Differential expression of human chorionic gonadotropin (hCG) glycosylation isoforms in failing and continuing pregnancies: preliminary characterization of the hyperglycosylated hCG epitope

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Abstract

Human chorionic gonadotropin (hCG) glycoforms change as pregnancy progresses. We have developed an antibody (B152) which can measure a hyperglycosylated early pregnancy isoform of hCG. This putative hyperglycosylated form of hCG arises very early in pregnancies and is rapidly replaced by an isoform that predominates for the remainder of the pregnancy. The profiles of these hCG glycoforms are measured as a ratio of values of two immunometric assays. The profiles of these ratios differ between pregnancies which persist and those which will experience early failure.

In this report, daily urine hCG isoform ratios from donor eggs (no exogenous hCG pretreatment), in vitro fertilization pregnancies were profiled and analyzed from the first day following embryo transfer (ET). Significant differences were found between continuing pregnancy and pregnancy loss throughout days 5–20 post-ET. When hCG isoform ratios were analyzed from the first day of detectable hCG, pregnancy loss could be predicted in the case of a single fetus both during the 5- to 10-day time segment (P=0·018) and the 10- to 15-day time segment (P=0·045). When single and multiple fetus pregnancies were analyzed together significance was approached in the 10- to 15-day time period (P=0·058).

In a second population of pregnant women who conceived naturally, in whom urine samples were collected at approximately weekly intervals to either term birth or clinical spontaneous abortion, the ratio could discriminate between miscarriages and normal term pregnancies (P=0·043). In later pregnancy, the ratio of hCG isoforms declined more rapidly in miscarriages than in term pregnancy.

Antibody B152 was produced using a choriocarcinoma-derived hCG (C5), which was hyperglycosylated at both N- and O-linked sites and was 100% nicked at position β47–48. Western blot analyses supported the assay results showing that early pregnancy urine does not contain nicked C5-like hCG. Also, the early pregnancy hCG appeared to be the same size as later pregnancy hCG as judged by SDS gel electrophoresis. A series of Western blot analyses and immunoassays conducted with the samples either non-reduced or reduced showed that B152 is directed to a linear epitope located in the COOH-terminal peptide region of the β subunit. This indicated that only the O-glycan groups and not the N-linked glycans are part of the antibody epitope.


Introduction

We have previously documented that the expression of various urinary human chorionic gonadotropin (hCG) molecular forms differs between naturally conceived pregnancies, which carry to term, and those destined for early pregnancy loss (Elish et al. 1996, O’Connor et al. 1998). These differences encompass both the relative production of hCG metabolic fragments and the expression of hCG hyperglycosylated isoforms. We now report preliminary studies which indicate that the temporal pattern of these isoforms differ significantly between sustaining pregnancies and those which will fail.

This early pregnancy hCG isoform declines rapidly, being replaced by the presumably less highly branched hCG glycoform of a mature pregnancy (Kovalevskaya et al. 1997).
The replacement of the early hCG isoform(s) by those of later pregnancy is a continuous process. Both families of hCG isoforms appear to be present throughout pregnancy, but at substantially different concentrations. Since no one hCG assay is 100% specific for a particular hCG isoform we have developed a method to present the relative amounts of hCG isoforms as a ratio of hCG concentrations measured by two assays. One has primary affinity for early pregnancy hCG (B152-B207*, asterisk means radiolabeled detection antibody) and the other (B109-B108*) has primary affinity to the later isoform(s). We believe that the ratio of the two assays most accurately reflects the continuous transition of hCG isoforms as pregnancy progresses. The present paper expands these studies to characterize the expression of a family of putative hyperglycosylated hCG isoforms in in vitro fertilization (IVF) normal pregnancy and IVF early pregnancy loss (EPL) as well as throughout normal pregnancy compared with later miscarriages.

Although it was reported earlier that B152 is directed to carbohydrate and peptide backbone structure, characterization of its recognition patterns was not completed. For example, it appeared that both N-linked as well as O-linked carbohydrates may be involved in the recognition site (Birken et al. 1999). Furthermore, it was assumed that B152 was a conformation-directed antibody. In the following studies we report that B152 is directed to a linear epitope within the βCOOH-terminal region (CTP) of hCG or hCG free β subunit (hCGβ).

Materials and Methods

Patient samples

Set no. 1. Urine samples from patients undergoing ovum donation The program for oocyte donation was reviewed and approved by the Institutional Review Board and the Medical Ethics Committee of Columbia University, College of Physicians and Surgeons, Columbia-Presbyterian Medical Center.

Upon completion of required pre-screening examinations and matching with an approved donor, oocyte donors underwent ovarian hyperstimulation with human menopausal gonadotropins and were monitored by transvaginal ultrasound and serum estradiol levels. When follicle size reached 18–20 mm, hCG was given to trigger ovulation. Thirty-six hours later, transvaginal oocyte aspiration was performed under ultrasound guidance. Recipients were synchronized to an oocyte donor using a standard regimen consisting of oral micronized estradiol and intramuscular progesterone, as previously described (Sauer et al. 1995). Recipients with residual ovarian function underwent pituitary down-regulation with a gonadotropin-releasing hormone agonist to render them functionally agonal prior to initiating hormone replacement. Following fertilization, three to five embryos were transcervically transferred at 72 h to the recipient’s uterus. Pregnancy was confirmed by serum hCGβ 9 and 12 days following embryo transfer (ET). Prescribed hormonal replacement was continued through 12 weeks of gestation.

Twenty-two IVF donor egg recipients who underwent an ET without pretreatment with exogenous hCG (i.e. donor egg) collected urine every morning starting the day after ET. A total of 542 samples was collected and analyzed in this study. The ten patients who did not have detectable hCG (presumably because the embryo failed to implant into the endometrium) were excluded from the analysis. Twelve women were positive in the hCG assays (399 samples). Five of the twelve subjects had EPL, four had a singleton pregnancy and three had multiple fetuses (two or three).

For those twelve patients who had a positive value of hCG (>2 × least detectable dose) the range of the collection period was 7–48 days with a median collection duration of 34 days. One hCG-positive patient collected urine during days 1 to 20 and 39 to 60. The maternal age in this group of patients was in the range 33–50 years with a median of 42.8 years.

Two urine samples (50 ml each) from the same pregnant woman (43 years old) who underwent IVF procedure were used for immunoblotting. One sample was collected at the 31st day post-ET, the second at the 45th day post-ET.

Set no. 2. Normal pregnancy urine samples Two hundred and fifteen samples from ten normally conceived pregnancies were collected at an interval of 1–2 weeks throughout complete pregnancies (i.e. terminated in a live birth) and were compared with eleven miscarriages (37 samples). Sample collection started from weeks 3 to 9 (median 6-13 weeks) from the last menstrual period (LMP).

Urine samples were collected at Columbia-Presbyterian Medical Center under an Institutional Review Board-approved protocol.

Materials

Antibodies B152 and B151 were developed to a choriocarcinoma hCG (C5) (Birken et al. 1999), while B201 and B207 are directed to hCGβ core fragment (hCGβcf) and hCGβ (Krichevsky et al. 1988, O’Connor et al. 1988). CTP103, developed to synthetic peptide β residues 109–145 (Krichevsky et al. 1994a) was described earlier. Anti-mouse IgG antibody was from DAKO (Carpinteria, CA, USA). Sequencing grade modified trypsin (13 U/µg) was obtained from Promega (Madison, WI, USA). ECL Western blot reagents and Hyperfilm-ECL were purchased from Amersham International (Amersham, Bucks, UK). Polyvinylidene difluoride membrane (PVDF) and other materials related to electrophoresis and


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electrophoretetic transfer were obtained from Bio–Rad (Richmond, CA, USA). Other reagents were from Fisher (Pittsburg, PA, USA).

The non-nicked hCGβ (preparation 863), isolated from the CR127 preparation of hCG, was used in immunoblotting (Birken et al. 1993). Choriocarcinoma hCG C5 (designated 673 in this report), a 100% nicked hCG having extra carbohydrate moieties, purified from the urine of a choriocarcinoma patient (Elliott et al. 1993), was supplied by Dr L Cole (University of New Mexico, NM, USA). Hormones used for B151–B152* assay cross-reactivity characterization were: hCG C5 (preparation prep. 510), hCG C5 free β (prep. 544), hCG nicked (prep. 656), hCG CR129, hCGβ CR117, hCGβ cf (prep. 455), hCGα CR119.

Immunometric assays

Immunometric assay construction has been described previously (O’Connor et al. 1988). The methodology used to analyze urine samples has also been completely described (Kovaleskaya et al. 1999a). It is based on the combination of two hCG solid–phase immunometric assays: B109–B108* and B152–B207*. Intra-assay variation was 3% for B109–B108* assay and 5% for B152–B207* assay, inter-assay variation was 10% for B109–B108* assay and 11% for B152–B207* assay. Sensitivity (least detectable dose) defined as ±2 SD from the zero calibrator, was 1 fmol/ml for B152–B207* assay. Sensitivity (least detectable dose) for the B109–B108* assay, 4 fmol/ml for the B152–B207* assay and 11 fmol/ml for the B151–B152* assay.

Urinary non-nicked hCG was purified from the CR127 preparation of pooled normal pregnancy hCG and was used as a standard in both B109–B108* and B152–B207* assays. For the newly described B151–B152* assay, hCG C5 was used as a standard and 10 µg/ml B151 was used for microtiter plate coating.

Statistical analysis

Set no. 1. Urine samples from patients undergoing ovum donation For statistical analysis, the trajectory of the daily log-transformed B152/B109 ratio from the day following ET (first analysis) or from the 1st day of detectable hCG (second analysis) through the 30 succeeding days were compared between the IVF normal pregnancy group and the IVF EPL group using mixed–effects models (Laird & Ware 1982). Based on graphical inspection of outliers, mean ratio trajectories and their distributions, the ratio was log transformed to improve linear time trend and normality. The mixed–effects model was used because (a) heterogeneity of an individual trajectory can be modeled through a random slope representing the time trend of the ratio, (b) serial correlation within repeated measures of the log-ratio can be modeled, and (c) incomplete data due to pregnancy loss (i.e. unequal number of sampling points) can be used, thus avoiding any imputation and utilization of all available data. To capture the observed time trend, we divided the sampling period into time intervals (e.g. 1st to 5th days, 5th to 10th, 10th to 15th, 15th to 20th and so on), and employed the piecewise regression method (Montgomery & Peck 1982) so that the differential time trend within each time interval can be modeled. Significant interaction of the time trend for a given interval and experimental group implies a differential time trend between two groups within the time interval. Although the mixed–effects modeling approach allows us to model inter– and intrapatient variability based on incomplete data, we are aware of the limitation of statistical inference based on a relatively small sample size including lack of statistical power and generalizability of findings.

Set no. 2. Normal pregnancy urine samples Log-transformed measurement profiles (log B152/B109 ratio, logB152, logB109) were analyzed by the mixed–effects model consisting of random intercept, random slope against gestational age of subject, outcome (i.e. pregnant and spontaneous abortion) and interaction between time trend (i.e. slope) and outcome.

Partial purification of early pregnancy urine samples prior to SDS–PAGE

Two specimens (50 ml each) were obtained from pregnant woman at gestation days 31 and 45 from ET who underwent IVF procedures. The specimens were adjusted to neutral pH with 1 M Tris–HCl buffer (pH 8·0) and allowed to precipitate overnight at 4 °C. The supernatant was filtered through a 0·45 micron filter (Nalgene, Rochester, NY, USA) and concentrated in a 3000 cutoff Centricon ultrafilter YM–3 (Amicon, Danvers, MA, USA) to 3 ml. Each urine concentrate was desalted on a Sephadex G–25 column (Pharmacia, Piscatway, NJ, USA) in 0·05 M ammonium bicarbonate and then lyophilized. The dried powder was suspended in 1–2 ml column buffer (0·05 M ammonium bicarbonate) and then size-fractionated on Superose 12 (30 cm × 1 cm; Pharmacia) tandem columns (two in series). Fractions eluting where the hCG standard eluted were pooled and lyophilized. These fractions were used for the SDS gel electrophoresis and blots.

Separation of subunits of C5 (673)

Fifty micrograms hCG C5 were incubated in 4 M guanidine acetate, pH 4 for 2 h at 37 °C, and the dissociated subunits were separated on a Pharmacia Smart system using a uRP C2/C18 PC 3·2/3 column. Column buffers were 0·1% trifluoroacetic acid (TFA) in water (A) and the 0·1% TFA in 100% acetonitrile (B); flow rate was 0·24 ml/min; gradient: 0% B 10 min; 50% B at 40 min, linear...
gradient. The nicked β peak elutes prior to α and both peaks were separately pooled and dried.

Reduction and alkylation of hCGβ and choriocarcinoma hCG C5 free β subunit
Lyophilized hCG C5 free β subunit (20 µg) and hCGβ preparation 863 (87 µg) were reduced, alkylated, and separated on reverse phase HPLC as described earlier (Birken et al. 2001).

SDS-PAGE, Western blot and dot-blot analysis
SDS gels were run according to the procedure of Laemmli (1953). Proteins in the gel were transferred electrophoretically according to the directions provided by the manufacturer (Bio-Rad). Blots were visualized by autoradiography according to the manufacturer’s directions (Amersham ECL reagent and Kodak Hyperfilm–ECL).

Liquid phase radioimmunoassay competition studies
Liquid phase radioimmunoassay was conducted as described (Birken et al. 1999). Preparation of all competitor forms of hCG have been described earlier (Rosa et al. 1984, Puett & Birken 1989, Birken et al. 1991, 1993, 2000). Freshly made water solutions (calibrated in molarity by amino acid analysis) were diluted with carrier protein-containing buffer.

Protein determination
Hormone concentrations were determined by amino acid analysis on a Beckman 6300 according to the manufacturer’s instructions.

Creatinine
Urine hCG values were normalized to creatinine, measured in a 96-well microtiter plate format by a procedure adapted from Taussky (1954).

Table 1 Comparison of the log-transformed B152/B109 ratio time trend in IVF early pregnancy and IVF early pregnancy loss (EPL) urine samples (from the day after embryo transfer (ET))

<table>
<thead>
<tr>
<th>Time interval (days from the day after ET)</th>
<th>All IVF normal pregnancy vs IVF EPL</th>
<th>IVF normal pregnancy (single fetus) vs IVF EPL</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>F (df,df)</td>
<td>P value</td>
</tr>
<tr>
<td>5–10</td>
<td>4.28 (1,263)</td>
<td>0.0395</td>
</tr>
<tr>
<td>10–15</td>
<td>7.21 (1,263)</td>
<td>0.0077</td>
</tr>
<tr>
<td>15–20</td>
<td>6.37 (1,263)</td>
<td>0.0122</td>
</tr>
<tr>
<td>20–25</td>
<td>0.74 (1,263)</td>
<td>0.3900</td>
</tr>
</tbody>
</table>

df, degree of freedom.

Results

Analysis of daily urine collection from IVF patients (set no. 1)
The apparent values of hCG (normalized to creatinine) in the B152 assay were higher than in the B109 assay in both IVF normal pregnancy and IVF losses (Fig. 1). However, the proportion of each isoform presented as the B152/B109 ratio changed significantly both as a function of gestational age and pregnancy outcome. As shown in Fig. 1, in normally continuing IVF early pregnancy the B152/B109 ratio declined rapidly as pregnancy progressed (Fig. 1 insert C1). In contrast, the B152/B109 ratio exhibited a more shallow decline in IVF EPL (Fig. 1 insert D1). Table 1 presents the outcome of the statistical analysis of the B152/B109 ratio commencing with the day after ET. It was determined that, for both single fetus pregnancies and single and multiple fetuses combined, significant differences of the B152/B109 ratio were observed between the IVF normal pregnancy and IVF pregnancy loss group. These differences maintained significance throughout days 5–20 from the day after ET.

Upon assaying the urine samples which had been started from the day after ET we found that the day of first detection of a positive hCG value in urine varied. The mean day of the first detection was 6.75 days from ET (range was 5–10 days). When the time trend analysis commenced with the first day of detectable hCG, a statistically significant difference in the B152/B109 ratio between IVF normal pregnancy for both multiple and singleton fetuses analyzed together and IVF EPL was only approached (P=0.058) in the time interval of the 10th to the 15th days (Table 2). There was, however, a significant difference in the 5– to 10-day and the 10– to 15-day intervals between IVF normal pregnancy (single fetus) and IVF EPL.

There was no statistically significant difference in maternal age between the IVF normal pregnancy group (mean was 45.3 years) and the IVF EPL group (mean was 41.5 years) (P=0.242).
Analysis of weekly collections through normal pregnancy and spontaneous abortions (set no. 2)

The B152/B109 ratio profiles for naturally conceived normal pregnancy and miscarriages throughout pregnancy are presented in Fig. 2.

Statistical data analysis revealed a significant differential time trend between the profiles of normal pregnancies and spontaneous abortion on log (B152/B109 ratio) \( [F(1,62)=4·25 \quad P=0·043] \). The B152/B109 ratio declined with gestational age but the rate of decline was faster in the spontaneous abortion case than in normal pregnancy. Levels of the B109 hCG isoform became progressively higher in the normal pregnancy group as compared with the loss group \( [F(1,62)=6·89 \quad P=0·011] \).

Figure 1 hCG isoform profiles in IVF patients (normal pregnancy and EPL) urine. hCG was measured by the B152-B207* assay (A) and by the B109-B108* assay (B); hCG isoforms (C) and the B152/B109 ratio (insert C1) in normal IVF pregnancy; hCG isoforms (D) and the B152/B109 ratio (insert D1) in IVF EPL. Note that the ratio in IVF EPL (insert D1) has a shallow slope as compared with that of IVF continuing pregnancy (insert C1). crt, creatinine.

Construction, characterization and application of immunometric assay B151-B152* specific for C5 (choriocarcinoma hCG)

This immunoassay was constructed using monoclonal antibody B151 as a capture and monoclonal antibody B152 for detection. The B151 antibody recognizes the nicked epitope of C5 as well as the nicked epitope on hCG from normal pregnancy (Birken et al. 1999, Kovalevskaya et al. 1999b). The B152 antibody recognizes a hyperglycosylated epitope of C5 and a putatively similar structure present in an early pregnancy hCG isoform (Birken et al. 1999, Kovalevskaya et al. 1999a). The B151–B152* assay, which recognizes C5 choriocarcinoma hCG (nicked hyperglycosylated hCG), is quite specific. Cross-reactivity with nicked hCG is 9%, with hCG (CR129) it is 2%, with
hCG C5 free β subunit it is 1% and cross-reactivity with all other tested analytes (hCGβ, hCGβcf, hCGδ) is under 1%.

On the basis of the ratio of hCG isoforms detected by the B152-B207* and B109-B108* assays for intact hCG we found a similarity between early pregnancy hCG and choriocarcinoma hCG C5 (Kovalevskaya et al. 1999a). When we used B151 as a capture and analyzed early pregnancy urine samples in the B151-B152* assay we did not detect a C5-similar hCG to any appreciable extent. The conclusion was that early pregnancy hCG is not nicked, in contrast to C5 which is 100% nicked. Immunoblot experiments are described below which also support this finding.

**Characterization of the B152 epitope by Western blot analysis and by competitive liquid phase immunoassay**

Dot blots were used to compare four different antibodies by the relative intensity of immunostaining of reduced and non-reduced hCGβ and choriocarcinoma hCG C5 (Fig. 3). Two antibodies known to require a conformational epitope were compared with one antibody made to a synthetic peptide that was directed to a linear determinant in the β-COOH region (CTP). The conformational-directed antibodies, B201 and B207, were both generated to the hCGβcf, and both also recognize hCGβ (Krichevsky et al. 1988, 1994b). The linearly directed antibody CTP103 was made to a synthetic peptide.

### Table 2 Comparison of the log-transformed B152/B109 ratio time trend in IVF early pregnancy and IVF EPL urine samples (from the 1st day of hCG detection)

<table>
<thead>
<tr>
<th>Time interval (days from the first hCG detection)</th>
<th>All IVF normal pregnancy vs IVF EPL</th>
<th>IVF normal pregnancy (single fetus) vs IVF EPL</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>F (df,df)</td>
<td>P value</td>
</tr>
<tr>
<td>1–5</td>
<td>1·16 (1,214)</td>
<td>0·2819</td>
</tr>
<tr>
<td>5–10</td>
<td>2·57 (1,214)</td>
<td>0·1104</td>
</tr>
<tr>
<td>10–15</td>
<td>3·63 (1,214)</td>
<td>0·0582</td>
</tr>
<tr>
<td>15–20</td>
<td>1·88 (1,214)</td>
<td>0·1714</td>
</tr>
</tbody>
</table>

**Figure 2** B152/B109 ratio throughout pregnancy. Urine samples were collected from ten normally pregnant patients (215 samples) and from eleven patients with miscarriages (37 samples) weekly. Regression curve and 95% confidence interval are shown for normal pregnancy (r²=0.76) and for miscarriages (r²=0.57).

**Figure 3** Comparison of B152 immunoreactivity towards non-reduced and reduced forms of choriocarcinoma hCG C5 (673) and non-nicked hCGβ (863) with monoclonal antibodies CTP103, B201 and B207. R indicates reduced forms of 673 and 863.
composed of residues 109–145 of hCGβ and appears to bind this mobile region of hCG not visible on the crystal model of hCG (Krichevsky et al. 1994a, Lathorn et al. 1994, Wu et al. 1994). It is clear from Fig. 3 that B201 and B207 stain non-reduced hCGβ (863) and non-reduced hCG C5 (673) much better than reduced forms of these same molecules. The choriocarcinoma hCG C5 was used for the generation of B152 (Birken et al. 1999). In contrast, B152 and CTP103 bind much better to reduced hCGβ and reduced choriocarcinoma hCG C5 as compared with non-reduced forms. Thus, B152 most closely resembles the binding properties of antibody CTP103 which is known to be directed to a linear epitope.

**Examination of the isoforms recognized by B152 in early pregnancy urine**

Choriocarcinoma hCG C5 was reported to be hyperglycosylated at both its N- and O-linked carbohydrate moieties (Elliott et al. 1997). Extra carbohydrate makes this hCG isoform migrate slower in SDS gels than standard late first trimester pregnancy hCG. Likewise, if early pregnancy hCG isoforms are similarly hyperglycosylated, they too should migrate more slowly. All samples were boiled in SDS so we were visualizing dissociated β chains. Figure 4A shows that choriocarcinoma C5 hCGβ (from 673) is only slightly larger than the standard hCGβ (863) on these blots but that the 31st day hCGβ (rich in the B152 hCG isoforms as determined by measurement of a B152/B109 ratio of 3:4) as well as the 45th day hCGβ (lower in the B152 isoform as indicated by a B152/B109 ratio of 0:5) are both similar in migration position to standard mid-pregnancy hCGβ (863). Reduction of these samples by mercaptoethanol also did not show a migration difference between standard hCGβ and the forms of hCGβ in 31st day and 45th day gestation urine (Fig. 4B). It was noted on these blots of reduced samples that no nicked forms of β were stained except for the nicked β subunit present in choriocarcinoma hCG C5 (673). Reduction greatly increased the intensity of immunoblot staining of hCG isoforms, as shown by the dot blots of Fig. 3, making it difficult to discern differences in band positions of the reduced isoforms in Fig. 4.

The β subunit of nicked hCG, e.g. hCG C5, can be cleaved into two halves by simply reducing the disulfide bridges. Separation of reduced hCG C5 on SDS gels separates the N-glycosylated amino terminal half from the O-glycosylated COOH terminal half. In reduced hCG C5 (673), antibody B152 (Fig. 4B) as well as CTP103 (not shown) recognize a single band at approximately 20 kDa molecular weight, and we assumed this band is β_{48-145}. Preparation of purified, reduced and alkylated β_{48-145} confirmed its staining by B152. However, this does not prove that B152 can recognize the COOH-terminus (β_{115-145}) alone.

Therefore, we performed a series of liquid phase radioimmunoassay studies using radiolabeled hCG C5 as a tracer and a variety of hCG isoforms as competitors (Fig. 5). This study demonstrated that hCG C5 is the most potent competitor in liquid phase with itself as tracer as one would expect, since it is the immunogen for antibody B152. Free C5 β was the next most potent. The other reagents were derived from standard pregnancy hCG, which is not hyperglycosylated, resulting in their lower potencies when displacing labeled hCG C5 from B152.
were hCG, hCG C5, hCG C5

The existence of hCG glycoforms has been documented for some time (Fein et al. 1980, Wide & Hobson 1987, Skarulis et al. 1992, Nemansky et al. 1998). Most of this work was either based on isoelectric focusing patterns which are not feasible for general measurements or else by carbohydrate analysis on a few specimens. All current hCG immunoasays participate in the recognition of these isoforms to an undetermined extent since standard preparations of hCG glycoforms have not been available. We have developed the first immunometric system capable of quantifying the relative content of early pregnancy isoforms as compared with late first trimester hCG isoforms (Kovalevskaya et al. 1999a). Our methodology allows for the measurement of two isoform families recognized by their differential affinity for the antibodies used in the construction of two immunometric assays. One of the assays (B152–B207*) has primary affinity for the hyperglycosylated hCG isoform which predominates in very early pregnancy (first 6 weeks). The second assay (B109–B108*) has primary affinity for the hCG isoforms of later pregnancy. The ratio of values obtained in these two assays allows us to visualize an immunochemical profile of the progression of hCG isoforms throughout pregnancy.

As can be observed from Fig. 1C, the B152 hCG isoform rises rapidly in IVF early normal pregnancy, preceding the increase of the B109 hCG isoform. Figure 1(insert C1) presents the ratio of the two isoforms, showing the shift in the relative proportion of the two isoforms as gestation proceeds. Figure 1D presents the apparent hCG isoform concentrations in IVF EPL and Fig. 1 (insert D1) illustrates the B152/B109 ratio in IVF EPL. The original analysis of the data started from the day of ET (Table 1), and resulted in an improved discrimination over that when the data were analyzed from the first day of positive hCG detection in both assays (Table 2).

It is important to note that elimination of these first points in the latter approach had the effect of diminishing an observed difference between the IVF continuing pregnancy group and the IVF loss group – namely that the loss ratio profile (log ratio vs time) lags behind that of the continuing pregnancy group, i.e. if the loss group ratio is shifted forward by approximately 5 days the profiles assume greater similarity.

The observed decrease in the power of discrimination between the IVF pregnancy and IVF EPL groups consequent to the two statistical approaches (i.e. analysis from day of ET vs day of first hCG detection) appears to be, at least in part, due to the partial loss of the lag time observed in the IVF EPL group as compared with the IVF pregnancy group as a result of the normalization of the two profiles to the day of hCG detection. The profiles of the two groups (log ratio vs time) become roughly parallel when this frame shift is eliminated. It appears that the biological activity or transformation, perhaps cytotrophoblast cell differentiation, which is signified by the change in isoform ratio, can be at least partially accounted for by its occurrence at a later time in EPL gestations.

Discussion

hCG isoforms in early and later pregnancy urine

However, it is clear that hCGβ and reduced, carboxymethylated hCGβ (RCM β) which is a permanently denatured, linearized molecule, are equipotent, demonstrating that the B152 does not require a conformational epitope. In addition, the similar potencies of free β115–145 indicate that most or all of the B152 epitope is in this region. Removal of sialic acid from RCM β diminishes its potency somewhat but it can still compete, demonstrating that sialic acid is a minor component of the epitope of B152.

These studies by liquid phase competition were further supported by direct comparisons of calibrated reference solutions using the B152–B207* two-site assay (Fig. 6). The solutions, molarity calibrated by amino acid analysis, were hCG, hCG C5, hCG C5, hCGβ and desialylated hCG. The comparative binding curves agreed well with the liquid phase competition curves of Fig. 5 (except that reduced and alkylated and free CTP solutions could not be tested in the B152–B207* assay since they are not recognized by the detection antibody). hCG C5 is the highest affinity standard followed closely by its free β subunit. All other reference solutions bind with lower affinity and desialylated hCG binds very similarly to that of naturally sialylated hCG.

Figure 6 Binding curves for hCG-related molecules in the solid phase two-site immunometric assay B152-B207*.

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Figure 6 Binding curves for hCG-related molecules in the solid phase two-site immunometric assay B152-B207*.

However, it is clear that hCGβ and reduced, carboxymethylated hCGβ (RCM β) which is a permanently denatured, linearized molecule, are equipotent, demonstrating that the B152 does not require a conformational epitope. In addition, the similar potencies of free β115–145 indicate that most or all of the B152 epitope is in this region. Removal of sialic acid from RCM β diminishes its potency somewhat but it can still compete, demonstrating that sialic acid is a minor component of the epitope of B152.

These studies by liquid phase competition were further supported by direct comparisons of calibrated reference solutions using the B152–B207* two-site assay (Fig. 6). The solutions, molarity calibrated by amino acid analysis, were hCG, hCG C5, hCG C5, hCGβ and desialylated hCG. The comparative binding curves agreed well with the liquid phase competition curves of Fig. 5 (except that reduced and alkylated and free CTP solutions could not be tested in the B152–B207* assay since they are not recognized by the detection antibody). hCG C5 is the highest affinity standard followed closely by its free β subunit. All other reference solutions bind with lower affinity and desialylated hCG binds very similarly to that of naturally sialylated hCG.

Discussion

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Discussion

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The existence of hCG glycoforms has been documented for some time (Fein et al. 1980, Wide & Hobson 1987,
Our prior studies of hyperglycosylated hCG excretion throughout pregnancy (103 samples at irregular intervals) indicated an inflection point in the regression curve around 29 weeks of gestation (Kovalevskaya et al. 1999a, Fig. 2). The present study, with more samples (215 from ten subjects) at regular intervals, fails to show this inflection. The B152/B109 ratio decreases to low levels after 22 weeks of gestation and remains essentially constant until birth (Fig. 2).

Although significant differences in the behavior of the hCG isoform ratio between clinical spontaneous abortion and naturally conceived normal pregnancy were found, the pattern of change did not mimic or coincide with that of IVF EPL. This may indicate that different cellular factors controlling the expression of hCG isoforms are operative as gestation proceeds.

C5-like hCG in early pregnancy urine

In a previous study we reported that nicked hCG is not a significant constituent of early pregnancy urine (Kovalevskaya et al. 1999b). For that study we used the B151-B604* assay which recognizes nicked hCG originating from normal pregnancy hCG. Now we have constructed the B151-B152* assay which specifically recognizes choriocarcinoma hCG C5 (100% nicked). Both B151 and B152 antibodies were developed to choriocarcinoma hCG C5 (Birken et al. 1999). We applied this assay to IVF early pregnancy urine and did not detect hCG by this assay.

Comparison of results from the B151-B152* and the B152-B207* assays allows us to conclude that the similarity between early pregnancy hCG and choriocarcinoma hCG C5 is not due to nicking but to hyperglycosylation.

Epitope recognized by B152

We used B152 as a capture antibody in a two-site assay and a common, anti-native hCGβ antibody as the detection antibody. The detection antibody, B207, is clearly directed to a conformational determinant as shown by the dot-blot analysis (Fig. 3). B152 binds preferentially to reduced, denatured hCGβ (Fig. 3) and binds specifically to the COOH-terminal half of the hCGβ (Fig. 4). Liquid phase competition assays using radiolabeled hCG C5 as a tracer indicate that B152 is directed to a linear epitope since it binds to reduced and alkylated hCGβ. In liquid phase assay, B152 recognized hCGβ and reduced and alkylated hCGβ equivalently. This result differs from the blot analysis, which showed better recognition of the reduced hormone form. Presumably this is due to additional changes induced by binding of the reduced protein to the membrane surface. The liquid phase studies also showed that free β115–145 competed as well as hCGβ, proving that B152 is a linearly directed antibody with all or most of its binding within this CTP region.

Conformationally directed hCGβ antibodies did not bind to alkylated hCGβ to a significant extent in either liquid or solid phase immunonassays. Liquid phase radiomunnoassay showed that the extent of sialylation did not affect B152 binding significantly. These findings were further substantiated by examining binding of the amino acid analysis-calibrated hormone solutions by the B152-B207* two-site assay. The lack of the effect of sialic acid content was substantiated. Our earlier report demonstrated that carbohydrate was part of the B152-recognized site by the use of hCG isoforms which were not hyperglycosylated but identical in all other respects (Birken et al. 1999).

Antibody B152 was made to a choriocarcinoma form of hCG which exhibited both hyperglycosylated N- and O-linked carbohydrate moieties. In our earlier report it was not clear if both types of carbohydrates were involved in the B152 epitope and this determinant was presumed to be conformational (Birken et al. 1999). The studies reported here have shown that the determinant is linear and similar to that COOH-terminal part of hCGβ which is recognized by CTP antibodies, except for its preference for hyperglycosylated O-linked carbohydrate groups. (The O-linked glycans of hCGβ are located at serine residues 121, 127, 132, and 138.)

The O-linked glycosylation process appears altered in the trophoblast cells in early placenta, resembling the type of sugar synthetic pathway operative in trophoblastic malignancy. Although many types of cancerous cells tend to synthesize truncated O-glycans rather than more complex types, this is not the case for trophoblastic malignancies (Brockhausen 1999, Kobata & Takeuchi 1999). hCG-secreting malignancies display more complex glycans, some of which are not sialylated or reduced in sialic acid content compared with normal pregnancy hCG.

Acknowledgements

This work was supported by grant ES07589 from the NIEHS and the Office of Research on Women’s Health, NIH, grant RR00645 from NCRR, NIH, and grant AG13783, NIH. The authors express their appreciation to Dr Cole (University of New Mexico) for providing choriocarcinoma hCG C5. We gratefully acknowledge the contribution of Alana Shear and Rebecca Dean (Columbia-Presbyterian Medical Center) in the performance of IVF specimen collection.

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Chorionic gonadotropin (hCG), intact hCG, and hCG beta fragment from standard reference preparations and raw urine samples.

Endocrinology 133 1390–1397.


Received in final form 22 October 2001

Accepted 30 October 2001