Detection of an oestrogen receptor-like protein in human meningiomas by band shift assay using a synthetic oestrogen responsive element (ERE)

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Summary When a ligand binding or enzyme immunoassay is used, meningiomas are found to be rich in progestin receptors (PR) whereas oestrogen receptors (ER) are virtually undetectable. A protein that can bind to an oestrogen responsive element (ERE) was detected in meningiomas, by using the band-shift assay. The binding of ER to the ERE is inhibited by the anti-ER monoclonal antibody ER-P31, which is directed to the A/B domain of the ER, indicating that the binding protein is an ER-like protein.

Female sex hormones may be involved in the etiology of human meningioma. Following the first report in 1979, on the putative presence of oestrogen receptors (ER) in human meningiomas (Donnell et al., 1979), several research groups, including our own (Blankenstein et al., 1983; Martuza et al., 1985) have tried to confirm these findings. Using Scatchard plot analysis (Blaauw et al., 1986) or enzyme immunoassay (Blankenstein et al., 1987) the majority of meningiomas were found to be ER negative. The same experiments revealed that meningiomas contain high amounts of progestin receptors (PR). Since in classical female sex steroid target tissues like breast and uterus, the synthesis of PR is ER dependent the finding that (at the protein level at least) meningiomas are rich in PR, yet virtually devoid of ER initiated further research.

We recently reported on the detection of mRNA coding for the wild type receptor by using the reversed-transcriptase-PCR technique (Koehorst et al., 1992). Apart from the wild type (wt) mRNA we also found two alternatively spliced mRNAs. One variant is missing exon 4 and the other is missing exon 7 (Koehorst et al., 1993).

The present study was designed to evaluate the presence in meningiomas of ER or ER-like proteins which could induce PR synthesis. We hypothesised that if the ER (at concentrations below the detection limit of the conventional receptor assay) or ER-like protein (escaping detection by lack of ligand binding) is responsible for the induction of PR synthesis in meningiomas at least a protein should be present which can bind to the oestrogen responsive element (ERE) of the PR-gene. The aim of the present investigation, therefore, was to test if such an ERE-binding protein is present in meningioma. To this end the binding of meningioma cytosol components to a synthetic ERE was investigated with the band shift assay. This assay is a very sensitive method to detect ERE binding proteins. Scott et al. (1991) could detect as little as 0.1 fmol of DNA bound ER from whole cell extracts containing 50 μg protein.

We found evidence for the existence of a protein that binds to the ERE and is recognised by the specific anti-ER antibody ER P31.

Materials and methods

Tissues

Human meningioma tissue was placed on ice immediately after removal from the patient. Representative specimens were frozen at -80°C until they were used for cytosol preparation or receptor assay. The MCF7 breast cancer cell line was routinely grown with 10% foetal calf serum (Gibco BRL, Paisley, UK) with RPMI 1640 (Gibco BRL, Paisley, UK). Cells were harvested by scraping with a rubber policeman in PBS.

Biochemical assay of steroid hormone binding

The ER and PR content of the tumours were measured as described previously (Blankenstein et al., 1983) by the ligand binding assay according to the guidelines of the EORTC, Breast Cancer Cooperative Group (EORTC, Breast Cancer Cooperative Group, 1980).

Cytosol preparation for ERE-binding experiment

Approximately 400 mg of fresh tissue powder was added to 1.5 ml of ice cold extraction buffer (0.4 M KCl, 10 mM Tris (pH 7.4), 1 mM EDTA, 10 mM monothioglycerol, 10% glycerol (v/v)) or 20 x 10^6 MCF7 cells were dounced in 1.5 ml ice cold extraction buffer. The resulting homogenate was centrifuged for 5 min at 4°C at 3,600 r.p.m. The supernatant was centrifuged again in a swing out rotor at 132,000 g for 10 min at 4°C. Twenty-five μl of the clear supernatant was used for protein determination by the Coomassie Brilliant Blue method (Biorad, Richmond, CA, USA).

Band shift assay

As probe for the band shift assay, we chose the ERE from the vitellogenin A2 gene which has been shown to have high regulatory potential in transient transfection of ERE-TKCAT reporter construction into oestrogen responsive MCF7 cells (Klein-Hitpass et al., 1986). Both strands corresponding to the ERE were synthesised as 31 (5'-GATCCGTACGGTACAGCTGATGGATC-3'; palindrome is underlined) base oligonucleotides using an Applied Biosystems DNA synthesiser (San Jose, CA, USA). Equimolar amounts of the two strands were annealed in buffer (10 mM Tris, 1 mM EDTA (pH 8.0)) by heating to 95°C and cooling to room temperature during a period of 2 h. The double stranded ERE oligomer was end-labelled using [β-32P] ATP (Amersham, Amersham, UK) and T4 polynucleotide kinase (Boehringer Mannheim, Mannheim, Germany). Five μl of the supernatant containing 20–30 μg protein (for MCF7 containing 2 μg of protein) were added to 15 μl incubation buffer (100 mM Tris (pH 8.0), 1 mM EDTA, 10% glycerol (v/v), 2 μg poly(dI-dC)) and placed on ice.

15 min 1 ng 32P-labelled double-stranded ERE oligomer was added and the reaction mixture incubated for 20 min at 20°C. The protein-ERE complexes were separated by electrophoresis through 6% acrylamide (19:1, acrylamide/bis) gels using a buffer consisting of 50 mM Tris, 50 mM boric acid, 1 mM EDTA (pH 8.0). The gels were run at 180 V for 3 h, vacuum dried and autoradiographed. The ER-containing human breast cancer cell line MCF7 was used as a positive

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control for ERE binding.
To confirm the integrity of the protein isolated from the tumours, a band shift assay was performed using a oligonucleotide which contained the Sp1 binding consensus sequence (5'-CAAAAGTCTGGCCGCGGATCAAG-3'; consensus is underlined). For the competition binding analysis in which the specificity of the ERE was proven, a oligonucleotide was used which contained the consensus of the progesterone responsive element (PRE)(5'-CCAAA-GTCAGAACACGTGTCGATCAAG-3'; palindrome is underlined and differences with palindrome of ERE are in bold underlined).

Results

The ERE from the vitellogenin A2 gene which was used in the band shift assay has proven specificity in oestrogen binding experiments (Klein-Hitpass et al., 1988; 1989; Klock et al., 1987).

The specificity of binding of the ERE oligonucleotides to ER in crude, high salt extracts from MCF7 and meningioma is shown in Figure 1. The intensity of the ER-ERE complex decreases with increasing ratios of unlabelled/labelled ERE, while the intensity of the ER-ERE complex is unaffected with increasing molar excess of PRE (Figure 1a and b). When MCF7 cytosol was run together with the meningioma sample a single, prominent band co-migrating with that obtained from MCF7 whole cell extract was observed. We attribute this result to the presence of ER or an ER-like protein in the meningioma tissue.

To identify the ER in the ER-ERE complex, cytosols were incubated with the anti-ER monoclonal antibody ER-P31 (MEDAC, Hamburg, Germany). This antibody inhibits the binding to the ERE of the ER from MCF7 cells as well as the ER-like protein in the two meningiomas and the ER in solid breast tumours (Figure 2). After we had proven the specificity of this ERE oligonucleotide under our experimental conditions, we tested 15 meningiomas with different ER and PR concentration. In all meningiomas but one, including the meningiomas which were ER negative in the ligand binding assay, we could detect a protein which can bind to ERE oligonucleotide (Figure 3, upper panel). The ERE-protein complex from the meningiomas co-migrates with the MCF7 ER-ERE complex even in gels run for longer times (data not shown). To check the integrity of the isolated protein all the meningiomas were tested for Sp1 binding and all except one were found to be positive (Figure 3, lower panel). This meningioma extract also failed to show ERE-binding. We also checked the ER-ERE binding in meningiomas and the normal meninges from two patients. An ER-ERE complex was detectable in meningiomas, the normal meninges were negative for ER-ERE binding. However the normal meninges was also negative for Sp1 binding and therefore these results do not allow conclusions to be drawn on differences between normal and tumour tissue.

Figure 1 Binding specificity of the ERE oligonucleotide in crude, high salt extracts from MCF7 cells and meningioma tissue. Competition binding analysis was performed in which extracts from MCF7 cells a, and meningioma tissue b, were incubated with 1 ng of 32P-labelled ERE and increasing amounts of unlabelled ERE as specific competitor or unlabelled PRE as nonspecific competitor as reflected in the molar ratio indicated above each lane.

Figure 2 Anti-ER monoclonal antibody ER-P31 reduces ERE-protein complex in crude, high salt extracts. The extracts were first incubated with 5 μg antibody ER-P31 or nonspecific antibody (anti-C peptide) at 4°C for 18 h. After administration of 1 ng labelled ERE, the mixture was incubated for 30 min at room temperature and separated on a 6% gel. Extracts from MCF7 cells, an ER+/PR+ and ER-/PR meningioma tissue, and ER+/PR+ breast cancer were used. The ER and PR levels are given as fmol/mg protein. (F: protein-free ERE).
The abundant presence of PR in meningioma and the absence of ER measured by ligand binding assay or enzyme immunoassay prompted us to search for the ER or ER-like protein in meningioma using other techniques. We have already detected mRNA coding for the ER in meningioma (Koehorst et al., 1992). Besides the wild type transcript two variants were detected, one missing exon 7 and the other missing exon 4 (Koehorst et al., 1993). If the wt transcript or the variants are responsible for the PR synthesis, we hypothesise that a protein has to be present which can bind to an ERE.

We analysed DNA-binding activity to the ERE in human meningiomas by a gel mobility assay. The promotor regions of the PR gene are under investigation to define the functional ERE in this gene. So far two functional promotor regions have been described to be important for the induction of PR synthesis by ER. These two regions showed oestrogen inducibility in transient co-transfection experiments with vectors expressing the human ER, although no 'classical' ERE was detected in these promotor regions (Kastner et al., 1990). The sequences which are responsible for this ER inducibility are not well defined. Since no classical ERE is present in the promotor region of the PR gene and as the functional ERE in these regions is not well defined we
OESTROGEN RECEPTOR-LIKE PROTEIN IN HUMAN MENINGIOMA

is known however that the band shift assay is not always in agreement with the ligand binding assay. Foster et al. (1991) tested 79 breast tumours. The which 55 showed that the band shift assay was in agreement with the hormone-binding assay. In 13 tumours the hormone binding assay was negative whereas the band shift assay was positive. In two cases a ER-negative/PR-positive tumour was tested, in both cases the band shift assay was positive (Foster et al., 1991). These two cases have resemblance with the phenotype of meningiomas which are mostly also ER-negative/PR-positive.

In meningiomas an ER or ER-like protein is detectable by the band shift assay but not detectable by ligand-binding (Blaauw et al., 1986) or by enzyme immunoassay (Blankenstein et al., 1987). This may be due to the sensitivity of the assay (Scott et al., 1991).

Another explanation could be a mutation in the hormone-binding domain of the ER so that such an ER variant would not be detected by the ligand binding assay and probably also not in the immunoassay. For example in ER–/PR + breast tumours an ER variant was detected missing exon 5. This variant was constitutively active in a trans-activation assay (Fuqua et al., 1991). We detected in meningiomas two variant ER mRNA species (Koehorst et al., 1993). One variant with a major deletion in the ligand binding domain was an alternatively spliced product missing exon 7. This variant has no hormone independent transcriptional activity as shown by McGuire et al. (1991). Therefore this variant can probably not account for the apparently autonomous PR synthesis in meningiomas. The variant missing exon 4 codes for ER missing amino acid 254–365. Exon 4 includes the last part of the DNA binding domain, the hinge region and the first hundred amino acids of the ligand binding domain. The two zinc fingers of the DNA binding domain are intact so DNA binding can probably occur but no oestadiol will be bound by this variant because a great part of the ligand binding domain is missing. In addition, the highly positively charged region situated between amino acid 251 and 271 is missing. This sequence is very important for the formation of the non-DNA binding 8–9 S ER complexes, which bind heat shock protein 90 (hsp 90) (Chambraud et al., 1990). A variant missing amino acids 251–271 such as the variant missing exon 4 can probably not bind hsp 90 and therefore is always in the 4–5 S DNA binding form. This variant could play a role but whether it is capable of transactivation as well as its relationship to the ER-like protein described here remain to be investigated.

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