Functional Expression of Microsomal and Mitochondrial Cytochrome P-450 (d and SCC) in COS-7 Cells from Cloned cDNA

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ABSTRACT. Using full length cDNA introduced into COS-7 cells, two species of P-450 with entirely different physiological functions have been expressed in enzymatically active form. One is P-450d, which is known to reside in the microsomes of rat hepatocytes where it acts as a drug-metabolizing enzyme; the other is P-450(SCC), which catalyzes the conversion of cholesterol to pregnenolone in the rate-limiting reaction of steroidogenesis in mitochondria of adrenal cortex cells. Northern blot and immunoblot analyses revealed that the mRNA and protein of these P-450 species were efficiently produced in COS-7 cells. The protein contents amounted to nearly 0.1% of the total cell protein as estimated from immunoblotting and low temperature CO difference spectra. The subcellular localization of the products indicated that they were correctly sorted to the microsomes and mitochondria, respectively. We have succeeded in eliciting most of the activity of the expressed microsomal P-450d by reconstruction with NADPH-cytochrome P-450 reductase, while the optimal conditions for the mitochondrial enzyme in the COS cells remain to be studied.

These results show the applicability of the COS-7 expression system to investigations of the functions of members of the P-450 superfamily whose cDNA has been newly isolated.

Cytochrome P-450 constitutes a superfamily of heme-proteins which are believed to have evolved from a common ancestor. They catalyze the monooxygenase reaction of xenobiotics, such as drugs, carcinogens and environmental pollutants, as well as endobiotics, such as steroids, fatty acids, and prostaglandins (1, 2). This versatility of the enzyme system is probably due to the large number of P-450 species and their different and sometimes broad substrate specificities. The multiple molecular forms of P-450 have several properties in common, such as size, structure, and immunologic and biochemical characteristics, which makes it difficult to separate one form from another biochemically. Recent application of molecular cloning technology to the field of P-450 has overcome this difficulty and brought about clear-cut identification of many forms of P-450 (2, 3). However, there seem to remain a considerable number of uncharacterized P-450 species, especially the minor forms in higher organisms. When a cDNA clone for a certain species of P-450 whose function is not known is isolated by cross-hybridization with the cDNA of a known P-450, it is prerequisite for characterization of the new species to establish a suitable expression system for the cDNA information. The expression system may also be helpful for investigating the structure-function relationship in P-450 molecules by site-directed mutagenesis and for analyzing causative mutations in defective genes of genetic diseases related to the P-450 superfamily.

In the present study we attempted to express two P-450 species [P-450d and P-450(SCC)] with different functions and localizations in the monkey kidney cell line COS-7 by transfection with on expression vector (pKCR) carrying their cDNA sequences. The P-450 molecules expressed in COS cells exhibited their correct cellular localizations and functional activities, although their activities seemed suboptimal. Our laboratory has reported a successful application of this system to the analysis of defective genes in congenital P-450c21 deficiency (4).

MATERIALS AND METHODS

Enzymes and chemicals. Restriction endonucleases, DNA polymerase (Klenow fragment), T4 DNA ligase, and bacterial restriction enzymes were purchased from Takara Shuzo Co. (Kyoto, Japan). Other chemicals were purchased from Sigma Chemical Co. (St. Louis, MO).
alkaline phosphatase were purchased from Takara Shuzo Co. (Kyoto, Japan). Proteinase K was obtained from Boehringer-Mannheim (Penzberg, West Germany). Tryptsin, glucose-6-phosphate, and glucose-6-phosphate dehydrogenase were obtained from Sigma (St. Louis, MO, U.S.A.). NADPH was obtained from Oriental Kobo Co. (Tokyo, Japan). [α-32P]dCTP (2000–3000 Ci/mmol) was obtained from The Radiochemical Center (Amersham, England); [2-3H]-estradiol-17 (25.3 Ci/mmol), and [7-3H]-pregnenolone (22.6 Ci/mmol) were from INEN Research Products (Boston, MA, U.S.A.).

Construction of plasmids. Yeast expression vectors containing P-450d or SCC cDNA were kindly supplied by Dr. T. Shimizu (Tohoku University). A cDNA insert of P-450d was obtained by HindIII digestion of pAAH5-P-450d, and that of P-450(SCC) was by Eco RI digestion of Yepl1PT-P-450(SCC) (5). The expression vector pKCRH2 for COS cells was kindly supplied by Drs. M. Mishina and S. Numa (Kyoto University) (6). The P-450d cDNA insert was ligated to the unique HindIII site of pKCRH2. Prior to ligation of the P-450(SCC) cDNA insert to the Eco RI site just flanking the HindIII site, another Eco RI site of pKCRH2 located at the 5'-terminal side of the SV40 promoter region was disrupted by partial digestion with Eco RI; the sticky ends were filled with the Klenow fragment, and the resultant blunt ends were ligated.

Transfection of COS cells. COS-7 cells were a kind gift from Dr. Y. Gluzman (7). Introduction of pKCRP-450 expression plasmids into COS-7 cells was by the calcium phosphate method as previously described (8). The efficiency of transfection by the method was greater than by the DEAE-dextran method under our conditions as judged from the amounts of mRNA expressed.

The night before transfection, COS-7 cells were passaged from a confluent 100-mm culture dish (Corning 2520) by trypsinization. Transfection was performed 4 h after the medium was replaced by 5 ml of the fresh one. Calcium phosphate-DNA precipitates were developed for 10 min on ice from a 15-μg aliquot of DNA per dish in 400 μl of 50 mM Hepes buffer (pH 7.10) containing 280 mM NaCl, 1.5 mM Na2HPO4, and 0.121 M glycine, 20% ethanol onto a nitrocellulose filter. The protein-transferred filter was blocked in a 1:4 mixture of FCS and PBS-TX100-BSA solution [10 mM sodium phosphate (pH 7.5), 0.15 M NaCl, 0.05% Triton X-100, 0.5% BSA] at room temperature for 1 h. Then the filter was incubated at room temperature for 1 h with the first antibody solution, containing anti-P-450c IgG or anti-P-450(SCC) IgG. The second antibody solution [consisting of 0.05 mg/ml peroxidase-conjugated goat anti-rabbit IgG (CAPPEL) in PBS-TX100-BSA] was applied to the filter, and it was incubated at room temperature for 1 h. The immunoreactive bands on the filter were stained by incubation in 50 mM Tris-HCl (pH 7.5), 0.4 mg/ml 4-chloro-1-naphthol, and 0.01% H2O2. During each interval of the incubation, the filter was washed three times in PBS-TX100.

In order to quantitate the intensity of the immuno-stained bands, the stained filter itself or the photograph of the filter was subjected to reflectometry with a Shimizu CS-910. Known amounts of purified P-450d (14) and P-450(SCC) (15) were used as standards.

Fractionation of subcellular components. COS cells harvested from 4 dishes (4.8 mg protein) were homogenized by brief ultrasoundization in 6 ml of 0.25 M sucrose solution containing 5 mM Mops-NaOH (pH 7.5), 1 mM EDTA, 10 μg/μl of peptatin A, and 10 μg/μl of leupeptin. The homogenate was centrifuged differentially (16) to obtain subcellular fractions: P0.7 (“nuclei”: 0.7k G, 10 min), P8 (“mitochondria”: 8k G, 10 min), P120 (“microsomes”: 120k G, 60 min), and S120 supernatant (“cytosol”: 120k G, 60 min).

Succinate-cytochrome c oxidoreductase (SCOR) activity was measured as a marker of mitochondria (17). The reaction mixtures (2.0 ml) contained 50 mM potassium phosphate (pH 7.4), 1 mM KCN, 20 mM sodium succinate, 25 μM cytochrome c, and a suspension of one of the subcellular fractions (1/4 portion of each pellet from cells of 4 dishes). The time course of absorbance at 550 nm was measured and the initial velocity was calculated. As a marker enzyme of microsomes, glucose-6-phosphatase activity was measured (18) in the reaction mixture (0.5 ml) containing 40 mM glucose-6-phosphate, 50 mM Tris-Malate [pH 6.8], and one of the subcellular fractions (1/3 portion of each fraction). Each mixture was incu-
bated at 30°C for 30 min; then the reaction was stopped by addition of TCA solution. Liberated phosphate was determined by the method of Fiske and SubbaRow (19).

Subcellular fractions from COS cells were digested with proteinase K as described (20).

Low temperature CO-reduced difference spectrum. The mitochondrial and microsomal fractions [P8 and P120 suspended in 0.25 M sucrose, 5 mM Mops-NaOH (pH 7.5), 3–4 μg/μl of protein] from COS-7 cells were reduced with an excess amount of dithionite (21). One-half of the fraction was then saturated with carbon monoxide. Fifty μl of 60% glycerol in PBS was added to 50-μl aliquots of CO-saturated and non-treated fractions, respectively. Co difference spectra were recorded with a High-Sensitivity Spectrometer (Union Giken Co.) in which a liquid N2 Dewar bottle with quartz glass windows for the light path was settled (22, 23). P-450cam was used as the standard for estimation of the chromophore concentrations in the sample solution.

Reconstruction of microsomal P-450d with exogenous NADPH-cytochrome P-450 reductase. Purified P-450 reductase (Fp) (24) was a generous gift from Dr. T. Iyanagi (Tsukuba University). The amount of Fp was determined by measurement of absorbance at 450 nm using the molar absorbance coefficient of 1 × 10^4 cm−1 M−1.

The reconstruction was performed essentially according to the procedure by Miwa and Lu (25). Sodium cholate (1.2 mg) was added to the microsomal fractions (P120) (440 μg of protein), and the solution was subjected to brief sonication. Then 440 pmol of Fp was added to these fractions. This amount corresponds roughly to a 1 : 10 molar ratio relative to the amount of P-450d in each fraction, estimated from the low temperature CO-reduced difference spectra. The volumes of these fractions were then adjusted to about 270 μl by addition of dialysis buffer [50 mM potassium phosphate (pH 7.4), 0.1 mM EDTA, 0.1 mM DTT]. Finally, to equilibrate the system, they were dialyzed three times at 4°C for 12 h against 6 liters of the dialysis buffer. The dialyzed microsomal fractions were subjected to the activity measurements. The above procedure was essential for efficient reconstruction of P-450d and exogenous Fp, as indicated in Results.

Assay of expressed P-450 activities. Expressed P-450d monooxygenase activity was assayed with estradiol as a substrate (26). The reaction mixture (500 μl) contained 5 nmol or 100 nmol of [2-3H]-estradiol-17 (0.5 μCi), 3 mM NADPH, 40 mM glucose-6-phosphate, 1 unit of glucose-6-phosphate dehydrogenase, 70 mM potassium phosphate (pH 7.4), and the microsomal fraction (440 μg of protein). The mixture was incubated at 37°C for 60 min, and the reaction was stopped by addition of 1 ml of 1.5 N HCl. After centrifugation (3k rpm, 20 min), the supernatant of the reaction mixture was applied to a Pasteur-pipette column packed with about 3 ml of XAD-2 resin equilibrated with H2O. The column was washed first with 7 ml of H2O and then with 8 ml of methanol. The estradiol 2-hydroxylation reaction was accompanied by formation of 3H2O which was eluted first with H2O, while the unreacted labeled estradiol was later eluted with methanol. The radioactivity of tritium in the collected H2O fractions, methanol fractions, and 1.5 N HCl precipitate was measured with a liquid scintillation counter LS-3801 for calculation of the substrate conversion ratio.

P-450(SCC) enzyme activity was detected by measurement of pregnenolone released into the culture medium of the COS cells transfected with the P-450(SCC) expression vector. RIA was performed for determination of the pmol level of pregnenolone as described previously (27). On the fourth day after transfection, 20 nmol of 25-hydroxycholesterol was added to 10 ml of medium in a culture dish, and the cells were incubated for another 24 or 120. One milliliter of this medium was left standing at room temperature for 30 min in order to equilibrate with [7-3H]-pregnenolone (3000 cpm) which served as an internal standard for recovery calculation. It was then extracted with 10 ml of methylene chloride. The organic phase was evaporated by passing N2 over it and dissolved in 100 μl of HPLC solvent (4% isopropanol/hexane). The wall of the plastic tube used for evaporation was rinsed with another 100 μl of the same solvent; the combined 200 μl was dried again and re-dissolved in 50 μl of the solvent. It was then subjected to HPLC on a silica-60 column and eluted with the same solvent. The radioactive fractions were collected and dried. This purification procedure was essential to eliminate cross-reaction of the pregnenolone-antiserum with 25-OH-cholesterol which might have been in the extract solution.

The obtained residues were dissolved in 700 μl of RIA assay buffer consisting of 0.1 M sodium phosphate (pH 7.0), 0.9% NaCl, 0.1% sodium azide, and 1 mg/ml egg albumin. The solution was left standing for at least 1 h at room temperature to improve solubility. The tritium radioactivity in 200 μl of this purified-extract solution was counted for estimation of the recovery, and the remaining 500 μl was applied to RIA using sheep antiserum SLB SE-0000-53 against pregnenolone (1/1000 dilution with the assay buffer). One hundred μl of the diluted serum and 100 μl of [7-3H]-pregnenolone (3 × 10^4 dpm) dissolved in the assay buffer were added to the 500 μl of HPLC-purified extract solution and the mixture was left standing at 4°C overnight. Then free pregnenolone was removed by addition of 200 μl of charcoal-dextran solution (0.625% Norit A, 0.0625% dextran). After incubation at 4°C for 20 min, the mixture was centrifuged (5k rpm, 2 min), and the tritium radioactivity in the supernatant was counted. A standard measurement was performed in exactly the same way using known amounts of cold pregnenolone.

RESULTS

The transient expression system mediated by the plasmid DNAs possesses the advantages of efficiency of transfection and freedom from possible effects of the integration site in the genomic sequence (28). In COS cells, the pKCR-expression plasmid insures abundant transcription (6) (Fig. 1a). Marked hybridization signals
could be seen 2 days after transfection. Co-transfection of P-450d and P-450(SCC) cDNA into COS-7 cells led to production of the respective mRNAs with almost equal efficiency. Both the cDNAs gave a single band of mRNAs at 19.5 S, which is consistent with their full lengths.

The left half of Fig. 1b shows the bands of the P-450d protein stained with P-450c antibody which is cross-reactive to P-450d (11). The mixture (1:2 by weight) of standard P-450d and P-450d from transfected cells gave a single band, indicating that the transfected cells produced an immunoreactive protein which is electrophoretically indistinguishable from authentic P-450d. The amount of expressed P-450d in the cell was estimated from the band intensity to be 0.1% of the total cell proteins (20 pmol P450d/mg cell protein) 3 days after transfection.

On the other hand, the P-450(SCC)-cDNA transfected cells produced two immunoreactive bands against anti-P-450(SCC) antibody, (the right half of Fig. 1b). Since the P-450(SCC) cDNA contained the full-length sequence coding for the P-450(SCC) precursor, we assume that the larger band corresponds to P-450(SCC) precursor and the smaller to the mature form. This figure also indicates that a mixture of standard P-450(SCC) purified from bovine adrenal cortex and P-450(SCC) from the transfected cells [P-450(SCC) Mt] gave two bands, of which the smaller one contained P-450(SCC) proteins from the two origins, showing that the smaller protein expressed by the COS cells had the same molecular weight as the authentic mature form. The amount of the product was approximately

Fig. 1. Expression of mRNA and protein from cDNA of P-450d and P-450d (SCC). a. mRNA-blot hybridization analysis of transfected COS-7 cells. Cells were harvested 2 days after transfection and 5 μg of extracted total RNA was analyzed. Lanes 1 and 4 (d), total RNA from cells transfected with PKCRP-450d (15 μg); lanes 2 and 5 (d+SCC), PKCRP-450d (15 μg) plus PKCRP-450d (SCC) (15 μg); lanes 3 and 6 (SCC), P-450d (SCC) (15 μg). The nitrocellulose filter of lanes 1–3 was hybridized with nick-translated P-450d cDNA and the filter of lanes 4–6 with P-450d (SCC) cDNA. The hybridized filters were washed, then exposed to Fuji-RX film for 2 h at -80°C. The positions of RNA markers are shown to the left of the autoradiogram. b. Immunoblot analysis of expressed P-450d and P-450d (SCC) in COS-7 cells. Lane 1 (BSA+cyc. c), bovine serum albumin (2 μg) and cytochrome c (2 μg) as molecular weight markers; lane 2 (P-450d std.), purified P-450d (10 ng); lane 4 (d-COS), whole cell lysate (20 μg) of cells transfected with PKCRP-450d; lane 3 (Mix), a mixture of purified P-450d (5 ng) and d-COS (10 ng); lanes 5 and 10 (H2-COS), whole cell lysate (20 μg) transfected with PKCRH2; lane 6 [P-450d (SCC) std.], purified bovine P-450d (SCC) (40 ng); lane 8 (SCC-COS Mt), mitochondrial fraction (20 μg) from cells transfected with PKCRP-450d (SCC); lane 7 (Mix), a mixture of purified P-450d (SCC) (20 ng) and SCC-COS Mt (10 μg); lane 9 (SCC-COS), whole cell lysate (20 μg) transfected with PKCRP-450d (SCC). Protein-blotted filters of lanes 1–5 were incubated with anti-P-450c antibody, followed by incubation with peroxidase-conjugated goat anti-rabbit IgG and with 4-Cl-1-naphthol solution. The notation "anti-P-450c" at the bottom of the filter represents only its reactivity. In lane 2, the faint band just above the dense band of P-450d indicates contamination with P-450c. The filter of lanes 6–10 was incubated with anti-P-450d (SCC) antibody and treated in the same way.
0.2% of the total mitochondrial protein (40 pmol/mg) when harvested 3 days after transfection. The band intensity of mitochondria of the plasmid-transfected COS cells was higher than that of the whole-cell lysate, indicating the mitochondrial localization of the product.

As shown in Fig. 2, the maximum amount of the mRNA for P-450d was achieved on the third day after transfection. P-450d protein also increased similarly to reach a maximum on the third or fourth day after transfection. Most of the mRNA disappeared after 1 week, whereas the cells at this time still retained half or more of the maximal level of P-450d protein.

The concentration of mRNA for P-450(SCC) could be equal to that of P-450d, as deduced from Fig. 1a. The amount of P-450(SCC) protein expressed from its vector reached a maximum value on the fourth or fifth day as indicated by the immuno-stained filter of Fig. 2; this corresponded to about 0.1% of the whole cell protein. Similar to the case with P-450d, more than half of the maximal concentration remained after 1 week; this disappeared 2 weeks later.

The intracellular distribution of expressed P-450d and P-450(SCC) was examined by fractionating the components of transfected COS cells. As shown in Fig. 3, the distribution of P-450d protein was consistent with that of a microsome marker enzyme (G-6-Pase), while the distribution of the P-450(SCC) mature form was consistent with that of a mitochondrial marker enzyme (SCOR). The mitochondrial localization of the mature form was demonstrated more clearly when each fraction was treated with proteinase K, which selectively digests the precursor form. The resistance of the mature form suggests that it was localized on the matrix-side surface of the inner mitochondrial membranes. The precursor form, on the other hand, appeared to reside on the outer surface of all membranous fractions (P0.7, P8, and P120), and was thus sensitive to the proteolytic digestion.

The content of the expressed P-450d in microsomes was also determined from the CO difference spectra at liquid nitrogen temperature. The molar extinction coefficient at 448 nm for the protoheme of P-450 was estimated from the known amount of P-450cam to be $1.88 \times 10^3 \text{mM}^{-1}\text{cm}^{-1}$, about 20 fold higher than at room temperature.

The spectrum in Fig. 4 clearly shows a positive peak.
Fig. 3. Subcellular localization of expressed P-450d and P-450 (SCC). Transfected cells were homogenized and fractionated into four fractions (P0.7, P8, P120, S120) as described in Methods. Each fraction was used for marker enzyme activity assay and for immunoblotting. a. The cross-hatched bars show the mean values ±S.D. (standard deviation) of observed succinate cytochrome c oxido-reductase activities (SCOR) as a mitochondrial marker. The blank bars show the values of glucose 6-phosphatase activities (G-6-Pase) as a microsome marker. Each datum is the arithmetic mean of duplicate or triplicate determinations. As indicated, the purification calculated from the increment of each marker activity is much higher for mitochondria than for microsomes (7 fold for mitochondria at P8, but 2 fold for microsomes at P120), suggesting a difference in protein content between microsomes and mitochondria in COS-7 cells. b. Immunoblotting of subcellular fractions from PKCRP-450d transfected COS-7 cells. The fraction for each lane contains 20 μg of protein. c. Immunoblotting of the fractions from pKCRP-450d (SCC) transfected COS-7 cells. d. The same samples as c were subjected to proteinase K (70 μg/ml) digestion for 120 min at 15°C, then analyzed by immunoblotting.

Fig. 4. Low temperature CO-reduced difference spectrum of microsomes from PKCRP-450d transfected cells. The spectra were recorded as described in Methods. Microsomes from pKCRP-450d transfected COS-7 cells (d-COS Ms) and from pKCRH2 transfected cells (H2-COS Ms) were used.
450d activity were performed with estradiol-17, which is known to be one of the best substrates for P-450d.

The activity of Estradiol 2-hydroxylase in microsomes of the transfected cells was lower than that expected from the results of immunoblotting and CO difference spectroscopy. From the activity of 2.34 pmol·mg⁻¹·min⁻¹ in the microsomes, the amount of active P-450d enzyme is estimated to be 0.17 pmol/mg, two orders of magnitude smaller than that calculated from the results of the immunoblotting. When assayed after the d-COS microsomes were reconstructed with P-450 reductase (Fp, 1 : 10 mol/mol), the measured hydroxylase activity increased remarkably to a full activity of 211.8 pmol·min⁻¹·mg⁻¹, from which the P-450d content was calculated to be 15.3 pmol/mg on the assumption that the turnover rate of pure P-450d is 13.8 min⁻¹. This value corresponds to 0.08% of the microsomal total protein. Since the reconstruction procedure usually yielded about 56% recovery (33), we concluded that almost all of the expressed P-450d protein was active enzyme, but was limited in interaction with the reductase.

The activity of expressed P-450(SCC) is also shown in Table 1B. It was, however, extremely limited; at least 100 h of incubation of the cells transfected with P-450 (SCC) cDNA was required for detection of the product. Intracellular cholesterol or a large amount of serum cholesterol contained in the culture medium did not seem to be available to the enzyme, whereas exogenous 25-hydroxycholesterol was utilized as a substrate.

**DISCUSSION**

In addition to E. coli (29) and yeast (5, 30), the COS cell, a cultured mammalian cell, provides a useful system for producing protein products from cloned cDNAs or genomic DNAs (4, 31). Since all eukaryotic P-450s are microsomal or mitochondrial proteins and need an electron transport pathway for their activities, the host cells used for the expression of these proteins may have to have such intracellular compartments for the stability and activity of the expressed proteins. These requirements are considered to be fulfilled in the COS cell.

In the present report we have demonstrated that two types of P-450, P-450d and P450(SCC), which were expressed in COS-7 cells were correctly sorted out and localized in their normal intracellular compartments. The expression of P-450 cDNAs in mammalian cell cultures has also been reported recently by other laboratories (32-34) with essentially the same results. Both types of P-450 functioned as oxygenase, indicating the existence of electron transport chains within microsomes and mitochondria in COS-7 cells, although the electron supply seemed to be limited.

As shown in Fig. 3, the full length cDNAs of P-450d and P-450(SCC) did carry the information for their intracellular localization, which was functional even in the non-drug-metabolizing, non-steroidogenic COS-7 cells. While the mature form of expressed P-450(SCC) was located in the mitochondrial compartment, its precursor was found in all membraneous fractions (Fig. 3). The relative amount of precursor among the membrane fractions varied form preparation to preparation. It is known that many eukaryotic mitochondrial proteins which are encoded by the nuclear genome are synthesized on cytoplasmic polyosomes in precursor forms and then translocated into mitochondria to form mature proteins (35, 36). Accordingly, the cytosol may contain a pool of precursors for mitochondrial proteins which are in equilibrium between their continuous inflow and outflow (37). It is highly likely that most of the precursors found in the membrane fractions of the COS-7 cells (Fig. 3) originated from the cytosolic pool and that

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**Table I. Catabolic activity of expressed p-450.**

<table>
<thead>
<tr>
<th>A. Effect of P-450 Reductase (Fp) Incorporation on Catalytic Activity of Expressed P-450d</th>
<th>Esr 2-hydroxylase Activitya (pmol/mg/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H2</td>
<td>0.04 &gt;d</td>
</tr>
<tr>
<td>d</td>
<td>2.34</td>
</tr>
<tr>
<td>d + Fp</td>
<td>8.20</td>
</tr>
<tr>
<td>d + Fp (RC)</td>
<td>211.8</td>
</tr>
</tbody>
</table>

| B. Pregnenolone Production by COS-7 Cells Transfected with P-450 (SCC)-cDNA | cpm boundb | Cold PRG added (pmol/ml) |
| --- | --- | |
| CT0 | 5370 | 0 |
| CT2 | 4200 | 2 |
| CT5 | 3850 | 5 |
| H2 + 250H-ChL | 5940 | 0.1> |
| SCC | 5520 | 0.1> |
| SCC + 250H-ChL | 4150 | 2.1 |

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a Estradiol 2-hydroxylase activity of microsomal fraction was measured. Averages of two independent determinations are shown. "RC" indicates reconstructed samples.
b By means of competitive binding of hot pregnenolone with cold pregnenolone to be estimated, RIA was performed as described in Methods. Radioactivities of [7-H]-pregnenolone bound to the specific antibody are indicated. In the control experiments (CT0-CT5), the indicated amount of cold pregnenolone was added to 1.0 ml of fresh medium, which was then extracted, purified by HPLC, and applied to RIA. The cpm values of control experiment are averages of four determinations. Those of the experiments with transfected cells are averages of two or three (SCC + 250H – ChL).
c Values were obtained from the standard curve of the control experiment shown above (CT0-CT5). COS-7 cells were transfected and incubated for 120h as indicated. The pregnenolone produced in 1.0 ml of the medium was determined.
d x>, less than x.
during the cell fractionation they attached aberrantly to the membrane fractions because of their lipophilic nature.

The time course of the precursor and the mature form of P-450(SCC) did not show a typical precursor-product relationship (Fig. 2) and may suggest that through translocation of steroidogenic P-450(SCC) to mitochondria is somehow hindered in the COS cells either due to uncontrolled synthesis of the protein from the transfected expression vector or to lack of some specific factor for translocation.

All of the P-450d polypeptide obtained in the present expression system appears to be active enzyme with the same properties as the enzyme naturally occurring in rat hepatocytes (Fig. 4, Table 1). The same situation could hold true for P-450(SCC), although the heme incorporation could not be ascertained due to the high background noise in the CO difference spectrum. As shown in Fig. 3, P-450(SCC) precursor is correctly processed and transported to the mitochondrial inner membrane.

Since there must exist abundant, newly synthesized protoheme in the matrix space, it seems probable that all the P-450(SCC) could incorporate the protoheme moiety to acquire enzymatic activity. Needless to say, however, one cannot ignore the possibility that the incorporation of protoheme into the apoprotein of P-450(SCC) is somehow inhibited in this expression system.

The rate of reactions catalyzed by the expressed P-450d is limited by the amount of coupled NADPH-cytochrome P-450-reductase (Fp) in microsomes. The observation that the addition of purified Fp protein to microsomes and the subsequent reconstruction was sufficient to achieve efficient coupling of P-450d to the NADPH-Fp electron transport chain suggests that co-transfection with the expression vectors of P-450d and Fp might be a more convenient procedure. It has been reported that two microsomal P-450s, P-450(17) (34) and P-450 (C21) (4), from steroidogenic cells were active enough to permit detection of their products in culture medium within several hours without supplying any of the reductase when COS cells were transfected with an expression plasmid containing the respective cDNAs. Although the activities of these P-450s could be increased by supplementing with Fp, these observations suggest that the intrinsic Fp of COS cells is sufficient to allow detection of the enzymic activity if substrate accessibility and product stability are ensured.

The COS-7 cells harboring the expressed P-450(SCC) were found to convert exogenous 25-hydroxycholesterol into pregnenolone, whereas endogenous cholesterol or that of the medium seemed hardly to be available to P-450(SCC), as already described (33) (Table 1). Such a preference of substrates is consistent with the hypothesis that COS cell mitochondria lack the postulated "substrate translocator" (38) which would exist in steroidogenic mitochondria. Other lines of evidence suggest that the mitochondrial membrane lipids themselves might be a significant factor in activation of P-450(SCC) (39). Moreover, a recent report has demonstrated that certain peptide factors may activate the mitochondrial steroidogenic system in rat Leydig cells (40). Non-steroidogenic cells like COS cells may lack these components supporting the steroidogenic system.

In spite of some limitations in practical use, we have proved that the COS cell expression system is useful for production and characterization of cloned P-450 cDNA and genomic DNA (data not shown). Our laboratory has already successfully applied this system to analysis of causative mutations in defective P-450 genes of the inherited P-450 C21-deficient disease (4).

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