Placenta Expressing the Greatest Quantity of Bisphenol A Receptor ERRγ among the Human Reproductive Tissues: Predominant Expression of Type-1 ERRγ Isoform

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Estrogen-related receptor γ (ERRγ), one of the 48 human nuclear receptors, has a fully active conformation with no ligand. We recently demonstrated that ERRγ binds strongly bisphenol A (BPA), one of the nastiest endocrine disruptors, and thus retaining ERRγ's high basal constitutive activity. A report that BPA accumulates in the human maternal-fetal placental unit has led us to hypothesize that a large amount of ERRγ might exist in the human placenta. Here we report evidence that placenta indeed expresses ERRγ exceptionally strongly. We first ascertained the presence of nine different ERRγ mRNA variants and the resulting three ERRγ protein isoforms. By real-time PCR, we estimated the relative amount of ERRγ mRNA using total RNA extracts from human reproductive tissues. Placenta was found to express ERRγ extremely highly. Among the three ERRγ protein isoforms, placenta exclusively expresses the type-1 isoform, which possesses additional 23-mer amino-acid residues at the N-terminus of the ordinary ERRγ. This N-terminal elongation was found to elevate by approximately 50% the basal constitutive activity of ERRγ as evidenced in the luciferase reporter gene assay. The present results suggest that BPA accumulates in the placenta by binding to ERRγ.

Key words: alternative splicing, bisphenol A receptor, estrogen-related receptor γ, placenta, real-time PCR.

INTRODUCTION

Bisphenol A (BPA), 2,2′-bis(4-hydroxyphenyl)-propane, is one of the highest volume chemicals produced worldwide as a starting material for polycarbonate plastics and epoxy resins. Long known as an estrogenic chemical, BPA is suspected of interacting with human estrogen receptor ER (1, 2) or acting as an antagonist for a human androgen receptor (AR) (3, 4). However, BPA’s binding to ER and AR and its hormonal activity are extremely weak: 1,000–10,000 times weaker than with natural hormones.

Based on the idea that BPA may interact with nuclear receptors (NRs) other than ER and AR, we screened a series of nuclear receptors and eventually explored estrogen-related receptor γ (ERRγ) as the BPA target receptor. BPA was found to bind strongly to ERRγ with high constitutive basal activity (5–7). BPA’s binding to ERRγ was further demonstrated by X-ray crystallographic analysis of the complex between BPA and ERRγ (8, 9).

In our efforts to explore the genuine characteristics of ERRγ as a BPA receptor, we have noticed the presence of several different ERRγ mRNA isoforms. NRs often possess a number of mRNA isoforms produced by alternative splicing to exhibit functions in a tissue-specific or developmental stage-specific manner (10, 11). However, little is known about the in vivo physiological functions of those splicing variants, and even the variants’ tissue distributions are poorly understood.

BPA as an endocrine disruptor poses the worrisome threat of low-dose effects on reproductive and developmental processes in humans (12). To ensure the presence of ERRγ mRNA isoforms in human reproductive organs and brains, we attempted to quantify the total amount of ERRγ mRNAs and then the amount of each mRNA isoform. Here we report evidence that the human placenta expresses ERRγ mRNA extremely highly, and that the class of isoforms is type-1 ERRγ.

MATERIALS AND METHODS

cDNA Cloning—To confirm the presence of eight reported isoforms of ERRγ mRNA, we cloned cDNA by...
using human pancreas and skeletal muscle. These total RNA samples (Clontech, Mountain View, CA, USA) were reverse-transcribed by using the forward primer of ERRγ-R1 (5′-GAAGAGCTGCATCTGTCGGTCTG-3′) and the enzyme SuperScriptII™ RNase H− Reverse Transcriptase (Invitrogen, Carlsbad, CA, USA) at 42°C.

To confirm that all clones had inconsistent 5′-UTR sequences, the forward primers were designed separately, depending on the unique structure of each exon (Table 1). As reverse primers, ChERRγR1 and ChERRγR2 were used in the first and the nested PCRs, respectively. As for the amplification of ERRγ-1 cDNA, the first PCR was carried out using a primer set of ChERRγ1F/ChERRγR1 and the enzyme Pfu Turbo® Hotstart DNA Polymerase (Stratagene, La Jolla, CA, USA). The second PCR was performed by using PLATINUM® Taq DNA polymerase (Invitrogen) with another primer set of ChERRγ1F/ChERRγR2 and the product from the first PCR as a template. For amplification of all other ERRγ, PCR was carried out by using the same method. Sequence analysis was carried out on CEG8800 Genetic Analysis System (Beckman Coulter, Fullerton, CA, USA).

Real-Time PCR—The total RNA samples extracted from brains (adult and fetal) and various different reproductative tissues (ovary, uterus, placenta, prostate and testis) were purchased from Clontech, Stratagene and Biochain (Hayward, CA, USA). Each total RNA sample (1μg) was reverse-transcribed by using SuperScriptII™ (Invitrogen) and oligonucleotide ERRγ-R2 and the enzyme SuperScriptII™ RNase H− Reverse Transcriptase (Invitrogen, Carlsbad, CA, USA). The total RNA samples extracted from placenta and kidney lysates were purchased from ProSci (Poway, CA, USA). These lysates (20μg each) was electrophoresed on 10% polyacrylamide gels. After electrophoresis, gels were electro-blotted onto Hybond-P (GE Healthcare, Waukesha, WI, USA). The blot was incubated overnight in the presence of the anti-ERR γ monoclonal antibody. ERRγ protein was visualized by chemi-luminescence (GE Healthcare) using HRP-conjugated secondary antibody (Jackson ImmunoResearch, West Grove, PA, USA). To discriminate a positive band from negative ones, negative staining controls were performed without the first antibodies.

Table 1. The oligonucleotide sequences of primers used for cDNA cloning of a series of ERRγ mRNA isoforms.

<table>
<thead>
<tr>
<th>Name of primers</th>
<th>Oligonucleotide sequences</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primers for amplification of ERRγ-1 cDNA</td>
<td>ChERRγ-R1F 5′-CGTTGGTCTCTTCAGAAGATTTGG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R1a 5′-GAAGAGCTGCATCTGTCGGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R2b 5′-TTTCCACAGTAGAAGATGGGAAG-3′</td>
</tr>
<tr>
<td>Primers for amplification of ERRγ-2 cDNA</td>
<td>ChERRγ-R2F 5′-GAAGAGCTGCATCTGTCGGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R2-adF1 5′-CGTTGGTCTCTTCAGAAGATTTGG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R2-adF2 5′-GCTCACCTCCATTTACCTGTCGGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R1a 5′-GAAGAGCTGCATCTGTCGGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R2b 5′-TTTCCACAGTAGAAGATGGGAAG-3′</td>
</tr>
<tr>
<td>Primers for amplification of ERRγ-2-gig cDNA</td>
<td>ChERRγ-2-gigF1 5′-CCACACACCTTCCATGATTTGG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-2-gigF2 5′-GCTCACCTCCATTTACCTGTCGGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R1a 5′-GAAGAGCTGCATCTGTCGGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R2b 5′-TTTCCACAGTAGAAGATGGGAAG-3′</td>
</tr>
<tr>
<td>Primers for amplification of ERRγ-3 cDNA</td>
<td>ChERRγ-3F1 5′-GCGGTCCTCCATTTAGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-3F2 5′-GCTCACCTCCATTTACCTGTCGGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R1a 5′-GAAGAGCTGCATCTGTCGGTCTG-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-R2b 5′-TTTCCACAGTAGAAGATGGGAAG-3′</td>
</tr>
<tr>
<td>Primers for insertion/deletion confirmation of exon K</td>
<td>ChERRγ-F 5′-CAAGATCTGATCTCTGACACA-3′</td>
</tr>
<tr>
<td></td>
<td>ChERRγ-LR 5′-CAAGATCTGATCTCTGACACA-3′</td>
</tr>
</tbody>
</table>

aThe antisense reverse primer R1 has the same nucleotide sequence.
bThe antisense reverse primer R2 has the same nucleotide sequence.

Real-time PCR was performed on a capillary-type LightCycler™ rapid thermal cycler system (Roche Diagnostics, Mannheim, Germany). Reactions were completed in a 10μl solution mixture and SYBR Green Realtime PCR Master Mix (Toyobo, Tokyo). For normalization, the mRNA gene (gpdh) of the enzyme glycerolaldehyde-phosphate dehydrogenase was amplified as an internal standard. The assay includes the steps of denaturation at 95°C for 1 min, annealing at 61°C for 3 s and extension at 72°C for a variable time, depending upon the size of products. The product specificity was always confirmed by agarose gel electrophoresis and routinely estimated by the melting curve analysis. To depict the standard curves for quantitative real-time PCR, a 10−1-fold series of dilutions of each plasmid with the same DNA sequence was simultaneously amplified. Quantification of mRNA was achieved using LightCycler software (version 3.5). Standard curves had a correlation coefficient (r2) of 1.00, linear over a sample concentration range, and mean square error values of 0.03–0.08 were involved.

Western Blotting Analyses—Western blotting was used to detect ERRγ protein isoforms from human kidney and placenta. ERRγ-specific mouse monoclonal antibody was purchased from Perseus Proteomics (Tokyo). The human placenta and kidney lysates were purchased from ProSci (Poway, CA, USA). These lysates (20μg each) were electrophoresed on 10% polyacrylamide gels. After electrophoresis, gels were electro-blotted onto Hybond-P (GE Healthcare, Waukesha, WI, USA). The blot was incubated overnight in the presence of the anti-ERR γ monoclonal antibody. ERRγ protein was visualized by chemi-luminescence (GE Healthcare) using HRP-conjugated secondary antibody (Jackson ImmunoResearch, West Grove, PA, USA). To discriminate a positive band from negative ones, negative staining controls were performed without the first antibodies.

Reporter Gene Assay for ERRγ Types 1 and 2—Type I and type II ERR γ fragments were cloned into the vector pCDNA3.1(+) (Invitrogen). As an ERR response element (ERRE)-luciferase construct, 3× ERRγ/pGL3 was used as described previously (7). HeLa cells were maintained in Eagle’s MEM medium (Nussui, Tokyo) with 10% (v/v) fetal bovine serum at 37°C. HeLa cells were transfected with 3μg of luciferase reporter gene (pGL3/3× ERRE), 1μg of the expression plasmid of the wild-type of either type I or type II ERR γ and 10ng pSEAP-control plasmid as an internal control by Lipofectamine Plus reagent (15μl/ml, Invitrogen). Approximately 24 h after transfection, cells were harvested and plated into 96-well plates at a concentration of 5×104 cells/well. The cells were then treated with varying doses of chemicals, BPA (Tokyo Kasei Kogyo, Tokyo) and 4-OHT (Sigma-Aldrich, St. Louis, MO, USA), diluted with 1% BSA/PBS (v/v). After 24h, luciferase activity was measured by using the Luciferase assay reagent (Promega, Madison, WI). SEAP activity was assayed by using Great EscA PeTM SEAP assay reagent (Clontech) according to the Fluorescent SEAP Assay protocol. Light emission was measured on a microplate reader Wallac 1420 ARVOx (Perkin Elmer, Turku, Finland). Cells treated with 1% BSA/PBS were

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used as a vehicle control. Each assay was performed in duplicate and repeated at least three times.

RESULTS

Confirmation and Classification of Alternative Splicings of ERRγ Gene—To date, three independent investigations have revealed six alternative splicing sites for the human ERRγ mRNA gene and eight different ERRγ mRNA variants (13-15). However, there is no systematic study to unify these results, and thus it is unclear whether or not all of these variants are present simultaneously in one species, such as humans. Thus, we first attempted to confirm the full-length sequences of cDNAs derived from the eight ERRγ mRNA splicing variants.

To amplify each ERRγ mRNA isoform—ERRγ1, ERRγ2, ERRγ2-gig and ERRγ3—PCR was conducted successfully by using a series of forward sense and 5′ reverse antisense primer sets was designed for each mRNA isoform (Table 1) and commercially available pancreas and skeletal muscle cDNAs. As a result, seven of the eight splice variants reported were definitely identified, and skeletal muscle cDNAs. As a result, seven of the eight splice variants reported were definitely identified, but we could not identify ERRγ3 in this study. Although we carefully searched many other human tissues, the mRNA corresponding to ERRγ3 was not detected. Instead, we identified another novel variant ERRγ2-bcd (accession number AB362218). It should be noted that these variants have variable nucleotide sequences of the 5′-UTR, with no structural changes of ERRγ2 protein.

We now know nine splicing variants in total. It should be noted, however, that these variants afford or produce three distinctly different protein isoforms (Fig. 1), as the variants are classified into three mRNA isoforms: ERRγ1, ERRγ2 and ERRγ3. Here, type-2 mRNA ERRγ2 consists of seven subclasses of splicing variants: ERRγ2-df, ERRγ2-def, ERRγ2-di, ERRγ2-d, ERRγ2-ad, ERRγ2-bcd and ERRγ2-gig (Fig. 1C), although all of these variants produce the same receptor protein molecule of ERRγ2. The variants are constituted from 15 distinct exons, A–O, coded in the human genomic DNA in the very broad region of chromosome 1 (about 1,000 kbp) (Fig. 1B). The heterogeneity at the 5′-UTR is due to the presence of alternative transcription start sites and alternative splicing sites.

The type-1 protein isoform ERRγ1 has an additional 23-mer elongated N-terminal sequence. Type-2 mRNA genes, ERRγ2 isoforms, include six variants containing the exon D-based fragment (designated d) in the 5′UTR. ERRγ2-ad, where ad indicates that the exons a and d are involved in this order, was first isolated from the human fetal brain library by Eudy et al. (13). On the other hand, ERRγ2-gig has been found only in the skeletal muscle cDNA library (14). The mRNA ERRγ3-bcf gene producing ERRγ3 has recently been reported by Kojo et al. (15), although we could not identify this gene in the present study. The ERRγ3 protein isoform has a deletion of 39-mer amino-acid residues in the DNA-binding domain of ERRγ2, resulting in an incomplete construction of the DNA binding site. In addition, it was found that ERRγ mRNAs each have two alternative polyadenylation isoforms (13).

Quantitative Analysis of ERRγ mRNA Genes as a Whole by Real-Time PCR—By means of real-time PCR, the total expression amount of the human ERRγ mRNA genes was estimated to amplify the region common to all the splicing variants. We designated the hERRγ-whole mRNA segment. We did confirm that there is no contamination of the genomic DNA in these RNA samples, since we could not amplify any cDNA when we used the primer sets directly for the samples.

To analyse the hERRγ-whole mRNA gene by real-time PCR, the primer set of sense hERRγ-wholeF and antisense hERRγ-wholeR was utilized (Table 2). For the quantification of each ERRγ splicing variant, real-time PCR was carried out by using a series of primer sets.

Fig. 1. Structural constitution of ERRγ mRNA isoforms and ERRγ protein isoforms. (A) Structure of ERRγ protein isoforms. ERRγ1 has 23-mer amino-acid extension at the N-terminus of ordinary ERRγ, namely ERRγ2. ERRγ3 has a 39-mer amino-acid deletion in the DNA binding domain. (B) Structural constitution of exon and intron of human ERRγ genomic gene. Alphabetic letters A–O each indicate an independent exon. (C) Structural constitution of exons in nine ERRγ mRNA isoforms. Closed arrowheads indicate the position of an AUG initiation codon, and open arrowheads indicate the position of a termination codon in the ERRγ open reading frame.
whole mRNA and respective mRNA isoforms, with the cDNAs prepared from each total RNA sample of exon region. Both sense and antisense primers in the independent ERR (Fig. 2, Table 2). These primers were designed not to amplify the ERRγ sequence on the genomic DNA, setting both sense and antisense primers in the independent exon region.

Real-time PCR was carried out at least three times for the cDNAs prepared from each total RNA sample of human adult kidney, placenta, ovary, uterus, prostate, testis and brain, as well as from the human fetal brain. We determined the number of molecules of the whole mRNA and respective mRNA isoforms, with 1 × 10^5 molecules of gapdh mRNA being the internal standard. For example, the molecular number of the hERRγ-whole mRNA of adult brain was 942 per 1 × 10^5 gapdh mRNA (Table 3). The fetal brain had 739 molecules (about 80% of the molecular number of adult brain).

In kidney, hERRγ-whole mRNA had a molecular number of 4,369, approximately 4.6-fold that of the adult brain (Table 3). Among the reproductive organ tissues, the placenta was found to express the highest number of hERRγ-whole mRNA molecule, 4,544, ~5% greater than that of kidney and thus about 5-fold that of the adult brain. This very high expression of hERRγ mRNA genes in the placenta was of course found to exceed those of other reproductive organs: 151 in ovary, 167 in uterus and 571 in testis (Fig. 3). The second highest amount was in the prostate (1,637) (Table 3). Thus, the amount in the placenta was approximately 3-fold greater than that in the prostate. These results suggest that ERRγ plays a very significant role in the placental functions. This may indicate adversely that the placenta is potentially the most affected by BPA.

**Table 2. The nucleotide sequences of the primers used for the quantification of whole ERRγ mRNA and each ERRγ mRNA isoform.**

<table>
<thead>
<tr>
<th>Name of primers</th>
<th>Oligonucleotide sequences</th>
<th>Length of products (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primers for quantification of ERRγ-whole mRNA</td>
<td>hERRγ-wholeF 5'-CAGAATTCACAAAGATCTGACAC-3', hERRγ-wholeR 5'-GCTTTGAACGTGACACGTGC-3'</td>
<td>148</td>
</tr>
<tr>
<td>Primers for quantification of ERRγ1 mRNA</td>
<td>hERRγ1F 5'-GCACATGATCTGAGAGCTGTTGACCG-3', hERRγ1R 5'-GCTTTGAACGTGACACGTGC-3'</td>
<td>215</td>
</tr>
<tr>
<td>Primers for quantification of ERRγ2 mRNA</td>
<td>hERRγ2F 5'-TGTTTGAACGTGACACGTGC-3', hERRγ2R 5'-GCTTTGAACGTGACACGTGC-3'</td>
<td>183, 300, 338, 359</td>
</tr>
<tr>
<td>Primers for quantification of ERRγ2-gig mRNA</td>
<td>hERRγ2-gigF 5'-GCCACCATCAGACTGTTGACCGGAGACG-3', hERRγ2-gigR 5'-GCTTTGAACGTGACACGTGC-3'</td>
<td>339</td>
</tr>
<tr>
<td>Primers for quantification of ERRγ2-bcd mRNA</td>
<td>hERRγ2-bcdF 5'-GTGTTTGAACGTGACACGTGC-3', hERRγ2-bcdR 5'-GCTTTGAACGTGACACGTGC-3'</td>
<td>208</td>
</tr>
<tr>
<td>Primers for quantification of gapdh mRNA</td>
<td>hgapdhF 5'-CAGAATTCACAAAGATCTGACAC-3', hgapdhR 5'-GCTTTGAACGTGACACGTGC-3'</td>
<td>107</td>
</tr>
</tbody>
</table>

*These antisense primers have the same nucleotide sequence. The number of products depends on the number of alternative splicing sites in the particular region amplified.*
Table 3. The results of real-time PCR quantification of ERRγ-whole mRNA and its subtypes in human tissues.

<table>
<thead>
<tr>
<th>Tissues</th>
<th>ERRγ-whole mRNA</th>
<th>ERRγ mRNA isoforms (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Type-1</td>
<td>Type-2</td>
</tr>
<tr>
<td></td>
<td>d-series   gig</td>
<td>bcd&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>Brain (adult)</td>
<td>942 ± 24</td>
<td>73.5 ± 9.8</td>
</tr>
<tr>
<td>Brain (fetal)</td>
<td>739 ± 195</td>
<td>11.1 ± 2.8</td>
</tr>
<tr>
<td>Kidney</td>
<td>4369 ± 1276</td>
<td>94.7 ± 1.2</td>
</tr>
<tr>
<td>Pancreas</td>
<td>2742 ± 798</td>
<td>7.1 ± 1.0</td>
</tr>
<tr>
<td>Skeletal muscle</td>
<td>247 ± 48</td>
<td>798 ± 4.0</td>
</tr>
<tr>
<td>Placenta</td>
<td>4544 ± 1572</td>
<td>45.2 ± 18.3</td>
</tr>
<tr>
<td>Prostate</td>
<td>1637 ± 217</td>
<td>98.9 ± 0.7</td>
</tr>
<tr>
<td>Testis</td>
<td>571 ± 93</td>
<td>72.6 ± 8.1</td>
</tr>
<tr>
<td>Ovary</td>
<td>151 ± 50</td>
<td>93.1 ± 3.7</td>
</tr>
<tr>
<td>Uterus</td>
<td>167 ± 127</td>
<td>4.3 ± 2.3</td>
</tr>
</tbody>
</table>

aThe amount of mRNA was calculated as the number of molecules per 1.0 × 10<sup>5</sup> gapdh mRNA molecules. bThe analysis with specific sense and antisense primers, both of which were set in the same exon C (Fig. 2) was originally designed to measure the total amount of c-containing ERR mRNAs including ERRγ-2-bcd and ERRγ-3-bcf. Since the amount of ERRγ-3-bcf that gives ERRγ-3 was negligible, the measurement gave the amount of only ERRγ-2-bcd that affords ERRγ2.

Fig. 3. Quantitative real-time PCR for estimation of ERRγ mRNA expression in human brains and reproductive organs. The gapdh mRNA gene was used as an internal control. The copy number per 1.0 × 10<sup>5</sup> gapdh mRNA was estimated for ERRγ-whole mRNA in each tissue. The error bars indicate SEM.

Information such as age and sex is given for samples provided from each agent.

It should be noted that in the placenta, the ERRγ1 mRNA isoform is accounted for 98.9% of the quantity of all mRNA isoforms (Table 3). This predominance of the type-1 mRNA isoform is very surprising and unique among the human tissues. All other human tissues express ERRγ2 as a major mRNA isoform. It is likely that ERRγ1 produced from the ERRγ1 mRNA isoform may play a crucial and central role in the placenta.

Reproductive tissues other than placenta expressed the ERRγ2 mRNA isoform much more than ERRγ1. The content of ERRγ2 mRNA isoform was estimated as 79.6% in prostate, and 81.9% and 95.7% in ovary and uterus, respectively (Table 3). Since the expression level of ERRγ2-gig mRNA was almost negligible in these tissues (Table 4), estimated ERRγ2 mRNA isoform includes exclusively six subclasses of splicing variants (d-series that contain: ERRγ2-df, ERRγ2-def, ERRγ2-di, ERRγ2-d, ERRγ2-ad and ERRγ2-bcd) (Fig. 1C). As for the testis, however, a prominent inconsistency was found. It contained about 11% of the ERRγ2-gig mRNA isoform. Furthermore, a discrepancy between the sum of all mRNA isoforms and the ERRγ-whole mRNA—the sum of total numbers of ERRγ mRNA isoform molecules was clearly smaller (by about 10%) than the total molecular number of ERRγ-whole mRNA (data not shown)—strongly suggested the presence of mRNA isoform(s) other than those measured.

The adult and fetal brains were found to have almost the same isoform constitutions (Table 3). The expression ratios of ERRγ1 and ERRγ2 mRNAs respectively were 30.2% and 69.8% in the fetal brain and 26.1% and 73.9% in the adult brain. Thus, brain is the tissue in which type-2 ERRγ mRNA is expressed predominantly. In kidney, the expression levels of ERRγ1 and ERRγ2 mRNAs were estimated to be approximately 11.1% and 88.9%, respectively. Pancreas was also one of the tissues in which type-2 ERRγ mRNA was expressed predominantly, 4.0% ERRγ1 and 96.0% ERRγ2. The sum of ERRγ1 and ERRγ2 mRNAs occupied approximately 100% in these tissues.

The constitutional ratio of ERRγ1, ERRγ2(d-series) and ERRγ2-gig mRNAs in skeletal muscle was unique (Table 3). Their expression ratios were 45.2%, 7.1% and 47.7%, respectively. Skeletal muscle was found to be the tissue in which type-2 ERRγ2-gig mRNA is expressed very highly.
In the estimation of the tissue distributions of $ERR_{\gamma}$ mRNA splicing variants, another important issue is to calculate the actual amount of each mRNA isoform. As shown in Fig. 4, $ERR_{\gamma}1$ mRNA isoform was most abundant in placenta, the amount of this isoform (4,544 (98.9% as in Table 3 = 4,494 molecules) being exclusive and predominant. The kidney is the tissue in which the $ERR_{\gamma}2$ mRNA isoform is richest (3,884). Pancreas (2,632) and prostate (1,306) are also abounding in $ERR_{\gamma}2$. As for the $ERR_{\gamma}2-gig$ mRNA isoform, skeletal muscle (118) and testis (63) contain it relatively highly.

Western Blotting Detection of $ERR_{\gamma}$ Protein Isoforms—
To verify the amount of type-1 $ERR_{\gamma}$ protein isoform expressed in the placenta, Western blotting was carried out for the lysates of the human placenta and kidney. Kidney was selected as a reference because of its expected high expression (90%) of the type-2 $ERR_{\gamma}$ protein isoform (see above). $ERR_{\gamma}$ protein isoforms were detected by using a monoclonal antibody specific for the N-terminal region (1–100) of $ERR_{\gamma}2$, because this antibody can detect all three isoforms: $ERR_{\gamma}1$, $ERR_{\gamma}2$ and $ERR_{\gamma}3$. The calculated molecular weight (51,313) of type-1 $ERR_{\gamma}$ is larger than that of type-2 standard $ERR_{\gamma}$ by ~2,700, apparently due to the N-terminal addition of 23-mer amino-acid residues.

The lysate isolated from kidney exhibited an intense protein band of 49 kDa, corresponding to the molecular weight of type-2 standard $ERR_{\gamma}$. This band accompanied a faint band of 51 kDa, which corresponds to the molecular weight of type-1 $ERR_{\gamma}$ (Fig. 5). When we examined the lysate isolated from the placenta, the band of 51 kDa was markedly detected, as shown in Fig. 5. This result very clearly proves the consequences observed for the mRNA isoforms. It is evident that the mRNA variants expressed indeed produce a consequent protein and that the placenta expresses the type-1 $ERR_{\gamma}$ protein isoform predominantly and exclusively.

Transcription Activity of Type-1 and Type-2 $ERR_{\gamma}$ Isoforms in the Reporter Gene Assay—$ERR_{\gamma}$ is a constitutively active nuclear receptor that exhibits a high basal activity with no ligand. In the present study, we examined the reporter gene activity of type-1 and type-2 $ERR_{\gamma}$ isoforms by means of the luciferase reporter gene assay using HeLa cells. To normalize for transfection efficiency, we simultaneously carried out the SEAP assay (16). When the type-1 $ERR_{\gamma}$ isoform was compared with the type-2 standard $ERR_{\gamma}$ isoform, the constitutive activity level of type-1 $ERR_{\gamma}$ was found to be about 50% higher than that of type-2 $ERR_{\gamma}$. As shown in Fig. 6, the type-2 $ERR_{\gamma}$ isoform exhibited significantly elevated constitutive activity (210% of the basal activity). Type-1 $ERR_{\gamma}$ isoform also exhibited considerably elevated constitutive activity (260%), a notably higher level than that of type-2. The results indicate that the N-terminal

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Fig. 4. Quantitative real-time PCR for estimation of $ERR_{\gamma}$ mRNA expression and percentage of the constitutions of $ERR_{\gamma}$ mRNA isoforms in human brains (A), standard tissues (kidney, pancreas and skeletal muscle) (B) and reproductive organs (C). The circled area shows the total expression amount of $ERR_{\gamma}$ mRNA in each tissue. The area of each color region in the circle is proportional to the expression amount of the $ERR_{\gamma}$ mRNA isoforms. Blue, purple, yellow and red areas indicate $ERR_{\gamma}1$, $ERR_{\gamma}2(d\text{-}series)$, $ERR_{\gamma}2\text{\text{-}gig}$ and $ERR_{\gamma}2\text{\text{-}bcd}$ mRNA isoforms, respectively. The expression rates of each $ERR_{\gamma}$ mRNA isoform were calculated against the sum of the copy numbers of $ERR_{\gamma}$ mRNA isoforms.
23-mer elongation has a distinct effect on the reporter gene transactivation activity of ERR. 4-OHT deactivated the ordinary standard type-2 ERR as reported (7, 17), diminishing the basal activity of ERR by up to 60–80% at a concentration of 10 μM (Fig. 7A). These were exactly revealed for the type-1 ERR isoform, ERR1. BPA, on the other hand, showed no effect on the basal constitutive activity of ERR even at a concentration of 10 μM, completely preserving ERR’s high constitutive activity (Fig. 7A). The inverse agonist activity of 4-OHT for ERR1 was reversed or inhibited by BPA in a dose-dependent manner (Fig 7B).

This reversing activity of BPA, namely, inverse antagonist activity of BPA, was revealed originally for the type-2 ERR isoform (7).

DISCUSSION

Extremely High Expression of ERR mRNA in the Placenta—In the present study, using commercially available human gene samples of reproductive organ
For accuracy in the quantification of hERRγ-whole mRNA, we repeated the real-time PCR. By using a new set of sense and antisense primers set at the 3’ terminal region, the results eventually obtained were almost the same as those of the first quantification shown in Fig. 3 (data not shown). For further confirmation, we tested internal controls other than gapdh mRNA. Those include the mRNA genes of human β-actin, ubiquitin C, sdha (succinate dehydrogenase complex, subunit A) and hprt1 (hypoxanthine phosphoribosyl-transferase 1). The results for the amount of ERRγ mRNA were almost the same as those obtained by the quantification using gapdh mRNA (data not shown). These further evidenced that the ERRγ mRNA expression in the human placenta is extreme, and the highest among the tissues examined.

Other reproductive organ tissues, such as ovary, uterus and testis, also express ERRγ mRNA, but at very low levels: 3.3%, 3.7% and 12.6% that of the placenta, respectively. Compared to these tissues, the considerably high expression of ERRγ mRNA in the prostate should be noted. The prostate had the second highest amount of ERRγ mRNA, approximately 36% of that in the placenta.

**Predominant Expression of Type-1 Isoform of ERRγ mRNA in the Placenta**—Nuclear receptors usually have several mRNA and protein isoforms by alternative splicing mechanisms, resulting in the exhibition of their functions in a tissue-specific or developmental stage-specific manner (10, 11). Unfortunately, there is little understanding not only of the physiological functions of splicing variants, especially in vivo, but also of their tissue distributions in the majority of tissues throughout the body.

All transcript ERRγ mRNA variants consist of several distinct exons coded on human genomic DNA in the very broad region of chromosome 1 (about 1,000 kbp). As shown in Fig. 1B, the exons are thought to be distinguished between those in a variable region (A~1) and those in a consistent region (Δ~0). The latter includes almost all of the open reading frames of ERRγ, while 1~3 exons are selected from the variable region to form a 5’-UTR. Including alternative polyadenylation mechanisms, the gene expression of ERRγ appears to be severely regulated in a post-transcriptional manner. The sequence difference of the 5’-UTR should bring about a different translational efficiency, depending on the stability of mRNA produced and the presence of some upstream open reading frames (uORF) (18).

Type-1 ERRγ isoform (ERRγ1) has an additional 23-mer amino-acid residue extension at the N-terminus, and exhibits about 50% increased basal constitutive activity relative to that of ERRγ2 (Fig. 6). Although this isoform possesses exactly the same structure of its ligand-binding domain as other isoform types, its activation mechanism may differ from those of other isoforms by having this 23-mer N-terminal elongation. This might bring about unique tissue-specific function(s) in the placenta.

**High Concentration of BPA in the Placenta due to High Expression of Type-1 ERRγ**—BPA is an industrial chemical, and exposure to it is now widespread (19, 20). We know now that BPA binds to ERRγ about 100 times greater than to ERα and ERβ, and that BPA exhibits a distinct inhibition activity against 4-OHT in ERRγ.
Chief Expression of Bisphenol A Receptor ERRγ in Placenta

In the present study, we did know that there are three isoforms possessing precisely the same ligand-binding domain (LBD). It should be noted that BPA can bind to all these ERRγ isoforms in any tissue.

This strongly suggests that the concentration of BPA in human tissues is directly proportional to the amount of ERRγ proteins. In our estimation in the present study involving reproductive organ tissues, the expression of ERRγ mRNA is largest in the placenta and second largest in the prostate. The BPA concentration in the placenta has been reported to be approximately five times higher than that in maternal and fetal plasma (21). It is now quite reasonable to believe that the high concentration of BPA is due to the large amount of ERRγ protein isoforms, almost entirely type 1, in the human placenta.

The Effects of BPA Accumulation or Binding to Type-1 ERRγ on Placental Functions—The placenta receives nutrients, oxygen, antibodies and hormones from the mother’s blood and removes waste. It forms the placental barrier, which filters out some substances that could harm the fetus. However, some substances, including BPA and viruses, are not filtered out, suggesting that the placenta does not act as a barrier against BPA (21, 22). In addition to transferring gases and nutrients, the placenta also has metabolic and endocrine activity. It produces estrogen, relaxin, and human chorionic gonadotrophin, progesterone and somatomammotropin, all of which are important in maintaining pregnancy and the large amounts of glucose and lipids in the maternal blood. It is now evident that the placenta expresses BPA receptor ERRγ very highly. What would happen with the accumulation of BPA in the placenta?

Placenta in BPA-administered mice during pregnancy was reported to be abnormal (23), directly decreasing the number of embryos. In addition, almost all mouse neonates exposed to BPA were dead within 3 days after birth. Thus, BPA might disrupt the placental functions directly or indirectly, and might affect the mortality of neonates through indirect exposure of embryos. These are likely mediated through the BPA receptor ERRγ2, at least in part. BPA administration has also been reported to significantly increase the weight of the uterus and the number and fertilization quality of sperm (24). DNA microarray analysis has shown that BPA administration increases the mRNAs of some nuclear receptors in mouse placenta (25). These results suggest that BPA affects the transcriptional regulation in the placenta or other reproductive organs through certain particular transcription factors.

Based on the fact that BPA strongly binds to ERRγ, the abnormality and probable change of gene expression in the placenta are likely accompanied by BPA binding to ERRγ. In this study, we demonstrated that the ERRγ1 mRNA gene expresses almost fully in the placenta, and that the resulting type-1 ERRγ receptor is noticeably more potent than the resulting type-2 ERRγ receptor that expresses dominantly in other tissues. BPA would sustain unnecessarily this very high basal constitutial activity of type-1 ERRγ receptor in the placenta.

Other Human Tissues with High Expression of ERRγ mRNA—Adult and fetal brains, kidney and pancreas were the tissues in which ERRγ expresses significantly and considerably highly. In these tissues, type-2 ERRγ mRNA is expressed predominantly, while the expression levels of ERRγ3 is almost negligible. The expression ratios of ERRγ1 and ERRγ2 mRNAs respectively varied 4–30% and 69–95%. As compared to the type-2 ERRγ isoform (ERRγ2), type-1 isoform ERRγ1 has an additional 23-mer amino-acid elongation at the N-terminus. Although ERRγ1 exhibits about 50% increased basal constitutive activity relative to that of ERRγ2 (see above), physiological functions in these tissues have never been clarified nor analysed for both ERRγ1 and ERRγ2.

As for the prostate, the mRNA gene was for the most part (~80%) ERRγ2. Various effects of estrogenic chemicals including BPA have been reported for the prostate. For example, acceleration in the proliferation rate of prostate epithelium during fetal life was noted to disrupt permanently the cellular control systems and to predispose the prostate to disease in adulthood (26, 27). The effects of estrogens on the prostate, or the effects of their involvement in prostate cancer development and benign prostatic hyperplasia, are likely mediated through their ERs. In addition to the androgen receptor, which plays a central role in the normal development and neoplastic growth of the prostate gland, estrogens have long been suggested to play synergistic or distinct roles in the same processes. However, studies from ER knockout mouse models have shown neither ERRα nor ERRβ affects the targeted disruption of prostatic phenotype and function (28). This strongly suggests the involvement of one or more nuclear receptors other than ER and that ERRγ is a probable candidate for involvement in prostatic growth and development. ERRγ might play regulatory roles in normal and neoplastic prostatic cells by sharing similar ER-mediated pathways or acting independently.

CONCLUSION

The present study provides a valuable blueprint of ERRγ mRNA expression and important clues to understanding BPA’s low-dose effects in humans. For instance, although the issue of bioavailability of parent BPA in humans has been contentious, the present results strongly suggest that the BPA concentration is proportional to the expression amount of ERRγ. There are scientific debates over whether or not low doses of BPA can have developmental or reproductive effects in humans. Now, it is clear that ERRγ, the receptor of BPA, abounds in brains and reproductive tissues such as the placenta and prostate.

Strong expression of ERRγ as a possible receptor of BPA in both the placenta and the fetal brain could have important implications for newborns. ERRγ is also a probable candidate for involvement in prostatic growth and development. However, the physiological roles of ERRγ are poorly understood at the moment. It is thus important to clarify such physiological functions and characteristics of ERRγ. Moreover, it is crucial to examine the content and extent of which BPA may influence these roles.
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CONFLICT OF INTEREST
None declared.

REFERENCES