LIGHT MICROSCOPE RADIOAUTOGRAPHY:_ Analysis of the radioactive content of liver sections by radioautography (Fig. 6) revealed a loss of radioactivity at late time intervals (30–60 min). Identical studies carried out on colchicine-pre-treated rats (Fig. 6) showed a lower radioactive content in liver at 2 to 10 min which progressively increased to levels higher...
subcellular Fractionation

The temporal accumulation of silver grains over lipoprotein-containing vesicles in colchicine-treated rats (Figs. 8 and 10) was in apparent contradiction to our previous observations (26) where it was found that colchicine treatment inhibited the accumulation of $^{125}$I-oPRL into purified Golgi fractions. Therefore, in order to resolve this apparent discrepancy, we fractionated homogenates into L, P, and S fractions at various times after $^{125}$I-oPRL injection into normal or colchicine-treated rats. The radioactive content of liver homogenates from colchicine-treated rats was lower at 5 min postinjection than the radioactive content in control animals (Table I) in accordance with our previous data (26). Moreover, the rate of loss of radioactivity in liver homogenates from colchicine-treated animals was less than that from controls (Table I). At early time intervals, the majority of the homogenate radioactivity was recovered in microsomes (P, Fig. 11). At later time intervals (30 and 60 min), radioactivity accumulated in the cytosol fraction (S, Fig. 11) isolated from control animals but not in the cytosol fraction isolated from the colchicine-treated rats. Rather, colchicine treatment accumulated radio-label in the L fraction at the later time intervals with a slight diminution observed for the microsomal fraction.

DISCUSSION

The utility of the in vivo radioautographic method employed herein has been demonstrated by the extensive mapping studies

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smooth membranous body compartment was noted (Fig. 8) which decreased with longer time intervals. The compartment scored as lipoprotein-containing vesicles became increasingly labeled with time (Fig. 8). The location of the lipoprotein-filled vesicles was markedly altered by colchicine treatment. In normal animals these vesicles were found within the Golgi region often closely apposed to a Golgi complex stack (Fig. 9). In contrast, the colchicine-treated animals labeled hormone accumulated in lipoprotein-containing vesicles which were close to the sinusoidal surface of the hepatocyte (Fig. 10).

FIGURE 5 Distribution of filamentous silver grains about the hepatocyte plasmalemma at 2 min after the injection of $^{125}$I-oPRL. The perpendicular distance of the geometric center of every grain was measured to the nearest hepatocyte plasmalemma as described previously in Bergeron et al. (5). On the left-hand ordinate is indicated the number of grains scored in histogram form as a function of their distance from the nearest hepatocyte plasmalemma (abscissa). On the right-hand ordinate the data are plotted in cumulative form. 93% of the grains were within 2 µm of the hepatocyte cell membrane and the first 50% of these within 1,792 Å. By direct scoring of the 266 silver grains, 87.3% were observed over hepatocytes with 9.8% over endothelial cells and the remaining grains over the lumina of sinusoids. Of the grains over hepatocytes, 75.7% were scored as plasmalemmal, 14.2% over cytoplasm, 7.6% over cytoplasmic vesicles, 6.1% over mitochondria, 1.3% over stacked Golgi sacculles, 0.5% over the endoplasmic reticulum and 0.5% over lipid drops.

By light microscope radioautography (Fig. 7 a-c), a peripheral location of silver grains at 2 min postinjection was revealed over liver sections from control animals. At 10 and 30 min, grain density was decreased and displaced towards the Golgi/bile canalicular region of the cell. By contrast, in colchicine-pretreated rats, grain density over liver sections remained high at 10 and 30 min postinjection (Fig. 7 d-f) and the location of the silver grains remained at the cell periphery.

ELECTRON MICROSCOPE RADIOAUTOGRAPH Y: The distribution of fine grains in electron microscope radioautographs was altered with time in the normal rats. Thus, at 10 min the ratio of fine grains within 2 µm of the sinusoidal plasmalemma as compared to those within 2 µm of the bile canaliculus was 2.04:1. By 20 min this ratio changed to 0.27:1. By contrast, fine grains were mainly observed within 2 µm of the sinusoidal plasmalemma at all time intervals for colchicine-treated rats. At 20 min, the ratio was 1.57:1 and similar values were observed for other time intervals (data not shown).

Direct scoring of structures beneath fine grains was carried out for each time interval (Fig. 8). Remaining grains were mainly over the cell membrane, cytoplasm, and mitochondria. No temporal difference was noted for cell cytoplasm and mitochondria and no effect of colchicine was observed on the distribution of grains over these compartments. Representative examples of grains scored as endoplasmic reticulum, smooth membranous bodies (an heterogeneous compartment), lipoprotein-filled vesicles, and secondary lysosomes are noted in Figs. 9 and 10 for normal and colchicine-pretreated females, respectively.

An early peak in the proportion of silver grains over the
Filamentous grains at 2 min revealed a distribution about the hepatocyte cell membrane with 93% within 2 µm. Of these, 50% were within 1,792 Å. Under identical development conditions, a line-source of 125I-thyroglobulin revealed 100% of the grains at 14,400 Å from the center of the line and 50% within 1,568 Å (5). Thus, as for 125I-insulin (5), 125I-oPRL is found close to the hepatocyte cell membrane at 2 min after injection. The peripheral location of the grains was also noted by direct scoring. However, few grains were observed over specific regions of the plasmalemma (e.g., coated pits) even by the fine grain method of silver grain development (1,4% of grains at 2 min in coated vesicles, data not shown). The attribution of silver grains to the various domains of the hepatocyte cell surface (sinusoidal, lateral, and bile canicular) revealed a distinct lack of labeling of the bile canicular membrane but with labeling of the sinusoidal and lateral surfaces which approximated published estimates of their surface areas (8). Thus, as for insulin (8), prolactin seems to be initially bound to receptors dispersed randomly over the sinusoidal and lateral surfaces of the hepatocyte.

Internalization

The radioactive content of liver as assessed by light microscope radioautography was observed to decrease markedly by 20–30 min after injection, presumably due to hormone degradation. The location of the prolactin label was also observed to change quite markedly with time. Thus under the light microscope most silver grains were observed peripherally at 2 min but at 10 and 30 min were observed to cluster near the Golgi complex/bile canicular region of the hepatocyte.

Electron microscope radioautography revealed the progressive accumulation of silver grains over lipoprotein-filled vesicles in the Golgi region of the cell. These lipoprotein-filled vesicles were often in close association with the Golgi apparatus although the Golgi complex stacks were themselves unlabeled. These observations are in general accord with our previous in vivo studies on insulin internalization in which we observed 125I-insulin accumulation into the Golgi region and into Golgi elements which were largely acid-phosphatase negative (8). Lipoprotein-filled structures in the Golgi region have been previously considered as secretory-vesicular elements of Golgi

of receptor sites for a variety of peptide hormones (e.g., 2, 3, 5–9, 31–38).

In the present study on the prolactin receptor of liver, competition for binding of 125I-oPRL was effected by unlabeled oPRL in the target tissue of interest (liver) but not in a site (renal proximal convoluted tubule) known to be nonspecific for other hormones (6, 38). The sex specificity of the binding (Fig. 1) attested to the identification of the binding site as the lactogen receptor (17, 19, 29).

### FIGURE 7
Light microscope radioautographs of liver sections from rats perfused at 2, 10, and 30 min after injection of 125I-oPRL (393 × 10^6 dpm; 114 µCi/µg) into normal adult females (a, b, and c) and females pretreated with colchicine (d, e, and f). In sections from normal animals, silver grains are observed (arrows) over the cell periphery at 2 min (a). By 10 (b) and 30 min (c) silver grains are clustered (arrows) over the bile canicular region. In colchicine-pretreated animals, silver grains are observed at the cell periphery at all time intervals (arrows). The nuclei (N) of hepatocytes and sinusoids (S) are indicated. × 800.
vesicles by some investigators (e.g., references 13, 14). However, A. Novikoff (24) and P. Novikoff and Yam (25) consider lipoprotein-containing structures on the trans aspect of the Golgi stack in liver to be part of the distinct organelle GERL. Most recently, these investigators have noted the internalization of ferritin-labeled lactose into lipoprotein-filled GERL structures as well as lipoprotein-filled secretory vacuoles and secondary lysosomes (16). Chao et al. (11) have also documented the internalization to the Golgi region of the ligand LDL into structures which they called "multivesicular-body like" but which would have been scored as lipoprotein-containing vesicles by the criteria of our present study on prolactin internalization. Therefore lipoprotein-filled vesicles close to Golgi stacked cisternae appear to accumulate a variety of ligands besides the hormone insulin (8) and prolactin (present study).3

Colchicine Treatment

In an attempt to ascertain with more accuracy the identity of the lipoprotein-containing vesicle, we carried out a parallel internalization study with colchicine-treated female rats. Similar colchicine pretreatment of rats has been documented by Redman et al. (30) to result in a block in the secretion of lipoproteins and other plasma proteins by hepatocytes. Morphologically, this block in secretion was reflected by an accumulation of lipoprotein-filled secretion vacuoles at the sinusoidal front of the cell (30) and an increased accumulation of secretory proteins in Golgi fractions isolated from liver homogenates of colchicine-treated rats (30). Our previous biochemical studies have indicated a decrease in the labeling of Golgi fractions isolated from liver homogenates of colchicine-treated rats (30). Most recently, these investigators have noted the internalization to the sinusoidal side of the plasma membrane following colchicine treatment. In the studies of Redman et al. (30) as well as Bergeron et al. (4), lipoprotein-filled structures were characterized as secretory in nature based on subcellular fractionation studies. The same isolated Golgi fractions with lipoprotein-filled vesicles as their most prevalent component have previously been shown by us to accumulate i.v. injected 125I-oPRL (17) in normal rats.

The present studies are therefore in apparent contradiction to our previous work in which colchicine treatment was shown to inhibit the accumulation of 125I-oPRL into lipoprotein-filled vesicles in Golgi fractions (26). As galactosyl transferase relative specific activities and recoveries were identical for Golgi fractions isolated from control or colchicine-treated rats (26), then the labeled lipoprotein-containing structures observed in the present work (Fig. 10) may represent a subpopulation of lipoprotein-containing vesicles which do not accumulate in microsomes and are not enriched in galactosyl transferase marker enzyme.

To investigate this possibility, we fractionated liver homogenates into L, P, and S fractions following standard protocols (12, 20). Colchicine treatment inhibited the accumulation of radioactivity into microsomes (Fig. 11) in confirmation of our previous results (26). Furthermore colchicine treatment augmented the accumulation of radioactivity in the L fraction and inhibited markedly the accumulation of radioactivity in S (Fig. 11). Based on this limited data, it is suggested that the labeled lipoprotein-containing vesicles in colchicine-treated rats may be equivalent to the "unique" vesicle previously identified in the total L fraction (20) and in the L, subfraction of purified lysosomes (27).

Radioactivity in the S fraction has previously been shown to reflect largely degraded hormone (27), and the inhibition by colchicine is consistent with the present morphologic (Fig. 6) and biochemical (Table I) studies indicating a retention of radioactive content in liver following colchicine treatment.

Lysosome and Endosome Involvement

Using identical definition for lysosomes as used most recently by Hamers et al. (16), namely a thick membrane and partial or complete halo, we observed a maximum of 18.7% of silver grains over secondary lysosome and secondary lysosome-like structures, with no difference noted by colchicine treatment. From 10 to 30 min, the proportion of silver grains over lysosomes was about 15% of the total grains.

A peak in the smooth membranous body compartment was noted at the early time intervals. The smooth membranous body compartment was heterogeneous and consisted of small (50–100 nm diameter) vesicles and heterogeneous larger structures (up to 200 nm diameter). At 2 min most of the silver grains (75%) in the smooth membranous body compartment of normal animals was in the smaller vesicular structures. At later intervals a greater proportion of grains was observed over the larger structures in this compartment. The small vesicles observed in the present study corresponded in size and temporal labeling to the small smooth-surfaced vesicles which accounted for the labeling of the Golgi heavy fraction in previous studies assessing 125I-oPRL internalization by subcellular fractionation (17, 26). Our combined in vivo studies of the present work and previous subcellular fractionation data (17, 26) therefore indicate a sequence of internalization involving initial binding of hormone to the sinusoidal and lateral surfaces of the plasma membrane followed by internalization into small, smooth-surfaced intracytoplasmic vesicles and temporary accumulation within vesicles with a lipoprotein content.

Further analysis of the effect of colchicine on prolactin accumulation and comparison to the effect of chloroquine (27)
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Portions of this work were presented at the Canadian Society for Clinical Investigation (Montreal) and published in abstract form (Bergeron, J. J. M., R. Rachubinski, R. Sikstrom, and B. I. Posner, 1978, 


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125I-labeled insulin in hepatocytes from intact rat liver. Proc. Natl. Acad. Sci. USA. 76:2809-2807.


TABLE I

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<th>Time</th>
<th>Control</th>
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<td>5</td>
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<td>64.2 ± 5.0</td>
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<td>44.1 ± 4.5</td>
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*125I-oPRL (31.25 X 10^6 dpm, 142 μCi/μg) was injected into each of three animals for each time interval and experiment indicated. Livers were rapidly excised and placed in ice-cold 0.25 M sucrose and homogenized. Radioactive content of liver homogenates from control rats at 5 min after injection averaged 2.05 X 10^6 dpm/g liver. All other values have been expressed as a proportion of this 5-min value for both control and colchicine-treated rats.

should help in the elucidation of the molecular mechanisms of receptor recycling and perhaps hormone action.

* That colchicine treatment may have affected receptor recycling can be inferred from the data in Fig. 6 and Table 1 in which hormone binding at 2 min (Fig. 6) and 5 min (Table 1) is lower in colchicine-pretreated rats than in the normal animals. However, the continued internalization of 125I-oPRL in the colchicine-treated animals (Fig. 8) argues against a tight coupling between the phenomena of endocytosis and exocytosis, the latter completely inhibited by identical colchicine treatment (30).

FIGURE 11 Distribution of radioactivity in subcellular fractions isolated from liver homogenates (Table I) at indicated time intervals after injection of 125I-oPRL. The conditions of 125I-oPRL injection and preparation of liver homogenates are indicated in the legend to Table I. The percent distribution of radioactivity ± SEM (n = three animals) in each of the fractions is indicated for each time interval and expressed as a proportion of the radioactive content of parent liver homogenates. Remaining radioactivity was in the "nuclear-mitochondrial" pellet, and total recoveries were 101.3 ± 2.4% of the radioactive content of the parent homogenates. Protein contents of homogenates were 229.9 ± 4.6 mg protein • g^-1 liver wet wt for control rats and 231 ± 4.5 mg • g^-1 liver for liver homogenates from colchicine-treated rats. Protein values for fractions isolated from these homogenates were I fraction, 4.52 ± 0.12; 4.6 ± 0.2; P, 47.5 ± 1.3; 44.6 ± 1; S, 89.8 ± 1.1; 87.3 ± 1.3 mg protein • g^-1 liver for fractions from control and colchicine-treated rats, respectively.

FIGURE 10 Electron microscope radioautographs of liver sections from colchicine-treated rats at 20 min after 125I-oPRL injection. Silver grains are over lipoprotein-containing vesicles (LPV) in the top panel, over a secondary lysosome (LYS) in the center panel (arrow), and over an amorphous dense body (ADB) and smooth membranous body (SMB) in the lower panel. The sinusoidal plasmalemma (SPML), bile canaliculus (BC), and mitochondrion (M) are indicated. × 30,000.