Role of Specific Response Elements of the \textit{c-fos} Promoter and Involvement of Intermediate Transcription Factor(s) in the Induction of Sertoli Cell Differentiation (Transferrin Promoter Activation) by the Testicular Paracrine Factor PModS* \\

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Role of Specific Response Elements of the c-fos Promoter and Involvement of Intermediate Transcription Factor(s) in the Induction of Sertoli Cell Differentiation (Transferrin Promoter Activation) by the Testicular Paracrine Factor PModS*

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ABSTRACT

A mesenchymal-epithelial cell interaction exists in the testis between the Sertoli cells that form the seminiferous tubule and the mesenchymal-derived peritubular myoid cells that surround the tubule. Analysis of the mesenchymal-epithelial interactions between these cells revealed the local production of a mesenchymal factor, PModS. PModS modulates the differentiated functions of Sertoli cells in vitro, including stimulation of the iron-binding protein transferrin (Tf). Previous results have indicated that PModS-induced Tf gene expression involves the activation of immediate early genes. One of the immediate early genes was identified as c-fos. The importance of c-fos was demonstrated in the current study when a c-fos antisense oligonucleotide was found to inhibit the ability of PModS to induce the expression of a Tf promoter-chloramphenicol acetyltransferase (CAT) construct. The regulation of c-fos by PModS was investigated with various CAT constructs containing segments of the c-fos promoter, such as the serum response element (SRE), sis-inducible element (SIE), CAMP response element (CRE), and phorbol ester/TPA response element (TRE), transfected into cultured Sertoli cells. PModS has no effect on CAMP response element-CAT or TRE-CAT, suggesting that PModS does not act through stimulation of CAMP and protein kinase C pathways. PModS was found to activate the c-fos SRE-CAT construct and the SIE-CAT construct. A construct containing both SIE and SRE was stimulated to the same degree as either element alone. Gel mobility shift assays using nuclear extracts from PModS-stimulated Sertoli cells and a radiolabeled SRE oligonucleotide resulted in retarded mobility of a DNA-protein complex. A gel shift with a SRE oligonucleotide containing an ETS domain resulted in a unique intermediate transcription factor(s) that regulates downstream Sertoli cell differentiated functions, such as Tf expression. These studies have initiated an investigation of the transcriptional regulation of Sertoli cell differentiation. (Endocrinology 136: 3046–3053, 1995)
Treatment of Sertoli cells with the tyrosine phosphorylation inhibitor, genistein, suppresses PModS-stimulated Tf production (5). Therefore, the actions of PModS on Sertoli cells appear to be in part through an activation of tyrosine kinase(s). Regulation of Sertoli cell differentiation by PModS on a molecular level has also been investigated. An inhibitor of translation, cycloheximide, was used to determine whether PModS could directly stimulate Tf messenger RNA (mRNA) expression or whether intermediate protein synthesis was involved. Cycloheximide inhibited PModS-induced Tf mRNA synthesis, indicating that activation of immediate early genes is required for stimulation of Tf expression (6). Northern analysis of potential immediate early genes demonstrated that PModS stimulates c-fos mRNA synthesis, but has no effect on two other nuclear proteins c-jun and C/EBP. The protein c-fos is a protooncoprotein that is transiently stimulated by a variety of mitogens and differentiation factors (7). Time-course data show that c-fos mRNA levels increase within minutes after PModS treatment and subsequently decline to basal levels after 2 h. In contrast, PModS-induced increases in Tf mRNA do not occur until 2 h, then peak at approximately 12 h, and are maintained for up to 5 days in culture. Treatment of Sertoli cells with a c-fos antisense oligonucleotide partially suppresses PModS-enhanced Tf protein production (6). These data indicate that c-fos is involved as an immediate early gene in response to PModS. The current study was designed to analyze PModS activation of the c-fos promoter and relate this to the previous data on signal transduction and Sertoli cell differentiation. Activation of the Tf promoter was investigated as a downstream event involved in Sertoli cell differentiation.

**Materials and Methods**

**Cell preparation and culture**

Sertoli cells were isolated from the testes of 20-day-old rats by sequential enzymatic digestion (8), with a modified procedure described by Tung et al. (9). Decapsulated testicular fragments were digested first with trypsin (1.5 mg/ml; Gibco-BRL, Gaithersburg, MD) to remove the interstitial cells and then with collagenase (1 mg/ml type I; Sigma Chemical Co., St. Louis, MO) and hyaluronidase (1 mg/ml; Sigma). Sertoli cells were then plated under serum-free conditions in 24-well Falcon plates at 1 × 10^6 cells/well. Cells were maintained in a 5% CO₂ atmosphere in Ham’s F-12 medium (Gibco-BRL) at 32°C. Sertoli cell cultures were treated as described in Results, after 48 h of culture and transfection. Cultures were treated with FSGI (100 ng/ml, ovine FSH-I-16, National Pituitary Agency), (Bu)₃cAMP (1 mM), 10% calf serum, or greater than maximally effective concentrations of PModS(S300) (25 µg/ml) or PModS(C4) (25 ng/ml).

Peritubular cells were obtained from the collagenase digestion supernatant, as described by Skinner et al. (2), after the tubule segments had sedimented. Peritubular cells were plated in Ham’s F-12 medium containing 10% calf serum and grown to confluence. Cells were then subcultured and plated at 25% confluence. When subcultured cells were confluent, they were washed with serum-free medium. The cells were subsequently cultured in serum-free medium for up to 4 weeks with 48- to 72-h medium collections.

Fresly collected serum-free conditioned medium from the peritubular cells was treated with phenylmethylsulfonylfluoride (25 µM final concentration) and benzamidine (0.1 mM final concentration) and centrifuged at 10,000 × g for 15 min at 4°C to remove cell debris. When necessary, medium was stored at −20°C. Conditioned medium was concentrated 100-fold by ultrafiltration with an Amicon system (Amicon Corp., Lexington, MA) and a 3000 mol wt exclusion limit membrane.

**PModS preparation**

PModS was obtained from concentrated peritubular cell-conditioned medium, as previously described (2). An ammonium sulfate precipitate of concentrated conditioned medium was applied to a size-exclusion Sephacryl S300 column (Pharmacia, South San Francisco, CA). The active peak of the S300, determined by bioassay of Sertoli cell Tf production, was collected and applied to a chromatofocusing column (Pharmacia). The pH 5.7-6.8 pool was applied to a 1 × 15-cm heparin-Sepharose affinity column. Eluted proteins were applied to two successive C4 reverse phase columns (Vydac, Hesperia, CA) and eluted with a linear gradient from 25–60% acetonitrile. The partially purified PModS(S300) pool, termed PModS(S300), and the more highly purified PModS(C4) fraction, termed PModS(C4), were stored at −70°C before use in the presence of 1 mg/ml BSA.

**Reporter gene constructs**

**c-fos promoter.** The chloramphenicol acetyltransferase (CAT) reporter plasmids (PBL-CAT2) with the thymidine kinase minimal promoter (10) containing either the cAMP response element (CRE) (12-base pair (bp) 5′-flanking sequence from the transcriptional start site of the human c-fos promoter with a 5′-TGACGTTT-3′ sequence at −60 bp) or TPA/activating protein 1 (API) response element (TRE; 5′-TGCCTCA-3′), or the serum response element (SRE; 5′-CAGGATGCCAATTAAGACATC-3′) sequences of the c-fos promoter were generated: the entire c-fos promoter was generously provided by Dr. Jeff Holt (Vanderbilt University, Nashville, TN). The cis-inducible element (SIF)-CAT reporter plasmid was constructed by cloning a synthetic 15-bp oligonucleotide (CAGTCGGTTAATACT) into HindIII and SphI sites on either ends into PBL, CAT2 plasmid. The SIE/SRE construct was prepared by cloning the 15-bp SIE oligo into HindIII/SphI sites of the SRE/CAT plasmid.

**Tf promoter.** The CAT reporter plasmid containing −581 bp (PUC8 CAT) and the human GH reporter plasmid containing −3.0-kilobase (kb) sequences of the mouse Tf (mTf) promoter (12) were generously provided by Dr. G. Stanley McKnight (University of Washington, Seattle, WA). The CAT reporter plasmids containing −3.0-kb mTf promoter and its deletions were constructed as follows: 1) The −3.0-kb mTf CAT construct was made by ligating a −3.0-kb BamHI-BamHI digest to the BamHI site in PGL2-CAT plasmid (Promega). 2) The 2-kb upstream HindIII-HindIII fragment in the 3-kb mTf promoter was ligated in the HindIII site of −581-bp mTf-CAT to obtain the 2.6-kb mTf-CAT plasmid. 3) The −1.6-kb mTf CAT plasmid was derived from the −3.0-kb mTf CAT (no. 1 above) by digesting out the upstream −1.4-kb PsIl fragment. 4) To prepare the upstream 1-kb mTf-CAT (1-kb mTf-Tf-CAT), the 3-kb mTf plasmid was digested with HindIII-PstI. The 1-kb fragment (1.6–2.6 kb) was then cloned into a PBL-CAT2 plasmid.

Constructs for each of the plasmids that did not contain the specific response element or promoter fragment (i.e. promoterless plasmid) were also generated. In all cases, experiments were performed to examine the actions of various treatments on these promoterless plasmids. As discussed in Results, these promoterless plasmids generally did not respond to the treatments. If a small response was observed, the data are shown in the figures or stated in Results.

**Transfection**

Sertoli cells, cultured for 48 h, were transfected with a reporter gene construct by the calcium phosphate method coupled with hyperosmotic shock (10% glycerol), as previously described (13). Cells were treated 2 h after transfection, and unless otherwise stated, CAT activity was determined after 48 h in culture.

**CAT assay**

Assay of CAT activity was performed as follows. Medium was removed from the wells, and the cells were washed once with PBS. One hundred microliters of 1 × cell lysis buffer (Promega Corp., Madison, WI) was added to each well, and the cells were disrupted by shaking vigorously. After incubation at room temperature for 10 min, 250 µl of chloramphenicol acetyltransferase (CAT) reporter plasmid was added to each well, and the cells were disrupted by shaking vigorously. After incubation at room temperature for 10 min, 250 µl of chloramphenicol acetyltransferase (CAT) reporter plasmid was added. The mixture was incubated at 37°C for 2 h and then assayed for CAT activity as follows. Medium was removed from the wells, and the cells were washed once with PBS. One hundred microliters of 1 × cell lysis buffer (Promega Corp., Madison, WI) was added to each well, and the cells were disrupted by shaking vigorously. After incubation at room temperature for 10 min, 250 µl of chloramphenicol acetyltransferase (CAT) reporter plasmid was added. The mixture was incubated at 37°C for 2 h and then assayed for CAT activity.
Antisense procedure

Sertoli cells. The Sertoli cells were isolated as described above and cultured in 137-mm petri dishes (Falcon Plastics, Oxnard, CA). The cells were treated with 500 µM 2,5-dimethylfucose for 1 h at room temperature. After five washes under the 48 h before harvest of the cells for CAT assay.

Gel mobility shift assay

Gel shift assays were performed with nuclear extracts of isolated Sertoli cells. The Sertoli cells were isolated as described above and cultured in 137-mm petri dishes (Falcon Plastics, Oxnard, CA). The cells were treated at 48 h in culture either FSH, PModS(S300), or 10% calf serum. After 72 h, the cells were scraped off the petri dishes and washed once with PBS. The nuclear extracts of these cells were then prepared as described by Guillou et al. (14). The probes used in gel retardation assays were the SRE (5'-GGATGTCCATATTAGGACACATCTG-3') and the ETS-SRE element (SRE with 5'-CAGGAT sequence) of the c-fos promoter; the SIE (5'-CATITCCCGTAAATC-3') located 25 bp upstream of the c-fos SRE (15), and API dimer (5'-TTAGTCATGAGTCA-3'). The complementary oligonucleotides were 5'-end labeled with [32P]ATP using polynucleotide kinase. Of these fragments, two gave a positive gel shift. The restriction fragments SE1 (200 bp) and SE2 (180 bp) were located at -2.4 and -1.9 kb, respectively, on the 3-kb mTf 5'-flanking region. These two Tf promoter fragments (i.e., SE1 and SE2) were routinely isolated, dephosphorylated, and end labeled with [γ-32P]ATP for gel shift analysis.

The gel retardation assay used was a modification of the protocol described by Garner and Rezvani (15). The final reaction volume of 20 µl was 15% approximately 200-1 fragments were isolated, dephosphorylated, and end labeled with [γ-32P]ATP using polynucleotide kinase. These fragments, two gave a positive gel shift. The restriction fragments SE1 (200 bp) and SE2 (180 bp) were located at -2.4 and -1.9 kb, respectively, on the 3-kb mTf 5'-flanking region. These two Tf promoter fragments (i.e., SE1 and SE2) were routinely isolated, dephosphorylated, and end labeled with [γ-32P]ATP for gel shift analysis.

Antisense procedure

A c-fos antisense oligonucleotide, 3'-TACTACCGGTCA-5', was prepared and used, as previously described (16, 17). A c-fos sense oligonucleotide was prepared for use as a control. Transfected Sertoli cells were treated with either the antisense or sense oligomer (4 µM) starting 8 h after transfection and treated every 8 h for a total of 48 h before harvest of the cells for CAT assay.

Immunoblot procedure

A gel mobility shift assay was electrophoretically transferred to a polyvinylidene difluoride membrane (Immobilon Millipore, South San Francisco, CA) by electrophoresis in Tris-glycine buffer containing 20% methanol. The blot was then blocked with 5% nonfat milk (dissolved in 50 mM Tris (pH 7.4), 150 mM NaCl, and 0.05% Nonidet P-40) and incubated with an overnight incubation at room temperature. The wells were then discarded, and the buffer was collected in 1.5-ml microfuge tubes. Tubes were heated to 95 C for 10 min to inactivate endogenous acetylases and then spun at 12,000 x g for 10 min at 4 C to remove cell debris. An aliquot of cell extract (54 µl) was mixed with 65 µl 25 mM Tris (pH 8.0), 25 µg n-butylryl coenzyme A (5 mg/ml; Sigma), and 0.1 µCi [3H]chloramphenicol (0.1 µCi/µl; ICN, Costa Mesa, CA) and incubated overnight at 37 C. The mixture was extracted once with 300 µl xylene and back-extracted with 100 µl 25 mM Tris (pH 8.0). A 200-µl aliquot of the organic phase was counted in a scintillation counter to determine the relative amount of CAT activity.

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Statistical analysis

Each data point was converted to a percentage of the control value, and the mean and SEM from multiple experiments were determined, as indicated in the figure legends. Data were analyzed by analysis of variance, using the SPSS Statistical Package (SPSS, Chicago, IL).

Results

To confirm that c-fos is involved at a molecular level in PModS stimulation of Tf, a CAT reporter gene construct containing -2.6 kb of the Tf gene promoter was transiently transfected into cultured Sertoli cells. Sertoli cells were isolated from 20-day-old rat testes and cultured under serum-free conditions unless otherwise stated. After transfection, the cells were left untreated (control) or were treated with FSH, a partially purified PModS preparation from a size-exclusion column termed PModS(S300), a combination of PModS(S300) and the antisense c-fos oligonucleotide (S300+ANTI), or a combination of PModS(S300) and the sense c-fos oligonucleotide (S300+SENSE) for 48 h before analysis of the cell extracts for CAT activity. FSH, S300, and S300+SENSE activated the Tf CAT construct (Fig. 1). A control promoterless plasmid (pCAT-basic) was slightly stimulated by PModS(S300) treatment. Treatment of the transfected cells with the antisense oligonucleotide inhibited PModS(S300)-induced CAT activity (Fig. 1). These data confirm that c-fos is involved in PModS-stimulated Tf gene expression.

c-Fos promoter analysis

To determine how PModS activates c-fos gene expression, CAT reporter gene constructs containing different regulatory

![Figure 1](image-url)
elements of the c-fos promoter were generated, including the CRE, TRE, SIE, and SRE. A schematic representation of the locations of these elements within the c-fos promoter is shown in Fig. 2A. These constructs were transiently transfected into cultured Sertoli cells. After transfection, the cells were treated with FSH, 10% calf serum, or more highly purified PModS preparations from the C4 reverse phase column or 10% calf serum, (Bu)_2cAMP, PModS(S300), or more highly purified PModS preparations from the C4 reverse phase column termed PModS(C4) for 48 h. The CRE of the c-fos promoter was activated by both FSH and (Bu)_2cAMP 400% above the control level (P < 0.05), which demonstrated that the reporter construct can be activated (data not shown). Treatment with PModS(S300) did not increase CRE-CAT activity above the control level (Fig. 2B). Treatment of the c-fos TRE-transfected cells with FSH resulted in a 300% stimulation above the control level (P < 0.05), which demonstrated that the reporter construct can be activated (data not shown). Treatment of the transfected Sertoli cells with PModS(S300) does not stimulate the TRE construct (Fig. 2B). A SRE construct was used that included the 5'-flanking 5-bp ETS domain ETS-SRE sequence of the c-fos promoter. Activation of the c-fos SRE construct was accomplished by treatment with 10% calf serum as a positive control to demonstrate that the reporter construct can be activated (data not shown). Treatment of the transfected Sertoli cells with PModS(S300) did not stimulate the TRE construct (Fig. 2B). A SIE construct was also stimulated by PModS(S300) and PModS(C4) (Fig. 2B). Interestingly, a construct containing both the SIE and SRE was stimulated to the same or lesser degree as that containing either element alone. The entire 400-bp c-fos promoter construct was stimulated approximately 900% over the control level by PModS(S300) (data not shown). The control plasmids not containing a promoter were not influenced by any of the treatments (data not shown). Purified PModS had a similar action as the partially purified PModS(S300). This indicates that stimulation of c-fos expression by PModS is in part mediated through activation of the SRE and SIE.

To extend these observations, gel retardation/mobility shift assays were performed. Nuclear extracts were obtained from Sertoli cells cultured in the absence (control) or presence of PModS(S300) or 10% (vol/vol) calf serum as a positive control. A gel shift with an oligonucleotide to Oct-1 was used to check the quality of the nuclear extracts (Fig. 3). All extracts resulted in a similar level of Oct-1 gel shift. To confirm that c-fos was induced, a gel shift with an API oligonucleotide was performed, because the c-fos-jun complex binds to the API site. Incubation of the labeled API oligonucleotide with nuclear extracts of Sertoli cells treated with PModS(S300) or serum resulted in the formation of a DNA-protein complex with retarded mobility (i.e. gel shift). The observed gel shift was specific for API, because it could be effectively competed by unlabeled API (Fig. 3). This confirms the induction of c-fos by both PModS(S300) and serum.

Activation of SRE of the c-fos promoter was also investigated with a gel mobility shift assay using a SRE oligonucleotide (Fig. 4A). PModS(S300) induced a gel shift that was competed with excess unlabeled SRE. Serum was used as a positive control and also promoted a gel shift. To assess whether PModS may promote binding of a ternary complex, an ETS-SRE sequence was used in a gel shift. The ETS domain is a 5'-CAGGAT flanking sequence of the SRE that is needed for the binding of ETS oncogene-like proteins and is required for the formation of a ternary complex with the serum response factor (SRF) that binds to the SRE. PModS(S300) promoted a distinct shift with the ETS-SRE that was not present in control nontreated cells or in serum-treated cells (Fig. 4A). PModS(S300) promoted two distinct gel shifts, whereas serum promoted a diffuse shift below that of PModS(S300). These gel shifts were displaced with excess unlabeled oligonucleotide (data not shown). Therefore, PModS(S300) induced protein binding to the SRE and ETS-SRE oligonucleotides. An alternate response element, located 25 bp from the SRE, is the 25 bp OCT1 site. The data are representative of five different experiments.

![Fig. 2](image)

**Fig. 2.** a. Schematic representation of the c-fos promoter and the locations of the CRE, SRE, SIE, and TRE in promoter-CAT constructs. The designation for the promoter constructs is listed at the left, and that for the plasmid used at the right. b. Regulation of the c-fos promoter constructs (CRE, TRE, SIE, and SRE) in Sertoli cells cultured in the absence (control) or presence of PModS(S300) (■) and PModS(C4) (■). Data are expressed as a percentage of the control and are presented as the mean ± SEM from six different experiments performed in replicate.

![Fig. 3](image)

**Fig. 3.** Gel mobility shift assay with 32P-radiolabeled Oct-1 and API oligonucleotides. Nuclear extracts from Sertoli cells cultured in the absence (C) or presence of PModS(S300) (P) or 10% calf serum (S) were used. The DNA-protein complexes were electrophoretically separated on 5% polyacrylamide gels, then dried and autoradiographed. The data are representative of five different experiments.
upstream from the SRE, is the SIE, which is also responsive to serum. A SIE oligonucleotide gel shift demonstrated that PModS(S300) promotes a SIE gel shift (Fig. 4B). Again, serum was used as a positive control and promoted the anticipated gel shift. This SIE gel shift was specific and could be competed by excess unlabeled SIE oligonucleotide. Previously, a SIE gel shift has been shown to involve three shifts, termed A, B, and C. PModS was found to promote all three forms, with A (lowest) being the predominant (Fig. 4B). The gel shift data presented confirm the reporter gene experiments previously discussed. Therefore, PModS appears to act through the SRE and SIE to promote c-fos gene expression.

**Tf promoter analysis**

To determine the location of response elements within the Tf promoter that are activated by treatment with PModS, four CAT constructs containing a 581-bp, 1.6-kb, 2.6-kb, or 3-kb upstream region of the Tf promoter were produced (Fig. 5A). FSH treatment of cells transfected with any of the constructs, except the 2.6- to 1.6-kb Tf PBL-CAT2, termed 1-kb Tf-Tk-CAT, stimulated CAT activity approximately 400% above control values, confirming that these constructs can be activated (data not shown; \(P < 0.01\)). PModS(S300) treatment induced activation of the 581-bp, 1.6-kb, and 3-kb Tf constructs (Fig. 5B) to a similar extent. The -2.6-kb Tf construct was stimulated to a greater extent by PModS(S300) (Fig. 5B). Interestingly, the PModS(C4) preparation stimulated primarily the -2.6-kb Tf construct. This indicates that an enhancer-like activity involved in PModS stimulation of Tf production is located between -1.6 and -2.6 kb, and a repressor is located between -2.6 and -3 kb of the promoter. To extend these observations, the 1000-bp region of the promoter between -1.6 and -2.6 kb was cloned into a Tk-CAT construct (PBL-CAT2, Fig. 5A), termed 1-kb Tf-Tk-CAT. PModS was found to activate this construct, but had only a slight effect on the control plasmid not containing the 1-kb Tf promoter (Fig. 6). PModS(C4) had an action similar to that of the partially purified PModS(S300). The plasmid used contained a minimal thymidine kinase (Tk) promoter with a TATA box (Tk-CAT) that was required for PModS to activate this 1-kb Tf construct. In the absence of a minimal promoter with a TATA box, a similar construct was not activated (data not shown). Therefore, the ability of PModS to activate the enhancer elements in this Tf promoter fragment does require a minimal promoter containing a TATA box.

Although the region between 0 and -581 bp has been sequenced and mapped, little is currently known about the distal region of the Tf promoter. Therefore, the exact nature of these putative upstream regulatory elements is unknown. To initiate an examination of this region, the entire 2.5-kb fragment of the Tf promoter between -581 bp and -3 kb was digested with various restriction enzymes to generate 15 approximately 200-bp fragments. These fragments were used in gel mobility shift assays to identify potential PModS-
FIG. 6. Regulation of a 1-kb fragment of the TF promoter between −1.6 kb and −2.6 kb containing SE1 and SE2 inserted into a Tk-CAT plasmid and termed 1 kb Tf-Tk-CAT. Sertoli cells were cultured in the absence (C) or presence of PModS(S300) or PModS(C4). The actions of PModS(S300) on a control Tk-CAT plasmid are also shown. Data are expressed as a percentage of the value in control nontreated cells for each construct and are presented as the mean ± SEM from three different experiments performed twice. ***, Statistically significant difference from the control (P < 0.01).

responsive regions. Thirteen of these TF promoter restriction fragments did not cause a gel shift under control or PModS-stimulated conditions (data not shown). Two restriction fragments upstream of the mTF transcriptional start site located at −2.4 kb, designated SE1, and at −1.9 kb, designated SE2, did cause a gel shift after PModS stimulation (Fig. 7). The location of these Sertoli elements (i.e. SE1 and SE2) on the Tf promoter is diagrammatically shown in Fig. 5A. To determine whether either SE1 or SE2 is a potential AP1-binding site, a competition experiment was carried out with excess unlabeled API oligonucleotide. The unlabeled API was not able to displace the SE1 or SE2 complex. Therefore, the gel shift appears not to be due to a c-fos/jun-binding API site. The SE2 fragment was unable to form the DNA-protein complex with the nuclear extracts of Sertoli cells treated with serum (Fig. 7). PModS(S300) caused a gel shift with both SE1 and SE2 that was competed with excess unlabeled oligonucleotide (Fig. 7). Excess SE2 DNA fragment did not compete for the SE1 complex, nor did the SE1 compete for the SE2 complex. These two, apparently distinct, non-AP1-containing regions of the TF promoter appear to be influenced by PModS actions on Sertoli cells.

To confirm that c-fos does not associate with the PModS-induced SE1 and SE2 gel shifts, an immunoblot with a c-fos antiserum of the gel shifts was performed (Fig. 8). Although a gel shift was detected with both SE1 and SE2, c-fos was not detected in the DNA-protein complex (Fig. 8). A positive control API gel shift did contain c-fos in its DNA-protein complex (Fig. 8). These observations imply that c-fos regulates an intermediate transcription factor(s) that influences the TF promoter at SE1 and SE2 (Fig. 9).

Discussion

The ability of the antisense c-fos oligonucleotide to inhibit PModS-induced activation of the TF promoter in Sertoli cells transfected with the −2.6-kb Tf-CAT reporter construct correlates with its ability to inhibit Tf protein secretion (6). These data demonstrated that c-fos is involved in PModS stimulation of Tf expression by cultured Sertoli cells. To further investigate the actions of PModS on a molecular level, the effects of PModS on both the c-fos and TF promoters were examined.

c-Fos promoter analysis

The responses of Sertoli cells treated with PModS and transfected with the reporter constructs containing the different regulatory elements of the c-fos promoter correlated with previous data obtained on the pharmacology of PModS. Treatment of cultured Sertoli cells with PModS did not affect cAMP levels (4), nor was it able to stimulate the reporter construct with the CRE of the c-fos promoter. The inability of PModS(S300) treatment to activate the TRE supports the observation that PModS does not stimulate either calcium mobilization or phosphoinositide turnover (5), both markers of protein kinase C activation. The inability of PModS to influence the TRE and AP1 (i.e. FAP) site at −290 bp of the c-fos promoter supports previous findings that PModS actions do not involve calcium mobilization or protein kinase C activation (5). Only the SRE and SIE were stimulated by treatment with the PModS(S300) and the highly purified PModS(C4) preparations. Interestingly, the presence of both SRE and SIE was not additive; the effect was the same or lesser than that of either element alone. Therefore, the ability of PModS to induce c-fos expression appears to be through the activation of SRE and SIE.

The SRE is a 29-bp region within the c-fos promoter. A 67-kilodalton protein has been isolated that binds to the SRE, the SRF (18). Evidence indicates that the binding of SRF to
SRE requires its phosphorylation (19–21). The actions of PModS require tyrosine phosphorylation events (5). The DNA-binding domain of SRF is the region necessary for dimerization and interaction with other proteins to form ternary complexes (18). Because SRF is ubiquitously expressed, cell-specific activation of the c-fos SRE by SRF appears to be mediated by other ternary complex protein factors. The binding of the ternary complex to the SRE requires the presence of the adjoining 5′-side of the SRE-binding site, the ETS domain binding motif. The protein kinase C-dependent and -independent pathways target different parts of this ternary complex (22). The expression and/or phosphorylation of these proteins may be involved in mediating cell-specific responses by the SRE. The SRE gel shift data presented demonstrate that PModS induced a gel shift similar to that of the positive control with serum treatment. Under less stringent conditions, a SRE gel shift was detected in control cells, but a more abundant gel shift was present in PModS-treated Sertoli cells (data not shown). The identity of the PModS-induced SRE-binding protein as SRF or other binding proteins remains to be elucidated. Interestingly, PModS induced a unique gel shift with the ETS-SRE that was not observed in control nontreated cells or in serum-treated cells. Therefore, PModS appears to promote the binding of a protein complex (e.g., ternary complex) that activates the SRE within the c-fos promoter. The formation of these ternary complexes may enable a common regulatory element, such as the SRE, to have cell-specific activity using ubiquitously expressed DNA-binding proteins. Alternatively, different or unique binding factors may be another method by which a common regulatory element may differentially activate cell-specific gene expression. In support of this proposal, several other DNA-binding factors have been identified that interact with the SRE. The 62-kilodalton SRE direct binding factor (p62DBF) may be related to the MAPF1 protein isolated from yeast (23). This factor is phosphorylated in vivo and mediates muscle-specific expression of α-actin. SRE-zinc-binding protein is a member of the C2H2 zinc finger family (24). High affinity binding is achieved through seven tandemly repeated zinc finger motifs. Rat NFIL-6 is a C/EBP-related factor whose binding to the SRE is mutually exclusive with SRF (25). Further analysis of activation of the SRE by PModS will investigate the potential involvement of known ternary complex factors, unique ternary complex factors, or novel SRE-binding proteins.

A second regulatory element that appears to be involved in PModS stimulation of c-fos is the SIE. It is located 25 bp upstream of the SRE. The SIE is activated by binding of the SIF, which has three forms, A, B, and C, and contains a phosphotyrosine residue (11). Treatment of human epidermoid carcinoma cells with epidermal growth factor and human hepatoma cells with interleukin-6 induces the binding of SIF-A, whereas treatment of epidermoid cells with interferon-γ-induced binding of SIF-C (11). This is also an example of the ability of different binding proteins to activate a common regulatory element and stimulate a gene-specific response. PModS induced a SIE gel shift that appears to involve predominantly SIF(A), but also had detectable SIF(B) and SIF(C) gel shifts. PModS was also found to activate a SIE reporter construct. Interestingly, the effects of both the SIE and SRE were not additive in response to PModS. PModS appears to influence the c-fos promoter at both the SRE and SIE. Whether the SIE can compensate and/or regulate the SRE response remains to be elucidated.

**Tf promoter analysis**

The downstream effects of PModS-induced c-fos expression were investigated with use of the Tf gene as a differentiated marker of Sertoli cells. The gene expression of Tf was inhibited by an antisense c-fos oligonucleotide, indicating that actions of PModS on the Tf promoter are indeed a downstream c-fos event. The regions controlling liver-specific transcription of the human Tf gene are composed of multiple positive and negative acting elements, mostly interacting with DNA-binding proteins present in either human or rat liver nuclear extracts (26, 27). The regulation of Tf gene expression in unstimulated Sertoli cells has been shown to be distinct from that of liver gene expression (28). Although the first 581 bp of the human Tf promoter (hTF) and mTF promoter have little similarity, they both contain a CRE site,
which was found to be responsive to FSH in the current study. The results with the CAT-mTf deletion mutants suggest that an enhancer activity is located between -2.6 and -1.6 kb for PModS and a repressor between -2.6 and -3.0 kb. Analysis of restriction enzyme fragments of the Tf promoter between -581 bp and -3kb identified two distinct domains, designated SE1 and SE2, that bind specific nuclear proteins from Sertoli cells stimulated with tMonds. Interestingly, these Sertoli response elements (*i.e.*, SE1 and SE2) are located within the apparent enhancer activity between 2.6 and -1.6 kb of the Tf promoter. Treatment with serum also induced a similar DNA-protein complex with SE1. The complex formed with SE2 was specific to PModS(S300), because treatment with serum or cAMP (data not shown) could not induce the gel shift. The lack of competition of SE1 and SE2 with API suggested that these sites were independent of API and the binding of a c-fos-jun complex. The c-fos immunoblot confirmed the absence of c-fos in the SE1 and SE2 DNA-protein complexes. Therefore, PModS induced Tf gene expression may involve both c-fos-dependent and -independent response elements. It is postulated that c-fos regulates the expression or activities of an intermediate transcription factor(s) that subsequently binds the SE1 and SE2 regions. The general mechanism is currently proposed, in that PModS, through activation of the SRE and SIE, induces the immediate early response gene c-fos, which then influences an intermediate transcription factor(s) that regulates downstream Sertoli cell differentiated functions such as Tf gene expression. Future studies will involve elucidation of the precise mechanism by which PModS activates the SRE and SIE of the c-fos promoter and Sertoli cell-specific response elements (*e.g.*, SE1 and SE2) of the Tf promoter. The possibility that Sertoli cell-specific trans-acting factors and ternary complex factors as well as unique response elements may be involved in the Sertoli cell-specific activation of these promoters is being investigated.

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