Sex hormone-binding globulin is synthesized in target cells

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Abstract

Sex hormone-binding globulin (SHBG) is a multifunctional protein that acts in humans to regulate the response to steroids at several junctures. It was originally described as a hepatically secreted protein that is the major binding protein for sex steroids in plasma, thereby regulating the availability of free steroids to hormone-responsive tissues. SHBG also functions as part of a novel steroid-signaling system that is independent of the classical intracellular steroid receptors. Unlike the intracellular steroid receptors that are ligand-activated transcription factors, SHBG mediates androgen and estrogen signaling at the cell membrane by way of cAMP. We have reviewed the current state of knowledge on the SHBG gene and the role of SHBG in steroid signaling (we shall not address its function as a plasma-binding protein).

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The sex hormone-binding globulin (SHBG) gene

The SHBG gene (Fig. 1) is located on chromosome 17p13.1, only 30 kb away from the p53 tumor-suppressor gene, and within a region known to undergo allelic deletions and mutations in a large variety of tumors (Cousin et al. 2000). Its proximity to p53 raises the unaddressed question of whether genomic events that alter the SHBG locus might also lead to changes arising in hormone-dependent cancers, e.g. breast and prostate. This question arises because, as we shall review, it is clear that SHBG is synthesized in these two tissues. Two major SHBG transcripts are known, each originating from a different promoter (minor SHBG transcripts have received little attention and will not be discussed here) (Gershagen et al. 1987, 1989, 1991, Hammond et al. 1987, 1989, Joseph et al. 1991, Bocchinfuso et al. 1992, Bocchinfuso & Hammond 1994, Hammond & Bocchinfuso 1996, Janne et al. 1998). The first major transcript encodes a precursor for the secreted (plasma) form of SHBG, and was originally described in the liver (SHBG_L) (Que & Petra 1987), while the second encodes a protein of unknown function and was originally described in the testis (SHBG_T) (Hammond et al. 1989).

SHBG_L

SHBG_L is encoded by eight exons, ranging in size from 90 to 208 bp. With the exception of a 733 bp intron separating exons 6 and 7 (which perhaps contains alternative splicing regulatory elements), the remaining introns are relatively small (133–331 bp). SHBG_L is under the transcriptional control of a TATA-less promoter which possesses multiple protein-binding sites, including those for hepatocyte nuclear factor-4 and SP-1 (Janne & Hammond 1998, Hogeveen et al. 2001). The nascent SHBG_L transcript encodes a precursor protein with a 29 amino acid, lysine-rich signal peptide (encoded within exon 1 and part of exon 2) at its amino terminus. The mature, secreted form of SHBG in human plasma lacks this signal peptide and circulates as a glycosylated, 92·5 kd homodimer (Khan et al. 1985, Hammond et al. 1986, Englebienne et al. 1987, Danzo et al. 1989, Grishkovskaya et al. 2000) containing two steroid-binding sites (Avvakumov et al. 2001).

SHBG_T

The second major transcript, SHBG_T, is regulated by an uncharacterized promoter that lies upstream of the SHBG_L promoter (Hammond & Bocchinfuso 1996). SHBG_L and SHBG_T differ in their 5’ sequences and in the absence of exon 7 in SHBG_T. The complete 5’ end sequence of SHBG_L has not been reported; the incomplete sequence contains an initial, long open reading frame wherein the first ATG start codon does not appear until the shared
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SHBG-mediated steroid signaling through the SHBG receptor

The current view of SHBG function differs dramatically from the way in which it was originally conceptualized, e.g. to regulate the concentration of certain free steroids in plasma. Although of undeniable importance, this original concept has been substantially broadened by the realization that SHBG is also part of a signal transduction system for steroids at the cell membrane.

The SHBG receptor

An active role for SHBG in steroid signaling was suggested initially by the discovery of specific, high-affinity binding sites for SHBG on uterine endometrial cell membranes (Strel’chyonok et al. 1984), isolated prostatic cell membranes (Avvakumov et al. 1985) and human placenta (Avvakumov et al. 1985). Subsequently, SHBG binding was also demonstrated in MCF-7 breast cancer cells (Frairia et al. 1991, Porto et al. 1992b, Fissore et al. 1994), normal breast (Frairia et al. 1991, Fortunati et al. 1992a) and epidermis (Guéant et al. 1991, Felden et al. 1992, Porto et al. 1992a, Krupenko et al. 1994), but not with striated muscle, colonic epithelia, or lymphocytes (Avvakumov et al. 1985, Felden et al. 1992, Fortunati et al. 1992a,b, Frairia et al. 1991, Porto et al. 1992a,b, Krupenko et al. 1994). The binding properties of SHBG are consistent with the presence of a specific RSHBG on cell membranes, and the biochemistry of SHBG–RSHBG binding is well characterized. Foremost, RSHBG only binds steroid-free SHBG. All steroids that bind to SHBG inhibit the binding of SHBG to RSHBG; the magnitude of the inhibition is directly proportional to the magnitude of the association constant for the steroid–SHBG interaction (Fig. 2) (Hryb et al. 1989, 1990). Once bound to RSHBG, SHBG binds steroids with affinities equal to SHBG that is in solution (Hryb et al. 1990). The SHBG domain, or at least a portion of it, that interacts with RSHBG has been localized to a ten amino acid stretch (TWDPEGVIFY) (Khan et al. 1990) encoded within exon 3. This region is shared between SHBG1 and SHBG1, and is the most highly conserved portion of the molecule, both across species (Khan et al. 1990) and in related proteins, e.g. protein S, laminin A, merosin, and Drosophila crumbs protein (Gershagen et al. 1987, Khan et al. 1990, Joseph & Baker 1992). Although there is a substantial body of knowledge about RSHBG, its structure remains elusive; the RSHBG gene has yet to be identified and characterized.

Steroid activation of cAMP through RSHBG

Our current conception of SHBG–RSHBG–steroid signaling is shown in Fig. 3. As discussed above, a specific sequence of events is necessary to initiate signaling through RSHBG, binding of unoccupied SHBG to RSHBG on the cell membrane, followed by binding of steroid to the SHBG–RSHBG complex, thereby activating it. Activation of RSHBG induces the synthesis of cAMP which, in turn, triggers downstream signaling and initiates genomic effects through the activation of promoters containing cAMP responsive elements (Nakha et al. 1990, Rosner et al. 1992). These events occur too rapidly to be affected either by the dissociation of SHBG–RSHBG, seen subsequent to binding of the agonist, or by the transcriptional activation of classical steroid hormone receptors.
a dose-related decrease in the binding of SHBG to RSHBG. Whether or not a steroid is an agonist of RSHBG-mediated cAMP signaling (Nakhla et al. 1998), G-protein diol (3α,17β diol) is active in this system at physiologic concentrations, was previously thought to be an inactive metabolite of dihydrotestosterone (DHT). Other steroids that bind SHBG with high affinity, e.g. DHT, testosterone, and 2-methoxyestradiol, are not agonists, but instead antagonize the effects of 3α-diol. On the contrary, DHT is an agonist for SHBG–RSHBG in both the LNCaP prostate cancer cell line (Nakhla et al. 1990) and in cultured human placenta (Queipo et al. 1998). Not surprisingly, the degree to which agonists induce cAMP through RSHBG appears to vary with cell type. For instance, the fractional increase in cAMP in cultures of human (Nakhla et al. 1994) and canine prostate (Nakhla et al. 1995) far exceeds that seen in LNCaP cells. It should not be lost sight of that, in both LNCaP cells (Nakhla et al. 1990) and placenta (Queipo et al. 1998), SHBG in the absence of steroid causes a modest increase in cAMP. Although the relationship between steroidal structure and affinity for SHBG has been examined in some detail (Cunningham et al. 1979, 1981), those studies shed no light on whether a given steroid might be an agonist or antagonist in the SHBG–RSHBG system.

Furthermore, inhibitors of the transcriptional activation of the estrogen receptor and androgen receptor (AR) do not affect the cAMP response, supporting the independence of this pathway.

RSHBG appears to be coupled to a G-protein. There is a dose-related decrease in the binding of SHBG to RSHBG after incubation of the receptor preparation with the non-hydrolyzable GTP analogue, guanylyl-5’-imidodiphosphate (Nakhla et al. 1999), a phenomenon typical of the behavior of receptors coupled to G-proteins. In addition, in COS-1 cells, which express a functional RSHBG expression of dominant negative mutants of the G-protein α-subunit (Otsawa & Johnson 1991), cause a decrease in RSHBG-mediated cAMP signaling (Nakhla et al. 1999).

Steroids that bind to SHBG act as either agonists or antagonists of RSHBG-mediated signaling. Furthermore, whether or not a steroid is an agonist of RSHBG-mediated signaling appears to be dependent on cell type. In the prostate, two steroids, estradiol and 5α-androstan-3α,17β diol (3α-diol) are potent agonists (Nakhla et al. 1990, 1995). In fact, 3α-diol, which is active in this system at physiologic concentrations, was previously thought to be an inactive metabolite of dihydrotestosterone (DHT). Other steroids that bind SHBG with high affinity, e.g. DHT, testosterone, and 2-methoxyestradiol, are not agonists, but instead antagonize the effects of 3α-diol. On the contrary, DHT is an agonist for SHBG–RSHBG in both the LNCaP prostate cancer cell line (Nakhla et al. 1990) and in cultured human placenta (Queipo et al. 1998). Not surprisingly, the degree to which agonists induce cAMP through RSHBG appears to vary with cell type. For instance, the fractional increase in cAMP in cultures of human (Nakhla et al. 1994) and canine prostate (Nakhla et al. 1995) far exceeds that seen in LNCaP cells. It should not be lost sight of that, in both LNCaP cells (Nakhla et al. 1990) and placenta (Queipo et al. 1998), SHBG in the absence of steroid causes a modest increase in cAMP. Although the relationship between steroidal structure and affinity for SHBG has been examined in some detail (Cunningham et al. 1979, 1981), those studies shed no light on whether a given steroid might be an agonist or antagonist in the SHBG–RSHBG system.

**Figure 2** Inhibition of the binding of 125I-SHBG to the soluble RSHBG by steroids (from Hryb et al. 1990). Soluble RSHBG was added to a constant amount of 125I-SHBG and varying concentrations of either cold SHBG or the steroids indicated. Incubations were for 40 h at 37 °C, which achieved steady-state binding. The receptor is stable for this period of time at this temperature. SDS gel electrophoresis, followed by autoradiography of both receptor-bound and free 125I-SHBG, after 40 h, showed the 125I-SHBG to be unmetabolized. Each steroid caused an inhibition in the binding of 125I-SHBG to the soluble SHBG receptor. Further, their inhibitory potency (K_i) is in precisely the same sequence (DHT >> 2-methoxyestradiol > 2-MethoxyE2 >> testosterone > estradiol > methyltrienolone > cortisol) as the tightness of their association (K_A) with SHBG. Indeed, the relative (to testosterone) ability of each steroid to inhibit the binding of SHBG to its receptor (K_i(testosterone)/K_i(steroid)) was almost identical to its relative SHBG-binding activity (K_A(steroid)/K_A(testosterone)). These results, taken together with double reciprocal plots (not shown), show that the inhibitory effects of the steroids were due to the interaction between them and SHBG (non-competitive inhibition), and not between the steroid and the RSHBG. Thus, liganded SHBG must have a conformation which prevents it from binding to its receptor.

**Biologic effects of steroid signaling through RSHBG**

*Induction of prostate specific antigen (PSA) in prostate cells*

Delineation of the biologic effects of SHBG signaling through RSHBG has lagged behind our understanding of the biochemical analysis of its signaling pathway. Details regarding the downstream effects of steroid signaling through SHBG exist, but are not extensive. A downstream event of potential biologic importance is the intersection of this pathway with an AR-mediated event, the activation of the PSA gene and secretion of its translational product (Nakhla et al. 1997). The human PSA gene possesses an androgen response element in its promoter, and is transcribed upon activation of the AR in prostate cells. Prostate explants secrete PSA when treated with DHT; however, they do not when treated with estradiol, which does not bind to the AR. When such explants were treated first with SHBG, and then with estradiol, they produced PSA at concentrations similar to those seen when they were exposed to DHT. Furthermore, inhibitors of estrogen receptor activation did not block estradiol–SHBG–RSHBG-mediated PSA induction, whereas inhibitors of AR activation did. These results indicate that estradiol–SHBG–RSHBG initiates ligand-independent activation of PSA secretion.

*Cell growth*

RSHBG signaling affects growth in two different cell lines, with opposite results. It decreases the estrogen-mediated growth of the human breast carcinoma cell line, MCF-7.
Figure 3 The SHBG signaling system. In its steroid-free configuration, SHBG binds to $R_{SHBG}$ on cell membranes, forming a bipartite complex (SHBG–$R_{SHBG}$). SHBG, already bound to a steroid, non-competitively inhibits the binding of SHBG to $R_{SHBG}$. However, within minutes after exposure of SHBG–$R_{SHBG}$ to a steroid agonist, e.g. estradiol (Nakhla et al. 1990, 1994) or 5α-androstan,3α,17β-diol (Nakhla et al. 1995), a tripartite complex (steroid–SHBG–$R_{SHBG}$) forms that activates adenylyl cyclase, leading to the generation of the second messenger, cAMP.
Figure 4  SHBG expression in normal prostate (from Hryb et al. 2002) (A) In situ hybridization (× 400). A 5 μm human prostate section was processed using the Biogenex (San Ramon, CA, USA) super sensitive in situ hybridization kit. RNase activity was blocked and the section was incubated with a 521 bp human SHBG cDNA probe (prepared by PCR incorporation of biotin-14-dCTP). After heating and incubation, slides were developed using the ABC method (ABC elite system; Vector Labs, Burlingame, CA, USA), using DAB as the substrate, and counterstained with hematoxylin. Photographs were taken with a 35 mm camera mounted to a BX60 microscope and digitized.

(B) Immunohistochemistry (× 400). A 5 μm human prostate section was fixed and incubated overnight at 4 °C with a rabbit anti-SHBG polyclonal antisera (64–4), generated in our laboratory. The section was developed by the avidin–biotin complex (ABC) method using DAB as the substrate, and counterstained with hematoxylin. Photographs were taken as above.
(Fortunati et al. 1996), whereas the human prostate cancer cell line, ALVA-41, has its growth enhanced by both estradiol and DHT in the presence of SHBG–RSHBG (Nakha & Rosner 1996). These results mirror the effect of cAMP in each of the two cell lines. Whether SHBG–RSHBG–induced cAMP elevation is solely responsible for these observations, or whether other factors involved in growth regulation play a role, remains to be investigated. Furthermore, these are cancer cell lines; whether signaling through RSHBG has the same effects on normal breast and prostate epithelial cells is not known. On a very speculative note, if this relationship exists in normal cells, SHBG might be considered a tumor-suppressor gene in breast cancer, and agonists of the SHBG–RSHBG Pathway might be used to suppress the malignant phenotype, while antagonists of SHBG–RSHBG signaling might be useful in prostate cancer, where inhibition is wanted.

**Localized expression of SHBG in hormone-responsive tissues**

The presence of SHBG in cells that respond to sex steroids has been examined in a number of laboratories. Early immunohistochemical studies, using rabbit polyclonal antisera, showed SHBG antigen in both the prostate and breast (Bordin & Petra 1980, Tardivel-Lacombe et al. 1984, Sinnecker et al. 1988, 1990, Meyer et al. 1994, Germain et al. 1997). However, whether SHBG was delivered to these cells through the plasma or was locally expressed remained a question. Indeed, although all the antisera were raised using highly purified SHBG, there was no proof that this intracellular antigenic activity was SHBG, rather than a related antigen.

More recently, SHBG mRNA has been demonstrated in a number of non-hepatic tissues and cell lines (Larrea et al. 1993, Misao et al. 1994, 1997, Moore et al. 1996, Murayama et al. 1999). Although the data in the cell lines that stain for SHBG protein, and show SHBG mRNA by RT-PCR and/or by Northern blotting, are convincing, the conclusions based on experiments using human tissue sections are ambiguous. With one exception (Noe 1999), studies showing the tissue mRNA did not show the protein, and those demonstrating the protein did not show the mRNA. In the one exception, Noe (1999) detected both the protein (immunostaining) and the mRNA by RT-PCR in human Fallopian tubes. However, no studies were presented to ascertain whether the mRNA was translated, e.g. the possibility remained that the mRNA was not translated and the protein arrived via the plasma. Although it is possible to demonstrate the causal relationship between an mRNA and its protein in cell lines, this cannot be done in tissue sections. The strongest inferential evidence that is possible, under these circumstances, is to show that the mRNA (in situ hybridization) and the protein exist in the same cells. Thus, we (Hryb et al. 2002) undertook an examination of human prostate and breast tissue sections by *in situ* hybridization and immunocytochemistry. In the prostate, cells that expressed SHBG mRNA (Fig. 4A) also stained for SHBG protein with a monospecific, polyclonal rabbit anti-SHBG (64–4) (Fig. 4B) or monoclonal antibodies (data not shown). Comparable results were obtained for breast tissue (authors’ unpublished observations). While we cannot dismiss internalization of plasma SHBG as at least a partial source of the immunoreactive SHBG in these studies, it is likely that locally produced SHBG is the major species in these cells. If so, regulated SHBG synthesis and secretion in the breast and prostate could affect intracellular free steroid concentrations and participate in RSHBG signaling independent of plasma SHBG. These results raise a number of important new questions. (1) Does locally expressed SHBG affect intracellular steroid signaling pathways or act in an autocrine or paracrine manner through RSHBG? (2) Does SHBG participate in crosstalk between epithelial and stromal cells, as SHBG is predominantly expressed in the former and RSHBG is predominantly expressed in the latter? (3) Do perturbations of SHBG expression in cancer cells, through allelic deletions, contribute to the malignant phenotype and, if so, can agonists or antagonists of RSHBG signaling serve as useful therapeutic agents?

In summary, the portrait of SHBG as a monofunctional plasma steroid-binding protein has changed to that of one with multiple functions. It appears that it not only participates in steroid signaling at the cell membrane, but that the regulation of its synthesis and secretion in target cells offers new possibilities for the local modification of steroid hormone effects.

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