The NAD(P)H cytochrome b$_5$ oxidoreductase, Ncb5or (previously named b5+b5R), is widely expressed in human tissues and broadly distributed among the animal kingdom. NCB5OR is the first example of an animal flavohemoprotein containing cytochrome b$_5$ and cytochrome b$_5$R reductase domains. We initially reported human NCB5OR to be a 487-residue soluble protein that reduces cytochrome c, methemoglobin, ferricyanide, and molecular oxygen in vitro. Bioinformatic analysis of genomic sequences suggested the presence of an upstream start codon. We confirm that endogenous NCB5OR indeed has additional NH$_2$-terminal residues. By performing fractionation of subcellular organelles and confocal microscopy, we show that NCB5OR colocalizes with calreticulin, a marker for endoplasmic reticulum. Recombinant NCB5OR is soluble and has stoichiometric amounts of heme and flavin adenine dinucleotide. Resonance Raman spectroscopy of NCB5OR presents typical signatures of a six-coordinate low-spin heme similar to those found in other cytochrome b$_5$ proteins. Kinetic measurements showed that full-length and truncated NCB5OR reduce cytochrome c actively in vitro. However, both full-length and truncated NCB5OR produce superoxide from oxygen with slow turnover rates: $k_{\text{cat}} = -0.05$ and $-1 \text{ s}^{-1}$, respectively. The redox potential at the heme center of NCB5OR is $-108 \text{ mV}$, as determined by potentiometric titrations. Taken together, these data suggest that endogenous NCB5OR is a soluble NAD(P)H reductase preferentially reducing substrate(s) rather than transferring electrons to molecular oxygen and therefore not an NAD(P)H oxidase for superoxide production. The subcellular localization and redox properties of NCB5OR provide important insights into the biology of NCB5OR and the phenotype of the Ncb5or-null mouse.

The NAD(P)H cytochrome b$_5$ oxidoreductase, Ncb5or (previously named b5+b5R), was first identified in human cell lines and tissues by mining the expressed sequence tag data base (1). This is the first example in animals of a flavohemoprotein containing a cytochrome b$_5$-like domain at the NH$_2$ terminus and a cytochrome b$_5$ reductase (b5R)-like domain at the COOH terminus (1). NCB5OR is encountered in a broad range of vertebrates and invertebrates and is expressed in a variety of cells and tissues. Similar to the classical single-domain cytochrome b$_5$ and b5R counterparts, the cytochrome b$_5$ domain in rat NCB5OR has a six-coordinate b-type heme (2), and the b5R domain binds flavin adenine dinucleotide (FAD) and NAD(P)H prosthetic groups as found in all members of the ferredoxin-NADP$^+$ reductase superfamily (3). The classical cytochrome b$_5$ and b5R proteins form a complex and reside on endoplasmic reticulum (ER) membranes with $\Delta\Delta$ stearoyl-CoA desaturase to catalyze fatty acid desaturation. To anchor on ER membrane, both cytochrome b$_5$ and b5R single-domain proteins use a stretch of hydrophobic residues at their COOH- and NH$_2$-end, respectively, and b5R has additional N-myristoylation (4). Two pairs of charged residues are responsible for the proper docking between these two proteins to allow electron flow from the b5R to cytochrome b$_5$ protein (5). In contrast, NCB5OR is a soluble flavohemoprotein, and its hinge apparently mediates the docking between the cytochrome b$_5$ and b5R domains (1). Moreover, the hinge sequences show a well-conserved CHORD and SGT1 (CS) motif as found in p23/HSP20, SGT1, CHP, and melusin, where it functions as a mediator for protein–protein interaction (6). NCB5OR expressed in Escherichia coli reduces a variety of artificial substrates in vitro, including cytochrome c, ferricyanide, methemoglobin, and molecular oxygen (1). Because of its
ubiquitous expression and capability for superoxide production, we speculated that NCB5OR may function as an oxygen sensor (7).

Since our original publication on human Ncb5or, new sequence information has been generated from genome data bases suggesting the presence of an upstream start codon (8). This new start codon would result in a full-length transcript encoding 54 additional amino acid residues at the NH terminus of the truncated NCB5OR. There has been significant interest regarding whether NCB5OR could produce superoxide in vivo. To clarify these issues, we report in this article the identification of the Ncb5or gene family in animals by bioinformatic analysis. We also confirm by Western analysis that endogenous NCB5OR in mouse tissues is indeed the full-length protein. We show here by both confocal microscopy and subcellular fractionation that endogenous NCB5OR colocalizes with calreticulin, an ER marker. Both full-length and truncated recombinant NCB5OR proteins have been generated in a bacterial system. Kinetic measurements have been performed to analyze their NAD(P)H oxidase activity in superoxide production, as well as their NAD(P)H reductase activity in cytochrome c reduction. These studies of cellular localization, enzymology, and redox properties have considerably enhanced our understanding of the biological function of NCB5OR and the phenotype of the Ncb5or-null mouse.2

EXPERIMENTAL PROCEDURES

Fractionation of Subcellular Organelles—Equal-sized livers from wild-type or Ncb5or-null mice (5-week-old females)2 were rapidly excised, rinsed briefly with phosphate-buffered saline, and minced using a sterile razor blade. Tissue slices were transferred to a 50-mL tube, washed with ~30 mL of phosphate-buffered saline, and allowed to settle to the bottom. This step was repeated once, and the tissue was then resuspended in 7.5 mL of homogenization medium (0.25% sucrose, 1 mM EDTA, and 10 mM HEPES, pH 7.4) containing Complete protease inhibitor mixture (Roche Applied Science) at the recommended concentration. Cells were disrupted by Dounce homogenization (15 turns) and then centrifuged at 3000 × g for 15 min to obtain a post-nuclear supernatant. The post-nuclear supernatant for each sample was transferred to a polyallomer centrifuge tube (Seton Scientific) and centrifuged at 100,000 × g for 40 min to pellet light organelles. This pellet was gently resuspended in homogenization buffer and made up to a volume of 0.6 mL with a density of 5% Nycodenz (5-N,2,3-dihydroxypropylacetamido)-2,4,6-triido-N,N-bis(2,3-dihydroxypropyl)-iso-propylphthalamide; Accurate Chemical). Each sample was loaded on top of a 9-mL continuous 10–30% Nycodenz gradient and then centrifuged for 18 hr at 100,000 × g in an SW41 rotor on a Beckman L8-55M ultracentrifuge. Twelve 0.75-mL fractions were collected by tube puncture. Aliquots of each fraction (40 μL) were loaded onto a 10% denaturing gel for SDS-PAGE and Western blot analysis.

Western Blot Analysis—Primary antibodies were rabbit polyclonal antibodies against NCB5OR NH-terminal sequence (NH2-LN-VPQAKPAFPSQGCOOH; Genomed Synthesis Inc.), cytochrome c (H104; Santa Cruz Biotechnology), or calreticulin (PA3-900; Affinity BioReagents), all at a 1:1000 dilution. Horseradish peroxidase-conjugated goat anti-rabbit IgG (1:1000; Pierce) was used as the secondary antibody, and enhanced chemiluminescence detection was performed with SuperSignal West Dura Extended Duration Substrate (Pierce).

Two-photon Confocal Microscopy of Hep72 Cells—Hep72 cells (American Type Culture Collection) were grown to subconfluence on poly-lysine-coated glass coverslips. They were fixed by ice-cold methanol/acetic acid (1:1) for 10 min (~20 °C) and then blocked with 3% bovine serum albumin in phosphate-buffered saline. Primary antibodies were rabbit polyclonal antibody against human NCB5OR (1:200 dilution; Genomed Synthesis Inc.) and chicken polyclonal antibody against human calreticulin (1:100 dilution; Affinity BioReagents). Secondary antibodies were the Alexa-568-conjugated goat anti-rabbit IgG (1:400 dilution; Molecular Probes) and the Cy2-conjugated rabbit anti-chicken Ig (1:200 dilution; Jackson Immunoresearch). Visualization was performed with a Nikon E1000 microscope (Nikon, Düsseldorf, Germany), equipped with a charge-coupled digital camera (Spot RT Color; Vision Systems, Puchheim, Germany) and the image acquisition software EZ2000 (Coord, Utrecht, Netherlands). Two-photon images of single immunostained cells were captured by a converted Nikon TE300 microscope employing a Ti:sapphire laser that was tuned to 850 nm and delivered 110-fs pulses at a repetition rate of 76 MHz and a 590–650 nm Alexa (a 500–530 nm (Cy2) emission bandpass filter). Two different fluorescence channels were recorded simultaneously. Image stacks (512 × 512 × 64 pixels) were obtained by optical sectioning with a 60× water objective (Nikon, Plan Apochromat DIC H, 1.2 NA). The spatial data were three dimensionally reconstructed from image stacks with the Application Visualization System (Waltham, MA). To enable a comparative analysis of the different fluorescent structures, the fluorescence emission was normalized. Areas of equal intensity of fluorescence emission were rendered as isosurfaces and encoded in false colors (red and blue).

Confocal Microscopy of Transfected COS7 Cells—Subconfluent COS7 cells were transfected with expression constructs of human full-length or truncated NCB5OR (8 μg DNA/8 million cells on a 10-cm dish) with LipofectAMINE Plus reagents (Invitrogen) in Dulbecco's modified Eagle's medium containing 2% fetal bovine serum for 5 h. Cells were then grown in 10% fetal bovine serum medium overnight and split into 6-well plates to grow on glass cover slides. Subconfluent cells were treated as described previously before labeling with antibodies (1). Primary antibodies against the rabbit polyclonal antibody against human calreticulin (1:25 dilution; Affinity BioReagents) and mouse monoclonal antibody 9E10 against c-myc epitope (1:100 dilution; Sigma). Secondary antibodies were the fluorescein isothiocyanate conjugated goat anti-rabbit IgG (1:200 dilution; Jackson Immunoresearch) and the rhodamine-conjugated goat anti-mouse IgG (1:200 dilution; Roche Applied Science). Confocal microscopy was performed using a Zeiss LSM 510 system (Carl Zeiss, Inc.) at Harvard Center for Neurodegeneration and Repair. The confocal images of transfected COS7 cells were obtained on a Zeiss Axioscope 2 microscope with a 560–615 nm (rhodamine) and a 505–530 nm (fluorescein isothiocyanate) emission bandpass filter. Two different fluorescence channels were recorded sequentially with a 63× oil objective (1.4 NA).

Preparation of Recombinant NCB5OR—The full-length human and mouse NCB5OR, as well as truncated human NCB5OR, were prepared with a bacterial expression system (1). The coding sequence was cloned into NdeI and BamHI cloning sites of the pET19b vector (Novagen), and NCB5OR proteins were produced in BL21(DE3) host cells under isopropyl-β-D-thiogalactopyranoside induction at 8 °C to ensure proper folding. The poly-His NCB5OR was purified to homogeneity with nickel-nitrilotriacetic acid chelation (Qiagen) plus ion-exchange Sepharose Q column chromatography (Amersham Biosciences), all performed at 4 °C. NCB5OR holo-proteins, with a A660/A274 of ~1.0 (explained below), were eluted from Q column with a salt gradient starting with a low-salt buffer (20 mM potassium phosphate, pH 8.0) containing 150 mM NaCl and concentrated with YM10 centrifron (Amicon). Small aliquots were made, flash-frozen in liquid nitrogen, and stored at ~80 °C. All NCB5OR samples used in biochemical assays are 90% pure on SDS-PAGE gel and contain ~5% holo-protein with a >~85% NADH-reducible heme content (explained below). Pure cytochrome b6-core proteins with an A420/A660 of ~2.5 were generated by using the same system.

UV-Visible Spectroscopy for Recombinant NCB5OR—Approximately 0.5–1.0 μM (heme) NCB5OR protein in 0.3 mL of low-salt buffer (see above) was analyzed as follows on a Hitachi dual-beam U-2000 spectrophotometer at room temperature. Spectrum 1, a wavelength scan from 650 to 250 nm, was recorded for the oxidized NCB5OR as purified and nonreducing conditions (see above). NCB5OR was reduced instantaneously upon mixing with 3 μL of 10 mM NADH (0.1 mM final concentration) within the sample cuvette, and spectrum 2 was recorded immediately. The difference spectrum [2–1] represents “NADH-reducible” heme. A few granules of solid dithionite powder were added to the above sample to generate spectrum 3. The difference spectrum [3–1] including both NCB5OR protein, NADH product, cytochrome b6. The concentration of heme can be calculated from its difference spectrum, either [2–1] or [3–1], with ε420 to 410 = 185 m$m^3$/cm *1 (1), or from spectrum 1 of 100% oxidized NCB5OR with ε410 = 138 m$m^3$/cm *1. All NCB5OR proteins were analyzed for NADH-reducible heme contents and for their heme/protein ratios. The percentage of NADH-reducible heme was calculated from: $f_{heme,\text{reduced}} = \frac{\epsilon_{420} - \epsilon_{410}}{\epsilon_{410}}$. This value reflects the content of NCB5OR among the total heme proteins (NCB5OR+55), as well as the electron flow within NCB5OR, which is intrinsically determined by protein folding. Pure and fully native NCB5OR flavohemoprotein is expected to have

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100% NADH-reducible heme. The heme/protein ratio, as calculated from their corresponding absorbance A100 and A280, was used to estimate the total heme content of proteins. Pure NCB50R holo-protein has an A100/A280 ratio of ~1.0, indicating one heme per protein. This value has been confirmed by our reverse phase HPLC analysis, which simultaneously quantifies prothetic groups and polypeptides of the same protein (see below).

Reverse Phase HPLC—Approximately 500 pmol of (heme) recombinant human NCB50R (truncated) were loaded onto a Discovery C8 column (15 cm × 4.6 mm; 5 μm; Supelco) connected to a Ranin Rabbit HP system (Varian). The FAD, heme, and polypeptides were eluted with a water and acetonitrile gradient containing 0.1% trifluoroacetic acid and monitored at 220 nm with Dynaxmax UV-C detector (Varian). The absolute amounts of FAD, heme, and polypeptides were determined from individual peak area against standard curves of 320–8400 pmol of FAD and heme (Sigma) and 4–50 μg of bovine serum albumin (Pierce). Peak area from the chromatogram was integrated with Dynamax Method Manager 1.4.3 (Varian) and used to quantify FAD, heme, cytochrome b5, and NCB50R fractions. All major peaks were collected, lyophilized, and analyzed by reverse phase HPLC (Figs. 2 and 3).

Reverse phase HPLC analysis was performed as described above. The reverse phase HPLC profile of NCB50R holo-protein is shown in Fig. 2A. The major peak was the intact protein, identified as NCB50R by its NH2-terminal 15-mer sequence (NH2-LNVPSQ-WVFRQAGPGPSQQ-COOH).

Kinetics of Superoxide Production—The spectrophotometric-based steady-state kinetics apparatus is specifically designed for this experiment (see Fig. 6A). Oxygen (O2) and nitrogen (N2) gases (BOC gases) were mixed to the desired concentration using a computerized gas mixer (Envionics Series 200) before being delivered at 120 ml/min to the gas-tight reaction vessel (The Custom Glass Shop) with Viton tubing (Masterflex L/S 16; Cole-Parmer). Sample dissolved oxygen concentrations and temperature were monitored using a temperature compensated oxygen and temperature electrode and meter (OM2000 from Cameron Instrument Company or InoLab OxI DO WTW from Fisher Scientific). Before use, the electrode was zeroed by exposure to 100% N2 gas and oxygen (O2) gas or room air. The UV-visible spectrophotometer was interfaced via RS-232C to a Macintosh IIsi computer running FutureBasic II (STAZ Software, Inc.) to provide remote control, data acquisition, and data storage. All chemicals were from Sigma. Stock solutions of 10 mM NADH, 1 mM NAD or 10 mM NADPH, 1 mM NADP and 2000 unit/15 μl superoxide dismutase in 50 mM potassium phosphate buffer, pH 7.8, were freshly prepared and kept on ice until use. The reaction mixture was prepared on ice immediately before individual kinetics runs. Samples included 10–40 nM (heme) for full-length NCB50R or 0.5–1 μM (heme) for truncated NCB50R plus 1000 units of superoxide dismutase in a final volume of 1.5 ml of 50 mM potassium phosphate buffer, pH 7.8. The reaction mixture was equilibrated to the desired dissolved O2 concentration at 20 °C with gentle stirring before initiating the reaction with 15 μl of NADH/NAD or NADPH/NADP solution. The final reaction mixture was stirred for an additional 5 s and subsequently delivered under gas pressure to the flow cell (10-mm path length; Starna). A340 was measured as a function of time from 0 to 600 s in 6-s intervals. Data analysis was performed on a Mac G4 computer (Apple running Excel (Microsoft) with Solver). ΔA340/dt was fit to a linear function, the slope of which was used to calculate the initial velocity of NAD(P)H consumption or superoxide production, vo, using $\text{vo} = \frac{(\text{initial concentration of substrate}) \times \text{time}}{\text{final concentration of substrate}}$. The Michaelis-Menten equation to determine $K_m$ and Vmax and to calculate $K_m$ and the enzyme efficiency $k_{cat}/K_m$. For all calculations, 100% O2 = 1.3 mbar (20°C, 1 atm).

Kinetic Measurement of Cytochrome c Reduction—The spectrophotometric-based steady-state kinetics apparatus, similar to that outlined in Fig. 6A, was used to monitor cytochrome c reduction. The reaction vessel and gas control devices were omitted in the NAD(P)H reductase assay for the following reasons. When compared with the rate of cytochrome c reduction directly by NCB50R, the rate as contributed from superoxide, a product from the NAD(P)H oxidase activity of NCB50R, was negligible under ambient air. The measurement was initiated immediately after NADH or NADPH was added to a quartz cuvette and mixed thoroughly with 25 nM (heme) NCB50R plus oxidized cytochrome c (Sigma) in a final volume of 0.3 ml of low-salt buffer (20 mM potassium phosphate, pH 7.6, 50 mM KCl). To measure the $K_m$ for cytochrome c, the substrate was mixed at a final concentration of 0–40 μM with 0.1 mM NADH or NADPH, and the absorbance change at 540 nm was recorded. To measure the $K_m$ for NADPH or NADP, the reduction was mixed at a final concentration of 0–10 μM with 20 μM cytochrome c, and the absorbance change at 550 nm was recorded. Data analysis was performed as described above. The $v_o$ versus [substrate] curve was fitted to the Michaelis-Menten equation to generate kinetic constants. vo is defined as the rate of electron transfer in all calculations.

$$\text{vo} = \frac{dA_{340}/dt \times \text{c}}{\text{c}} \quad \text{when [cytochrome c] is monitored}$$

$$\text{c}_{\text{340}} = 20 \text{ mM} \times \text{cm}^{-1} \text{ \text{g}}\text{.}$$

(Eq. 1)
Measurement of Oxidation-Reduction Potentials—Potentiometric titrations were performed as described previously (12). 20–50 pmol (10–30 nM) of human NCB5OR (truncated) or cytochrome \(b_5\) core proteins in 1.8 ml of 100 mM KCl, 50 mM MOPS, pH 7.0, were used for each measurement. This buffer was supplemented with Complete proteinase inhibitor mixture (Roche Applied Science) to prevent degradation of the sample during the analysis (3–4 h). Spectra were recorded between 600 and 520 nm at a series of electrode potentials, using a Perkin-Elmer Lambda 18 spectrophotometer. An Orion 720A meter (Orion Research Inc.) was used to measure the half-cell potential relative to a saturated calomel reference electrode. The potential was adjusted by the addition of \(<5 \mu l\) aliquots of solutions of sodium dithionite (reductive titrations) and potassium ferricyanide (oxidative titrations). The accuracy of the apparatus was checked by titration of a 5 mM solution of phenosafranin (\(E = 252\) mV). The following mediators were used at 10 \(\mu M\): phenazine methosulfate; phenazine ethosulfate; anthraquinone; anthraquinone 2-sulfonate; anthraquinone 2,6-disulfonate; 2-hydroxy-1,4-naphthoquinone; and 2,3,5,6-tetramethylphenylenediamine. Pyocyanine was added at 6 \(\mu M\). The degree of reduction of NCB5OR or cytochrome \(b_5\) core was estimated from the height of the absorbance band at 558 nm. The values of absorbance were plotted against electrode potential, and the best fit to the data points was calculated using the GraphPad Prism software package (San Diego, CA) using the Nernst equation.

\[
u = -2 \times \frac{d\Delta u}{dt} \times \varepsilon\]

when [NAD(P)H] is monitored

\(\varepsilon_{240} = 6.22 \text{ mM}^{-1} \text{ cm}^{-1}\)

\(\text{Eq. 2}\)
RESULTS

Ncb5or Gene Family—The Ncb5or gene, first cloned from human liver cells (1), has now been identified in all animals tested, including mammals, chicken, *Xenopus*, fish, tunicate, *Drosophila*, nematodes, sea urchin, and algae. The amino acid sequences of all currently identified NCB5OR proteins are aligned in Supplementary Fig. 1. As reported previously by our group (1), NCB5OR proteins contain a cytochrome *b*₅-like domain at the NH₂ terminus (H₁₁₀₁₁₁₁₀ residues), a *b*₅R-like domain at the COOH terminus (H₁₁₀₁₁₁₁₀ residues), and a hinge (H₁₁₀₁₁₁₀ residues) linking the two domains. The primary sequence of the cytochrome *b*₅ domain is very conserved in all NCB5OR proteins. Likewise, the *b*₅R domains share conserved FAD- and NAD(P)H-binding sites. NCB5OR family members can be distinguished by their overall structural motifs, as well as their signature sequences, which are different from those of the closely related cytochrome *b*₅ and *b*₅R. Namely, NCB5OR is one residue shorter between the two axial histidines in the cytochrome *b*₅ domain and fifteen residues shorter between FAD2 and NAD(P)H1 sites in the *b*₅R domain (Fig. 1 in Ref. 1).

NH₂ Terminus of Endogenous Ncb5or—All Ncb5or genes currently identified in animals have an additional start codon upstream of the one previously assigned. This start codon was not detected during our initial cloning for 5′-untranslated region sequence by rapid amplification of cDNA ends-PCR. Moreover, our previous sequence alignment of 5′-untranslated region presented no consensus among all expressed sequence tag entries available at that time.

Our bioinformatic analysis supports the assignment of an upstream start codon. The Ncb5or genes of zebrafish-Singapore and algae have only the 5′ start codon. Their second putative start site is absent due to a single-base change resulting in an amino acid substitution from methionine to leucine in zebrafish-Singapore or from methionine to valine in algae (shown in Supplementary Fig. 1). In addition, the Kozak sequences in all vertebrate Ncb5or genes have a better match to the optimal sequences at the 5′ ATG site (Fig. 1) because, according to a survey of 699 vertebrate genes (13), A/G is preferred over C/T at the −3 site.

To confirm this NH₂-terminal assignment, we have performed Western blot analysis to detect endogenous NCB5OR with a polyclonal antibody against the NH₂ terminus of NCB5OR. We took advantage of the Ncb5or-null mice that have been recently generated by gene-specific targeting.² By comparing equal amounts of protein samples from wild-type and Ncb5or-null mice, a protein band of the expected size was present in wild-type liver, but not in Ncb5or-null tissue (Fig. 2). The same pattern was observed in mouse pancreata with the same antibody (data not shown) or a polyclonal antibody against the whole mouse NCB5OR.² Therefore, endogenous NCB5OR is translated from the upstream initiation site in these tissues.

Subcellular Localization—Our initial studies of subcellular localization of NCB5OR were based on confocal microscopy on COS7 (monkey kidney) cells expressing exogenous human truncated NCB5OR with a c-myc epitope tag. To define the subcellular localization of endogenous NCB5OR in more detail, we have employed two different methods. As shown in Fig. 2,
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after the removal of nuclei and cytosol, fractionation of intact subcellular organelles of mouse liver on a continuous density gradient established that endogenous mouse NCB5OR is present in isolated fractions that are enriched with the ER marker calreticulin (lanes 4–8) but is not found in the fractions enriched with the mitochondrial marker cytochrome c (lanes 1–3). Because recombinant NCB5OR is soluble, endogenous NCB5OR is likely to reside in the lumen of isolated ER.

In parallel, we have performed two-photon confocal microscopy on endogenous human NCB5OR in HepG2 cells (Fig. 3A), as well as confocal microscopy on exogenous human NCB5OR in COS7 cells (Fig. 3B), to study the colocalization of NCB5OR with ER. Using antibodies against calreticulin and NCB5OR or its epitope tag, we have colocalized NCB5OR with calreticulin in both HepG2 and COS7 cells at the same perinuclear space as we previously reported (1). Because the highest red fluorescence is surrounded by the light blue limits of the ER signals (Fig. 3A), NCB5OR most likely resides within the ER. The same distribution and ER colocalization pattern is observed for both truncated and full-length NCB5OR (Fig. 3B). Both calreticulin and NCB5OR signals are antigen specific because there is no cross-reaction between each second antibody and the primary antibody against the other antigen (data not shown). Similar confocal experiments using antibody against cytochrome c have shown no colocalization between mitochondria and NCB5OR in either HepG2 or COS7 cells (data not shown). This finding agrees with Western blot results on intact subcellular organelles as shown in Fig. 2.

**Stoichiometry of NCB5OR**—To study the biochemical properties of NCB5OR, we have generated high-quality recombinant NCB5OR from a bacterial overexpression system as described previously (1). All our recombinant NCB5OR are native flavohemoproteins containing both FAD and heme. These prosthetic groups have been identified by comparing their retention time with that of FAD and heme standards on reverse phase HPLC (Fig. 4). The color and lack of Coomassie Blue staining of FAD and heme peaks further confirm their identities. Flavin mononucleotide has a longer retention time and thus can be easily distinguished from FAD. NCB5OR proteins have been routinely purified to near homogeneity containing ~85% NADH-reducible heme as estimated from Coomassie Blue staining and UV-visible spectroscopy. Our reverse phase HPLC analysis supports this conclusion.

**Heme Environment**—The heme environment in the native NCB5OR has been probed by resonance Raman spectroscopy. The 413.1-nm resonance Raman spectra of ferrous dithionite-reduced NCB5OR (human truncated) and its cytochrome b₅ core domain are shown in Fig. 5A, a and b, respectively. These spectra are indistinguishable from those obtained from the NADH-reduced NCB5OR proteins. The frequencies of core size marker bands, υ₂ and υ₅, are consistent with six-coordinate, low-spin ferrous heme. The vibrational bands in Fig. 5A have been assigned by analogy to cytochrome b₅ (14), flavocytochrome b₂ (15), and Fe²⁺(PP)IX(ImH₂) (16). The heme resonance Raman frequencies for NCB5OR and cytochrome b₅ are virtually identical, consistent with similar heme environments in the presence and absence of the reductase domain. The 202 cm⁻¹ bands in the NCB5OR and cytochrome b₅ spectra are tentatively assigned to the symmetric stretching mode, υ(His–Fe²⁺(His)), by analogy to 203- and 205-cm⁻¹ bands in the spectra of Fe²⁺(PP)IX(ImH₂) (16, 17) and ferrous flavocytochrome b₂ (15), respectively. Hence, in the ferrous form, NCB5OR and cytochrome b₅ appear to have Fe²⁺-His bond strengths typical of b-type cytochromes with neutral imidazole ligands. The band at 228 cm⁻¹ is in a frequency range consistent with axial imidazole ligands serving as H-bond donors (16). Thus it is possible that there are multiple forms of the low-spin six-coordinate heme.

The resonance Raman spectra for both ferric NCB5OR and cytochrome b₅ proteins (Fig. 5B) exhibit υ₂, υ₅, and υ₁₀ bands at frequencies consistent with six-coordinate, low-spin ferric hemes (17). Because FAD in aqueous solution does not exhibit strong Raman bands at 1372 cm⁻¹, the heme band at 1372 cm⁻¹ was used as an internal subtraction standard for generation of a “NCB5OR–b₅” difference spectrum. The bands shown in Fig. 5B, c correspond to the indicated Raman modes of oxidized flavin (15, 18, 19). These bands, which are attributable to the presence of oxidized flavin in the NCB5OR spectrum, account for the slight differences in the high-frequency regions of Fig. 5B, a and b. Thus, there are no apparent differences in the heme contributions to the spectra of NCB5OR and the cytochrome b₅ core domain. Given the breadth and low intensity of the band(s) near 200 cm⁻¹, no assignment of a symmetric His-Fe²⁺-His stretching band is suggested here. However, there is Raman intensity in the 200–230 cm⁻¹ region of the spectrum that may be attributable to modes with Fe-His stretching character.
Superoxide Production—A spectrophotometric assay was established to elucidate the kinetics of superoxide production by recombinant NCB5OR. Individual data points from seven independent series using human truncated NCB5OR and NADH, normalized by their respective $V_{\text{max}}$, are displayed in Fig. 6B. Using the Michaelis-Menten model, $k_{\text{cat}} = -1 \text{s}^{-1}$, and $K_m = -25 \text{mM} (O_2)$. Similar results were obtained when NADPH was used as the reductant (data not shown). As shown in Table I, human and mouse full-length NCB5OR proteins have almost identical kinetic constants: $k_{\text{cat}} = -0.05 \text{s}^{-1}$ and $K_m = -12 \text{mM} (O_2)$. Their $k_{\text{cat}}$ is 20-fold lower than that of truncated human NCB5OR. Because physiological enzymes have $k_{\text{cat}}$ values ranging from 1 to $10^6$ s$^{-1}$ (20), the $k_{\text{cat}}$ value of NCB5OR is either at the low limit of an enzyme-based reaction (truncated NCB5OR) or far beyond the low limit (full-length NCB5OR). The same conclusion can be reached by using their catalytic efficiency, $k_{\text{cat}}/K_m$ (20). Therefore, full-length NCB5OR by itself would not be expected to function as a NAD(P)H oxidase in vitro or in vivo.

Cytochrome c Reduction—We have used cytochrome $c$ to measure the NADH reductase activity of NCB5OR. As shown in Table I, both full-length and truncated NCB5OR are able to reduce cytochrome $c$ with NADH or NADPH. Overall, the two full-length NCB5OR proteins from human and mouse have very similar kinetic behavior. Each NCB5OR protein can use both NADH and NADPH as a reductant with similar catalytic efficiency.

The catalytic efficiency with artificial substrates, cytochrome $c$ (Table I), and ferricyanide (data not shown) for all the full-length and truncated NCB5OR proteins tested in this study is in the physiological range (20) and at least 68-fold higher than that with molecular oxygen. Therefore, both full-length and truncated NCB5OR function as a NAD(P)H reductase in vitro.

Redox Potential—We have measured the redox potential of the heme center in truncated human NCB5OR with potentiometric titrations. The midpoint potential was calculated to be $-108$ mV for truncated human NCB5OR (Fig. 7), as well as for its cytochrome $b_5$ core (data not shown). As shown in Table II, this value is almost identical to that of solubilized cytochrome $b_5$ on outer mitochondrial membrane ($-102$ mV). Despite sharing the same cytochrome $b_5$ global structure, both the cytochrome $b_5$ core of nitrate reductase and solubilized cytochrome $b_5$ on ER membrane have a similar value ($-0$ mV), whereas cytochrome $b_5$ core of yeast cytochrome $b_5$ has a value between $-10$ and $-30$ mV. Although the redox potential of the heme in NCB5OR is sufficiently high to permit its reduction by a number of biological reductants, in view of the marked similarity of this system to the classic cytochrome $b_5$ reductase/cytochrome $b_5$ system, it seems highly probable that the physiological reductant(s) of NCB5OR is NADH or NADPH, both of which have redox potentials around $-220$ mV (20).

**DISCUSSION**

Impact of NH$_2$ Terminus on NCB5OR Structure and Function—The NH$_2$ termini of NCB5OR from mammals to *Xenopus* have the same length and very similar amino acid sequences. As the comparison is extended into invertebrates, there is less similarity both in length and in primary sequence. Compared with the other regions of NCB5OR, the NH$_2$ terminus is most divergent, which implies a less important functional role. Nevertheless the NH$_2$-terminal exon clearly has functional significance when NAD(P)H-dependent superoxide production and cytochrome $c$ reduction are monitored. Truncated NCB5OR has a slightly higher catalytic efficiency than human and mouse full-length NCB5OR in both reactions. The only exception is the NADPH-dependent cytochrome $c$ reduction by mouse full-length NCB5OR.

The NH$_2$-terminal sequences appear to have no impact on the solubility or subcellular localization of NCB5OR. Both full-length and truncated recombinant NCB5OR proteins are soluble. The same distribution and ER colocalization pattern is observed for both truncated and full-length NCB5OR by confocal analysis. When overexpressed in COS7 cells, both forms of NCB5OR are found in the lysate, not in membrane fractions (data not shown).

**Lack of NAD(P)H Oxidase Activity**—Superoxide production by the full-length NCB5OR is not favored due to its extremely low $k_{\text{cat}}$ value. Moreover, the redox potential at the heme center of NCB5OR is at least $100$ mV higher than the values $-225$ and $-265$ mV (12) of the two heme centers in the respiratory burst.
oxidase cytochrome b<sub>558</sub>. This NAD(P)H oxidase is best known for its robust H<sub>2</sub>O<sub>2</sub> production with a k<sub>cat</sub> of ~300 s<sup>-1</sup> (29). Because E<sub>red</sub> = −160 mV for O<sub>2</sub> → O<sub>2</sub><sup>−</sup> (30), the superoxide production at the heme center of cytochrome b<sub>558</sub> is greatly favored thermodynamically, whereas reduction of oxygen to superoxide would not be favored at the NCB5OR heme center.

Until recently, neutrophil cytochrome b<sub>558</sub> was the only mammalian b-type cytochrome known to be capable of converting oxygen into superoxide as a natural product through six-coordinate low-spin hemes (31, 32). In cytochrome b<sub>558</sub>, the electron transfer from heme to oxygen is thought to be an outer-sphere process from the heme edge and not directly from the heme iron. A number of homologues of flavocytochrome b<sub>558</sub> have been discovered recently (the NAD(P)H oxidase family), and it is likely that some or all of them produce superoxide at the expense of NAD(P)H. These proteins are likely to play important roles in cell signaling (reviewed in Ref. 33). Other mammalian hemeproteins have also been shown to generate superoxide at low rates by autoxidation of heme; these include nitric-oxide synthase (34) and other members of the cyto-

**Table I** Kinetic constants of NCB5OR

| Substrate                  | Kinetic constants | Human full-length | Human truncated | Mouse full-length |
|----------------------------|-------------------|-------------------|-----------------|------------------|
| Molecular oxygen           | k<sub>cat</sub> (s<sup>−1</sup>) | 0.055             | 1.0             | 0.054            |
| Fixed [NADH]               | K<sub>m</sub> (μM) | 13                | 25              | 11               |
| Fixed [NADPH]              | K<sub>m</sub> (μM) | 7.0               | 2.5             |
| Cytochrome c               | k<sub>cat</sub> (s<sup>−1</sup>) | 0.004             | 0.04            | 0.005            |
| Fixed [NADH]               | K<sub>m</sub> (μM) | 3.0               | 4.0             | 3.2              |
| Fixed [NADPH]              | K<sub>m</sub> (μM) | 12                | 2.5             | 13               |
| Cytochrome c               | k<sub>cat</sub> (s<sup>−1</sup>) | 2.7               | 3.5             | 2.5              |
| Fixed [NADH]               | K<sub>m</sub> (μM) | 1.4               | 1.3             |
| Fixed [NADPH]              | K<sub>m</sub> (μM) | 2.8               | 7.2             | 2.7              |
| Cytochrome c               | k<sub>cat</sub> (s<sup>−1</sup>) | 3.0               | 3.5             | 3.5              |
| Fixed [cytochrome c]       | K<sub>m</sub> (μM) | 1.8               | 1.4             | 1.3              |
| NADH                       | k<sub>cat</sub> (s<sup>−1</sup>) | 2.4               | 2.4             | 6.4              |
| Fixed [cytochrome c]       | K<sub>m</sub> (μM) | 2.4               | 2.4             | 6.4              |
| NADPH                      | k<sub>cat</sub> (s<sup>−1</sup>) | 1.4               | 0.91            | 2.0              |
| Fixed [cytochrome c]       | K<sub>m</sub> (μM) | 1.7               | 2.7             | 3.2              |

**TABLE II** Redox potential of heme center in NCB5OR and related heme proteins

| Protein                                 | E (mV)   | Reference |
|-----------------------------------------|----------|-----------|
| NCB5OR                                  | −108     | This study|
| Human, whole                            | −108     | This study|
| Human, b<sub>5</sub> core               | −102     | 21        |
| Solubilized outer mitochondrial membrane b<sub>5</sub> | −102     | 21        |
| Human and rat, whole                    | −102     | 21        |
| Nitrate reductase                       | −3       | 24        |
| Spinach, b<sub>5</sub> core             | −10      | 25        |
| Green algae, b<sub>5</sub> core         | 4        | 26        |
| Solubilized ER membrane b<sub>5</sub>   | −31      | 27        |
| Human                                   | −10      | 28        |
| Hansenula anomala                       | −25      | 12        |
| Respiratory burst oxidase b<sub>558</sub> | −225, −265 | 12    |
| Human, 2 heme centers                   | −225, −265 | 12    |

Fig. 7. Oxidation-reduction potential measurements of recombinant NCB5OR. The absorbance at 558 nm (y axis) is used to monitor the reduced heme and plotted against electrode potential (x axis) to calculate the midpoint redox potential at the heme center of human truncated NCB5OR. The solid line represents the best-fit theoretical line for a single-species, one-electron transfer process with a midpoint potential of −108 mV. Triangles represent oxidative titrations with potassium ferricyanide; squares represent reductive titrations with sodium dithionite.
chrome P450 family, hemoglobin/myoglobin, and cytochrome $b_5$ (reviewed in Ref. 35). In all these cases, the production of superoxide is thought to be an unwanted side reaction. Because both rat and human NCB5OR have a six-coordinate low-spin heme as in other cytochrome $b_5$ proteins and, more importantly, the redox potential at the heme center is higher than that needed to favor the oxygen conversion to superoxide, it would not be expected for NCB5OR to function as an NAD(P)H oxidase. The NAD(P)H-cytochrome $c$ reductase activity of NCB5OR is likely mediated by electron channel through side chains of charged residues, similar to those found in classical single-domain cytochrome $b_5$ and cytochrome $c$ complex (36–38).

In the low frequency resonance Raman spectrum of NAD(P)H oxidase cytochrome $b_{550}$, the ferric Fe-His stretching frequency has been tentatively assigned to a band at 269 cm$^{-1}$. If this assignment is correct, it would be consistent with anionic (imidazolate) character of the axial His side chains (17). This is consistent with the relatively negative redox potential of cytochrome $b_{550}$ because anionic axial ligands generally stabilize the ferric state of the heme preferentially. By contrast, the resonance Raman spectra of NCB5OR and its cytochrome $b_5$ core domain only show the $v_{4b}$ bands of the ferric and ferrous hemes in this region of the spectrum. Although it is possible that a Fe$^{III}$-His band could be overlapping the 272 cm$^{-1}$ band, this would be a very high frequency and seems unlikely. It is more likely that the resonance Raman intensity between 200 and 230 cm$^{-1}$ is attributable in full or in part to vibrations with Fe-His stretching contributions. Interestingly, there seems to be little or no difference between the frequencies of these bands in the ferrous (Fig. 5A) and ferric (Fig. 5B) spectra. This apparent similarity in the Fe-His bond strengths of the ferrous and ferric hemes, as reflected by Fe-His stretching frequencies, is consistent with the more positive redox potential of $-108$ mV relative to that observed in the NAD(P)H oxidase cytochrome $b_{550}$. Thus, the lack of NAD(P)H oxidase activity may be correlated with relative invariance in the Fe(II) and Fe(III) axial ligand bond strengths in NCB5OR.

The coexistence of the cytochrome $b_5$ and cytochrome $b_5$ reductase activities in the same peptide begs the question of whether interaction of the heme and reductase domains or related with relative invariance in the Fe(II) and Fe(III) axial ligand bond strengths in NCB5OR and the cytochrome $b_5$ core domain suggest that the redox potential is an intrinsic property of the cytochrome $b_5$ domain. Hence, the thermodynamic driving force for heme electron transfer reactions and the immediate heme environment would appear to be insensitive to the reductase.

**ER Localization**—An important clue to understanding the biological function of NCB5OR is its subcellular localization. As a soluble protein, the full-length NCB5OR appears to reside in the lumen of ER, as documented by two independent methods. Because NCB5OR protein has a slow turnover in cells, its presence in the ER is unlikely to be due to rapid synthesis and degradation. NCB5OR is distributed closer to nuclei and more focused than calreticulin, probably because calreticulin, as a member of the ER molecular chaperones, can associate with incompletely folded protein intermediates in pre-Golgi compartments (39, 40).

The ER localization of NCB5OR is somewhat unexpected because the NH$_2$-terminal sequence in full-length NCB5OR lacks a canonical signal peptidase sequence. NCB5OR also lacks a COOH-terminal KDEL motif (41) as well as NH$_2$-terminal or COOH-terminal KXXX and KXXX motifs (42, 43). Some ER proteins lack these classic addressins (44). All vertebrate NCB5OR share a SQQR sequence just downstream from an NH$_2$-terminal hydrophobic stretch, consistent with the (S/T)-(Q/E)-(R/K) motif that directs glycosyltransferases to the Golgi (45). Alternatively, a thiol-mediated protein retention mechanism through Erp44 may be involved (46).

**Function of NCB5OR**—Based on circumstantial evidence that a cytochrome $b$-type NAD(P)H oxidase might regulate target gene expression through oxidative modification of hypoxia-inducible factor 1 (47–49), we initially suggested that NCB5OR might function as a candidate oxygen sensor. The truncated human NCB5OR has kinetic properties consistent with a weak oxidase with a $K_a$ of $\sim 2\%$, responding to oxygen in a physiological range. However, the markedly lower $K_a$ of full-length protein indicates that the endogenous NCB5OR cannot function as an NAD(P)H oxidase.

The biological function of NCB5OR remained elusive until we prepared mice with targeted inactivation of the gene. The predominant phenotypic defect is insulin-deficient diabetes. The pancreatic $\beta$ cell is particularly susceptible to ER stress (50). Thus, the phenotype of the Ncb5or-null mouse suggests that this enzyme plays a critical role in the ER stress response pathway. It is noteworthy that the phenotype of the Ncb5or-null mouse bears a striking resemblance to that of two mouse models of ER stress in the $\beta$ cell, the Akita mouse (Ins2 C96Y) (51) and the Perk-null mouse (52, 53). All three have normal pancreatic and $\beta$-cell development, but at age 4–6 weeks, they develop progressive loss of $\beta$-cell mass and insulin-deficient diabetes. Perinuclear space in various cells has been shown to stain heavily for hydroxyl free radicals (49), which correlates with the oxidative environment within ER lumen, a necessary condition for protein folding (54). As disulfides bonds are formed in the ER lumen, electrons are passed first to protein disulfide isomerase and then to Ero1, a soluble FAD-binding protein (9). Oxygen appears to be the terminal electron acceptor. Disulfide bond formation may account for about 25% of the superoxide generated in most cells (54). Oxidative stress in the perinuclear space in various cells has been shown to stain heavily for hydroxyl free radicals (49), which correlates with the oxidative environment within ER lumen, a necessary condition for protein folding (54). As disulfides bonds are formed in the ER lumen, electrons are passed first to protein disulfide isomerase and then to Ero1, a soluble FAD-binding protein (9). Oxygen appears to be the terminal electron acceptor. Disulfide bond formation may account for about 25% of the superoxide generated in most cells (54).

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