Regulation of thyrotropin receptor protein expression in insect cells

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ABSTRACT

Expression of large quantities of conformationally intact thyrotropin receptor (TSHR) is essential to understand the structure–function relationship of the receptor. We expressed three different constructs of full-length human TSHR in insect cells: (a) a TSHR cDNA lacking signal sequence (TSHR-ns), (b) a TSHR cDNA containing human TSHR signal sequence (TSHR-hs) and (c) a TSHR cDNA with baculovirus envelope protein encoded signal sequence gp-67 (TSHR-gp). No unique protein band, corresponding to any of these recombinant proteins, was visible upon Coomassie Blue staining after SDS-PAGE. However, Western blot using TSHR specific monoclonal antibody showed unique bands around 80, 100 and 100 kDa in TSHR-ns, TSHR-hs and TSHR-gp virus infected insect cells respectively. All three full-length TSHR proteins could neutralize the TSH binding inhibitory immunoglobulin (TBII) activity from sera of experimental animals. However, only glycosylated proteins (TSHR-hs and TSHR-gp) neutralized the TBII activity of sera from autoimmune thyroid patients, confirming the importance of glycosylation for patient autoantibody reactivity. Expression levels of full-length TSHR proteins were much lower than the levels of similarly produced corresponding ectodomains of TSHR proteins. Southern blot and Northern blot analyses showed that DNA and RNA levels in full-length TSHR virus infected insect cells were comparable to the levels found in cells infected with viruses encoding only the ectodomain of TSHR. These data suggest that full-length TSHR expression is very low and is regulated at the translational level.

INTRODUCTION

Thyrotropin receptor (TSHR) is the primary autoantigen in Graves’ disease and the disease is characterized by autoantibodies to TSHR (Nagayama & Rapoport 1992, Vassart & Dumont 1992, Kohn et al. 1995, Prabhakar et al. 1997). Large amounts of functional TSHR are essential to understand the structure–function relationship of the receptor as well as the pathogenesis of Graves’ disease. The full-length TSHR has been stably expressed in mammalian cells and binds TSH with high affinity (Nagayama et al. 1989, Perret et al. 1990, Chazenbalk et al. 1996) and is recognized by autoantibodies (Ludgate et al. 1990, Filetti et al. 1991, Tahara et al. 1991, Harfst et al. 1992a, Matsuba et al. 1995). However, the level of expression is low and is insufficient for purification. Moreover, attempts to overexpress full-length TSHR in insect cells have not been successful (Harfst et al. 1992a, Huang et al. 1993, Seetharamaiah et al. 1993, Misrahi et al. 1994, Chazenbalk & Rapoport 1995). In contrast, ectodomain of the TSHR (ETSHR) has been produced in large quantities using bacterial (Takai et al. 1991, Harfst et al. 1992b, Huang et al. 1992, Costagliola et al. 1994, Graves et al. 1995), baculovirus (Huang et al. 1993, Seetharamaiah et al. 1993, Chazenbalk & Rapoport 1995, Vlase et al. 1995, Seetharamaiah et al. 1997) and mammalian expression systems (Harfst et al. 1992c, Shi et al. 1993, Rapoport et al. 1996, Osuga et al. 1997, Costagliola et al. 1998, Da Costa & Johnstone 1998). In this report, we describe our efforts to express the full-length human TSHR in insect cells. We studied the effects of glycosylation of TSHR on autoantibody reactivity. To understand the basis for
the differential expression, we compared DNA and RNA levels in insect cells infected with recombinant virus encoding full-length TSHR and ETSHR. Our data strongly suggest that the level of expression is regulated at the post-transcription level.

MATERIALS AND METHODS

Source of human sera

Sera were collected from patients with either primary myxedema (p101 and p103) or Graves’ disease (p113–126). Patients were diagnosed using appropriate standard clinical and laboratory criteria. Sera were also obtained from healthy normal volunteers with no family history of thyroid autoimmunity. Sera were stored at −20 °C until use.

Cell culture

Sf9 cells (Gibco BRL, Gaithersburg, MD, USA) were maintained in TNM-FH medium (Sigma, St Louis, MO, USA) supplemented with 10% fetal bovine serum (Sigma) and antibiotic/antimycotic agents.

Generation of different recombinant baculoviruses

Figure 1 shows the different TSHR constructs used for the production of recombinant viruses. Construction of recombinant viruses encoding three different ectodomains of human (h) TSHR (ETSHR, TSHR-E and ET-gp) has been previously described (Seetharamaiah et al. 1997). For this study, we constructed three additional recombinant viruses, described below:

(a) TSHR-ns/PVL 1393: a full-length human TSHR cDNA lacking a signal sequence (TSHR-ns) was generated as follows. A cDNA encoding ETSHR lacking the signal sequence, representing nucleotides 64 to 1248, (Seetharamaiah et al. 1993) was digested with SspI to obtain 167 bp BamH1-SspI 5’ fragment. Similarly, 2474 bp full-length human TSHR cDNA (Tahara et al. 1991) was digested with SspI to obtain a 2144 bp SspI-EcoRI 3’ fragment. The 167 and 2144 bp cDNA fragments were subcloned into the BamH1–EcoRI site of the baculovirus transfer vector PVL 1393 (Pharmingen).

(b) TSHR-hs/PVL 1393: a full-length human TSHR cDNA including human TSHR signal sequence and 100 bp 5’ upstream sequence (Tahara et al. 1991) (2474 bp) was subcloned into the EcoRI site of the baculovirus transfer vector PVL 1393 (Pharmingen).

(c) TSHR-gp/pAcGP67B: a full-length human TSHR cDNA lacking the signal sequence, generated as described above, was subcloned into the BamH1–EcoRI site of the baculovirus transfer vector pAcGP67B (Pharmingen), which contains the baculovirus-encoded signal sequence gp67. This construct allows expression of TSHR as a fusion protein containing gp67 signal peptide.

Different recombinant viruses were generated by co-transfecting Sf9 cells with 0.5 µg of linearized Baculgold viral DNA (Pharmingen) and either 2 µg of TSHR-ns/PVL 1393 DNA, TSHR-hs/PVL 1393 DNA or TSHR-gp/pAcGP67B DNA, according to the manufacturer’s protocol. Media containing the recombinant viruses (TSHR-ns, TSHR-hs and TSHR-gp) were harvested after 5 days. The recombinant viruses were further amplified twice to obtain high titered viruses.

Expression of recombinant proteins

Monolayers of Sf9 cells were infected with different recombinant viruses. After 3–4 days, cells were harvested and lysed by vortexing in lysis buffer...
Neutralization of TSHR antibodies

A modified TSH binding inhibitory immunoglobulin (TBI) assay was used to test the ability of TSHR proteins to neutralize TSHR antibodies (Seetharamaiah et al. 1997, 1999). Rabbit antisera (50 µl) raised against ETSHR or ET-gp protein or ETSHR-derived peptide p367 (amino acids 22–41) for 2 h at room temperature before staining the Western blot. For large-scale protein production, Sf9 cells were grown in suspension cultures and processed as described previously (Seetharamaiah et al. 1993).

Titrations of recombinant virus

Virus titer was determined by end-point dilution as described by Summers and Smith (1987). Briefly, Sf9 cells were plated onto 96-well plates (2 × 10⁶ cells/well). After 2 h, different recombinant viruses were added to wells at dilutions ranging from 10⁻¹ to 10⁻⁸ and incubated at 26 °C for 7 days. Cells were monitored daily from day 2 for growth and infection. The highest dilution of virus at which at least 50% of the cells were infected was considered the end point.

Isolation and purification of total DNA and Southern blot analysis

The Sf9 cells infected with different recombinant viruses were used for the isolation of DNA at 48 h post infection. The culture media was removed and 5 ml of lysis buffer (0.03 M Tris–HCl pH 7.5, 0.01 M magnesium acetate and 1% NP-40) were added to the cells. The cells were kept on ice for 5 min with intermittent vortexing. The nuclei were pelleted at 2000 r.p.m./min, washed with cold PBS and repelleted. The pellet was resuspended in 4.5 ml of extraction buffer (0.1 M Tris pH 7.5, 0.09 M EDTA and 200 mM KCl) containing 200 µg/ml proteinase K and incubated at 50 °C for 1 h. The incubation was continued for another 2 h after the addition of 0.5 ml of 10% sarkosyl. The DNA containing aqueous phase was extracted twice with phenol/chloroform/isoamyl alcohol and precipitated with 10 ml of absolute ethanol. The DNA was pelleted at 2500 r.p.m. for 20 min and washed with cold 90% ethanol. The pellet was suspended in 0.1 M Tris-HCl pH 7.5, 5 µg/ml aprotinin and 0.5 µg/ml leupeptin, 5 µg/ml aprotilin and 0.5 mM phenyl-methyl-sulphonyl fluoride. The pelleted DNA was pelleted at 2500 r.p.m. and resuspended in 1 ml of salmon sperm DNA and 50% formamide). 1 kb cDNA encoding ETSHR was labeled with [α⁻³²P]dCTP by using prime labeling kit (Promega, Madison, WI, USA), added to the blot (5 × 10⁶ c.p.m./ml), and was hybridized for 16 h at 42 °C. The blot was then washed twice in 2 × SSC (1 × SSC in 0.15 M sodium chloride plus 0.015 M sodium citrate)–0.1% SDS at room temperature for 20 min and twice in 1 × SSC–0.1% SDS at 65 °C
for 20 min. The blots were exposed to Kodak X-AR film at −80 °C.

**Northern blot analysis**

Total RNA was isolated from Sf9 cells infected with different recombinant viruses using RNeasy kit (Qiagen, Santa Clarita, CA, USA) as recommended by the manufacturer. Ten micrograms of each RNA preparation were analyzed by electrophoresis on a 1.2% formaldehyde gel, stained with ethidium bromide, and photographed. The RNA was then transferred to a nylon membrane (S & S) by capillary method. The prehybridization, hybridization and autoradiography were carried out as described above for Southern analysis.

**RESULTS**

**Expression of TSHR proteins**

Expression of recombinant proteins was assessed by SDS-PAGE and Western blot analysis. Different recombinant virus infected insect cells were lysed in a lysis buffer, solubilized in SDS buffer and subjected to SDS-PAGE. As reported previously (Seetharamaiah et al. 1993, 1997), we observed major protein bands around 50 and 63 kDa in ETSHR and ET-gp virus infected insect cell extracts respectively (Fig. 2A, lanes 1 and 3). There was no apparent unique protein band corresponding to either TSHR-E (ectodomain with human signal sequence) or any of the full-length TSHR proteins. On an immunoblot using an ETSHR-specific monoclonal antibody (mAb 28) (Seetharamaiah et al. 1995) the 50 and 63 kDa proteins were identified as non-glycosylated and glycosylated forms of ETSHR protein respectively (Fig. 2B, lanes 1–3). Although protein bands corresponding to TSHR-E, and full-length TSHR proteins were not visible on the Coomassie Blue stained gel, they could be readily detected on a Western blot (Fig. 2B, lanes 2 and 4–6). A unique band, around 80 kDa, was detected on Western blot of cells infected with recombinant virus containing full-length TSHR without any signal sequence (TSHR-ns) (Fig. 2B, lane 4). The observed molecular mass of 80 kDa agrees closely with the expected size based on the predicted amino acid sequence (amino acids 22–764). Similarly, cells infected with virus containing full-length TSHR with either human or gp67 signal sequence showed a doublet around 95–100 kDa (Fig. 2B, lanes 5 and 6 respectively). This higher molecular mass of TSHR-ns and TSHR-gp proteins is most likely due to glycosylation of the protein. The reactivity of recombinant proteins in Western blot was specific and could be completely removed by pre-incubating mAb 28 either with the ETSHR protein or with an ETSHR-derived synthetic peptide (amino acid residues 22–41) (not shown).

![Figure 2](https://example.com/figure2.png) **FIGURE 2.** Expression of the human TSHR protein. Proteins from recombinant virus infected Sf9 cells were separated on a 10% SDS gel and stained with Coomassie Blue (A and D) or subjected to Western blot analysis using ETSHR-specific mAb 28 (B and E) or subjected to Western blot analysis using control mAb specific for Pemphigus vulgaris antigen (PVA) (C and F). For Western blot, significantly lower concentrations of ectodomains were used to obtain band intensities comparable with those of full-length TSHR proteins. In (A), (B) and (C), lanes 1–6 represent ETSHR, TSHR-E, ET-gp, TSHR-ns, TSHR-ns and TSHR-gp respectively. In (D), (E) and (F), lanes 1 and 2 represent ET-gp and control antigen PVA respectively. The left lane in each panel (M) represents molecular mass markers.

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TABLE 1. Inhibition of TBII activity. Serum samples (50 μl) from rabbits immunized with ETSHR protein, or with peptide 367 (amino acids 367–386) or with ET-gp or from patients with hypothyroidism (p101 and p103) or hyperthyroidism (p113–p126) were pre-incubated with 5 × 10⁶ insect cells producing either TSHR-ns, TSHR-hs, TSHR-gp or wild type (WT) polyhedrin protein and then tested for TBII activity in a radioreceptor assay. TBII values represent mean of duplicate determinations for each serum sample. A total of 10,000 c.p.m. of ¹²⁵I-bTSH was used per assay sample. Specific mean c.p.m. values for normal rabbit serum and normal human serum were 3350 and 3825 respectively. Specific mean c.p.m. values for test samples are given in parentheses.

<table>
<thead>
<tr>
<th>Serum sample</th>
<th>% TBII (c.p.m.)</th>
<th>% TBII (c.p.m.) after incubation with</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>anti-TSHR-ns</td>
<td>TSHR-hs</td>
</tr>
<tr>
<td><strong>Experimental</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anti-ETS-1</td>
<td>52·2 (1601)</td>
<td>2·8 (3694)</td>
</tr>
<tr>
<td>Anti-ETS-2</td>
<td>41·4 (1963)</td>
<td>−1·8 (3410)</td>
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<tr>
<td>Anti-ETS-3</td>
<td>39·1 (2040)</td>
<td>4·9 (3185)</td>
</tr>
<tr>
<td>Anti-ET-gp</td>
<td>85·3 (492)</td>
<td>15·1 (2844)</td>
</tr>
<tr>
<td>Anti-ET-gp2</td>
<td>80·5 (653)</td>
<td>11·0 (2981)</td>
</tr>
<tr>
<td><strong>Patients</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P101</td>
<td>84·3 (600)</td>
<td>85·1 (570)</td>
</tr>
<tr>
<td>P103</td>
<td>74·1 (991)</td>
<td>70·6 (1124)</td>
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<td>P113</td>
<td>30·6 (2654)</td>
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<td>P114</td>
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<td>P117</td>
<td>75·3 (945)</td>
<td>69·7 (1159)</td>
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<td>P118</td>
<td>50·3 (1901)</td>
<td>44·9 (2107)</td>
</tr>
<tr>
<td>P119</td>
<td>81·5 (707)</td>
<td>78·8 (810)</td>
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<td>P120</td>
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<td>22·8 (2953)</td>
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<td>P123</td>
<td>41·6 (2233)</td>
<td>39·5 (2314)</td>
</tr>
<tr>
<td>P124</td>
<td>48·0 (1989)</td>
<td>45·6 (2080)</td>
</tr>
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<td>P125</td>
<td>52·1 (1832)</td>
<td>54·8 (1729)</td>
</tr>
<tr>
<td>P126</td>
<td>85·2 (566)</td>
<td>87·4 (482)</td>
</tr>
</tbody>
</table>

shown). The specificity of reactivity of TSHR recombinant proteins with mAb 28 was further confirmed when mAb 28 reacted only with ET-gp (Fig. 2E, lane 1) but not with baculovirus-expressed Pemphigus vulgaris antigen (PVA) (Fig. 2D and E, lane 2) (Memar et al. 1996). Similarly, the mAb specific for PVA reacted with 130 kDa PVA (Fig. 2D and F, lane 2) and not with TSHR (Fig. 2C, lanes 1–6 and 2F, lane 1).

Reversal of TBII activity of experimental and patients’ sera

Earlier we had shown that all three ectodomains could neutralize the TBII activity of experimental sera, but only TSHR-E and ET-gp, the glycosylated ectodomains, could neutralize TBII in patients’ sera (Seetharamaiah et al. 1997). Therefore, to see whether glycosylated full-length protein is required for patient autoantibody reactivity, we tested the ability of full-length TSHR proteins to reverse the TBII activity of experimental antibodies in a modified RRA. As shown in Table 1, pre-incubating the α-ETS-1 or α-ET-gp or an α-peptide (amino acids 367–386) sera with either TSHR-ns, TSHR-hs or TSHR-gp protein resulted in almost complete neutralization of the antibodies and prevented them from blocking TSH binding to the TSHR. Encouraged by these results, next we tested these recombinant proteins for their ability to neutralize autoantibodies to TSHR in the sera of hypothyroid and hyperthyroid patients. As shown in Table 1, TBII activity remained unchanged in all 13 sera when they were pre-incubated with the non-glycosylated TSHR-ns protein or with proteins from control cells infected with wild-type baculovirus. In contrast, the glycosylated TSHR-hs and TSHR-gp proteins significantly neutralized the TSHR autoantibodies in 12 of the 13 sera tested. These data further confirmed earlier studies that glycosylation of the human TSHR is important for autoantibody reactivity (Rapoport et al. 1996, Seetharamaiah et al. 1997).

Replaccative ability of different recombinant viruses

Although we were successful in expressing full-length TSHR protein, the level of expression was very low compared with the levels of expression of
corresponding ectodomains of TSHR protein. Therefore, to see whether differences in replication potential could explain the differences in the level of protein expression, we tested the replicative ability of all six TSHR recombinant viruses. The titers of different recombinant viruses were comparable and ranged from $10^{-3}$ to $10^{-4}$. To compensate for small differences in viral titers, insect cells were infected with different recombinant viruses with the same multiplicity of infection (i.e. MOI of 1). Southern blot analysis, at 48 h post-infection, showed expected size DNA of almost equal intensity from all recombinant virus infected insect cells (Fig. 3) with the exception of cells infected with full-length TSHR cDNA containing human TSHR signal sequence (Fig. 3, lane 5).

Next, to see whether differences in the efficiency of transcription could account for different levels of protein expression, we carried out Northern blot analysis. Northern blot analysis of RNA from Sf9 cells infected with different recombinant viruses encoding extracellular domain of TSHR showed similar amounts of TSHR transcripts at all time points (Fig. 4A). However, cells infected with TSHR-hs virus showed consistently lower amounts of RNA compared with TSHR-ns and TSHR-gp virus infected insect cells at all time points (Fig. 4C). This is consistent with the lower levels of DNA seen in TSHR-hs virus infected cells.

**DISCUSSION**

A wide range of studies on thyroid require large quantities of relatively pure, functional TSHR. Because of lower levels of TSHR expression on thyrocytes, it has not been possible to purify the native receptor. Therefore, many laboratories, including our own, have attempted to generate recombinant TSHR protein (Takai et al. 1991, Harfst et al. 1992a,b,c, Huang et al. 1992, 1993, Seetharamaiah et al. 1993, 1997, Shi et al. 1993, Costagliola et al. 1994, Misrahi et al. 1994, Chazenbalk & Rapoport 1995, Graves et al. 1995, Matsuba et al. 1995, Vlase et al. 1995, Rapoport et al. 1996, Osuga et al. 1997). Unfortunately, to date, all efforts to produce high levels of full-length recombinant TSHR protein, using different expression systems, have failed (Harfst et al. 1992a, Huang et al. 1993, Seetharamaiah et al. 1993,

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**FIGURE 3.** Southern blot analysis of DNA from recombinant virus infected Sf9 cells. DNA was isolated, electrophoresed, transferred onto nylon membrane and probed as described under Materials and Methods. The autoradiogram was developed after exposure for 4 h. Lanes 1–6 represent ETSHR, TSHR-E, ET-gp, TSHR-ns, TSHR-hs and TSHR-gp respectively.

**FIGURE 4.** Northern blot analysis of RNA from recombinant virus infected Sf9 cells. Total RNA was isolated, electrophoresed, transferred into nylon membrane and probed as described under Materials and Methods. The enzymes used to digest the DNA are BamHI for ETSHR and ET-gp; EcoRI for TSHR-hs; and BamHI and EcoRI for TSHR-E, TSHR-ns and TSHR-gp.

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Misrahi et al. 1994, Chazenbalk & Rapoport 1995). In this study, like in an earlier study (Misrahi et al. 1994), we were able to express low levels of full-length hTSHR using a baculovirus expression system. Unlike the full-length protein, we could readily produce relatively higher levels of ectodomains of TSHR. These studies confirmed our earlier observation that lack of signal sequence or replacement of human signal sequence with gp67 signal sequence could lead to enhanced production of ectodomain of TSHR (Seetharamaiah et al. 1997). Therefore, we expressed full-length receptor protein either without signal sequence or with gp67 signal sequence. Neither of these modifications resulted in enhanced full-length protein production. Even using a sensitive Western blot staining the protein cannot be detected earlier than 48 h post-infection and the mRNA levels reached significant levels by 36 h post-infection. This suggests that the transcripts appear at least 12 h before we can detect the protein. As shown in Fig. 2, full-length TSHR protein could be detected around 48–72 h post infection, and the protein expression did not increase after 3 days. Glycosylated full-length TSHR proteins, although produced at low levels, were able to neutralize autoantibodies to TSHR in the sera of patients with autoimmune thyroid disease.

Because of very low levels of full-length protein expression, we wondered whether there were differences in the replicative ability of different recombinant baculoviruses. We determined the titer of different viruses at different intervals after infection and found that there were no significant differences. Next, we looked at the level of viral DNA and found that there was no correlation between the levels of viral DNA and protein. For example, all cells infected with recombinant viruses encoding different ectodomains exhibited similar levels of DNA and yet TSHR-E expression was lower. Similarly, the levels of TSHR-gp and TSHR-ns DNA were high relative to TSHR-hs and yet none of them was expressed at a higher level. These results indicated that differences in the DNA levels may not account for differences in the amounts of protein.

Level of protein expression is often reflected by the amount of mRNA present in the cell. Messenger RNA level is regulated either by the rate of transcription or by the stability of mRNA. To see whether differential regulation of TSHR specific transcripts could be responsible for differences in the levels of expression of full-length TSHR and their corresponding ectodomains, we performed Northern blot analysis. Northern analyses showed that there were considerable amounts of transcripts at all times in all cells irrespective of the recombinant virus used for infection. The levels of mRNA from full-length TSHR-expressing cells were comparable to the levels of mRNA in cells expressing corresponding ectodomains at all three time points, i.e. 36, 48 and 60 h post-infection, and still the full-length protein expression remained low. In contrast, ectodomain proteins reached maximal levels by 60–72 h. Our data showed that, in spite of stable mRNA levels, the levels of full-length TSHR protein expression did not increase over time. These data suggested that the protein expression is regulated at a step subsequent to transcription.

Length of non-coding upstream sequences can affect the level of protein production (Matsura et al. 1987). However, based on our current results it is unlikely that 100 bases of 5’ upstream non-coding sequence found in TSHR-hs significantly affected the protein production. This suggestion is based on the observation that there were no differences in the level of TSHR expression among different full-length TSHR constructs with or without the upstream sequence. More recently, studies using cell-free translation system have clearly shown that the length of poly(A) tail can have significant influence on translation by recruiting ribosome and allowing translation of uncapped mRNA (Gallie 1991, Iizuka et al. 1994, Tarun & Sachs 1995, Sachs et al. 1997, Preiss & Hentze 1998). These observations are consistent with observations in vivo where cap structure and poly(A) have been shown to have synergistic effect. We could rule out the influence of poly(A) tail on full-length TSHR protein production, since we used only cDNAs for protein expression.

In eukaryotic cells, cap structure at the 5’ end of the mRNA determines the amount and quality of the protein produced (Preiss & Hentze 1998). This is primarily accomplished by enhancing the translational initiation frequency and recruiting 40S ribosomal subunit to sites that are proximal to the 5’ end of the mRNA. However, some mRNAs contain internal ribosomal entry sites (IRESs) to which ribosomes can bind and influence translation (Pelletier & Sonenberg 1988). This occurs even when the 5’ is either blocked or uncapped. The cap structure may not account for differences in the levels of protein production because the 5’ ends of cDNAs are identical in corresponding ETSHR and full-length TSHR constructs. Again the 5’ ends of cDNAs used for generating all six recombinant viruses are identical and, therefore, differences in IRES, if any, should reside in the 3’ end of the cDNA. Another factor that controls the translational efficiency is the mRNA secondary structure (Londei 1998). The eukaryotic
initiation factor (eIF), eIF4F, is a complex of three polypeptides, eIF4E, eIF4A and eIF4G and facilitates the ATP-dependent unfolding of the mRNA secondary structure (Londei 1998). The eIF4G acts as an adaptor, which couples the cap-recognition function of eIF4E to the helicase function of eIF4A. If any of these initiation factors do not function optimally, then complex RNA secondary structure will be maintained, thus negatively affecting the translational efficiency. At the present time, we do not know how the TSHR protein production is translationally regulated. Therefore, understanding what role any of these structures play in the translation of TSHR might help us not only to produce larger quantities of full-length protein but, more importantly, the regulation of TSHR expression.

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