Zonal expression of the thyroid hormone receptor α isoforms in rodent liver

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Abstract

Many metabolic processes occur simultaneously in the liver in different locations along the porto-central axis of the liver units. These processes are often regulated by hormones, one of which is thyroid hormone which for its action depends on the presence of the different isoforms of the thyroid hormone receptor (TR). These are encoded by two genes: c-erbA-α encoding TRα1 and TRα2 and their respective Δ isoforms, and c-erbA-β which encodes TRβ1, TRβ2 and TRβ3. We recently found a zonal (pericentral) expression of and a diurnal variation in the TRβ1 isoform in rat liver. We were therefore also interested to see whether TRα1 and TRα2 expression showed similar characteristics. For this reason we raised both polyclonal and monoclonal antibodies against TRα1 and TRα2 isoforms and characterised these. Antibody specificity was tested using Western blots and immuno-
histochemistry in liver of TR isoform-specific knockout animals. Using these antibodies we found that the TRα1 and TRα2 isoforms are zonally expressed around the central vein in rat liver. The experiments show that the portal to central gradient of TRα1 is broader than that of TRβ1. Moreover, the expression of the TRα2 protein showed a diurnal variation with a peak in the afternoon when the animals are least active whereas no such variation was found for the TRα1 protein.

From our data it appears that both the TRα1 and TRα2 isoforms show a zonal distribution in liver. This finding, together with the observed diurnal rhythm, has major implications for interpreting and timing experiments concerning the TR and its downstream actions in liver. (Journal of Endocrinology (2003) 179, 379–385

Introduction

In the liver many metabolic processes occur simultaneously, but not all of them take place in every liver parenchymal cell. Instead, the various metabolic reactions occur in different locations along the porto-central axis of the liver units (Klinger et al. 1988, Jungermann & Katz 1989). For instance, lipolysis and glycolysis are predominant in liver cells around the central vein, whereas lipogenesis and gluconeogenesis are found in the area around the portal vein. This so-called metabolic zonation can be of a stable or dynamic kind which means that the expression of certain enzymes is either restricted to certain cells regardless of the metabolic or hormonal state, or the expression of the enzymes can expand or shrink along the porto-central axis depending on the metabolic state or time of the day. Enzymes with a stable distribution can be located pericentrically (like glutamine synthetase) or peripherally (like fructose 1,6-bisphosphatase (Eilers et al. 1995)). Similarly, enzymes with a dynamic distribution can be located pericentromitally (ornithine aminotransferase (Swick et al. 1970)) or peripherally (phosphoenolpyruvate carboxy-kinase, PEPCK (Bartels et al. 1990)).

Thyroid hormone has many diverse actions in the body. These actions depend on the presence of the different isoforms of the thyroid hormone receptor (TR) (Yen 2001). Thyroid hormone receptors belong to the nuclear receptor superfamily and act mainly as transcription factors. They are encoded by two separate genes: c-erbA-α which encodes TRα1 and TRα2 and their respective Δ isoforms, and c-erbA-β which encodes TRβ1, TRβ2 and TRβ3 isoforms. Thyroid hormone (triiodothyronine, T₃) signals its presence to the cell by binding to these TRs which then interact with so-called thyroid hormone response elements in the promoter of T₃-responsive genes and thereby activate or repress genes. TRs are differentially distributed in various tissues and during developmental stages, indicating distinct or specific functional
roles. TRα1 and TRα2 are mainly present in the central nervous system and muscle whereas TRβ1 is predominantly present in the liver. Studies using knockout mice have shown that certain actions of thyroid hormone are dependent on a particular receptor isoform (Amma et al. 2001) and the expression patterns of some of these genes overlap with that of the TRβ1 isof orm they depend on (H Gulberg, B Zandieh-Doulabi, D Forrest, O Bakker, B Vennström, unpublished observations).

We recently found a zonal (pericentral) expression of and a diurnal variation in the TRβ1 isof orm in rat liver (Zandieh Doulabi et al. 2002). Therefore, we were interested to see whether TRα1 and TRα2 expression showed similar characteristics. For this reason we raised antibodies against the TRα1 and TRα2 isoforms. We generated polyclonal and monoclonal antibodies against the TRα isoforms and characterised these. We tested antibody specificity using Western blots and immunohistochemistry in liver of TR isof orm-specific knockout animals.

Materials and Methods

Wistar rats (Harlan Sprague Dawley, Zeist, The Netherlands) were kept in a 12 h light/12 h darkness cycle with free access to food. All animal experiments were approved by our local Animal Welfare Committee. Lights were switched on at 0700 h. The liver of each animal was used for immunohistochemistry and Western blots.

Polyclonal antibodies

To produce anti-TR antisera, two linear synthetic peptides coupled to keyhole limpet haemocyanin (KLH) were made, namely amino acids 402–410 (NH2-EVFEDQEV-COOH) for human TRα1, and amino acids 425–442 (NH2-SLRGPVQLHQSPKPQQR-COOH) for TRα2. Using these peptides antisera were raised in New Zealand White rabbits and affinity purified by Eurogentec (Seraing, Belgium).

Monoclonal antibodies

The same KLH-conjugated linear synthetic peptides TRα1 and TRα2 described above were used for immunisations in mice with a total of four injections of 25 µg peptide each. The first injection (subcutaneously in complete Freund’s adjuvant) was seven weeks before death, the second and third injections (intraperitoneally in complete Freund’s adjuvant) were respectively two and four weeks after the first injection and finally the last injection was given intravenously. Five days after the last boost of immunisation, the spleen cells from the immunised BALB/C mice were fused with myeloma (SP20) by means of polyethylene glycol 1500 (Roche Molecular Biochemicals–Germany). The fused cells were plated on 24-well plates in hypoxanthine aminopterin thymidine (HAT)-DMEM/15% FCS/1% PenStrep (Sigma) in the presence of 5% hybridoma cloning factor (HCF; Sanvertech, Breda, The Netherlands). Fourteen days later the supernatants of the hybridomas were screened by immunohistochemistry. Positive clones were subcloned in 96-well plates in HAT medium by limiting dilution. The supernatants from the positive subcloned hybridomas were used for further screening on the liver of a rat killed at 1930 h and in nuclear or whole cell extracts on Western blots.

The selected clones were grown in bulk. The supernatant of each monoclonal antibody was precipitated with half the starting volume of saturated ammonium sulphate (Merck, Darmstadt, Germany), which was added dropwise and was then left stirring overnight at 4 °C. The next morning after centrifugation at 3000 g for 30 min, 20 ml ammonium sulphate were added at 4 °C for 6 h. After centrifugation for 30 min at 3000 g, the pellet was saved and dissolved in 0·1 volume (4 ml) of PBS. The concentrated antibody was dialysed overnight at 4 °C in a dialysis tube (cut-off 12 000 M.W., Thomas, USA).

A mouse monoclonal antibody isotyping kit (Hycult Biotechnology, Uden, The Netherlands) was used to establish the isotype of the monoclonal antibodies.

Western blots

Whole cell extracts (WCE) from livers were prepared by homogenising in 0·25 M sucrose containing complete protease inhibitor (Roche Molecular Biochemicals) using a homogeniser for 10 s at maximum speed. Protein concentration of the WCE was determined using Biorad reagent (Biorad, Germany). Subsequently, 25 µg protein were loaded on 10% SDS-PAGE gel. After electroblotting onto membrane (Protran BA45, Schleicher & Schuell, Dassel, Germany), the blots were blocked for 45 min in Tris-buffered saline (TBS) containing 1% (w:v) caseine (Roche, Germany) and 0·1% (w:v) Tween 20 (blocking buffer). Next, the blots were incubated for 2 h at 22 °C with the primary antisera (polyclonal antisera, 1:250 dilution and monoclonal antisera, 1:50 dilution). After 30 min incubation with secondary antibodies (1:20 000 dilution) in blocking buffer, the blots were washed and Lumilight² substrate (Roche Molecular Biochemicals) was added. The signals were visualised and quantified using a LumiImager (Roche Molecular Biochemicals).

Nuclei were isolated as previously described (Wiersinga et al. 1982). In short, 4 g liver were homogenised in Sol A (20 mM Tris.HCl, 0·25 M sucrose, 2 mM CaCl2, 1 mM MgCl2, and 5% (v/v) glycerol, pH = 7·6). The nuclei were pelleted by ultracentrifugation (45 000 g for 45 min at 4 °C) and washed with Sol A containing 0·5% Triton X-100 and resuspended in Sol B (200 mM Tris, 5% (v/v) glycerol, 0·25 M sucrose, 1 mM EDTA and 50 mM NaCl) and stored in liquid nitrogen. The amount of...
protein in this nuclear preparation was determined using the Biorad protein assay. For Western blots 50 µg protein equivalent of nuclei were used.

**Immunohistochemistry**

Both paraffin-embedded and frozen livers of several rats were used as tissue for immunohistochemistry. The frozen rat liver sections of 6 µm were fixed with different kinds of fixatives depending on the type of experiments: paraformaldehyde (PFA; Merck, Germany) 4% (w:v) in phosphate-buffered saline (PBS); pH=7-4), acetone, methanol or ethanol for 15 min at 4 °C. Paraaffin-embedded rat livers were fixed for 16 h at 4 °C in PFA sections (6 µm) and pasted on Superfrost/Plus (Menzel-Gläser, Germany) slides and dewaxed prior to immunostaining.

Sections were placed in TBS (pH=8·0) and microwaved for 10 min for the paraffin and 5 min for the frozen sections to unmask the antigenic epitope. After cooling to room temperature, the sections were placed in a blocking solution for 60 min (5% low fat milk (Campina, The Netherlands) and a detergent, either 0·1% (w:v) Triton X-100 in TBS at pH=8·0). After a washing period of 10 min in TBS, the sections were placed in alkaline phosphatase buffer (50 mM MgCl₂, 100 mM NaCl, and 100 mM Tris, pH=9·2) for 10 min. The first antibodies were diluted in the blocking solution, and incubated first for 1 h at room temperature and then overnight at 4 °C. The sections were washed with 0·5% (w:v) Triton X-100 in TBS (pH=8·0) for 10 min, and then incubated with the second antibody (conjugated with alkaline phosphatase) diluted with 0·5% (w:v) saponin (Sigma) diluted in TBS at pH=8·0). The first antibodies were diluted in the blocking solution for 60 min (5% low fat milk (Campina, The Netherlands) and a detergent, either 0·1% (w:v) Triton X-100 or 0·5% (w:v) saponin (Sigma) diluted in TBS pH=8·0). The antibody (Fig. 1B).

These blots were incubated with the polyclonal antibodies against TRα1 and TRα2 to check specificity. Position of the protein size markers is indicated alongside. (B) Western blots were prepared with whole cell extracts from wild-type (wt), TRα1−/−/β1−/− (α1−/β1−) and TRα2−/− (α2−) mice and Vaccinia-expressed TRα1 (α1Vac). These blots were incubated with the polyclonal antibodies against TRα1 and TRα2 to check staining with antiserum preadsorbed with antigenic epitope. After cooling to room temperature, the sections were placed in a blocking solution, and incubated first for 1 h at room temperature and then overnight at 4 °C. The sections were washed with 0·5% (w:v) Triton X-100 in TBS (pH=8·0) for 10 min, and then incubated with the second antibody (conjugated with alkaline phosphatase) diluted with 0·5% (w:v) saponin (Sigma) diluted in TBS at pH=8·0). The first antibodies were diluted in the blocking solution for 60 min (5% low fat milk (Campina, The Netherlands) and a detergent, either 0·1% (w:v) Triton X-100 or 0·5% (w:v) saponin (Sigma) diluted in TBS pH=8·0). The antibody (Fig. 1B).

**Results**

**Specificity of antibodies on Western blots**

**Polyclonal TRα1 and TRα2 antibodies** The polyclonal antibodies were typed as IgG type antibodies. In liver WCE and nuclear extracts a 47 kDa band was detected using the TRα1 polyclonal antiserum and a 58 kDa band was detected using the TRα2 polyclonal antiserum (Fig. 1A). As a control for the TRα1 polyclonal antiserum we used a Vaccinia-expressed TRα1. We also tested both our polyclonal antibodies on Western blots of liver WCE of animals where either the TRα1 or TRα2 was knocked out (Wikstrom et al. 1998, Ng et al. 2001). The TRα1 polyclonal antibody showed no signal at 47 kDa in liver WCE from a TRα1−/−/β1−/− (α1−/β1−) animal and the TRα2 polyclonal antibody did not show the 58 kDa band which was seen in the wild-type animals (Fig. 1B).

Furthermore no cross-reaction with TRα1 protein is found, since Vaccinia-TRα1 was not detected with this antibody (Fig. 1B).

The specificity of the antisera was further supported by negative pre-immune serum staining and by negative staining with antisera preadsorbed with antigenic...
peptide. Isoform specificity was also supported by the absence of cross-reactivity as examined by immunocytochemical staining of the TR peptides TR\(\alpha1\) (amino acids (aa) 402–410), TR\(\alpha2\) (aa 425–442), TR\(\beta1\) (aa 74–92) and TR\(\beta2\) (aa 131–145) fixed on nitrocellulose membrane.

**Monoclonal TR\(\alpha1\) and TR\(\alpha2\) antibodies** The monoclonal antibody clones were typed as IgM. The specificity of the monoclonal antibodies was tested on liver slices derived from mice lacking either TR\(\alpha1\) (TR\(\alpha1^{-/-}\)) or TR\(\alpha2\) (TR\(\alpha2^{-/-}/\beta1^{-/-}\)). As can be seen in Fig. 2 no staining was observed with the TR\(\alpha1\) monoclonal antibody in a TR\(\alpha1^{-/-}\) liver. This was not a general loss of TR since the TR\(\beta1\) monoclonal antibody was still able to detect the TR\(\beta1\) protein in the pericentral cells in the TR\(\alpha1^{-/-}\) animals (Fig. 2). TR\(\alpha2\) staining was absent in the TR\(\alpha2^{-/-}/\beta1^{-/-}\) animals. In these mice TR\(\alpha1\) was detected in a control experiment and found to be broader than in the wild-type mice. On Western blots the monoclonal TR\(\alpha1\) antibodies detected bands of similar size (47 kDa) to those detected with the polyclonal antibody but the signal was stronger in the nuclear preparation. With the TR\(\alpha2\) monoclonal antibody, a strong band of about 58 kDa was detected in the nuclear preparation at the same position as the bands found with the polyclonal TR\(\alpha2\) antibody.

As with the polyclonal antibodies, isoform specificity was also supported by the absence of cross-reactivity as examined by immunocytochemical staining of the fixed TR peptides TR\(\alpha1\), TR\(\alpha2\), TR\(\beta1\) and TR\(\beta2\) on nitrocellulose membrane.

**Immunohistochemistry**

In order to study the distribution of the TR\(\alpha1\) and TR\(\alpha2\) proteins in liver we set out to determine the optimal incubation conditions for our antibodies. The parameters of interest are microwave treatment before incubation with the first antibody, salt (NaCl) concentration and pH during the first antibody incubation. When performing immunohistochemistry on both frozen and paraffin embedded sections of the rat liver we found that microwave treatment diminished background and intensified staining (probably by unmasking the antigenic epitope) with both polyclonal and monoclonal antibodies on sections fixed with PFA. We therefore used PFA as fixative and microwave treatment in all immunohistochemistry experiments using both polyclonal and monoclonal antibodies.

To find the optimum salt concentration during incubation with the first antibody at pH 8 we performed experiments with both frozen and paraffin embedded sections by increasing the salt concentration from 0 mM to 600 mM NaCl. Increasing the NaCl concentration up to 400 mM had no effect on the staining in the case of both polyclonal and monoclonal TR\(\alpha2\). Surprisingly, the TR\(\alpha1\) monoclonal antibodies were very sensitive to the NaCl concentration. Since all different antibodies stained well at 150 mM NaCl this concentration was chosen in further experiments. Using this 150 mM NaCl concentration we next determined the pH optimum during incubation with the first antibody. For the TR\(\alpha1\) and TR\(\alpha2\) polyclonal antibodies and the TR\(\alpha2\) monoclonal antibody the pH optimum was around pH 8. However, for the monoclonal TR\(\alpha1\) antibody the pH optimum was around pH 5 (Fig. 3).

Since the monoclonal TR\(\alpha1\) antibody had a lower pH optimum we determined optimum salt concentration for this antibody at this pH and we found again that the antibody was very sensitive to the NaCl concentration. Moreover, at this pH background was reduced using saponin in the incubation mixture.

Taking these results together we decided to use the monoclonal antibodies in the experiments aimed at studying the distribution and diurnal rhythm of the TR\(\alpha1\) isoforms. In these experiments the TR\(\alpha1\) monoclonal antibodies were diluted 1:50 in blocking buffer, 0.5% (w/v) saponin, pH 5 with no additional salt and the TR\(\alpha2\)

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**Figure 2** Specificity of the TR\(\alpha1\) and TR\(\alpha2\) antibodies tested with immunohistochemistry. (Left panel) Slices from livers of wild-type (wt) or TR\(\alpha1^{-/-}\) mice were incubated with the monoclonal TR\(\alpha1\) antibody with the TR\(\beta1\) polyclonal as a control. (Right panel) The monoclonal TR\(\alpha2\) antibody was incubated with slices from livers of wild-type (wt) and TR\(\alpha2^{-/-}/\beta1^{-/-}\) mice using the TR\(\alpha1\) polyclonal antibody as a control. Magnification × 40.
monoclonal antibodies were diluted 1:50 in blocking buffer pH 8 with 150 mM NaCl.

Zonal distribution

Immunohistochemistry using the monoclonal antibodies on liver sections indicated that TRα1 and TRα2 were expressed around the central veins (Fig. 4). This expression pattern overlaps with that of the TRβ1 (Zandieh-Doulabi et al. 2002) and with the centrally expressed, T₃-responsive gene glutamine synthetase (GS), which we stained as a control on consecutive sections. However, the area of expression of the TRα1 and TRα2 proteins extends further along the porto-central axis than that of TRβ1 and GS.

Diurnal variation of TRα isoforms in rat liver

When rats were killed at different time points during the day we found different staining intensities for the TRαs in liver slices. Especially, TRα2 expression differed when the rats were killed in the morning compared with those killed in the afternoon. These results were confirmed using Western blotting where it can be seen that no change in expression level of TRα1 is found (Fig. 5A) but that there is a marked diurnal variation in the expression of the TRα2 protein (Fig. 5B) with a peak during the afternoon when the animals are least active. Immunohistochemistry showed that the expression of both TRα1 and TRα2 was of a stable kind.

Discussion

Here we report the generation of a set of antibodies, both polyclonal and monoclonal, directed against the thyroid hormone receptor α1 and α2 isoforms which are specific for these isoforms according to the criteria used by us (Western blots with wild-type and specific knockout WCE and staining on slices of wild-type and specific knockout livers). Since the monoclonal antibodies gave a better signal on the Western blots we performed our studies on the zonal distribution and the diurnal variation of the TRα1 and α2 isoforms with the monoclonal antibodies.

Several immunohistochemical studies have reported the presence of TR proteins in the liver of rodents and humans (Tagami et al. 1990, Strait et al. 1991, Macchia et al. 1992, Rodd et al. 1992, Falcone et al. 1994). Although not all hepatocytes were stained in these studies, none of these studies mention a topographical distribution of TR isoforms even though most known T₃-dependent genes are expressed in the liver in different zones. We showed recently that the TRβ1 protein was expressed in a rather small area around the central vein of rat liver and that its expression varies during the day (Zandieh-Doulabi et al. 2002). Moreover, certain TRβ1-dependent genes, for
example the low density lipoprotein-receptor or CYP7a (Gullberg et al. 2002) are expressed in the same area. Here we show that a similar stable zonal expression exists for two other TR isoforms, TRα1 and TRα2. Although these two isoforms are also expressed around the central vein, the extent of the zonal expression for both TRα isoforms is wider than that of the TRβ1. Interestingly, the TRα1 expression gets even broader in the TRα2/−/−/TRβ1/−/− mice probably as a result of the overexpression of TRα1 in these mice (Ng et al. 2001).

Our findings suggest the possibility that there may be genes solely dependent on TRα1 or TRβ1 whereas there will be another set which is specifically activated by one of the isoforms depending on its localisation along the portal-central axis of the liver unit. For instance, CYP7a is expressed mainly in the pericentral zone in conjunction with all receptors (Berkowitz et al. 1995, Brasil et al. 1995, Massimi et al. 1998) while a gene encoding PEPCK, for example, is expressed towards the periportal zone where the TRαs are present. Therefore, based on the localisation of PEPCK it can be suggested that TRα1 is the most likely candidate to confer T3 regulation on this gene.

From our results it is apparent that the expression of TRα2 shows a diurnal rhythm whereas that of TRα1 does not. This hints at the possibility of regulated splicing, a phenomenon known to occur in other systems (Nissim-Rafinia & Kerem 2002, Perrone-Bizzozero & Bolognani 2002). The diurnal expression of the TRβ1 protein is the reverse of that of the TRα2 protein, with maximum TRβ1 protein expression observed at the beginning of the dark period when the rats are nutritionally active.

Whether the rhythm is dependent on the suprachiasmatic clock or on food intake is at present unknown but the presence of a rhythm in the serum levels of thyroid hormones and in the expression of the TR isoforms together with the knowledge that thyroid hormone regulates many genes (especially metabolic ones) in liver (Feng et al. 2000) points to the fact that thyroid hormone may play a role in mediating the clock signal originating in the suprachiasmatic nucleus.

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Figure 5 Diurnal variation of the TRα1 and TRα2 isoform proteins. The figure shows the signal intensities (expressed relative to an internal standard) derived from Western blots of whole cell extracts of rat liver incubated with monoclonal TRα1 (A) or monoclonal TRα2 (B). Rats were killed (6 per time point) at the time points indicated in the figure. Lights were on from 0700 h to 1900 h. Data are expressed as means ± S.D. and significant differences (Mann-Whitney U test), when present, are indicated.
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