Molecular basis of aromatase deficiency in a 46, XX patient with mutation of arginine 550 to tryptophan in POR: Expanding the endocrine phenotype in PORD.

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Abstract

Context: Mutations in Cytochrome P450 oxidoreductase (POR) cause a form of congenital adrenal hyperplasia (CAH). We are reporting a novel R550W mutation in POR identified in a 46, XX patient with signs of aromatase deficiency.

Objective: Analysis of aromatase deficiency from R550W mutation in POR.

Design, setting, and patient: Both the child and the mother had signs of virilization. Ultrasound revealed the presence of uterus and ovaries. No defects in CYP19A1 were found, but further analysis with a targeted Disorders of Sexual Development NGS panel (DSDSeq.V1, 111 genes) on a NextSeq (Illumina) platform in Madrid and Barcelona, Spain, revealed compound heterozygous mutations c.73_74delCT/p.L25FfsTer93 and c.1648C>T/p.R550W in POR. WT and R550W POR were produced as recombinant proteins and tested with multiple cytochrome P450 enzymes at University Children’s Hospital, Bern, Switzerland.

Main outcome measure and Results: R550W POR showed 41% of the WT activity in cytochrome c and 7.7% activity for reduction of MTT. Assays of CYP19A1 showed a severe loss of activity and CYP17A1, as well as CYP21A2 activities, were also lost by more than 95%. Loss of CYP2C9, CYP2C19, and CYP3A4 activities was observed for the R550W-POR. Predicted adverse effect on aromatase activity as well as a reduction in binding of NADPH was confirmed.

Conclusions: Pathological effects due to POR R550W were identified, expanding the knowledge of molecular pathways associated with aromatase deficiency. Screening of the POR gene may provide a diagnosis in CAH without defects in genes for steroid metabolizing enzymes.

Précis

In a 46, XX patient with aromatase deficiency, a mutation in POR was identified, which inhibited aromatase activity, expanding the molecular pathogenesis of aromatase deficiency.
Introduction

Cytochrome P450 oxidoreductase (POR) is the redox partner of cytochrome P450 proteins located in the endoplasmic reticulum (1). POR deficiency (PORD, OMIM: MIM613571 and MIM201751) is a form of congenital adrenal hyperplasia, initially described in patients with altered steroidogenesis (2, 3) followed by several reports with a broad spectrum of disorders (4-6). In 2004, mutations in POR disrupting steroid biosynthesis were reported (2, 5, 6), and in subsequent reports, a range of POR mutations causing disorders of sexual development with and without bone malformation have been described (4, 7, 8). POR was identified by studies of Lu and Coon as a component of microsomal mixed oxidase system along with lipids and cytochrome P450 proteins (9). POR transfers redox equivalents from the reduced form of nicotinamide adenine dinucleotide phosphate (NADPH) to cytochrome P450 proteins for their catalytic activities (10). Also, POR can reduce heme oxygenase, cytochrome b5, and a range of small molecules and dyes like ferricyanide (FeCN), 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT), etc (1, 11). Due to its unique role in a range of metabolic processes, POR is an essential protein and POR knockout mouse are embryonically lethal (12). POR deficiency was initially identified as a disorder of sex development with ambiguous genitalia resembling some features of Antley-Bixler bone malformation syndrome. However, after findings of multiple genetic defects in POR from patients both with and without bone malformation features, PORD is now considered a separate genetic disorder (7).

Due to the involvement of POR in multiple reactions in the metabolism of both the steroids and drugs, exact effects of individual mutations in POR are hard to predict and require extensive characterization by enzymatic and biochemical analysis (2, 7, 13-20). Therefore, the exact mechanisms by which mutations in POR cause pathogenesis, remain mostly unknown. From the characterization of multiple mutations in POR by us as well as other laboratories, some general conclusions could be drawn. Mutations in POR that result in loss of flavin co-factors (FAD and FMN) show a severe loss of enzymatic activities and cause a lethal form of POR deficiency. Mutation located in the hinge region of POR that is necessary for FMN and FAD domain movements cause a phenotypically variable form of the disease and impact different P450 enzyme activities to different extents. Mutations located near the NADPH binding site of POR may impact the activities of multiple cytochrome P450 activities, but the impact on individual enzymes could be variable.

Aromatase is a 503 amino acid protein (NP_000094) encoded by the CYP19A1 gene (MIM10790, GeneID:1588, NCBI: NM_000103), located on chromosome 15 (15q21.2, GRCh38 15:51208056-51338597). CYP19A1 (EC: 1.14.14.14) regulates estrogen biosynthesis in humans (21) by converting androgens to estrogens (22-25). Aromatase is highly expressed in ovaries and plays a significant role in the regulation of the reproductive cycle in females (26). Some of the critical reactions catalyzed by...
aromatase include the conversion of androstenedione to estrone (E1), testosterone to 17β-estradiol (E2) and 16-hydroxytestosterone to estriol (E3) (21). The catalytic process of aromatization of androgens is multifaceted, and it comprises of three steps involving the transfer of three pairs of electrons from the reduced form of nicotinamide adenine dinucleotide phosphate (NADPH) and consumption of oxygen. Electron transfer from NADPH to aromatase is carried out by NADPH cytochrome P450 oxidoreductase (POR).

The first case of placental aromatase deficiency was reported by Shozu et al. (27), and aromatase deficiency manifests during fetal life in both sexes (28). Multiple genetic defects are associated with decreased aromatase expression in both male and female reproduction (29). In females, aromatase deficiency affects maternal virilization during gestation, abnormal external genitalia in the female fetus, normal ovarian development but increased polycystic ovary syndrome occurrence and virilization at puberty (30, 31). Impact of aromatase deficiency on growth, skeletal maturation, and bone homeostasis, as well as changes in insulin resistance and abnormal plasma lipid profile, have been described in earlier studies (32, 33).

Here we have performed a detailed characterization of a novel mutation in POR, R550W, that caused aromatase deficiency in a 46, XX patient and led to maternal virilization during pregnancy. By use of biochemical analysis of patient samples and production of a recombinant form of POR containing the R550W mutation, we have been able to characterize the different enzyme activities affected by the R550W mutation in POR. We investigated the effect of R550W mutation in POR on steroid metabolizing P450 enzymes CYP17A1 (17-hydroxylase activity), CYP21A2 (21-hydroxylase activity) and CYP19A1 (aromatase activity) and could show that CYP19A1 activity was severely affected by the R550W mutation in POR, explaining the cause of virilization observed the in 46, XX female child. Moreover, we also tested the effect of R550W mutation in POR on activities of four drug-metabolizing cytochromes P450 (CYP2C9, CYP2C19, CYP3A4 and CYP3A5) in addition to testing the direct metabolic activities of POR on small molecules (FeCN and MTT) and a soluble probe substrate cytochrome c, to identify different mechanisms that may affect POR due to R550 mutation. Our experiments identified two distinct mechanisms for pathogenicity of the POR mutation R550W: inhibition of CYP19A1 activity leading to virilization of the female child, and the overall reduced POR activities due to loss of flavin content, suggesting protein instability. These results expand our knowledge about the pathology of PORD and disruption of steroidogenesis resulting from mutations in POR.
**Case Report and Methods**

**Patient**

The patient was born at term in Malaga (Spain) in 2009, from the 1st pregnancy of non-consanguineous parents. The mother had presented signs of virilization from the 6th month of pregnancy (acne, hirsutism, clitoris growth, and voice deepening) that regressed progressively post-partum. Hormone analyses at post-partum day 5 revealed a highly increased testosterone (T) serum concentration that returned to normal female levels at the fourth postpartum month (Table 1). The child was explored at seven days: length 48 cm (-1.22 SDS), weight 2,920 g (-1.02 SDS), the external genitalia (almost complete labia majora fusion with scrotal aspect, no gonads palpable, genital tubercle of 1.5 cm with proximal urethral opening, considered as Prader type 3 ambiguous genitalia). At three months, clitoris hypertrophy had regressed. Initial (7 days of age) 17-hydroxy-progesterone (17-OH-P) and T were elevated and progressively diminished; T reached normal female levels from the 1st month, whereas 17-OH-P was still slightly elevated at seven months (Table 1). The karyotype was 46, XX, and 21-hydroxylase deficiency was ruled out since the 17-OHP increased at birth, but the values were below those seen in the classic form of 21-hydroxylase deficiency and showed a progressive reduction. Pelvic ultrasound revealed the presence of uterus and bilateral gonads. Transient intrauterine virilization was diagnosed due to maternal pregnancy luteoma or aromatase deficiency that would explain the transient virilization of the mother and the female child. The CYP19A1 gene was sequenced from the DNA of the child, but no pathogenic sequence variant was detected. The patient developed without any further significant clinical manifestations. A genitoplasty was performed at 13 months.

**Hormone assays**

Hormone assays were performed in Málaga (Hospital Carlos Haya), Spain. The 17-OH-P was determined with a commercial radioimmunoassay technique (Coat-A-Count. Siemens Healthcare Diagnostics Ltd. Frimley, Camberley, UK); dehydroepiandrosterone sulfate (DHEA-S), T, estradiol (E2) and cortisol were determined with automated electrochemiluminescent assays (Modular E, Roche Diagnostics GmbH Mannheim, Germany) and ACTH with an automated chemiluminescence immunoassay (LIAISON, DiaSorin, Saluggia, Italy).

**Genetic studies**

The study was approved by the Ethics Committee of Hospital Universitari Vall d'Hebron (CEIC), Barcelona, Spain (PR (IR) 23/2016) and parents provided informed consent. Genomic DNA was isolated from whole peripheral blood samples from the patient and her family using standard procedures. The DNA samples were analyzed with a custom-designed targeted Disorders of Sexual Development NGS panel (DSDSeq.V1, 111 genes, and three regulatory regions) using SeqCap EZ technology (Roche Nimblegen) and sequenced on a NextSeq500 platform (Illumina) platform. All
procedures were carried out according to the manufacturer’s instructions. Data analyses were performed
using VariantStudio V2.2.1 (Illumina) and in-house bioinformatic analysis as previously described (34).
The observed variants were evaluated using the Alamut Visual software v2.11 (https://www.interactive-
biosoftware.com/es/alamut-visual/), which includes following pathogenicity prediction tools: Mutation
Taster, PolyPhen, Align GVGD and SIFT) MutAssesor, Fasthmm, Vest and CADD V1.4
Spanish Variant Server (CSVs; CIBERER BIER, Valencia, Spain; http://csvs.babelomics.org/) were
consulted for allelic frequencies. Candidate variants observed by targeted panel (DSDSeq.V1) were
sequencing as well as segregation analysis in the family were validated and genotyped by Sanger
sequencing. Variant classification was performed according to ACMG recommendations (35).

Materials
Tris-base, NADPH, acetic acid, magnesium acetate, Sucrose, potassium phosphate EDTA, DTT,
glycerol, PMSF, and Benzonase were purchased from Sigma-Aldrich Chemie GmbH (Buchs,
Switzerland). Carbenicillin, FeCl₃, ZnCl₂, CoCl₂, Na₂MoO₄, CaCl₂, CuCl₂, M H₃BO₃ were purchased
from CarlRoth GmBH (Switzerland). Goat anti-rabbit antibodies labeled with infra-red dyes were from
LI-COR Bioscience Inc. (NE, USA). The RC-DC protein assay dye reagent was from Bio-Rad
(Hercules, CA). The anti-POR antibody was from GenScript (NJ, USA). BOMCC (Invitrogen Corp,
Carlsbad, CA, United States).

Expression of POR proteins in bacteria and membrane purification
The human POR WT and R550W mutant proteins (NCBI# NP_000932, UniProt# P16435) were
expressed in bacteria using heterologous gene expression (7, 20, 36, 37). The protocol for recombinant
expression of POR variants (N-23 form) and subsequent bacterial membrane purification was based on
our previous publications (7, 17, 18, 20, 38) with slight modification. Briefly, plasmid (pET15b)
containing cDNAs for WT or mutant POR were obtained from GenScript. The Escherichia coli
bacterial strain BL21(DE3) was transformed, and single colonies were selected for growth on
carbenicillin plate. For the main shake flask culture, autoinduction media consisting of terrific broth
supplemented with 40 mM FeCl₃, 4 mM ZnCl₂, 2 mM CoCl₂, 2 mM Na₂MoO₄, 2 mM CaCl₂, 2 mM
CuCl₂, 2 mM H₃BO₃, 0.5 mg/ml riboflavin, 100 µg/ml carbenicillin was used. The cells were grown at
37 °C to an optical density (OD) at 600 nm of 0.6, and then the temperature was reduced to 25 °C and
further grown for 16 h. The bacterial cells were collected by centrifugation, washed with PBS and
slowly stirred for 1 h at 4°C for in 50 mM Tris-acetate (pH 7.6), 0.25 M sucrose, 0.5 mM EDTA,
lysozyme (0.2 mg/ml), 1 mM PMSF and 20 U/ml endonuclease to prepare spheroplasts. The
spheroplasts were collected by centrifugation at 5000 x g for 20 min; and suspended in 50 mM
potassium phosphate (pH 7.6) containing 6 mM Magnesium acetate, 0.1 mM DTT, 20% (v/v) glycerol
and 1 mM PMSF; and disrupted by sonication. After centrifugation at 12000 × g for 15 min at 4 °C, the supernatant was centrifuged at 100000 × g for 90 min at 4 °C to collect membranes. Purified membranes containing POR were resuspended in 50 mM Potassium phosphate buffer (pH 7.8) and 20% (v/v) glycerol and kept at −70 °C. Protein concentration was measured by the RC-DC protein assay method (Protein Assay Dye Reagent, Bio-Rad, Hercules, CA) and POR content in membrane proteins was measured by western blot analysis.

**Western Blot Analysis of POR Content in the Bacterial Membranes**

Western blot analysis to determine POR content in membranes was done as described previously (57). Briefly, 1 µg of bacterial membrane proteins were separated on an SDS-PAGE gel and blotted on to polyvinyl difluoride (PVDF) membranes. Blots were first incubated with a rabbit polyclonal antibody against POR-WT from GenScript (GenScript, NJ, USA) at a dilution of 1:1000. We then used a secondary goat anti-rabbit antibody labeled with an infrared dye (IRDye 680RD, LI-COR Bioscience Inc., NE, USA) at a 1:15000 dilution. Signals were analyzed with the 700 nm fluorescent channel on an Odyssey Infrared Imaging System (LI-COR Bioscience Inc., NE, USA), and protein bands were quantitated using the Odyssey software (LI-COR Bioscience Inc., NE, USA). Purified wild-type POR was used as a standard for normalizing the POR content of each membrane preparation in all experiments described here, the normalized amount of POR content was used for POR-WT and POR-R550W protein.

**Small molecule Reduction Assay by POR-WT and POR-R550W**

Cytochrome c reduction by POR-WT or POR-R550W was performed as described previously by measuring the change in absorbance at 550 nm (ε=21.1 cm⁻¹ mM⁻¹) (39). In brief, varying concentrations of cytochrome c (1.3–40 µM) were mixed with membrane preparations containing 1 µg POR in 100 mM phosphate buffer (pH 7.6) in a total volume of 100 µl. The reaction was started by the addition of NADPH, and the change in absorbance at 550 nm was monitored over 6 minutes. The reaction was performed in 96-well plates, in triplicate, using a microplate reader (Spectramax M2e, Molecular Devices, Sunnyvale, CA). Data were fitted based on Michaelis-Menten kinetics using GraphPad Prism (GraphPad Software, La Jolla, CA USA) to determine the Vₘₐₓ and Kₘ. The MTT [3-(4,5-dimethylthiazol- 2-yl)-2,5-diphenyltetrazolium] reduction assay was carried out using different concentrations of MTT (3.9-500 µM), by measuring the rate of increase in absorbance at 610 nm (ε₆₁₀=11 mM⁻¹ cm⁻¹) (40). The reaction mixture consisted of bacterial membranes containing 1 µg POR in 100 mM phosphate buffer (pH 7.6) and 100 µM NADPH. Similarly, the ferricyanide reduction was measured as the rate of decrease in absorbance at 420 nm (ε₄₂₀=1.02 mM⁻¹ cm⁻¹). The reaction was started by adding 100 µM NADPH and the concentration of ferricyanide varied from 1.3 to 500 µM (41). Activities represent the mean of at least three replicates. For NADPH variation analysis,
cytochrome c (40 µM) and MTT (500 µM) were kept constant, and NADPH concentration was varied from 0.8-100 µM.

Flavin Content Analysis of WT and Mutant POR
Flavins were extracted by thermal denaturation of POR proteins and analyzed as described previously (42). POR-WT or POR-R550W (100 µg/ml) were denatured by heating at 95 °C for 10 min in the dark and flavins were separated by centrifugation at 14000 x g for 10 min from the precipitated protein. The fluorescence of the supernatants was measured at pH 7.7 and pH 2.6 to determine FMN and FAD ratio (excitation at 450 nm, emission at 535 nm).

Expression and Purification of human CYP19A1
The recombinant human CYP19A1 was expressed and purified, as described previously (43) with some modifications. Briefly, a single colony of E. coli BL21(DE3) transformed with expression vectors for CYP19A1, and molecular chaperones GroEL/GroES, was selected for protein expression. A 1:100 dilution of overnight growth culture was used to inoculate the autoinduction medium (terrific broth, 40 mM FeCl₃, 4 mM ZnCl₂, 2 mM CoCl₂, 2 mM CaCl₂, 2 mM CuCl₂, 2 mM H₂BO₃, 100 µg/ml carbenicillin and 50 µg/ml kanamycin). After 4 h of incubation at 25°C, 1 mM δ-aminolevulinic acid and 4 mg/ml arabinose (for induction of molecular chaperones GroEL/GroES) were added, and the culture was grown for another 20 h. Bacteria were then harvested, washed with PBS, and stored at -80 °C. Bacterial pellet was resuspended in 50 mM Tris-Acetate (pH 7.6), 250 mM sucrose, 0.5 mM EDTA, and 0.2 mg/ml lysozyme to prepare spheroplasts. For purification, spheroplasts were lysed using 10 x CellLytic B (Sigma-Aldrich) in a 100 mM potassium phosphate (pH 7.4) buffer containing 20% glycerol, 500 mM sodium acetate, 0.1 mM DTT, 0.1 mM EDTA and 1 mM PMSF. The supernatants after centrifugation were pooled and CYP19A1 was purified by Ni²⁺ metal-chelate chromatography at 4 °C using 200 mM histidine for elution. Eluted protein was dialyzed to remove histidine, and protein concentration was determined by RC-DC protein assay (Protein Assay Dye Reagent, Bio-Rad, Hercules, CA) using BSA as standard.

Cytochrome P450 CYP17A1 enzyme activity supported by WT and POR-R550W
Purified recombinant CYP17A1 was used to test the effect of POR-R550W to support the 17α-hydroxylase activity of CYP17A1. Bacterial membranes containing POR and purified CYP17A1 (CYPEX, Dundee, Scotland, United Kingdom) were reconstituted as described previously (7, 44-46) and 17α-hydroxylase activity of CYP17A1 was measured by using progesterone as substrate. The reaction mixture consisted of 30 pmol of CYP17A1, 60 pmol of POR, and ¹⁴C labeled progesterone (50000 cpm) in 50 mM potassium phosphate buffer (pH 7.4). The reaction mixture was supplemented with 10 mM magnesium chloride, 6 mM potassium acetate, and 1 mM reduced glutathione.
Progesterone concentration was varied from 0.3 µM to 5 µM for kinetic analysis, and the reaction was initiated by the addition of 2 mM NADPH and incubated for 60 min at 37 °C. Data were fitted based on Michaelis-Menten kinetics using GraphPad Prism (GraphPad Software, La Jolla, CA USA).

**Assay of cytochrome P450 CYP21A2 enzyme activity supported by WT and POR-R550W**

Purified recombinant CYP21A2 was used to test the effect of POR-R550W to support the 21-hydroxylase activity of CYP21A2. Bacterial membranes containing POR and purified CYP21A2 (CYPEX, Dundee, Scotland, United Kingdom) were reconstituted as described previously (7, 44-46) and 21-hydroxylase activity of CYP21A2 was measured using progesterone as the substrate. The reaction mixture consisted of 20 pmol of CYP21A2, 40 pmol of POR, and 14C labeled progesterone (50000 cpm) in 50 mM potassium phosphate buffer (pH 7.4). The reaction mixture was supplemented with 10 mM magnesium chloride, 6 mM potassium acetate, and 1mM reduced glutathione. Different concentrations (0.3–5 µM) of progesterone were used for kinetic analysis, and the reaction was initiated by the addition of 2 mM NADPH and incubated for 60 min at 37 °C. Data were fitted based on Michaelis-Menten kinetics using GraphPad Prism (GraphPad Software, La Jolla, CA USA).

**Cytochrome P450 CYP19A1 enzyme activity supported by WT and POR-R550W**

Purified recombinant CYP19A1 using the bacterial expression system was used to test the effect of POR-R550W to support the aromatase activity. Aromatase activity was measured by the tritiated water release assay based on an earlier method described by Lephart and Simpson (47) with some modifications (20, 44) using a reconstituted system and androstenedione (androst-4-ene-3,17-dione) as the substrate. Bacterial membranes containing POR and purified CYP19A1 was reconstituted in a 1:2 ratio. Reaction mixture consisted of 50 pmol of POR, 100 pmol of CYP19A1 and tritium labeled androstenedione ([1H-3H(N)]-androstene-3,17-dione; ~15,000 cpm) in 100 mM potassium phosphate buffer (pH 7.4) with 100 mM NaCl. Different concentrations (10–1000 nM) of androstenedione were used for kinetic analysis. The aromatase reaction was started by addition of 1 mM NADPH and was incubated for 60 min at 37 °C. After incubation, 0.8 ml of 5% charcoal/0.5% dextran solution was added to the reaction mixture. Reaction tubes were mixed by vortex and centrifuged at 15000 x g for 5 minutes. From each tube, 0.5 ml of supernatant was used for radioactivity measurement. Values obtained from duplicate experiments are expressed as mean ± S.E.M. (standard error of the mean). Data were fitted based on Michaelis-Menten kinetics using GraphPad Prism (GraphPad Software, La Jolla, CA USA). For NADPH kinetic analysis, androstenedione concentration was kept constant at 100 nM, and NADPH was varied from 62.5-1000 µM.

**Assay of cytochrome P450 CYP2C9 activity supported by WT and POR-R550W**

The activity of CYP2C9 supported by WT or mutant POR was tested using *in vitro* reconstituted system. It consisted of bacterial membranes containing WT/R550W POR, purified CYP2C9 (CYPEX, Dundee,
Scotland, United Kingdom) and purified cytochrome b5 at a ratio of 5:1:1 (48). The fluorogenic compound BOMCC (7-Benzoyloxy-4-trifluoromethylcoumarin) (Invitrogen Corp, Carlsbad, CA, United States) was used as substrate. 100 µL of assay mixture consisted of 5 µg DLPC (1,2-Dilauroyl-sn-glycero-3-phosphocholine), 3 mM MgCl₂ and 20 µM BOMCC in 100 mM Tris-HCl buffer (pH 7.4). 1 mM NADPH was added to start the reaction and fluorescence was measured on a Spectramax M2e plate reader (Molecular Devices, Sunnyvale, CA, United States) at an excitation wavelength of 415 nm and an emission wavelength of 460 nm for BOMCC.

**Cytochrome P450 CYP2C19 activity supported by WT and POR-R550W**

The activity of CYP2C19 supported by WT or POR-R550W was tested using the fluorogenic substrate EOMCC (Invitrogen Corp, Carlsbad, CA, United States). *In vitro*, CYP2C19 assays were performed using a reconstituted system consisting of WT/POR-R550W, CYP2C19 (CYPEX, Dundee, Scotland, United Kingdom) and cytochrome b5 at a ratio of 5:1:1 (48). It consisted of 2.5 µg DLPC and proteins (0.5 µM POR: 100 nM CYP2C19: 100 nM b5), 3 mM MgCl₂, 20 µM EOMCC in 100 µL of 100 mM Tris-HCl buffer (pH 7.4). The reaction was started by addition of 0.5 mM NADPH, and fluorescence was measured on a Spectramax M2e plate reader (Molecular Devices, Sunnyvale, CA, United States) at an excitation wavelength of 415 nm and an emission wavelength of 460 nm for EOMCC.

**Cytochrome P450 CYP3A4 Activity supported by WT and POR-R550W**

The activity of the major drug-metabolizing enzyme CYP3A4 supported by WT or POR-R550W was tested using the fluorogenic substrate, BOMCC, as described earlier (49). An *in-vitro* reconstituted system was used by mixing POR (WT or R550W), purified CYP3A4 (CYPEX, Dundee, Scotland, UK) and cytochrome b5 as described above at a ratio of 5:1:1 (48). The final assay mixture consisted of proteins (1 µM POR: 200 nM CYP3A4: 200 nM b5), 3 mM MgCl₂, 5 µg DLPC and 20 µM BOMCC in 100 µL of 100 mM Tris-HCl buffer (pH 7.4). 1 mM NADPH was added to start the reaction and progress of the reaction was monitored by fluorescence Spectrophotometer (Spectramax M2e plate reader; Molecular Devices, Sunnyvale, CA) with sample excitation at 415 nm and emission at 460 nm.

**Cytochrome P450 CYP3A5 activity supported by WT and POR-R550W**

The activity of CYP3A5 supported by WT or POR-R550W was tested using the *in-vitro* reconstituted system as described above. The purified CYP3A5 (CYPEX, Dundee, Scotland, United Kingdom), WT/mutant POR, and cytochrome b5 were mixed at a ratio of 1:5:1 (48). The assay mixture (100 µL) consisted of 5 µg DLPC and proteins (1 µM POR: 200 nM CYP3A5: 200 nM b5), 3 mM MgCl₂, 20 µM BOMCC in 100 mM Tris-HCl buffer (pH 7.4) and the reaction was started by addition of NADPH to 1 mM final concentration. Fluorescence was monitored for 1hr.
3D protein models

To study the potential impact of the mutation on structure, 3D models of POR-WT and POR-R550W were prepared using previously published model building protocols.

Statistical Analysis of results

Data are shown as mean, standard errors of the mean (SEM) in each group or replicates. Differences within the subsets of experiments were calculated using Student’s t-test. P values less than 0.05 were considered statistically significant.

Results

Identification of compound heterozygous variants in POR in CYP19A1 deficiency

Custom-designed targeted DSD NGS panel (DSDSeq.V1) revealed compound heterozygous variants in POR (NM_000941.2): c.73_74delCT/p.(L25Ffs*93) and c.1648C>T/p.(R550W) in the patient (46, XX karyotype). Sanger sequencing confirmed the segregation, the mother carried for the frameshift c.73_74delCT/p.(L25Ffs*93) and the father for the missense variant, c.1648C>T/p.(R550W) (Figure 1). A younger normal brother carried the c.1648C>T p.( R550W) variant. No further variants of interest were identified in the DSDSeq.V1 panel. At 8 years, the patient is prepubertal, and growth is normal (height 0.1 SDS and weight -0.41 SDS). No skeletal anomalies were detected, and the baseline hormonal analysis revealed normal DHEA-S, cortisol, T, and ACTH while 17OH-P was elevated (Table 1). Recommendations related to the stress situation needing glucocorticoid therapy were given to parents.

Both POR variants found in patient are observed as rare variants in genome databases

The POR variants identified in the patient in this report have been observed as rare variants in the genome databases (Table 2). The variant c.73_74delCT/p.(L25Ffs*93) (rs782696006) has a population minor allele frequency (MAF) of 0.00008125 (GnomAD), is classified as likely pathogenic following recommendations of ACGM, and due to truncation would generate a non-functional POR protein which would be unable to support the aromatase activity. We have reported a study of structural stability and sequence conservation analysis to find out which POR variants may be potentially disease-causing (50).

Human POR is a 680 amino acid protein that has evolved from ferredoxin and ferredoxin reductase-like domains to form a single redox protein in eukaryotes. The Arginine 550 residue studied here is highly conserved across species. The variant c.1648C>T/p.(R550W) (rs782551496) which has a MAF of 0.00004671 (GnomAD) was classified as a variant of uncertain significance (VUS) and predicted to be pathogenic with multiple pathogenicity prediction tools. Another variant at the same position, R550Q only had data available for the Global population. (Table 2).
Arginine 550 is located near NADPH binding site in POR and mutation R550W creates protein instability

To differentiate between the POR-WT and POR-R550W, we performed in-silico mutagenesis using the x-ray crystal structure of human POR. Human POR has distinct domains for the binding of NADPH/FAD and FMN (Figure 2A). The FMN binding domain interacts with the redox partners and is required for electron transfer to partner proteins. The redox equivalents for the electron transfer are provided by NADPH which is used as a substrate by POR and converted to NADP. The R550 residue is not directly at the surface of POR but is located near the NADPH binding pocket (Figure 2B). In the WT-POR, the arginine 550 residue forms hydrogen bonds with threonine 529 to stabilize the NADPH binding domain (Figure 2C). Mutation of arginine 550 to tryptophan results in loss of hydrogen bond interactions and destabilization of the POR protein. Binding of NADPH has also been associated in providing stability to POR and disruption of NADPH binding due to mutation of arginine 550 to tryptophan is predicted to cause protein instability.

The R550W mutation in POR results in reduced flavin content

To differentiate the conformational changes and effects of POR mutation R550W on flavin binding, we evaluated the relative flavin content since the activity of POR may be affected by the changes in the binding of cofactors FMN and FAD. As compared to WT POR, both the FMN and the FAD-binding was affected due to R550W mutation. As compared to WT, the FMN and FAD content of POR-R550W was 63.4% and 68.7% respectively. This suggests that POR-R550W affects both FMN as well as FAD binding to POR (Figure 3). Since the R550 residue is not directly involved in either the FAD or the FMN binding, loss of flavins suggest an overall instability of protein due to R550W mutation. It has been shown previously that binding of NADPH may affect the FAD/FMN binding in POR, and therefore, R550W mutation in POR may create a conformational change that is less favorable for the binding of FAD as well as FMN.

Small molecule Reduction Activity of POR is adversely affected by the R550W mutation.

To study the effect of the R550W mutation on POR activity, we expressed WT and POR-R550W in E. coli as N-23 form and purified bacterial membranes. The ability to transfer electrons from NADPH to cytochrome c, MTT or ferricyanide by POR-WT or POR-R550W was tested. The POR-R550W mutation had a much lower capacity to reduce cytochrome c, MTT, and ferricyanide (Table 3, Figure 4A-C). Compared to WT POR, the R550W variant lost ~60% cytochrome c reduction activity and ~70% of ferricyanide reduction activity. The MTT reduction activity (Table 3, Figure 4B) was also severely affected by the R550W mutation, with only 8% residual activity observed compared to WT-POR. The loss of activities with ferricyanide indicates disruption of electron transport from NADPH to FAD and loss of cytochrome c and MTT reduction activities indicates disruption of electron transfer.
between the two flavins and from FMN to the redox partners. Conformational instability due to an R550W mutation might be affecting domain movements and transfer of electrons from NADPH to FAD, FMN, and acceptor molecules. Since Ferricyanide reduction can proceed even without a functional FMN domain in POR, loss of Ferricyanide reduction activity could either be due to a disruption of electron transfer from NADPH to FAD or the loss of FAD-binding as reported above and a combination of these two factors. The cytochrome c and MTT reduction by POR requires a functional FMN domain in addition to the intact NADPH/FAD-binding domain. However, we have seen MTT reduction activity to be a better indicator of the effect on steroid metabolizing enzymes, and here the loss of MTT reduction activity was much severe compared to a loss of cytochrome c reduction activity.

CYP17A1- 17α-hydroxylase Activity

After the observation of lower activities of the R550W mutant of POR towards the reduction of small molecule substrates, we sought to examine the effect of arginine to tryptophan at the position 550 in POR on the activities of several steroid metabolizing cytochrome P450 enzymes that are dependent on POR for their enzymatic reactions. Towards this goal, we first examined the effect of POR-R550W on the 17-hydroxylase activity of CYP17A1, which is considered the qualitative regulator of steroidogenesis in human and guides the formation of steroids towards different pathways. The POR-R550W lost almost all 17α-hydroxylase activity of CYP17A1 (Figure 5A, Table 3). For the POR-R550W variant, the apparent Vmax was reduced by ~95% without affecting the apparent Km as compared to WT POR. The POR variant R550W showed only 3% residual activity in supporting CYP17A1 as compared to WT POR.

The 21-hydroxylase activity of CYP21A2 is severely affected by the R550W mutation in POR

The CYP21A2 is involved in the regulation of corticosteroids and is a critical enzyme in the steroid metabolism in human. Defects in CYP21A2 cause the standard form of congenital adrenal hyperplasia and mutation in POR may resemble CYP21A2 deficiency due to adverse effects of a malfunctioning POR on the activities of CYP21A2. The POR-R550W showed highly reduced CYP21A2 activity (Figure 5B, Table 3). The apparent Vmax of POR-R550W was reduced by ~98%. The POR variant R550W showed only 3 % residual activity in supporting CYP21A2 as compared to WT POR.

POR mutation R550W causes severe disruption of the aromatase activity of CYP19A1

Since the patient showed symptoms of aromatase deficiency as indicated by genital virilization, we were particularly interested in examining the role of R550W mutation in POR on the CYP19A1 activity. Compared to other cytochromes P450 studied in this report, CYP19A1 requires additional steps of electron transfer from POR to complete a single reaction cycle, and therefore, is predicted to be severely affected by changes in POR that affect electron transfer from POR to CYP19A1. The POR-R550W showed a severe effect on CYP19A1 activity (Figure 5C, Table 3). The apparent Vmax of POR-
R550W was reduced to 16% of WT-POR. A significant loss of CYP19A1 activity confirmed the diagnosis of aromatase deficiency and genital virilization in the patient.

**CYP2C9 Activity**

We tested the activity of CYP2C9 supported by the WT and R550W variant of POR in a reconstituted system. Compared to WT POR activity of CYP2C9 supported by R550W variant of POR was severely reduced to 2.5% of the WT-POR (Figure 6A). This almost complete loss of activity compared to WT POR indicated a severe effect on drug metabolism supported by CYP2C9.

**CYP2C19 Activity**

The activity of CYP2C19 was tested with both the WT and R550W variant of POR. We found that in CYP2C19 assays, the R550W variant of POR showed only 2.9% of the WT POR activity (Figure 6B). The effect of R550W variation in POR on CYP2C19 was also severe, indicating a highly reduced capacity of drug metabolism reactions by CYP2C19.

**CYP3A4 Activity**

The R550W variant of POR showed only 10.2% activity in CYP3A4 assays as compared to the WT-POR enzyme (Figure 6C). The loss of activities for drug-metabolizing cytochrome P450 enzyme-CYP3A4 by the POR variant R550W indicates problems with POR-P450 interactions which seem to be different for different cytochrome P450 partners.

**CYP3A5 Activity**

The R550W variant of POR had only 31.4% of the WT activity in CYP3A5 assay (Figure 6D). This was different from the results obtained for the CYP3A4 activity assays, indicating there are differences in the interaction of POR with these two closely related cytochrome P450 proteins. The effect of R550W variation in POR on CYP3A5 was not as strong as in case of CYP3A4 activity, but the loss of activity was still more than 50%, indicating a reduced capacity of drug metabolism reactions mediated by CYP3A5. CYP3A5 is involved in the metabolism of tacrolimus, an immunosuppressant drug used during organ transplants which has a narrow therapeutic index.

The R550W mutation in POR affects NADPH binding.

The structural analysis of the R550W mutation indicated an effect on NADPH. We sought to examine the impact of R550W mutation in POR on the utilization of NADPH using three different assays (Figure 7). For examining the role of NADPH concentration in reactions supported by POR, the substrate concentrations (Cytochrome c, MTT, and aromatase substrate androstenedione) were kept constant, and NADPH concentration was varied. In the cytochrome c reduction assay, a fivefold increase in Km for NADPH was observed for the POR-R550W compared to the WT-POR (Figure 7A).
A similar effect was also observed using MTT reduction assay with a 4.8-fold increase in Km for NADPH for the POR-R550W compared to WT POR (Figure 7B). We also performed the CYP19A1 assay with varying concentrations of NADPH and observed a severe loss of CYP19A1 activity for the POR-R550W (Figure 7C). In the CYP19A1 assay, a direct evaluation of NADPH oxidation could not be made as the final product was the metabolism of androstenedione into estrone, but the loss of protein-protein interaction coupled with the loss of flavins in POR-R550W would severely limit the activity of CYP19A1 due to multiple interactions required for completion of one catalytic cycle.

Discussion

In this report, we have investigated the cause of CYP19A1 deficiency manifested not by any changes in CYP19A1 itself but by a novel mutation in POR. A comprehensive examination of multiple cytochrome P450 enzymes encompassing both the steroid as well as drug-metabolizing activities was performed, which showed severe inhibition of multiple enzyme activities due to the R550W mutation in POR. Severe inhibition of CYP19A1 activity due to the R550W mutation in POR corroborates our previous hypothesis of both the virilization of the patient as well as maternal virilization during pregnancy due to the mutation in POR affecting CYP19A1 activity. The results described here also show that in severe cases of POR deficiency, the steroid as well as drug-metabolizing cytochrome P450 enzyme activities, that are dependent on POR, may be affected.

Remarkably, although the R550 residue is not directly involved in either the FAD or FMN binding in POR, a loss of one-third of flavin content, for both the FAD as well as FMN was observed. Previously we have shown that mutations in POR may cause protein instability which may ultimately lead to loss of flavin cofactors. A reduction in flavin content by itself does not explain the severe loss of activities of all cytochrome P450 enzymes studied. Consistent with this, some of the small molecule substrates of POR, that can be reduced through FAD (FeCN) or requiring FMN (MTT) were only moderately affected, indicating protein-protein interactions play a vital role in the overall effect of R550W mutation in POR. The CYP19A1, requiring several electron transfer steps for its metabolic reactions, is affected to a greater extent compared activities of POR towards small-molecule substrates. The overall effects of R550W mutation on steroid metabolism may be more complicated due to inhibition of several enzyme activities including CYP17A1 and CYP21A2, both of which catalyze critical steps in human steroidogenesis (51, 52).

The patient, an individual with 46, XX DSD, presented signs of prenatal aromatase deficiency as indicated from strong maternal virilization during pregnancy with highly elevated serum T levels at delivery that returned to normal, newborn ambiguous genitalia with elevated T for the female sex, and slightly elevated 17OH-P that progressively diminished. CYP19A1 gene analysis was normal, and it was not until seven years later that compound heterozygous POR gene variants were detected by using
a panel of candidate genes for DSD in DNA samples from DSD patients lacking a molecular diagnosis. Clinical and biochemical control at prepubertal age (8 years) revealed normal skeletal growth, normal baseline adrenal function (cortisol and ACTH), although with an elevated 17OH-P.

The allele c.73_74delCT/p.Leu25Phefs*93 is predicted to be a null allele resulting from a premature stop codon truncated at amino acid position 93 out of 680 residues in POR protein. Therefore, the protein encoded by the c.1648C>T/p.Arg550Trp allele would be responsible for the patient phenotype. Prenatal and neonatal phenotype predicted a severe aromatase deficiency, whereas the elevated 17-OH-P, although with normal cortisol and ACTH, may have indicated the presence of mild 21-hydroxylase and 17,20-lyase deficiencies. The mutation Arg550Trp is located in the NADPH binding region of POR. Computational analysis predicted instability in the NADPH binding region of POR by R550W mutation due to the disruption of hydrogen bonding, which may affect cytochrome P450 activities to a higher degree than small-molecule partners. Computationally predicted the negative effect on several cytochrome P450 activities as well as binding of NADPH were confirmed by experiments using recombinant proteins. These results suggest a pathological effect of POR R550W and a diagnosis of PORD in the patient with p.Arg550Trp/p.Leu25PhefsTer93 in POR. The 46, XX patient and her mother presented prenatal aromatase deficiency; at prepubertal age and without steroid hormone replacement therapy growth is normal and baseline adrenal function is normal (cortisol and ACTH) although the increased level of 17-OH-P demonstrates a mild 21-hydroxylase deficiency. It is predicted that aromatase deficiency will manifest at pubertal age, most probably needing sex hormone replacement therapy and interfering with future fertility while adrenal function will need to be monitored.

From the therapeutic perspective, supplementation with steroids is a viable and commonly used option to treat POR deficiency, especially when the loss of exact metabolites can be measured in serum or urine of patients by use of GC-MS or other methods (8, 53, 54). However, long term effect of multi steroid supplementation may have unknown side effects on normal growth and development, and therefore, an accurate characterization of individual variations in POR is required to characterize the effects of different steps in steroid metabolism and tailor the therapeutic interventions based on personalized steroid and enzyme activity profiles (55). A detailed characterization of the effects of POR mutation R550W on multiple enzymes described is a step towards this goal of personalized medicine.

Acknowledgments

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The datasets generated during and analyzed during the current study are not publicly available but are available from the corresponding author on reasonable request. Restrictions apply to the availability of data generated or analyzed during this study to preserve patient confidentiality or because they were used under license. The corresponding author will on request detail the restrictions and any conditions under which access to some data may be provided.
References


53. Shackleton C, Pozo OJ, Marcos J 2018 GC/MS in Recent Years Has Defined the Normal and Clinically Disordered Steroidome: Will It Soon Be Surpassed by LC/Tandem MS in This Role? Journal of the Endocrine Society 2:974-996


Table 1. Hormone analyses of the patient and her mother. ACTH: adrenocorticotropic hormone (pg/ml); DHEA-S: dehydroepiandrosterone sulphate (µg/dL); 17-OH-P: 17-hydroxy-progesterone (ng/dL); cortisol (µg/dL); T: testosterone (ng/dL); E2: estradiol (ng/dL).

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<th>ACTH</th>
<th>DHEA-S</th>
<th>17-OH-P</th>
<th>Cortisol</th>
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<td>(8 years)</td>
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<td>2.8-85.2</td>
<td>7-56</td>
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Table 2: Population distribution of POR variations reported in this study.

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<th>Sample size</th>
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<th>Alt Allele</th>
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<td>34162</td>
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Table 3: Kinetic parameters for activities of cytochrome c, MTT, ferricyanide reduction, and CYP17A1, CYP21A2 and CYP19A1 activities supported by POR-WT and POR-R550W. For the cytochrome c, MTT and Ferricyanide reduction assay, the NADPH concentration was fixed at 100 µM and varying concentrations of the substrate were used for the analysis. For the conversion of androstenedione to estrone (CYP19A1 activity), NADPH was fixed at 1 mM, and variable concentrations (10–1000 nM) of androstenedione were used. For the conversion of Progesterone to 17OH Progesterone (CYP17A1 activity) or Progesterone to 21OH Progesterone (CYP21A2 activity), NADPH was fixed at 2 mM, and variable concentrations (0.3–5 µM) of Progesterone were used. Data are shown as mean ± SEM of independent replicates (n=3 for small molecule reduction assays and n=2 for CYP assays).

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<th>Vmax (µmol/min/mg)</th>
<th>Km (µM)</th>
<th>Vmax/Km</th>
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<tr>
<td>POR-WT</td>
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<td>40.8</td>
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<td>2939±712</td>
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<th>Km (µM)</th>
<th>Vmax/Km</th>
<th>% WT</th>
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<td>POR-WT</td>
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<td>0.173</td>
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<td>POR-R550W</td>
<td>0.007±0.002</td>
<td>0.15±0.17</td>
<td>0.044</td>
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<td>CYP17A1; (Progesterone to 17OH-Progesterone)</td>
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<td>POR-WT</td>
<td>8.2±0.47</td>
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<td>POR-R550W</td>
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<td>CYP21A2; (Progesterone to 21OH-Progesterone)</td>
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Table 4: Effect of NADPH variation on activities supported by POR-WT and POR-R550W. To measure the effect of NADPH variation on activities of POR-WT and POR-R550W activities were measured using fixed concentrations of cytochrome c (40 µM), MTT (500 µM) and androstenedione (100 nM) while the NADPH concentrations in the assays were varied from 0 to 1000 µM. Data are shown as mean ± SEM of independent replicates (n=3).

<table>
<thead>
<tr>
<th></th>
<th>Vmax µmol/min/mg</th>
<th>Km (µM)</th>
<th>Vmax /Km</th>
<th>% WT</th>
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<td><strong>MTT reduction</strong></td>
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<tr>
<td>POR-WT</td>
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<td>81</td>
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<td><strong>CYP19A1 (Androstenedione to Estrone)</strong></td>
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Figure 1: Family tree and genetic analysis of POR sequences.
The DNA samples were analyzed with a custom-designed targeted Disorders of Sexual Development NGS panel (DSDSeq.V1, 111 genes, and three regulatory regions) using SeqCap EZ technology (Roche Nimblegen) and sequenced on a NextSeq (Illumina) platform. Sequence variations retained as candidates were verified by Sanger sequencing and DNA from parents were also analyzed as controls. (A) Compound heterozygous variants within the POR gene were identified in the proband (II-1): c.73_74delCT (p.Leu25Profs*93) located in exon 2, inherited from the mother (I-2), and a second variant c.1648C>T (p.Arg550Trp) in exon 13, inherited from the father (I-1). Parents are both carriers for each variant, and her brother (II-2) is a carrier for c.1648C>T (p.Arg550Trp). The arrow indicates the proband. N indicates normal allele. (B) Chromatogram of the variants in the POR gene (NM_000941) in this family. The arrows indicate the variant location.

Figure 2: Structural analysis of POR-R550W. (A). Location of R550W residue in POR. (B) R550 is not directly at the surface of POR. (C) Arginine 550 forms hydrogen bonds with threonine 529 to stabilize the NADPH binding domain. Its mutation to tryptophan results in destabilization of the POR structure.

Figure 3: Flavin content of POR-WT and POR-R550W. Flavin content of the POR proteins was analyzed by boiling the proteins under selective pH conditions. Relative fluorescence unit (RFU) of the flavins released from the POR variants measured at (A) pH 7.7 (FMN or F7.7) and (B) pH 2.6 (FAD or F2.6) are shown. The RFU of WT POR was fixed as a hundred percent. Data are shown as mean ± SEM of two experiments done in triplicates.

Figure 4: Cytochrome c, MTT, and Ferricyanide reduction assay with POR-WT and POR-R550W. (A) Cytochrome c (B) MTT (C) ferricyanide reduction assays were performed with the WT and POR-R550W variant. Kinetic assays were performed by monitoring the changes in absorbance at 550 nm for cytochrome c, 610 nm for MTT, and 420 nm for ferricyanide reduction. Data were fitted to the Michaelis-Menten kinetics model and analyzed using GraphPad Prism. The calculated Km and Vmax values are presented in Table 1. Data are shown as mean ± SEM of three independent replicates.

Figure 5: Enzymatic activities of steroid metabolizing cytochrome P450 supported by POR-WT or POR-R550W. A. Enzymatic activity of CYP17A1 supported by POR-WT and POR-R550W. Purified, recombinant CYP17A1 and enriched bacterial membranes containing POR proteins were reconstituted, and their activity to convert [14C] labeled Progesterone into 17OH-progesterone was tested by TLC and phosphorimager analysis. Data were analyzed using the Michaelis-Menten kinetics.
with GraphPad Prism. The calculated Km and Vmax values are shown in Table 3. B. Enzymatic activity of CYP21A2 supported by POR-WT and POR-R550W. The in-vitro reconstituted system was prepared by mixing purified CYP21A2, and bacterial membranes containing POR proteins and their activity to convert \([^{14}C]\) labeled Progesterone to 21OH-Progesterone was measured by TLC and phosphorimager analysis. Data were analyzed using the Michaelis-Menten kinetics with GraphPad Prism. The calculated Km and Vmax values are shown in Table 3. C. Enzymatic activity of CYP19A1 supported by POR-WT and POR-R550W. Bacterially expressed, purified, recombinant CYP19A1, and the enriched bacterial membranes containing POR proteins were mixed, and their activity to convert \([^{3}H]\) labeled androstenedione to estrone was tested by the tritiated water release assay. Data were analyzed using the Michaelis-Menten kinetics with GraphPad Prism. The calculated Km and Vmax values are shown in Table 3. Data are shown as mean ± SEM of two independent replicates.

**Figure 6: Activities of drug-metabolizing cytochromes P450 supported by POR-WT or POR-R550W.**

A. Activity of cytochrome P450 CYP2C9 supported by POR-WT and POR-R550W. Assay of CYP2C9 activity was performed to compare POR-WT and POR-R550W by using 20 µM BOMCC as a substrate. Activity with the WT POR was fixed as a hundred percent, and results are given as a percentage of WT activity. Data are shown as mean ± SEM from two experiments performed in triplicates. B. The activity of cytochrome P450 CYP2C19 supported by POR-WT and POR-R550W. Assay of CYP2C19 activity was performed to compare POR-WT and POR-R550W by using 20 µM EOMCC as a substrate. Activity with the WT POR was fixed as a hundred percent, and results are given as a percentage of WT activity. Data are shown as mean ± SEM from two experiments performed in triplicates. C. The activity of cytochrome P450 CYP3A4 supported by POR-WT and POR-R550W. Assay of CYP3A4 activity was performed to compare POR-WT and POR-R550W by using 20 µM BOMCC as a substrate. Activity with the WT POR was fixed as a hundred percent, and results are given as a percentage of WT activity. Data are shown as mean ± SEM from two experiments performed in triplicates. D. The activity of cytochrome P450 CYP3A5 supported by POR-WT and POR-R550W. Assay of CYP3A5 activity was performed to compare POR-WT and POR-R550W by using BOMCC as a substrate. Activity with the WT POR was fixed as a hundred percent, and results are given as a percentage of WT activity. Data are shown as mean ± SEM from two experiments performed in triplicates.

**Figure 7: Co-factor dependence of R550W mutation in POR.** Since the R550 residue is located near the NADPH binding site in POR, we sought to examine the role of NADPH variation and binding to R550W-POR. A. Assay of cytochrome c reduction activity with increasing NADPH concentration. In R550W-POR affinity for NADPH was diminished as indicated by a fivefold increase in KM for NADPH. B. Assay of MTT reduction activity with increasing NADPH concentration. Affinity for...
R550W-POR was also found to be lower as deduced from more than fourfold increase in Km for NADPH compared to WT-POR. C. CYP19A1 assay with increasing concentrations of NADPH. The POR-R550W showed a severe loss of activity in the CYP19A1 assay, which was more pronounced compared to fixed NADPH concentration assays shown in Figure 5C. In the CYP19A1 assays with varying NADPH concentration (Figure 7C), the activity using the POR-R550W was too low to allow a reliable estimation of kinetic parameters. Data are shown as mean ± SEM of three independent replicates.
Figure 1
Figure 2
Figure 3
Figure 4

A

POR-WT
POR-R550W

nmol/min/mg

Cytochrome c (µM)

0 10 20 30 40

B

POR-WT
POR-R550W

nmol/min/mg

Ferricyanide (µM)

0 100 200 300 400 500

C

POR-WT
POR-R550W

nmol/min/mg

MTT (µM)

0 100 200 300 400 500
Figure 5
Figure 6
Figure 7