

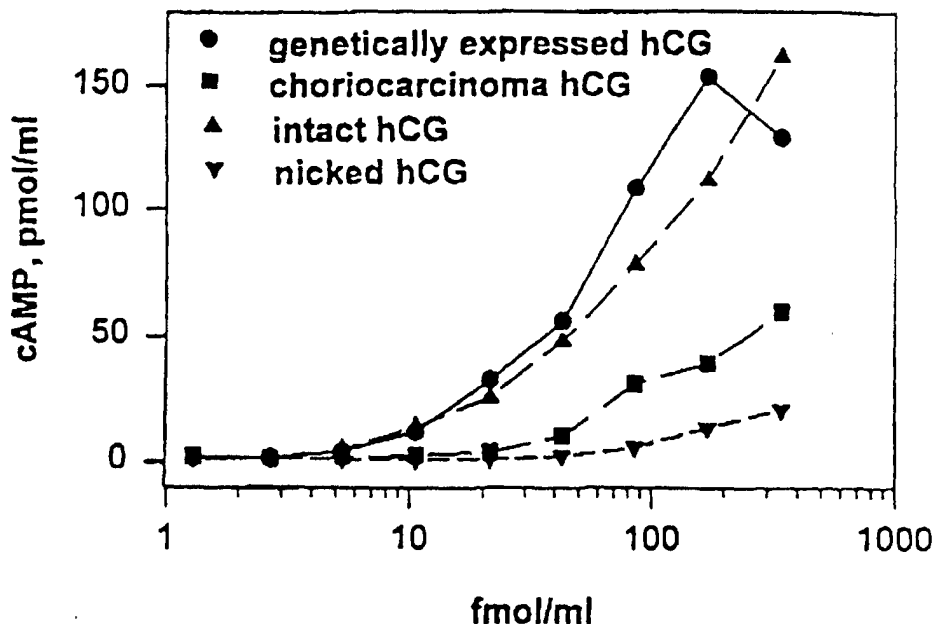


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(54) Title: METHODS FOR PREDICTING PREGNANCY OUTCOME IN A SUBJECT BY hCG ASSAY



## (57) Abstract

The present invention provides a method of predicting pregnancy outcome in a subject by determining the amount of an early pregnancy associated molecular isoform of hCG in a sample. The present invention further provides a method for determining the amount of early pregnancy associated molecular isoforms of human chorionic gonadotropin (hCG) in a sample. The present invention also provides a diagnostic kit for determining the amount of early pregnancy associated hCG in a sample. The present invention additionally provides an antibody which specifically binds to an early pregnancy associated molecular isoform of human chorionic gonadotropin. Finally, the present invention provides methods for detecting trophoblast or non-trophoblast malignancy in a sample.

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**METHODS FOR PREDICTING  
PREGNANCY OUTCOME IN A SUBJECT BY hCG ASSAY**

5 This application claims priority of and is a continuation-in-part of U.S. Serial No. 09/311,428, filed May 13, 1999, the contents of which are hereby incorporated by reference into this application.

10 The invention disclosed herein was made with United States Government support under National Institutes of Health Grant Nos. NIEHS ES-07589 and HD 15454. Accordingly, the U.S. Government has certain rights in this invention.

15 **Background of the Invention**

Throughout this application, various publications are referenced by author and date. Full citations for these publications may be found listed alphabetically at the end  
20 of the specification immediately preceding the claims. The disclosures of these publications in their entireties are hereby incorporated by reference into this application in order to more fully describe the state of the art.

25 Early pregnancy loss (EPL) is a widespread, but largely undiagnosed problem. In order to adequately diagnose and develop treatments for EPL it is essential to be able to detect and measure the rate of occurrence of EPL. This is critically important in epidemiological studies, some of  
30 which are related to exposures to known or suspected reproductive toxins in the workplace, in the environment or by personal use. These early pregnancy losses are often not recognized by women or physicians and are detected solely by the measurement of hCG in the urine at the time between  
35 implantation and expected menses. They are sometimes termed "chemical pregnancies" or "occult pregnancies." A landmark epidemiological study established that the incidence of EPL was 22% in a population of healthy women attempting to

conceive (Wilcox, A. J., et al., 1988). This investigation employed a very sensitive (0.01 ng/ml hCG) assay which detected only the intact hCG molecule with the unique beta subunit carboxyterminal peptide present.

5

There are multiple likely causes for EPL and clinical spontaneous abortion including genetic abnormality, immunological dysfunction, untreated infection or other unknown physiological problems. In addition, losses may be  
10 caused by failure of human chorionic gonadotropin (hCG) to induce adequate response at its target, the corpus luteum. This could result from inadequate hormonal potency. "Nicking" of the beta subunit in the loop 2 region of the molecule, specifically between residues 44-49, can reduce  
15 biopotency of hCG. Cleaved peptide bonds in this area of the molecule also exhibit reduced biopotency and reduced immunochemical recognition by monoclonal antibodies directed to the heterodimeric hormone (Cole, L. A., et al., 1991a; Cole, L. A., et al., 1991b; Puisieux, A., et al., 1990; Nishimura, R., et al., 1988; Nishimura, R. T., et al., 1989).  
20 Nicked forms of hCG were examined as possibly more prevalent in EPL situations and, at least in part responsible, for early pregnancy loss. Unfortunately many of the reports claiming that substantial concentrations of nicked hCG are  
25 produced during pregnancy, losses or successful pregnancies, are not accurate due to faulty assumptions regarding assay specificity (Wilcox, A. J., et al., 1988). Carbohydrate-modified hCG can also exhibit reduced biopotency. It is known that if the hCG has much reduced sialic acid content  
30 and its carbohydrate chains terminate in galactose, much hCG would be removed by the liver receptor for such altered glycoproteins (Braun, J. R., et al., 1996; Kawasaki, T. and G. Ashwell, 1996). The circulating life-time of asialo hCG is reduced and its *in vivo* potency is thereby low. Other  
35 carbohydrate changes also alter circulating half life;

glycoproteins terminating in sulfate-N-acetyl galactosamine are also extracted by a specific liver receptor and have reduced circulating lifetime (Baenziger, J. U., 1994; Fiete, D., et al., 1991).

5

At least two factors affect increased potency of hCG. First, it is known that a larger Stoke's radius will decrease clearance through the kidney glomerulus which generally clears proteins above an effective size of 70,000 very  
10 slowly. The effective size of urinary-isolated hCG is just at this borderline reduced clearance size. Generally, extra sugar content makes the hydrated radius of glycoproteins larger. It has been shown that by adding the hCG beta COOH-terminal peptide to hFSH or hLH, their circulating life-times  
15 greatly increased (Fares, F. A. et al., 1992; Matzuk, M. M., 1990). This addition was thought mostly due to the carbohydrate content of that peptide rather than simply the extra polypeptide size (Wilcox, A. J., et al., 1988).  
Second, increased negative charge of a protein will prolong  
20 its circulating time because of decreased renal clearance (Chmielewski, C. 1992, Quadri, K. H., et al., 1994; Maack, T., et al., 1985). This increased negative charge can be due to extra sialic acid or other negative groups, including sulfate such as is present on hLH and on the pituitary form  
25 of hCG (Birken, S., et al., 1996b). Changes which affect signal transduction at the receptor may also affect biopotency of hCG. It is known that deglycosylated hCG has much reduced receptor potency (Ravindranath, N., et al., 1992; Sairam, M. R., and L. G., Jiang, 1992; Browne, E. S.,  
30 et al., 1990; Sairam, M. R., 1989; Sairam, M. R., et al., 1988). Carbohydrate reduced forms of hCG also have reduced signal transduction (Amano, J., et al., 1990; Bahl, O. P., et al., 1995; Moyle, W. R., 1975).

35 According to the present invention EPL or recurrent spontaneous abortion is not due to an abnormal hCG form that

has reduced potency, such as nicked hCG. Instead, the present invention provides evidence that in successful outcome pregnancies women usually produce forms of hCG which are very highly potent in very early pregnancy; the standard 5 urinary reference preparations of hCG are less potent forms of the hormone produced later in pregnancy. The increased potency could be caused by a combination of factors from circulating half-life to increased receptor affinity or signal transduction or all of the preceding. Since hCG is 10 low very early in pregnancy, it is logical to find a more potent form of hCG on a molar basis to carry out its function until production levels rise as the trophoblastic cellular mass increases. The present invention describes molecular and immunological tools and methods including an antibody, 15 B152, described herein which recognizes the highly potent early pregnancy associated molecular isoforms of hCG. The determination of blood and urine profiles for the B152 hCG isoforms throughout healthy pregnancies can delineate the pattern of isoforms in successful pregnancies. These 20 isoforms can be measured by immunoassay alone, obviating the need to perform complex isoelectric focusing studies or other separation techniques. Additionally, the methods described herein are applicable to large numbers of samples.

**Summary of the Invention**

The present invention provides a method of predicting pregnancy outcome in a subject by determining the amount of an early pregnancy associated molecular isoform of hCG in a sample comprising: (a) contacting a sample with an antibody which specifically binds to the early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (b) measuring the amount of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample; and (c) comparing the amount early pregnancy associated molecular isoform of hCG in the sample determined in step (b) with either (i) the amount determined for temporally matched, normal pregnant subject(s) or (ii) the amount determined for non-pregnant subject(s), wherein the relative absence of the early pregnancy associated molecular isoform of hCG in the sample indicates a negative outcome of pregnancy for the subject.

20

The present invention further provides a method of predicting pregnancy outcome in a subject by determining the amount of an early pregnancy associated molecular isoform of hCG in a sample comprising: (a) contacting a capturing antibody which specifically binds to the early pregnancy associated molecular isoform of hCG with a solid matrix under conditions permitting binding of the antibody with the solid matrix; (b) contacting the bound matrix with the sample under conditions permitting binding of the antigen present in the sample with the capturing antibody; (c) separating the bound matrix and the sample; (d) contacting the separated bound matrix with a detecting antibody which specifically binds to hCG under conditions permitting binding of antibody and antigen in the sample; (e) measuring the amount of bound antibody on the bound matrix, thereby determining the amount of early

pregnancy associated molecular isoform of hCG in the sample;  
and (f) comparing the amount early pregnancy associated  
molecular isoform of hCG in the sample determined in step (e)  
with either (i) the amount determined for temporally matched,  
5 normal pregnant subject(s) or (ii) the amount determined for  
non-pregnant subject(s), wherein amounts of the early  
pregnancy associated molecular isoform of hCG in the sample  
similar to amounts of early pregnancy associated molecular  
isoform of hCG in temporally matched pregnant samples  
10 indicates a positive outcome, amounts of early pregnancy  
associated molecular isoform of hCG in the sample similar to  
amounts of early pregnancy associated molecular isoform of  
hCG in the non-pregnant samples indicates a negative outcome  
of pregnancy for the subject.

15

In addition, the present invention provides a method for  
determining the amount of early pregnancy associated  
molecular isoforms of in a sample comprising: (a) contacting  
the sample with an antibody which specifically binds to an  
20 early pregnancy associated molecular isoform of hCG under  
conditions permitting formation of a complex between the  
antibody and the early pregnancy associated molecular isoform  
of hCG; and (b) determining the amount of complexes formed  
thereby determining the amount of early pregnancy associated  
25 molecular isoform of hCG in the sample.

Further, the present invention provides a diagnostic kit for  
determining the amount of early pregnancy associated hCG is  
a sample comprising: (a) an antibody which specifically binds  
30 to an early pregnancy associated molecular isoform; (b) a  
solid matrix to which the antibody is bound; and (c) reagents  
permitting the formation of a complex between the antibody  
and a sample.

35 The present invention additionally provides an antibody which  
specifically binds to an early pregnancy associated molecular

isoform of human chorionic gonadotropin.

Further, the present invention provides a method for detecting non-trophoblast malignancy in a sample comprising:

5 (a) contacting a sample with an antibody which specifically binds to the early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (b) contacting the sample with a

10 second detection antibody which specifically binds to intact non-nicked hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (c) measuring the amount

15 of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample; and (d) comparing the amount of early pregnancy associated molecular isoform of hCG in the sample determined

20 in step (b) with the amount of early pregnancy associated molecular isoform of hCG in the sample determined in step (c), wherein a positive detection of early pregnancy associated molecular isoform detected in step (b) and a relative absence of the early pregnancy associated molecular isoform of hCG detected in step (c) indicates the presence

25 of non-trophoblast malignancy in the sample.

Finally, the present invention provides a method for detecting gestational trophoblast disease in a sample from a subject comprising (a) contacting a sample with an antibody

30 which specifically binds to the early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (b) contacting the sample with a second detection antibody which

35 specifically binds to intact non-nicked hCG without substantially cross-reacting with said antibody under

conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (c) measuring the amount of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample due to binding with the first antibody, and late pregnancy associated molecular isoform of hCG in the sample due to binding with the second antibody; (d) determining the ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated molecular isoform of hCG in the subject; and (e) comparing the ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated molecular isoform of hCG in the sample determined in step (c) over time, wherein a continuing high ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated molecular isoform of hCG in the sample determined in step (c) indicates the presence of gestational trophoblast disease in the subject.

**Brief Description of the Figures****Figure 1**

Bioassay for forms of hCG. This is data from recombinant CHO cells expressing the LH/CG receptor. The response factor is cAMP production. The x-axis is dose of one of four calibrated, pure hormones as described on graph legends. Expressed hCG has no nicks; choriocarcinoma hCG (C5) is 100% nicked; CR 127 was purified into a nick-free (non-nicked, intact) and nick-enriched fraction as shown.

**Figure 2**

Incidence (Panel A) and expression level (Panel B) of hCG-related molecules in the positive samples for each of the analyses measured (In early normal pregnancy, n=214; EPL cycles, n=49; and negative cycles, n=297).

**Figure 3**

Binding curves for three hCG types in the B152-B207\* assay (upper panel) and the B109-B108\* assay (lower panel).

**Figures 4**

Ratio of hCG isoforms measured by the B152-B207\* and B109-B108\* assays in normal pregnancy urine (n=103) at different gestational ages. (Regression curve and 95% confidence intervals are shown,  $r^2=0.79$ ). An inflection point in the curve occurs at approximately 29 weeks.

**30 Figures 5**

Box plot of the B152/B109 ratio for pregnancy matched serum/urine at 5-6 weeks of gestational age (n=12); or at 36-39 weeks of gestational age (n=11) and in JAR cell supernatant. Box extends to the 25<sup>th</sup> and 75<sup>th</sup> percentile. The upper and lower symbols indicate the

90<sup>th</sup> and 10<sup>th</sup> percentile respectively. A solid line inside the box marks the value of the 50<sup>th</sup> percentile.

**Figure 6**

5 Ratio of hCG isoforms measured by the B152-B207\* and B109-B108\* in urine of IVF patients (n=65). (Regression curve and 95% confidence intervals are shown,  $r^2=0.59$ ).

10 **Figure 7**

Immunoassay profiles of fractions from Superose 12 column chromatography of a pooled urine concentrate from pregnant women.

15 **Figure 8**

Liquid phase radioimmunoassays using antibody B151 (panel A) and antibody B152 (panel B). Radiolabeled choriocarcinoma hCG was used as tracer and calibrated (by amino acid analysis) solutions of pregnancy C5 chorioCG, hCG CR 127, hCG CR 127 non-nicked, and hCGn CR 127 were employed as competitors. Non-linear regression lines were plotted in logit transformed format.

25

**Figure 9**

Radio-Immunoassay (two-site) using antibody B152 as capture and B207 as radiolabeled detection reagent. Binding curves are shown for competitors, various competitor as detailed in methods and results. Each panel represents a separate assay in which all ligands were introduced in the same assay. Points were connected by straight lines although regression analysis (4 parameter logistic) indicated excellent fit to logistic or sigmoidal curve shape model. Panel A

30

35

and B represent two distinct assays with similar results. It is clear that this assay has greatest recognition of the nicked, choriocarcinoma hCG immunogen which is hyperglycosylated and binds similarly to nicked and non-nicked forms of hCG which contain the usual quantities of sugars. Reagent M1A is missing most of its beta COOH-terminal region, supporting a role of this region in the binding site of B152 (see panel B and discussion in text).

10

**Figure 10**

Enzymic-Immuno-metric (two-site) assay using antibody B152 as capture and peroxidase-labeled B4001 as detection antibody. A linear-linear plot of molar quantities of ligand added is plotted versus absorbance at 492nm which is the response factor from the peroxidase detection system. Eight different hormone forms were measured as ligands within the same assay as indicated in the legend. The two hyperglycosylated choriocarcinoma-derived hCG isoforms are both the most potent ligands (Figure 14). Potency correlated well with hyperglycosylation of ligand (see Figure 10 and text).

15

20

**25 Figure 11**

Characteristics of the Reagents Used to Define Antibody Specificity. The peptide and carbohydrate structures of the reagents used were determined earlier (26). The % nicked  $\beta$ -subunit refers to the proportion of molecules with cleavages (missing peptide bonds) in the region  $\beta$ 43 to  $\beta$ 48. The % tetrasaccharide core is the proportion of O-linked oligosaccharides with tetrasaccharide (vs. disaccharide) core structure, and the % sialic acid, is the proportion of O-linked structures with antennae terminated by sialic acid

30

35

residues. The proportion of triantennary N-linked oligosaccharides on  $\beta$ -subunit is given, as is the corresponding % sialic acid.

- 5       <sup>a</sup>.% sialic acid residues per sugar chain, N-linked on  $\beta$ .  
       <sup>b</sup>.% sialic acid residues per sugar chain, O-linked on  $\beta$ .  
       <sup>c</sup> The "CR" series of hCG reference preparations were made at Columbia University and were distributed internationally as reference materials for purified  
 10       hCG. CR 119 is also known as the 3<sup>rd</sup> international immunoassay reference preparation for hCG.  
       <sup>d</sup> ND is not done; NA is not applicable to that reagent.  
       <sup>e</sup> Less than 15% of the beta COOH-terminal region is present on this preparation.

15

**Figure 12**

Affinity Constants<sup>a</sup> Determined by Liquid Phase Competition Assays Using C5 as Tracer Ligand.

<sup>a</sup>Ka as L/M

- 20       <sup>b</sup>hCG CR 127 is an NIH-distributed hCG reference preparation produced at Columbia University.

**Figure 13**

- 25       Matrices of data for binding characteristics of different pairs of detection antibodies using B151 or B152 as capture antibody.

A. Relative Cross-Reactivities of Two Site Assay Using B151 as Capture Antibody

- 30       <sup>a</sup> labeled detection antibodies

<sup>b</sup> out of low range detection

<sup>c</sup> this particular assay format was applied in O'Connor et al (25).

35

B. Relative Cross-Reactivities of Two Site Assay Using B152 as Capture Antibody

The molar quantity of ligand required to produce binding equal to 50% of the maximum binding achieved by C5 was determined. Cross-reactivity shown in this figure as a percentage is calculated by dividing the molar quantity of the standard by the molar quantity of the other ligand at 50% maximum binding dose.

- 5
- 10
- 15
- \* labeled detection antibodies
  - † maximum binding represents the total quantity of radiolabeled detection antibody which can bind to the plate in the system described.
  - ‡ out of low range detection
  - § this particular assay format was applied in O'Connor et al (25).

**Figure 14**

20 Immunoreactivity of antigens in the B152 immunoradiometric assay. The dose-response curves used to provide data for this figure are shown in Figure 9. Each curve was fitted with 4-5 points. Slope and coefficient of determination ( $R^2$ ) were determined using a non-linear regression algorithm. Slopes were used as an indicator of antigen potency. Relative potency was estimated as the slope of antigens relative to the slope of C5 Choriocarcinoma hCG (the immunogen).

- 25
- 30
- \* Slope are from Figure 9 as calculated in Sigmaplot 4.01 by linear regression analysis. Units of slope are pmole/ml absorbance at 492nm.

**Figure 15 (A-D)**

35 Urinary isoform profiles and ratio in normal singleton pregnancies. ET day is a day from embryo transfer.

**Figure 16(A-C)**

Urinary isoform profiles and ratio in normal multiple birth pregnancy. ET day is a day from embryo transfer.

**5 Figure 17(A-E)**

Urinary isoform profiles and ratio in early pregnancy loss. ET day is a day from embryo transfer.

**Detailed Description of the Invention**

A method of predicting pregnancy outcome in a subject by determining the amount of an early pregnancy associated molecular isoform of hCG in a sample comprising: (a) 5 contacting a sample with an antibody which specifically binds to the early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (b) measuring the amount of complexes formed, 10 thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample; and (c) comparing the amount early pregnancy associated molecular isoform of hCG in the sample determined in step (b) with either (i) the amount determined for temporally matched, 15 normal pregnant subject(s) or (ii) the amount determined for non-pregnant subject(s), wherein the relative absence of the early pregnancy associated molecular isoform of hCG in the sample indicates a negative outcome of pregnancy for the subject. In an embodiment of the present invention, the 20 antibody is B152. Another embodiment of this invention is the early pregnancy associated molecular isoform of Hcg.

The hybridoma producing the B152 monoclonal antibody was deposited on February 3, 1998 with the American Type Culture 25 Collection (ATCC), 12301 Parklawn Drive, Rockville, Maryland 20852, U.S.A. under the provisions of the Budapest Treaty for the International Recognition of the Deposit of Microorganism for the Purposes of Patent Procedure. The hybridoma, was accorded ATCC Accession Number HB-12467.

30

According to one embodiment of this invention, step (a) further comprises a second antibody which specifically binds to hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex 35 between the antibody and the early pregnancy associated

molecular isoform of hCG. In an embodiment of this invention, the second antibody is B207. According to another embodiment of this invention, step (a) further comprises a second antibody which specifically binds to intact non-nicked  
5 hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG. In an embodiment of this invention, the second antibody is B108 or B109. In an embodiment of this  
10 invention, step (c) comprises comparing the amount of the early pregnancy associated molecular isoform of hCG determined in step (b) for said antibody with the amount determined in step (b) for the second antibody, wherein a high ratio of amounts determined for said antibody relative  
15 to the second antibody indicates a positive outcome of pregnancy for the subject, a low ratio indicates a negative outcome of pregnancy for the subject.

In yet another embodiment of this invention, step (c)  
20 comprises comparing the amount early pregnancy associated molecular isoform of hCG in the sample determined in step (b) with either (i) the amount determined for temporally matched, normal pregnant subject(s) or (ii) the amount determined for non-pregnant subject(s), wherein amounts of the early  
25 pregnancy associated molecular isoform of hCG in the sample similar to amounts of early pregnancy associated molecular isoform of hCG in temporally matched pregnant samples indicates a positive outcome, amounts of early pregnancy associated molecular isoform of hCG in the sample similar to  
30 amounts of early pregnancy associated molecular isoform of hCG in the non-pregnant samples indicates a negative outcome of pregnancy for the subject.

This invention also provides a method of predicting the  
35 likelihood of a negative pregnancy outcome in a female subject comprising: (a) contacting a sample from the subject with a

capture antibody which specifically binds to an early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (b) contacting any complex formed in step (a) with a labelled detection antibody under conditions permitting binding to the complex the capture antibody and the hCG isoform; (c) measuring the amount of labeled detection antibody bound to the complex so as to thereby determine the amount of the early pregnancy associated molecular isoform of hCG in the sample; and (d) comparing the amount early pregnancy associated molecular isoform of hCG in the sample determined in step (b) with the amount determined for a normal pregnant subject, wherein the relative absence of the early pregnancy associated molecular isoform of hCG in the sample indicates a negative outcome of pregnancy for the subject.

According to an embodiment of this invention, the sample is a urinary sample or a blood sample. In one embodiment of this invention, the sample is an aggregate sample taken from at least two consecutive days. In an embodiment of this invention, the sample is a spot urine sample, a first morning void urine sample, or an aggregate sample of the first morning void urine samples for at least two consecutive days. In one embodiment of this invention, the antibody is labeled with a detectable marker. In an embodiment of this invention, the detectable marker is a radioactive isotope, enzyme, dye, magnetic bead, or biotin. In a preferred embodiment, the radioactive isotope is  $I^{125}$ .

The present invention further provides a method of predicting pregnancy outcome in a subject by determining the amount of an early pregnancy associated molecular isoform of hCG in a sample comprising: (a) contacting a capturing antibody which specifically binds to the early pregnancy associated

molecular isoform of hCG with a solid matrix under conditions permitting binding of the antibody with the solid matrix; (b) contacting the bound matrix with the sample under conditions permitting binding of the antigen present in the sample with the capturing antibody; (c) separating the bound matrix and the sample; (d) contacting the separated bound matrix with a detecting antibody which specifically binds to hCG under conditions permitting binding of antibody and antigen in the sample; (e) measuring the amount of bound antibody on the bound matrix, thereby determining the amount of early pregnancy associated molecular isoform of hCG in the sample; and (f) comparing the amount early pregnancy associated molecular isoform of hCG in the sample determined in step (e) with either (i) the amount determined for temporally matched, normal pregnant subject(s) or (ii) the amount determined for non-pregnant subject(s), wherein amounts of the early pregnancy associated molecular isoform of hCG in the sample similar to amounts of early pregnancy associated molecular isoform of hCG in temporally matched pregnant samples indicates a positive outcome, amounts of early pregnancy associated molecular isoform of hCG in the sample similar to amounts of early pregnancy associated molecular isoform of hCG in the non-pregnant samples indicates a negative outcome of pregnancy for the subject.

25

An embodiment of this invention further comprises (a) removing of the sample from the matrix; and (b) washing the bound matrix with an appropriate buffer. In one embodiment of this invention, the capturing antibody is B152. In one embodiment of this invention, the detecting antibody is B207. In an embodiment of this invention, step (a) further comprises a second capturing antibody which specifically binds to intact non-nicked hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG. According to an

embodiment of this invention, the second capturing antibody is B108 or B109. In an embodiment of this invention, step (d) further comprises a second detecting antibody which specifically binds to hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG. In an embodiment of this invention, step (f) comprises comparing the amount of the early pregnancy associated molecular isoform of hCG determined in step (e) for said antibody with the amount determined in step (b) for the second antibody, wherein a high ratio of amounts determined for said antibody relative to the second antibody indicates a positive outcome of pregnancy for the subject, a low ratio indicates a negative outcome of pregnancy for the subject.

According to an embodiment of this invention, the sample is a urinary sample or a blood sample. In one embodiment of this invention, the sample is an aggregate sample taken from at least two consecutive days. In an embodiment of this invention, the sample is a spot urine sample, a first morning void urine sample, or an aggregate sample of the first morning void urine samples for at least two consecutive days. In one embodiment of this invention, the antibody is labeled with a detectable marker. In an embodiment of this invention, the detectable marker is a radioactive isotope, enzyme, dye, magnetic bead, or biotin. In a preferred embodiment, the radioactive isotope is  $I^{125}$ .

In addition, the present invention provides a method for determining the amount of early pregnancy associated molecular isoforms of in a sample comprising: (a) contacting the sample with an antibody which specifically binds to an early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform

of hCG; and (b) determining the amount of complexes formed thereby determining the amount of early pregnancy associated molecular isoform of hCG in the sample.

5 According to an embodiment of this invention, the antibody specifically binds a region of the early pregnancy associated molecular isoform of hCG comprising a carbohydrate moiety. In one embodiment of this invention the antibody is produced by a hybridoma cell line. In one embodiment of this  
10 invention the antibody is B152. In another embodiment the antibody specifically binds to the nicked form of human gonadotropin. The antibody is unlabeled or labeled with a detectable marker. In one embodiment of the invention the detectable marker is a detection antibody. In another  
15 embodiment the antibody specifically binds an epitope dependent upon peptide bond cleavage in beta loop 2. In still another embodiment the antibody binds the human gonadotropin at the beta COOH-terminal region.

20 Further, the present invention provides a diagnostic kit for determining the amount of early pregnancy associated hCG in a sample comprising: (a) an antibody which specifically binds to an early pregnancy associated molecular isoform; (b) a solid matrix to which the antibody is bound; and (c) reagents  
25 permitting the formation of a complex between the antibody and a sample. In an embodiment of this invention, the antibody is B108, B109 or B152. An embodiment of this invention further comprises control sample(s) normal pregnant sample(s), nonpregnant sample(s), or male sample(s).

30

According to an embodiment of this invention, the sample is a urinary sample or a blood sample. In one embodiment of this invention, the sample is an aggregate sample taken from at least two consecutive days. In an embodiment of this  
35 invention, the sample is a spot urine sample, a first morning void urine sample, or an aggregate sample of the first

morning void urine samples for at least two consecutive days. In one embodiment of this invention, the antibody is labeled with a detectable marker. In an embodiment of this invention, the detectable marker is a radioactive isotope, 5 enzyme, dye, magnetic bead, or biotin. In a preferred embodiment, the radioactive isotope is  $I^{125}$ .

The present invention additionally provides an antibody which specifically binds to an early pregnancy associated molecular 10 isoform of human chorionic gonadotropin.

In an embodiment of this invention, the antibody specifically binds to a region of the early pregnancy associated molecular isoform of human chorionic gonadotropin comprising a 15 carbohydrate moiety. According to one embodiment of this invention, the monoclonal antibody is B152. In an embodiment of this invention, a hybridoma cell (ATCC Accession No. HB-12467 ) is provided capable of producing monoclonal antibody B152. Another embodiment of this invention is the early 20 pregnancy associated molecular isoform of hCG recognized by the B152 monoclonal antibody.

Further, the present invention provides a method for detecting non-trophoblast malignancy in a sample comprising: 25 (a) contacting a sample with an antibody which specifically binds to the early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (b) contacting the sample with a 30 second detection antibody which specifically binds to intact non-nicked hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (c) measuring the amount 35 of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the

sample; and (d) comparing the amount of early pregnancy associated molecular isoform of hCG in the sample determined in step (b) with the amount of early pregnancy associated molecular isoform of hCG in the sample determined in step 5 (c), wherein a positive detection of early pregnancy associated molecular isoform detected in step (b) and a relative absence of the early pregnancy associated molecular isoform of hCG detected in step (c) indicates the presence of non-trophoblast malignancy in the sample.

10

According to an embodiment of this invention, the antibody is B152 or B109. In an embodiment of this invention, the detection antibody is B207 for B152 assay, B108 for B109 assay. In an embodiment of this invention, the non-15 trophoblast malignancy is ovarian malignancy or prostate malignancy.

According to an embodiment of this invention, the sample is a urinary sample or a blood sample. In one embodiment of 20 this invention, the sample is an aggregate sample taken from at least two consecutive days. In an embodiment of this invention, the sample is a spot urine sample, a first morning void urine sample, or an aggregate sample of the first morning void urine samples for at least two consecutive days. 25 In one embodiment of this invention, the antibody is labeled with a detectable marker. In an embodiment of this invention, the detectable marker is a radioactive isotope, enzyme, dye, magnetic bead, or biotin. In a preferred embodiment, the radioactive isotope is  $I^{125}$ .

30

In an embodiment, the present invention provides a method for detecting gestational trophoblast disease in a sample from a subject comprising (a) contacting a sample with an antibody which specifically binds to the early pregnancy associated 35 molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early

pregnancy associated molecular isoform of hCG; (b) contacting the sample with a second detection antibody which specifically binds to intact non-nicked hCG without substantially cross-reacting with said antibody under 5 conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; (c) measuring the amount of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample due to binding with 10 the first antibody, and late pregnancy associated molecular isoform of hCG in the sample due to binding with the second antibody; (d) determining the ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated molecular isoform of hCG in the subject; and (e) 15 comparing the ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated molecular isoform of hCG in the sample determined in step (c) over time, wherein a continuing high ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated 20 molecular isoform of hCG in the sample determined in step (c) indicates the presence of gestational trophoblast disease in the subject.

In an embodiment of this invention, the antibody is B152 or 25 B109. In another embodiment of this invention, the second antibody is B108 for B109 B207 for B152 assay. In an embodiment of the present invention, the gestational trophoblast disease is choriocarcinoma or hydatidiform mole.

30 According to an embodiment of this invention, the sample is a urinary sample or a blood sample. In one embodiment of this invention, the sample is an aggregate sample taken from at least two consecutive days. In an embodiment of this invention, the sample is a spot urine sample, a first morning 35 void urine sample, or an aggregate sample of the first morning void urine samples for at least two consecutive days.

In one embodiment of this invention, the antibody is labeled with a detectable marker. In an embodiment of this invention, the detectable marker is a radioactive isotope, enzyme, dye, magnetic bead, or biotin. In a preferred 5 embodiment, the radioactive isotope is  $I^{125}$ .

As described herein below, unexpected isoforms of hCG are produced during normal early pregnancy. Using an *in vitro* bioassay, it appears that these isoforms have enhanced 10 potency for signal transduction. These isoforms can be measured using the novel sensitive, immunoassay described herein. This can help predict pregnancy outcome where one cause of early pregnancy loss is failure to produce the isoform of hCG of higher potency produced by successful 15 pregnancies. This enables physicians to intervene to sustain a failing pregnancy. Identification of the nature of the hCG isoform required might provide the proper reagent needed to sustain pregnancy.

20 New antibodies for measurement of nicked forms of hCG described herein below were developed based on the hypothesis that forms of hCG, which have greatly reduced bioactivity, contribute to early pregnancy loss (EPL), due at least in part to diminished biopotency. Evidence was found that the 25 hCG that appears in EPL patients displays reduced biological activity. However, it was determined that the cause of the reduced bioactivity is not the presence of nicked hCG in EPL patients. Instead, the hypothesis is that patients that carry pregnancies forward produce an isoform of hCG with 30 enhanced bioactivity. The instant invention describes a unique immunochemical assay to measure this unexpected and previously uncharacterized isoform of early pregnancy hCG directly in clinical samples of blood and urine. One of the antibodies developed reacted against a nicked form of hCG 35 isolated from a choriocarcinoma patient, was not specific for

a nicked form of hCG but appeared to discriminate among carbohydrate variants of hCG. This antibody, designated B152, appears to preferentially bind hCG forms from choriocarcinoma patients. In studying the content of hCG isoforms during pregnancy, the unique and unexpected observation was made that B152 in the first four weeks of pregnancy measured much higher quantities of an isoform of hCG as compared to the standard hCG isoforms measured by the usual heterodimeric hCG assays exemplified by a previously described B109 based assay. In fact, in early pregnancy (days 9,10,11 postovulation) B152 measured as much as 20-fold more hCG, than did another monoclonal antibody, B109. Later in pregnancy, the B152 isoform declines and is lower in third trimester pregnancy urine than the standard isoforms measured by B109. A further striking observation was that in very early pregnancy, a high B152/B109 ratio correlates with a successful pregnancy outcome while a low ratio correlated with pregnancy loss. This discovery is important as the potentially overlooked isoforms of hCG described herein during pregnancy may be predictors of successful pregnancy outcome. Such an assay has wide medical applications and provides a clinician with opportunity to intervene very early in pregnancy if the assay indicated that the pregnancy appeared troubled.

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An antibody, designated B152, produced by the hybridoma cell accorded ATCC Accession number HB-12467, generated against a nicked form of hCG isolated from a choriocarcinoma patient, but not specific for nicked isoform hCG is able to discriminate among carbohydrate variants of hCG. B152 is specific for an early pregnancy associated molecular isoform of hCG, which in the first four weeks of pregnancy is measured at much higher quantities than the hCG standard isoforms measured by the usual heterodimeric hCG assays exemplified by a previously described B109 based assay. Later in pregnancy, the B152 isoform declines and is lower

in third trimester pregnancy urine than the standard isoforms measured by B109.

This invention is illustrated in the Experimental Details 5 section which follows. These sections are set forth to aid in an understanding of the invention but are not intended to, and should not be construed to, limit in any way the invention as set forth in the claims which follow thereafter.



Unappreciated isoforms of hCG in blood and urine very early in pregnancy may be more potent *in vivo* than the forms of hCG produced later in pregnancy. The absence of such isoforms may be one cause of early pregnancy loss. A sensitive and  
5 specific immunoassay system was designed and made to measure unique early pregnancy associated molecular isoforms (EPMI) of hCG. These isoforms, likely to differ by carbohydrate composition, are predictive of a successful pregnancy outcome. When these early pregnancy associated molecular  
10 isoforms of hCG are absent or present in low concentration, the pregnancy may be lost very early and be observed as only a "chemical" pregnancy. These hCG isoforms may resemble the forms of hCG produced in some choriocarcinoma patients from which the immunogen used to produce monoclonal antibody B152  
15 was derived as described herein below. The isoforms resemble those from trophoblastic disease not in terms of nicking or intact peptide chains but likely in carbohydrate content. The present invention describes that the molar ratio of B152 to B109 epitopes are predictive of a successful pregnancy or  
20 a loss. Three categories of pregnant patients were analyzed: (a) normal pregnant women, (b) women who experience recurrent abortions, (c) women undergoing embryo implantation.

It is possible to determine the hCG isoforms present in the  
25 blood and urine of women who have a history of recurrent spontaneous abortion and a similar analysis of women undergoing embryo implantation. The combined EPL and spontaneous abortion rate in healthy populations is 31%. Subjects who experience three consecutive recurrent  
30 spontaneous abortions have a 32% risk of sustaining another (Hill, J. A.; Anderson, D. J., 1990). In *in vitro* fertilization IVF pregnancy, the loss rate is 70% with non-donor sperm and 50% when donor sperm is used. Delineation of pregnancies with a negative outcome from pregnancies with  
35 a positive outcome can be based on differences in the

concentrations of EPMI hCG isoforms (i.e. as differences in the B152/B109 ratio in patients). In addition, specimens from gestational trophoblastic disease (GTD) can be used to discriminate between GTD and normal pregnancy.

5

## **Results**

### In vitro bioassay for hLH/hCG

An hCG bioassay was constructed employing CHO cells expressing functional human LH/CG receptor. Table 1  
10 illustrates the differences *in vitro* in biological activity between nicked and non-nicked hCG as measured by this assay. This system, has been used to evaluate the activity of pituitary and placental hCG (Birken, S., et al., 1996b). Preparations of hCG were tested for nicked and non-nicked  
15 molecular isoforms of hCG in a second recombinant bioassay system (Ho, H-H., et al., 1997). Similar results were obtained in both systems.

Normal pregnancy values compared with EPL values. Results  
20 indicated that nicked hCG is not a significant molar constituent of either early pregnancy or EPL. Data indicated that biological activity is not correlated with nicked hCG, but is instead ascribed to a form of hCG recognized by the B152 monoclonal antibody ATCC # HB-12467 an early pregnancy  
25 associated molecular isoform of hCG (EPMI hCG). It has been established that there is diminished hCG bioactivity associated with EPL as compared to early normal pregnancy (Ho, H-H., et al., 1997). Thus, diminished hCG biological activity is a factor in EPL as a consequence of a heretofore  
30 unappreciated isoform of hCG - an early pregnancy associated molecular isoform of hCG.

hCG Urinary Analytes. Metabolites of hCG and hLH were studied in a variety of states (Birken, S., et al., 1996a).  
35 One study indicated a 31% pregnancy loss (Zinaman, MJ, et

al., 1996) while another indicated a 17.4% rate of early pregnancy loss based on hCG assays (Elish, N. J., et al., 1996). It is known that hCG and hCG beta core can be readily transferred from the uterus to the circulation even in the absence of implantation (Chang, P. L., 1997). The molecular spectrum of hCG urinary analytes in EPL cycles, normal conceptive cycles and non-conceptive cycles has been evaluated. The study design and demographics of the investigation have been described (Elish, N. J., et al., 1996).

Briefly, three urine specimens per cycle, corresponding to days 9,10, 11, post calculated day of ovulation were collected and analyzed in a screening assay (the "combo") which simultaneously detects intact, non-nicked hCG, hCG free beta subunit, and hCG beta core fragment. Individual determinations for each of these analytes, as well as for nicked hCG, and the form of intact hCG detected by monoclonal antibody B152 (EPMI hCG) were performed on these specimens. In addition, since the concentration of luteal phase hLH urinary analytes is a concern because of cross-reaction in hCG assays, levels of intact hLH, hLH free beta subunit and hLH beta core fragment were determined in the normal pregnancy cycles and the non-conceptive cycles. Table 1 summarizes the characteristics of immunometric assays employed.

**TABLE 1.** Assay format and specificity

<b>Assay format</b>	<b>Primary analyte</b>	<b>% cross-reactivity with related analytes</b>
5 B109- B108*	intact non-nicked hCG	<1% <sup>b</sup>
B201- C104*	hCG free beta subunit (non-nicked+nicked)	1% hCG; 10% hCG nicked (pregnancy); <1% <sup>b</sup>
10 B210- B108*	hCG beta core fragment	2% hLH beta core fragment; <1% <sup>b</sup>
B151- B207*	hCG nicked	10% hCG nicked free beta subunit; 12% hCG non-nicked; 2% hCG free beta subunit; 2% hLH; 5% hLH free beta subunit; <1% <sup>b</sup>
B152- B207*	choriocarcinoma hCG (C5) and choriocarcinoma hCG free beta subunit	100% hCG nicked (C5); 190% hCG free beta nicked (from C5); 10% hCG nicked (pregnancy); 5% hCG free beta nicked (pregnancy); 7% hCG (pregnancy); 6% hCG free beta subunit; <1% <sup>b</sup>

	B406- A201*	hLH	<1% <sup>a</sup>
	B505- B503*	hLH beta core fragment	<1% <sup>a</sup>
5	B408- B409*	hLH free beta subunit	29% hLH; <1% <sup>a</sup>

<sup>a</sup>-(if not indicated) hLH, free beta hLH, hLH beta core fragment, hCG, free beta hCG, hCG beta core fragment;

10 <sup>b</sup>-(if not indicated ) the same as <sup>(a)</sup> plus nicked hCG and nicked free beta hCG (pregnancy).

The results indicate that nicked hCG does not constitute a significant mole fraction of urinary hCG immunoreactivity in  
15 either EPL or early normal pregnancy. In addition, there is a substantial excretion of hCG free beta subunit in some subjects in both pregnancy and EPL. Further, both EPL and normal pregnancy cycles variably express all of the measured analytes. Although both the incidence and level of  
20 expression are different between EPL's and normal pregnancy, there is no hCG related analyte unique to either state. There was, however, a clear difference between the hLH associated analytes in the control population (non-conceptive cycles) and the normal pregnancy group. Virtually all of the  
25 non-pregnancy cycles expressed hLH free beta subunit and hLH beta core fragment while only a third of the conceptive cycles had detectable levels of either analyte. Intact hLH proved to be a minor constituent of the hLH profile in both groups.

30

These findings demonstrate both the necessity of measuring hCG beta core fragment in the detection of EPL, and also of making sure that the hCG beta core assay does not cross-react with beta core hLH, which is demonstrated to be present  
35 in that part of the luteal phase where EPL measurements are performed. The data is summarized in Figure 2.

Statistical analysis was performed after transformation of analyte values to mole fractions so as to produce a more useful analysis due to the wide excursion of hCG analyte values among groups. The mole fraction data were evaluated 5 by discriminant analysis and by a mixed effects model incorporating LMP (last menstrual period date). The discriminant analysis was performed both with and without "outliers" (defined as values greater than two standard deviation from the mean) removed. Both approaches produced 10 similar results.

A quadratic discriminant analysis based on a cross-validation method in order to minimize bias correctly classified 91% of the normal pregnancy subjects and 80% of the EPL subjects. 15

The mixed effects analysis, testing for interactions between mole fraction of analyte and time since LMP found no significant time or group (EPL vs. normal) effects in the intact hCG assay. In the free beta subunit of hCG assay, 20 there is a significant group effect but no time trend. In both the hCG beta core fragment measurement and the B152 measurement, both the hormone levels and the time trend from LMP were significantly different between the EPL and pregnancy groups. This study produced several important 25 findings. It defined the spectrum of analytes which in both early pregnancy and EPL, thereby resolving the issue of which hCG analytes to measure in epidemiological studies in which EPL is the end point determination. More importantly, it illustrated for the first time that there are significant 30 differences both in the pattern of analytes and the time course of their appearance between early normal pregnancy and EPL. This observation facilitates very early prediction of a distressed pregnancy by urinary hCG measurements at a time which would permit therapeutic intervention.

Immunoreactivity of different forms of hCG in the two IRMA's  
(B152-B207 and B109-B108)

The relative binding of three different forms of hCG (urinary  
5 hCG, pituitary hCG and choriocarcinoma hCG C5) has been  
characterized in the two hCG assays (Figure 3). Urinary non-  
nicked hCG and pituitary hCG are recognized nearly equally  
well by the two IRMA's, while C5 recognition is quite  
different. The B152-B207\* assay is more sensitive to C5,  
10 which is to be expected because B152 antibody was developed  
and selected on the basis of higher affinity to C5. Urinary  
non-nicked hCG is purified from the CR127 preparation of  
pooled normal pregnancy hCG. Conversely C5 is recognized with  
lower affinity by the B109-B108\* assay, which has primary  
15 specificity for the hCG isoforms of later pregnancy.

We have developed a method to directly profile changes of hCG  
isoforms in serum or urine throughout pregnancy. Two IRMA's  
for hCG are employed, each based on monoclonal antibodies to  
20 different hCG epitopes. The B109-B108\* assay is a commonly  
used intact hCG assay to the heterodimeric-dependent epitope.  
A new assay, B152-B207\*, is most likely sensitive to the  
carbohydrate portion of hCG carboxyterminal peptide. The same  
standard non-nicked hCG was used in both assays. Non-nicked  
25 hCG was employed since the B109 assay reacts poorly with  
nicked forms of hCG while the B152 assay does not  
discriminate between nicked and non-nicked forms of the  
hormone. The B152 assay detected with greatly enhanced  
sensitivity hCG isoforms which appear earlier in pregnancy  
30 than isoforms measured by the B109 assay (O'Connor et al.  
1998). Prior to development of the new immunometric assay  
system described in this report, it was not possible to  
readily discern the changes in hCG isoforms from very early  
pregnancy to mid pregnancy. The only available procedure for  
35 examining these changes was isoelectric focusing of every  
patient specimen followed by immunoassay of every focused

fraction (Berger et al. 1993; Ulloa-Aguirre et al. 1990). The IEF pattern reflects the heterogeneity of the charged sugar, sialic acid which varies with the multi-antennary structures of the carbohydrate moieties in which sialic acid is the terminal sugar. Although we do not yet know the precise nature of the isoform epitopes being measured, the evidence for carbohydrate discrimination is based upon the hyperglycosylated structure of the immunogen, C5, used to develop the B152 monoclonal antibody and the antibody's reactivity with the hCG isoforms found in the JAR choriocarcinoma cell line. C5 hCG was isolated from a choriocarcinoma patient and has been thoroughly characterized as to its protein and carbohydrate content and structure (Elliott et al. 1997). It has been shown that C5 (and hCG from other choriocarcinoma subjects) differ in the protein moiety mainly by the presence of an increased number of nicked sites and by increased glycosylation relative to the hCG of normal pregnancy. In comparison with the hCG of normal pregnancy, choriocarcinoma derived hCG has increased fucosylation of the N-linked biantennary oligosaccharides in the beta subunit. In addition, the O-linked oligosaccharides in preparation C5 (a form of hCG produced from a single patient with choriocarcinoma) has a 100% tetrasaccharide core on the COOH-terminal region of the beta subunit. Normal mid pregnancy hCG has only 10-20% of this structure (Elliott et al. 1997). These observations, plus our own determination that the hCG synthesized by the JAR choriocarcinoma cell line provides a B152/B109 isoform ratio similar to that observed in early pregnancy, leads us to the conclusion that in very early pregnancy, the developing trophoblast secretes an isoform of hCG which resembles that produced in choriocarcinoma.

We have also tested recognition of pituitary hCG since its N-Asn carbohydrates differ somewhat from those of placental hCG, bearing a closer resemblance to those of hLH which have

both sialic acid and sulfate groups (Birken et al. 1996). The carbohydrate structure of the b COOH-terminal portion of pituitary hCG is not yet known. Since B152 did not recognize any substantial differences between pituitary and placental hCG (Figure 3), differences in N-Asn recognition are unlikely. In terms of the COOH-terminal carbohydrates, it appears that pituitary and placental hCG (mid-pregnancy isoforms) may be similar, assuming the O-linked carbohydrate on the C5 antigen is part of the epitope of B152.

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**Example 2: B152/B109 Ratio Predicts Pregnancy Outcome**

The B152/B109 ratio measured in urine samples throughout the pregnancy

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The relative concentrations of hCG isoforms in 103 normal pregnancy urine samples (5-39 weeks post last menstrual period - LMP) were determined by two immunometric assays (B152-B207\* and B109-B108\*). Both because of the wide range of hCG concentrations in different samples, even at the same gestational age, and because neither of the assays is totally specific for the two (or more) families of hCG isoforms present, we find that presenting the data as a ratio of the observed two isoform groups more clearly delineates the change in isoform content as pregnancy progresses. This calculated ratio is shown in Figure 4. In weeks 5-8 of pregnancy, the ratio of B152/B109 isoforms ranged between 6.2 and 1.3, indicating a predominance of the B152 isoform(s) in early pregnancy. During the 10 to 12 week period, the ratio ranged from 1 - 0.2, indicating that an inversion in hCG isoform content is occurring as pregnancy progresses. This decline in the ratio continues, ranging from 0.54 - 0.08 in the 15 - 18 week period and reaching an inflection point at 29 weeks. At that time, the ratio reached a value of around 0.06 after which the ratio displayed a rise to a range of 0.2 - 0.07 in the 37 - 39.5 weeks of gestation time period.

Statistical analysis involved fitting the log transformed ratio data to second and third order polynomial regression models. Since the third order term was not significant (likelihood ratio  $\chi^2(1)=1.32$ ,  $P=0.25$ ), the second order model was used ( $r=0.793$ ). The log B152/B109 ratio reached an inflection point at LMP=29 weeks, based on this model.

The B152-B207\* values reflect a measurement of the B152 isoform in terms of later pregnancy hCG equivalents, not in absolute quantities. It must be emphasized that the "absolute" concentrations measure in the B152 assay cannot be compared with the results of the B109 assay on an equimolar basis since the potency of the hyperglycosylated isoform is much higher in the B152 assay vis-à-vis the standard, i.e. normal later first trimester pregnancy hCG. The actual molar values of this isoform are on the order of tenfold less than those recorded in the assay. For this reason we have chosen not to analyze absolute molar quantities of the two analytes but only the ratio of the two measurements.

Even in normal pregnancy, the hCG values obtained vary widely according to the characteristics of the immunological reagents employed (Cole and Kardana, 1992; Cole et al. 1993). We hypothesize that the two assays described in this report primarily detect hCG isoforms at opposing ends of this spectrum, each primarily recognizing a subset of closely related molecules in the continuum of early to later pregnancy hCG molecular forms.

30

We have retained the use of normal pregnancy hCG as the standard in B152-B207\* assay, despite its decreased affinity in this antibody configuration. The reasons for this include the limited and unrenovable supply of C5 (which was isolated from the urine of a single patient) and the variability in

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data which would result from investigations using different standards. The consequences of this choice are that the early pregnancy hCG isoforms have markedly increased immunopotency over that of normal pregnancy and hence their 5 molar quantities are overestimated in this assay. We use this difference in affinity to our advantage by employing a ratio of the molar results of two assays (B152 and B109). Either assay taken alone obscures this change due to the wide excursion of hCG values which occur in normal pregnancy.

10

Others have documented progressive changes in hCG isoforms throughout pregnancy. Skarulis et al. found that the fucose content of both intact hCG and also its free beta subunit increased as pregnancy progressed (Skarulis et al. 1992). 15 Diaz-Cueto et al. investigating the isoelectric focusing pattern of circulating hCG throughout pregnancy, found that in early pregnancy, more than 80% of the hCG isoforms were acidic. This fraction decreased to less than half (47%) late in the third trimester (Diaz-Cueto et al. 1996). In 20 contrast, Wide and Hobson found that the hCG of early pregnancy was more "choriocarcinoma-like" by virtue of its greater biological activity than the hCG of normal pregnancy (Wide and Hobson, 1987). Fein et al., in a study which employed gel filtration determined that first trimester hCG 25 was a larger size than that of the third trimester. Treatment with exoglycosidases eliminated the size differential, indicating that the first trimester hCG was more highly glycosylated (Fein et al. 1980).

30 The B152/B109 ratio in matched serum/urine samples in the first and third trimesters of pregnancy compared with hCG from JAR cells.

The B152/B109 ratio in serum is analogous to that found in 35 matched urine samples and undergoes a similar change as pregnancy progresses (Figure 5). The B152/B109 ratio in the

cell supernatant from JAR cells (a choriocarcinoma derived cell line) was similar to that of early pregnancy.

The B152/B109 ratios of both serum and urine hCG concentrations are significantly higher in the first trimester as compared to the third trimester of normal pregnancies (Table 2). Significant differences between serum and urine hCG concentration ratios as well as log transformed ratios in early (5-6 weeks) and late (36-39 weeks) gestation were evaluated by paired t-tests (Table 3). In both the first and third trimesters, urinary B152/B109 ratios were significantly higher than serum ratios, indicating that there was a preferential clearance of the B152-recognized isoform into urine, regardless of the relative concentrations of the two isoforms.

**TABLE 2.**

Analysis of the B152/B109 ratio in serum and in urine in the first vs third trimesters of pregnancy.

Measure	T-test (df)	P
Serum, ratio B152/B109	t(11)=6.65	0.0001
Serum, log(ratioB152/B109)	t(23)=21.61	0.0000
Urine, ratio B152/B109	t(11)=4.64	0.0007
Urine, log(ratioB152/B109)	t(15.7)=16.85	0.0001

**TABLE 3.**

Analysis of the B152/B109 ratio in serum vs urine in the first and third trimesters of pregnancy.

Gestational age	Measure	Paired-t (df)	P
5-6 weeks	Ratio B152/B109 Log(ratioB152/ B109)	t(11)=3.25 t(11)=6.25	0.0077 0.0001
36-39 weeks	Ratio B152/B109 Log(ratioB152/ B109)	t(10)=5.47 t(10)=7.14	0.0003 0.0001

In urine samples from IVF patients (1-4 weeks post embryo transfer - ET) the B152/B109 ratio was again between 2-8 and decreased as pregnancy progressed (Figure 6), similar to that observed in natural conceptions. The effect of pregnancy duration with respect to outcome variables could best be represented by a linear or quadratic function. ANCOVA models including the second order week were fitted to the general equation: Outcome= (effect of time post ET)+(effect of diagnosis). After an appropriate ANCOVA model was determined, the least square means (adjusted for week post ET effect) were compared among the normal pregnancy, ectopic pregnancy and spontaneous abortion populations (Table 4). The log transformed values of both B109-B108\* and B152-B207\* measured hCG forms discriminated both ectopic pregnancy and spontaneous abortions from normal pregnancy ( $P=0.0001$ ). The ratio of the log transformed values discriminated abortion from normal pregnancy ( $P=0.016$ ). However, neither the ratio of B152/B109 nor the log of that ratio discriminated either of the pregnancy disorders from normal pregnancy.

20

A significant number of spontaneous abortions and ectopic pregnancies occur in IVF pregnancies. We did not find a difference in the ratio of the isoforms between either of

these two categories as compared to normal controls, possibly a consequence of low statistical power. However a significant difference was found between the B152 hCG isoforms levels in normal pregnancy and spontaneous abortion. This supports our previous finding in early pregnancy loss, where diminished or absent levels of the B152 isoforms characterized an early pregnancy loss (O'Connor et al. 1998).

**TABLE 4.**

IVF patients: analysis of covariance of hCG isoforms among normal pregnancy (np), ectopic pregnancy and spontaneous abortion as a function of gestational age.

Outcome	<sup>a</sup> Adjusted R <sup>2</sup>	<sup>e</sup> -F	F	<sup>d</sup> -Pairwise Difference
<sup>a</sup> Log (ratio B152/B109)	0.51	0.89	0.41	none
<sup>a</sup> Log (B109-B108*)	0.56	21.33	0.0001	np vs abortion & ectopic
<sup>b</sup> Log (B152)/log (B109)	0.45	4.34	0.016	np vs abortion
<sup>b</sup> Log (B152-B207*)	0.50	26.94	0.0001	np vs abortion & ectopic

- <sup>a</sup>- ANCOVA model with 2nd order polynomial coefficient (or parameter).
- <sup>b</sup>- ANCOVA model with only 1st order (linear) coefficient.
- <sup>c</sup>- Adjusted R<sup>2</sup> is a R<sup>2</sup> adjusting number of coefficients on the ANCOVA model so that comparisons of two R<sup>2</sup> with different ANOVA models with different number of coefficients are meaningful.
- <sup>d</sup>- "Pairwise difference" is based on t-test comparing the least-square means of outcome variables (after adjusting effect of week ET).
- <sup>e</sup>- Degree of freedom (df1, df2) for F-test are (2,82) for a model with only linear coefficient and (2,81) for a model with both linear and 2nd order coefficient.

HCG analysis of trophoblastic disease samples

Trophoblast disease serum (17 samples) and urine (28 samples) were obtained from patients post therapy and hence contained low hCG levels. Due to limited amounts of sample all of these specimens were run at a 1:10 initial dilutions. HCG levels in serum were low. The highest hCG concentration in serum was 202 fmol/ml in the B152-B207\* assay, with a corresponding value of 148 fmol/ml in the B109-B108\* determination. Six of seventeen samples in serum had detectable levels, with 4/6 having a higher value in the B152-B207\* assay. Of the 15/28 positive urine samples however, 14/15 had higher levels in the B152-B207\* assay than in the B109-B108\* assay, with the highest hCG value being 20000 fmol/ml in the B152-B207\* assay and 18715 fmol/ml in the corresponding B109-B108\* assay. Due to the small sample size, no statistical treatment was performed on this data, but even in these post-treatment patients the B152/B109 ratio was  $\geq 1$ , which corresponds to the early pregnancy hCG isoform ratio.

20

The specimen limitations discussed above precludes our reaching any definitive conclusion on the analysis of trophoblastic disease samples. However it appears as might be anticipated that the B152 assay is more sensitive than B109 assay in detecting hCG immunoreactivity in the blood and in the urine of trophoblastic disease patients, even after treatment.

Chromatography of First Week of Gestation Pregnancy Pool.

In order to determine whether the B152-B207\* assay recognized other forms of hCG associated immunoreactivity in addition to the intact hCG molecule, specimens were pooled. FPLC on tandem Superose 12 columns followed by immunoassay of the fractions for all of the characterized forms of hCG revealed that only the intact hCG molecule (or hCG free beta subunit) gave a signal in this assay (See Figure 8). There were no

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lower molecular weight fragments identified by the B152-B207\* assay. The hCG free beta analyte was measured in the 159 urines described in Figure 6 and was found to make a negligible contribution to over all hCG immunoreactivity in 5 these specimens.

Molecules recognized by monoclonal antibody B152 in urine and pituitary extracts. In order to define the nature of the hCG isoforms recognized by B152, high resolution gel filtration 10 columns of both pituitary extracts and postmenopausal urine concentrates were used (See Figure 8). The rationale for use of pituitary extracts is to determine cross-reactive molecules, specifically those which are glycosylated, which are plentiful in pituitary which contains the entire family 15 of glycoprotein hormones, hLH, hTSH, and hFSH as well as free subunits and the pituitary form of hCG. Two peaks are detected in both of these cases. Only one peak was detected in similar studies of pregnancy urine concentrates as described earlier. In the pituitary, it is likely that the 20 larger molecule is pituitary hCG (70K) while the smaller sized molecule is hLH. Since hLH is present at 100x or so as compared to pituitary hCG, the apparent similar concentration of immunoreactivity indicates that B152 has reduced cross-reactivity to hLH as compared to hCG. 25 Likewise, both hCG and hLH occur in postmenopausal urine, again with much more hLH than hCG and the B152 pattern is similar to that of the pituitary extract. These results show that B152 is generally hCG specific except for cross reactivity to hLH (as shown by standard cross-reaction 30 studies in Table I) and that its carbohydrate specificity is both to the protein portion as well as to the carbohydrate moieties of hCG (and to a lesser extent of hLH) since it does not react with the multitude of other glycoylated proteins present in the pituitary nor with those in postmenopausal 35 urine except for hCG or hLH-related molecules.

Serum and urine specimens were analyzed using two assays, B109-B108\* and B152-B207\*, which recognize the difference in molecular isoforms of hCG. See Table I. The *in vitro* bioassay for hLH/hCG is described above. (See Figure 1).  
5 Results are indicated in Figures 6-11. The immunometric assay employs 96-well microtiter plate technology. The coating antibody, at a concentration determined to provide the most satisfactory combination of sensitivity and range, is applied to the microtiter wells (Immulon IV, Dynatech  
10 Laboratories) in carbonate buffer (0.2M, pH 9.5). The plates are incubated with the coating solution at 4°C, overnight, then aspirated, washed with washing solution (0.05% Tween, 0.15N NaCl), and blocked with a 1% solution of BSA (three hours at room temperature). The BSA solution is aspirated  
15 and the appropriate hCG standards (200µL/well), in buffer B (PBS/0.1% bovine IgG/0.1% sodium azide), or in hCG free serum (Chemicon, Inc.), or hCG free urine, as appropriate to the specimen matrix, and specimens are added to the wells. The plates are sealed with plate sealers, and incubated  
20 overnight at 4°C. The controls, specimens, and standards are then aspirated, the plates washed 5 times with washing solution, and iodinated detection antibody in buffer B (200 uL/well, 100,000 cpm/well) added and incubated overnight at 4°C. The wells are again aspirated, washed 5 times with  
25 washing solution, separated and counted (Packard Cobra gamma counted). Values are interpolated from a smoothed spline transformation of the count data. This assay procedure, as well as assay validation has been previously reported (O'Connor, J. F., et al., 1988).

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Creatinine analysis, when urine values are normalized to creatinine, is performed in a microtiter plate format following a modification of the Taussky procedure (Taussky, H. H., 1954).

35

Descriptive statistical and graphical methods are used to measures of serum and urine samples from normal healthy pregnancies to identify the distributions a) between patient first trimester average B152 levels, B109 levels and 5 B152/B109 ratio; b) between patient variability in time to B152/B109 ratio reaching 1.00; and c) between patient variability in time to B152/B109 ratio declining by 1/3rd from first trimester maximum levels. The variability in the timing of the crossover in the ratio of these two analytes 10 provides an empirical basis from which to estimate the value of these markers as biochemical signatures of a viable third trimester fetus.

Comparison of the assay profile of healthy normal pregnancies 15 to those of unsuccessful pregnancies from failed IVF implantations, two non-parametric hypotheses are available: 1) the proportion of pregnancies in which the B152/B109 ratio falls below 1.00 is no different in healthy normal and unsuccessful IVF pregnancies; 2) the proportion of 20 pregnancies in which the B152/B109 ratio declines by 1/3rd from first trimester maximum levels is no different in healthy normal and unsuccessful IVF pregnancies. These hypotheses can be tested as a difference between two proportions. For example, a comparison of week 14 vs. week 25 9, week 13 vs. week 6, week 12 vs. week 5 or week 11 vs. week 4 pregnancies to show a reversal of the B152/B109 ratio in healthy normal pregnancies and unsuccessful IVF implantations, respectively. The power analyses apply to an outcome defined as the time at which the B152/B109 ratio 30 declines by 1/3rd from first trimester maximum levels, although this outcome would necessarily provide earlier detection of pregnancy failure than the reversal of the B152/B109 ratio. Patterns of results less discriminantly different from these indicate a rejection of the dichotomous 35 outcome of B152/B109 ratio reversal as a clinically meaningful marker of pregnancy failure.

Alternatively, the same two non-parametric hypotheses can be recast as parametric hypotheses by considering the timing of the biochemical events within the assay profile of healthy normal pregnancies and unsuccessful pregnancies from failed IVF implantations: 1) the time at which the B152/B109 ratio falls below 1.00 is no different in healthy normal and unsuccessful IVF pregnancies; 2) the time at which the B152/B109 ratio declines by 1/3rd from first trimester maximum levels is no different in healthy normal and unsuccessful IVF pregnancies. Of course, the objective is to provide an empirical basis from which clinicians may counsel their patients. Thus, it is important to adopt a logistic model for this component of the data analysis. With pregnancy success as the outcome, logistic models allow the estimation of the (symmetrical) hypothesis of increase in risk of pregnancy failure for each additional week where either the B152/B109 ratio has failed to decline by one third from first trimester baseline maximum values or the B152/B109 ratio has failed to become less than 1.00 (measured in 20 weeks). The logistic model enables specification of the time at which results indicate a particular pregnancy exceeds an a priori defined likelihood of failure, given assay data regularly available during pregnancy, and allows incorporation of other risks for pregnancy failure in the same data analytic framework to assess the relative contribution of threats to pregnancy loss. The Cox proportional hazard model may be used to examine predictors of the crossover rates. Mixed effects models can also analyze repeated measures of the B152/B109 ratios taken during entire 30 cycles. These models are particularly useful since they allow inclusion of incomplete and imbalance data (i.e. data with missing values and unequal timing of data collection), to estimate effects of time-varying covariates, to model dependency structure of repeated measures and to model 35 possible heterogeneity of the ratio measures within each experimental group.

B152 hCG isoforms isolated from early pregnancy urine and determination of their protein and carbohydrate structures.  
Using the already developed scheme of concentration and immunoaffinity extraction of urine, hCG molecules are  
5 isolated from urine collected from women in early pregnancy for both protein and carbohydrate analyses. According to one approach, molecules are isolated from HPLC fractions, digested with proteases before and after reduction of disulfide bonds, examination of the resultant peptides by  
10 mass spectrometry and/or sequence analysis, isolation of carbohydrate moieties after glycosidase digestions and determination of carbohydrate structures by a combination of specific glycosidases and retention times on specialized anion exchange columns as compared to know branch-chain  
15 oligosaccharide standards. In a similar approach, the final purification stage for the isolated hCG isoforms is SDS gel electrophoresis. Both protease digests and glycosidase digests are performed on the blotted and cutout band. This method results in greater purity of the protein and less  
20 artifactual errors due to contamination by carbohydrates which are not in the purified protein but are derived from outside contaminants.

Carbohydrate compositional analyses and oligosaccharide  
25 branched chain identifications. The MALDI TOF mass spectrometric method may be used to confirm oligosaccharide structures by using specific glycosidases on the glycopeptides and determining the change in molecular weight as the sugars are digested off the glycopeptide. Only the  
30 hCG beta COOH peptide can be expected to contain O-linked sugar moieties. These are of special interest since it is thought that B152 has significant reaction with this region. The structures of this region can be determined in a similar fashion using enzymes that specifically release O-linked  
35 glycans. The O-linked structures has been previously examined using standard reference pregnancy hCG (Cole, L. A.,

et al., 1985). The O-linked branched chain structure are determined by a similar strategy using the Dionex chromatographic system as well as specific glycosidases on the C-terminal glycopeptides and Mass Spectrometry. In one 5 study (Elliott, M. M., et al., 1997), these techniques were used to elucidate the carbohydrate structures of CR series hCG preparations (standard urinary pregnancy hCG) and compared them to the structures of patient samples such as C5 which was the immunogen employed to generate antibody 10 B152. It was found that C5 contained significantly more mono and tri-antennary (2X mono and 3X tri-structures than the CR preparations) on the N-Asn residues. It was also found that more tetrasaccharide structures were on the hCG COOH-terminal peptide O-Serine residues in the choriocarcinoma hCG isoform 15 than in the CR preparations.

Biological activity and metabolic clearance of hCG isoforms.

Biological activity is a function both of molecular structure and half-life in the circulation, which can be influenced by 20 structure. Alterations in carbohydrate/sialic acid content of the glycoprotein hormones are thought to be responsible for the changes in hCG biological/immunological activity observed throughout pregnancy. In addition, signal transduction at the receptor is influenced by the pI of the 25 hCG isoform and the presence or absence of carbohydrate. Thus, it is valuable to examine both receptor binding and biological activity *in vitro* and, in order to determine the mechanism of action, to distinguish receptor binding and signal transduction as well as relative potency of signal 30 transduction along with *in vivo* bioactivity determinants such as circulating half life. Studies, including clearance rates, are performed on B152 hCG isoforms of early successful pregnancy, hCG from third trimester pregnancy, and the reference urinary hCG preparation, CR 127.

**Example 3: B152 and B151 immunoreactivity in non-trophoblastic malignancy.**

With the exception of trophoblastic disease and testicular cancer, hCG is expressed in the blood of about 20% of patients with all other types of cancer (Hussa, R. O., 1987). HCG beta core fragment in the urine has a significantly higher level of expression, especially in gynecological malignancy. Since the B152 antibody was developed to a form of hCG produced in a malignancy, it was of interest to examine the expression of B152 and nicked hCG immunoreactivity (B151) in non-trophoblastic malignancy. Accordingly, blood and urine derived from men undergoing chemotherapy for prostate cancer or women for ovarian cancer were evaluated for the expression of hCG isoforms in plasma and urine. It is significant that in prostate cancer, B152 hCG immunoreactivity is found in the blood and urine of prostate cancer patients in instances when there is no hCG detected by B109-B108\*. In ovarian cancer patients evaluated, there is evidence of nicked hCG in the blood, even in the absence of both B109 and B152 immunoreactivity. Neither of the above groups demonstrated the presence of hCG immunoreactivity when the standard pregnancy derived hCG assay was employed. It is reassuring to find that nicked hCG, the existence of which has been documented by several investigators, can be found and reliably measured in a clinical setting.

**Experimental Discussion**

In the course of these studies, a potentially important new signal was observed in the urine of women early in pregnancy, namely an epitope of a form of hCG which may indicate the likely success of carrying a pregnancy. Likewise, absence of this signal may indicate that EPL will occur. Since EPL can be a very sensitive marker of environmental toxins (Hakim, R. B., et al., 1995) and is frequently used as an

epidemiological marker of exposure, the finding of this epitope provides a powerful tool for monitoring the safety of the environment. In addition, this assay facilitates increasing the success rate of IVF infertility programs since the predictive value of the new measuring system would rapidly indicate successful approaches. Described herein is the novel and completely unexpected finding that successful pregnancies display a high content of unique isoforms of hCG that are maintained for the first few weeks of pregnancy and then rapidly decline as pregnancy progresses. Based on properties of the immunoassay system, it is hypothesized that these hCG isoforms may be hyperglycosylated. This is a striking observation never reported nor suspected earlier. Carbohydrate analyses (Elliot, M., 1997) demonstrate that C5 hCG employed as immunogen for antibody B152, contains two times the monoantennary content and three times the tri-antennary content of branch chain sugars as compared to the CR series of natural pregnancy urinary hCG. In addition, the O-linked carbohydrates are mostly tetrasaccharide instead of disaccharide in C5 as compared to CR 127 hCG. (CR 127 hCG is similar to the WHO preparation, the third international hCG standard, which was CR 119 hCG, prepared by Canfield and Birken twenty years ago but still in use today) (Birken, S., et al., 1991a). B152 recognizes C5 hCG much better than nicked CR127 hCG or non-nicked CR 127 hCG (Birken, S., et al., 1993). In addition, JAR cell type hCG is known to contain a similar array of carbohydrate moieties. It was found to be recognized by B152 similar to the early pregnancy isoforms in healthy pregnancies. The observation that the hCG isoform produced by JAR cells in culture (B152/B109 ratio) is similar to that found in early pregnancy hCG isoforms supports the hypothesis that the production of a type of hCG with a particular glycosylation pattern is a prerequisite for a viable pregnancy. This glycosylation pattern is not characteristic of the hCG of later pregnancy.

A variety of pregnancy disorders are testable. One category of patients consists of those women who experience a high rate of recurrent abortions. Even in populations with no known fertility problems, the total rate of pregnancy loss is 32% (EPL plus clinically recognized abortion) (Wilcox, A. J., et al., 1988). The risk of recurrent abortion increases with the number of spontaneous abortions experienced in the past, reaching an incidence of 32% after three consecutive abortions. (Hill, J. A., and Anderson, D. J., 1990). Probable causes of recurrent spontaneous abortion, comprising genetic, infectious, hormonal imbalance, or immunologic factors can be established in less than 60% of all spontaneous abortions, leaving 40+% of spontaneous abortions with a completely unestablished etiology. These facts, taken together with reports establishing that the administration of exogenous hCG can be an effective therapy in subjects with a history of recurrent spontaneous abortion (Quenby, S., and Farquharson, R. G., 1994; Harrison, R. F., 1985) lends support to the hypothesis that a disproportionate production of the ineffective isoforms of hCG in early pregnancy is a causal factor in both early pre-clinical loss as well as in spontaneous abortion.

A second category includes women undergoing embryo transfer. These patients provide several distinct advantages: The patients undergoing this procedure are not treated with crude hCG preparations, making measurement of hCG isoforms easy and decisive since all hCG forms derive from the embryo none from any injected hCG preparations. Second, is the opportunity to monitor the nature of the isoforms from day 9 of a successful pregnancy. Third, is the ability to obtain large volumes of urine to purify the early pregnancy isoforms to determine their structures. Fourth, since pregnancy loss is from 50% to 70% in this population, the loss can be defined as due to lack of the essential hCG isoform recognized by B152 or due to other causes. Comparison of early pregnancies

in populations of women not undergoing *in vitro* fertilization procedures with those undergoing embryo implantation can, thus, assess whether pregnancy loss situations present similar or different patterns of hCG isoforms during the 5 process. The mechanism of pregnancy loss in the general population as compared with the much higher rate of embryo loss in IVF programs may be different. Additionally, it has been established that the hCG produced in choriocarcinoma has differences in carbohydrate structures, sialic acid content 10 and biological activity (Wide, L., and Hobson, B., 1987; Elliot, M., et al., 1997; Husa, R. A., 1987). Since both the B151-B604\* and B152-B207\* assays incorporate monoclonal antibodies raised against an immunogen derived from choriocarcinoma, specimens may be evaluated from patients 15 with gestational trophoblastic disease in order to determine whether the above assays recognize the hCG produced in these conditions with greater sensitivity and specificity than do assays based on the hCG of normal pregnancy, as is apparently the case for the hCG produced in testicular and ovarian 20 cancer.

There are few reports of changes of carbohydrate content of hCG-related molecules during pregnancy. Blithe and colleagues studied free alpha subunit of hCG whose 25 carbohydrate content differs from that of alpha within hCG by additional carbohydrate antennae and fucose. The carbohydrate of free alpha becomes increasingly complex in terms of more branches and higher content as pregnancy proceeds. It has also been reported that the quantity of 30 fucose increased in both hCG and in free alpha as pregnancy proceeded (Skarulis, M. C., et al., 1992). Thus, the literature indicates increasing content and complexity of carbohydrate of hCG and free alpha subunits. However, immunological data using the B152 monoclonal antibody, 35 implies a progression to simpler carbohydrate content during

pregnancy. Since the beta COOH-region's O-linked carbohydrates may be involved in the epitope recognized by B152, it is conceivable that the carbohydrate structures of this region may be altered in a different pattern from the 5 N-linked glycans studied by Blithe and colleagues (Skarulis, M. C., et al., 1992; Blithe, D. L., and Iles, R. K., 1995). Data from Skarulis et al. indicate that heterodimeric hCG may contain additional fucose but do not provide data that this late pregnancy hCG becomes hyperglycosylated as does free 10 alpha.

Other studies indicated that the forms of hCG during EPL likely differ in biological activity from those hCG isoforms in successful pregnancies (Ho, H.-H., et al., 1997). The in 15 vitro bioassay employed in those studies are unsuitable for large-scale studies and are not as reliable as the immunoassays described herein. Furthermore, it is likely that *in vivo* assays may give different results since *in vitro* and *in vivo* assays sometimes give completely disparate 20 results. In this case, *in vivo* and clearance assays are most important in order to identify whether the hCG isoforms are truly more potent in the whole animal and to identify the reasons for the increased potency. Thus *in vitro* and *in vivo* bioactivities of the early pregnancy isoforms of hCG are 25 highly significant.

Carbohydrate differences is a widely accepted explanation for variations in biological to immunological ratio such as the forms observed by various studies of EPL (Ho, H.-H., et al., 30 1997). Various studies (Grotjan, H. R. J., and Cole, L. A., 1989; Hoermann, R., 1997; Stanton, P. G., et al., 1993; Szkudlinski, M. W., et al., 1995, Thotakura, N. R., et al., 1994; Szkudlinski, M. W., et al., 1993), have shown that sialic acid differences are an explanation for such 35 heterogeneity in biological activities of glycoprotein

hormones. These studies have also confirmed the dogma that *in vitro* biological activities can yield the opposite results from *in vivo* studies because of altered metabolic clearance rates in the latter studies. Thus, more acidic (more highly  
5 sialylated) forms of gonadotropins are more biopotent in the whole animal because of prolonged circulating half-lives. The same molecules may appear less potent in *in vitro* assays due to greater acidity, greater negative sialic acid content. Hoermann et al. (Hoermann, R., et al., 1997) demonstrated the  
10 exclusion of many of the acidic circulating hormone forms from the urine, thus, prolonging their half-lives. The pI pattern of normal pregnancy as well as trophoblastic cancer hCG in serum is quite different from that of urine. Since the studies described herein indicate that EPL hCG isoforms  
15 have reduced *in vitro* biological activity, this finding cannot be explained solely by what is known of biological activity and sialic acid content. Early pregnancy isoforms recognized by monoclonal antibody B152 may be more potent *in vivo* by virtue of prolonged half-life they may then display  
20 increased signal transduction at the receptor as well. This may be explained by a hyperglycosylated form of hCG which is not hypersialylated. In this case, the extra sugar portion would help prolong circulating half-life of a more basic pI form of hCG which also has increased *in vitro* bioactivity.

25

**Example 4: Diagnosis of Gestational Trophoblast Disease.**

An important application of the B152 (early hCG isoform)/B109 (late hCG isoform) ratio analysis described herein above is in the very early (and facile) diagnosis of gestational  
30 trophoblast disease. Examples of gestational trophoblast disease include choriocarcinoma or hydatidiform mole. In normal pregnancy, the ratio of B152/B109 of the two isoforms of hCG rapidly decreases, eventually inverting. In gestational trophoblast disease including choriocarcinoma or  
35 hydatidiform mole, the ratio is initially higher than found

in normal pregnancy, but does not diminish during the course of the apparent pregnancy. This approach provides a highly sensitive and specific diagnostic marker for gestational trophoblast disease.

5

Other pregnancy disorders in which hCG levels are abnormally high or abnormally low include Down's syndrome or other aneuploid pregnancies, ectopic pregnancy, preeclampsia, and intra-uterine growth retardation. Because the hCG production  
10 in these conditions is quantitatively abnormal compared with normal pregnancy, an altered ratio of the hCG isoforms identified by B152 (early hCG isoform) and B109 (late hCG isoform) can be detected. Thus, the dual isoform analysis (B152/B109) further provides a method for diagnosing  
15 pregnancy disorders and gestational trophoblast disease.

### **Materials and Methods**

#### **Hormones**

The non-nicked hCG isolated from the CR127 preparation of hCG  
20 was used as a standard in both assays (Birken et al. 1993). The pituitary hCG was isolated as described (Birken et al. 1996). C5, a 100% nicked hCG having extra sugars on both N- and O-linked carbohydrate moieties, purified from the urine of a choriocarcinoma patient (Elliott et al. 1997), was  
25 supplied by Dr. Laurence Cole (Yale University School of Medicine). Although the C5 immunogen used in the development of B152 antibody was 100% nicked hCG isoform (i.e. had cleavages in the peptide backbone of loop 2 of the b subunit) the antibody did not discriminate nicked from non-nicked hCG  
30 (O'Connor et al. 1998).

The same serial dilutions of non-nicked hCG, pituitary hCG and C5 were used for binding characterization in hCG assays. Hormone concentrations of initial stock standards solutions  
35 were determined by amino acid analysis.

Immunoradiometric assays (IRMA)

The methodology used in the construction and validation of the B109-B108\* assay has been fully described elsewhere (O'Connor et al. 1988). The B152-B207\* assay has also been characterized (O'Connor et al. 1998). Both assays were performed with a slight modification of the published procedure: the capture antibody was adsorbed onto the wells of microtiter plates (Immulon IV, Dynatech, Chantilly, VA) by incubating a 5 µg/ml solution (B109-B108\* assay) or 25 mg/ml solution (B152-B207\* assay) in coating buffer (0.2 M bicarbonate, pH 9.5) overnight at 4 C. The coating antibody solution was aspirated, the plates washed (wash solution: 0.9% NaCl, 0.05% Tween 20) and blocked with a 1% solution of BSA in PBS with 0.1% sodium azide. Following incubation with the BSA solution (minimum 3 hours at room temperature) the blocking solution was removed, the wells again washed with wash solution and 200 µl/well of the appropriate hCG standards were added in phosphate buffer B (PBS with 0.1% bovine gamma globulin and 0.1% sodium azide). After overnight incubation at 4 C, the plates were again aspirated and washed. The 200µl (50,000 cpm-100,000 cpm) of <sup>125</sup>I- labeled antibody was added to the wells which were again incubated for 24 h at 4 C. The tracer was aspirated, the plates washed with wash solution, the individual wells placed in glass tubes and the radioactivity determined in a Packard Cobra gamma counter. Doses were determined by interpolation from a smoothed spline transformation of the data points.

All samples were stored frozen at -20 C prior to assay. Because extreme values of sample pH may interfere with antibody binding, the urine pH was adjusted with 1.0M Tris (pH 9.0), 50µl/ml urine prior to assay, so that the final pH was in the range of 7-7.4 (O'Connor et al. 1988). Intra-assay variation was 6% for both assays, inter-assay variation was 12% for B109-B108\* and 13% for B152-B207\* assays. Sensitivity

(least detectable dose, defined as +2SD from the zero calibrator, was 1 fmol/ml for the B109-B108\* assay and 2.2 fmol/ml for B152-B207\* assay.

## 5 Patients samples

Urine samples from IVF patients were a gift from Dr. L. Cole. They included spontaneous abortion (n=14, range of gestational age 1.8-4.1 weeks from ET- embryo transfer),  
10 ectopic pregnancies (n=7, gestational age 2.3-4 weeks) and normal pregnancy controls (n=65, encompassing the range 0.6 to 5.4 week from ET). Some of the normal pregnancy urine samples throughout the pregnancies were also obtained from Dr. Cole. Others were obtained from the clinical practice of  
15 collaborating physicians at Columbia Presbyterian Medical Center (CPMC) (Total n=103). Matched serum/urine samples from the first (n=12) and the third (n=11) trimesters were provided by Dr. Amalia Kelly at CPMC. Trophoblast disease serum (n=17) and urine (n=28) samples were obtained from Dr.  
20 Cole, but were collected by Dr. Edward Newlands (Charing Cross Hospital, London, UK). All specimen collection protocols were approved by the appropriate Institutional Review Board.

## 25 Statistical Analysis

Polynomial regression models of log transformed hormone ratios were used to describe the relationship between the change in ratio as a function of gestational age in normal  
30 pregnancy. A paired t-test was used to evaluate the relationship between matched serum and urine hormone ratios. Analysis of covariance (ANCOVA) was used to describe the time adjusted relationship of hormone values in ectopic pregnancy and spontaneous abortion to those of normal pregnancy.

### Urine Processing.

Twenty-four hour urine samples are collected from women undergoing embryo transfer as well as women in early natural pregnancy. The urine is refrigerated during the collection procedure. After delivery of the urine to the laboratory, sodium azide is added to 1g/liter. Women undergoing *in vitro* embryo transfer are not pre-treated with hCG. Thus, all hCG which appears in their blood or urine is derived from the embryo (except for the small amounts of pituitary hCG present in all people). Raw urine is freed from particles by centrifugation followed by Pellicon filtration through a 0.45 micron membrane. Next, the procedure is to concentrate the urine with a Pellicon (Millipore) system which concentrates as much as 30 liters to 500ml overnight (4°C) using a 3,000 MW cutoff membrane. Smaller volumes can be concentrated in less than two hours. Next, the urine is desalted and delipidated by passage through a large volume of Sephadex G25 in 0.1 M ammonium bicarbonate. This step greatly increases the binding of CG to immunoaffinity columns. The desalted urinary concentrate is next size fractionated on the Pharmacia HiLoad Superdex 200 and the hCG and hCG subunit peaks are identified by specific immunoassays (O'Connor, J. F., et al., 1994) and the appropriate fractions are pooled and dried. The hCG and hCG subunits are purified from the gel filtered urine concentrate by immunoaffinity on insolubilized hCG antibody columns as described but with the use of either 4M guanadine (0.1M tris acetate, pH 5) or ammonium thiocyanate as eluant to decrease loss of sialic acid from the hormone. Alternatively, hCG is purified by conventional chromatographic procedures, anion exchange and hydrophobic chromatography. The subunits are separated on reverse phase HPLC using a 0.01M sodium phosphate, pH 5 buffer and acetonitrile, after incubation in 4M guanadine, 0.1M tris acetate, pH 5. A third method is final purification and separation of the hCG subunits on SDS PAGE

electrophoresis followed by electroblotting to PVDF. The PVDF band can be subjected to protease digestion to release peptides and glycopeptides which can be separated on reverse phase HPLC in neutral pH 5 buffers.

5

Separation of Glycopeptides from Isolated hCG subunits. To facilitate isolation of the glycopeptides from the hCG subunits, the subunits are both tryptic digested and the products of digestion are separated on reverse phase HPLC  
10 (using a pH 5 buffer). This procedure results in removal of the large beta COOH-terminal peptide which contains O-linked sugars. It also releases small, non glycopeptides from both subunits (Pollak, S., et al., 1990, Birken, S., et al., 1987; Birken, S., et al., 1986). Next, the main disulfide-linked  
15 core of each hCG subunit, is reduced and carboxymethylated, and separated on reverse phase HPLC at pH 5. At this stage, large peptides are isolated, including the glycopeptides. Each separated glycopeptide is redigested with trypsin and re-separated on HPLC at pH 5. These glycopeptides are next  
20 employed for two different methods of sugar chain analysis. One method is the approach of releasing the oligosaccharides by enzymatic digestions using PNGase F for the N-linked glycans. The released glycans can be separated from the peptides by ethanol precipitation, desialyated with  
25 neuraminadase, and separated directly on a Dionex Carbopac PA-100 column. Oligosaccharide standards are available from Dionex, Oxford Glycosystems and other companies for calibrating column elution times for various glycans (Hardy, M. R., and Townsend, R. R., 1994, Rohrer, J. S., et al.,  
30 1993, Weitzhandler, M., et al., 1993; Townsend, R. R., et al., 1989). Confirmation of the released structures is obtained by performing carbohydrate compositional analysis of eluted glycan peaks as well as performing digestions with specific glycosidases and rechromatographing the modified  
35 glycan on the Dionex system (Hardy, M. R., and Townsend, R. R., 1994; Rohrer, J. S., et al 1993; Weitzhandler, M., et

al., 1993; Townsend, R. R., et al., 1989; Townsend, R. R.,  
et al., 1991; Townsend, R. R., et al., 1989; Hardy, M. R.,  
and Townsend, R. R., 1989; Townsend, R. R., et al., 1988;  
Hardy, M. R., et al, 1997; Hardy, M. R., and Townsend, R. R.,  
5 1988; Dionex, 1997; Spellman, M. W., 1990; Kumarasamy, R.,  
1990). The newly modified glycan can be observed to elute at  
the same time as the appropriate standard oligosaccharide  
and, in addition, the released monosaccharide can frequently  
be identified as well (Dionex, 1997). Structure  
10 determination is facilitated by the use of specific  
glycosidases for branch chain cleavage as well as for  
digestion of individual sugars from each of the branch  
chains. For example, Endo H cleaves high mannose type and  
hybrid oligosaccharide chains while glycosidase Endo F2  
15 cleaves biantennary complex types and PNAase F cleaves tri  
and tetra-antennary chains down to the N-Asn bond.

Competitive receptor binding and in vitro bioassay.

Bioassays are performed with recombinant-engineered CHO cells  
20 transfected with the human receptor to LH/CG Cells are  
maintained in Ham's F-12 medium, 4mM Glutamine, 400ug/ml G418  
(Gibco), 5% fetal calf serum, 100IU/ml penicillin, 100ug/ml  
streptomycin. The cells are removed from the flask surface  
by versene only.

25

A competitive receptor assay constructed as follows: The  
receptor binding assay mixture contains 100 ul of the  
appropriate dilution of serum/urine samples or hCG dilutions  
for standard curve, 100ul of <sup>125</sup>I-hCG (50,000-100,000cpm) in  
30 buffer A(PBS/0.1%BSA) and 100ul of CHO cells (2x10<sup>5</sup> cells in  
PBS). The mixture is incubated at 37°C with slight shaking  
followed by centrifugation for 10 minutes at 750xg. The  
supernatant is aspirated and the cell pellet is counted in  
gamma-counter.

35

In vitro bioassay. Transfected CHO cells are seeded (200,000 cells/well) into a 24 well plate in culture medium and incubated for 2-3 days until the cells reach confluence. Non-transfected CHO cells are included to monitor non-specific response. The medium is removed and replaced with medium containing 1 mM isobutylmethylxanthine with appropriate dilutions of tested serum or urine. The plates are incubated at 37°C for two hours. The supernatant is removed, and the wells washed with Hank's balanced salt solution. The intracellular cAMP is extracted with 95% ethanol, which is diluted 1:5, (or up to 1:40, depending on cAMP content) in assay buffer provided by the cAMP kit (New England Nuclear). cAMP assay is performed according to manufacturer's instructions. Response is normalized to well protein content (BCA protein assay kit, Pierce, Rockford, IL).

In vivo bioassay is determined by the uterine weight assay in immature female mice, following the procedure of Wide and Hobson (Wide, L., and Hobson, B., 1987). The mice are injected subcutaneously with one third of the total dose of gonadotropin on three consecutive days and killed 72 hours after the first injection. Uteri are dissected free from mesentery, fat and oviducts, blotted to remove intrauterine fluid and weighed to the nearest 0.1mg. Five to ten mice are used at each of these dose levels. The hCG standard preparation used is a nicked hCG. This material may be run concurrently with specimens isolated from first and third trimester pregnancy. Sham saline injection may be used as a control. The response signal is the log mouse uterine weight.

Clearance of hCG isoforms. The clearance of hCG is determined in the rat. Blood (200ul/sample) is obtained at 0, 120, 240, 360 and 480 minutes post injection, from an

indwelling catheter in an catheterized external jugular vein, following the procedure described by Newman et al. (Newman, C. B., et al., 1985) and Brown and Hedge (Brown, M. R., and Hedge, G. A., 1972). Briefly, adult male Sprague-Dawley rats (Charles River Laboratories, Wilmington MA), wt 175-225g, are given free access to food and water. Rats are handled for acclimatization for one week after arrival, and several days before the hCG infusion, the rats are cannulated under pentobarbital anesthesia. A 21 gauge stainless steel cannula is inserted into the one external jugular vein. The placement of the catheter allows for the collection of blood from the unrestrained, unstressed rat. After the animals have recuperated from the cannula implacement, an hCG isoform is injected (10 µg/ml sterile saline) through the cannulated vein. Blood samples are obtained at the four time intervals listed above. The blood is allowed to clot and the serum separated and stored at -80°C for immunometric assays specific for different hCG isoforms.

Clearance rate of the isoforms of hCG from the circulation of the rat are estimated by computer fitting the concentration data to an equation of the general form:  $\text{Concentration} = Ae^{-\alpha t} + Be^{-\beta t}$  at time  $t$ ;  $A$  and  $\alpha$  are parameters of the rapid component and  $B$  and  $\beta$  are parameters of the slow component. The metabolic clearance rate (MCR) is calculated as  $\text{MCR} = \text{Dose} / (A/\alpha + B/\beta)$  and the initial volume of distribution is calculated from  $V_d = \text{Dose} / (A + B)$ . The MCR is normalized to body weight for statistical analysis, which is performed using ANOVA with Duncan's range test for determination of significance (Cassals, J. W., et al., 1989).

Mice. The mouse species used in the experiments described herein are Balb/c mice, aged 12-20 weeks old and adult Sprague-Dawley rats of either sex. Mice used for the production of monoclonal antibodies through ascites and for

the determination of *in vivo* biological activity as described. Balb/c mice are used because hybridoma cell lines were developed using Balb/c splenocytes.

**EXPERIMENTAL DETAILS FOR THE SECOND SERIES OF EXPERIMENTS**

Human Chorionic Gonadotropin exists in blood and urine as a variety of isoforms one of which contains peptide bond cleavages within its beta subunit loop 2 is referred to as nicked hCG (hCGn). This hCG isoform appears to be more prevalent in the urine of patients with certain malignancies and possibly in some disorders of pregnancy. Until now, only indirect immunoassays could be used to quantify hCGn. We report the development of two monoclonal antibodies to a form of hCGn isolated from a choriocarcinoma patient. This hCG isoform was not only 100% nicked but also contained 100% tetrasaccharide-core O-linked carbohydrate moieties in its beta COOH-terminal region. Two-site immunometric assays have been developed using these new antibodies, B151 and B152. The former exhibits good specificity for hCGn independent of the source of the hCGn, that form excreted by choriocarcinoma patients or the form of hCGn from normal pregnancies. The latter antibody, B152, is sensitive to the carbohydrate moieties and possibly other differences in hCG isoforms but not for nicking of the beta subunit. These two immunometric assays provide potential novel diagnostic tools for direct measurement of hCG isoforms which could not be accurately quantified earlier before development of the assays using these newly generated antibodies.

25

Human Chorionic Gonadotropin (hCG) is a glycoprotein hormone produced by trophoblast cell of the placenta. The measurement of this hormone in blood or urine is the basis of all pregnancy tests. It is secreted by the trophoblast starting very early in pregnancy and functions to maintain steroid production by the corpus luteum until the placenta takes over that function later in pregnancy (1). There has been much recent interest in measurement of the hormone or its subunits or fragments for purposes other than the diagnosis of pregnancy such as monitoring of therapy for hCG-

secreting malignancies (1), tests for indication of Down syndrome or other genetically-abnormal pregnancies (2-11), ectopic pregnancies (12), etc. HCG appears in urine in a variety of forms including free subunits, heterodimeric hCG with peptide bond cleavages in loop 2 of its beta subunit (known as nicked hCG), free nicked beta subunits, and the beta core fragment (13-16). The nicked form of heterodimeric in hCG has been reported to be present in blood as well as urine and is known to have much reduced immunochemical recognition by some antibodies directed to heterodimeric hCG as well as greatly reduced biological activity (13, 16-17). There are reported associations between increased nicking of hCG with certain hCG-secreting malignancies (18,19).

There are no satisfactory direct immunoassays for the nicked form of hCG nor for the nicked form of free hCG beta subunit. All measurements to date have been conducted with subtractive assay procedures or immunoassays which include scavenger antibodies (20-22). We report here the development of two antibodies of distinct specificity, using as immunogen the form of hCG produced by a single individual with choriocarcinoma. This particular hCG isoform was 100% nicked and hyperglycosylated both in its N and O-linked carbohydrate moieties. One resultant antibody, B151, displays significant preference towards binding to its choriocarcinoma hCG immunogen and does not recognize the nicked characteristic. Immunoassays developed using B152 are of special interest in their enhanced recognition of hCG isoforms more prevalent in preeclampsia and Down syndrome (23,24). B152 also appears to detect an hCG isoform associated with healthy pregnancies as juxtaposed to those pregnancies destined to fail early which have little of this isoform (25).

**Results:**Characteristics of antibodies

A variety of hCG isoforms were employed to characterize the new antibodies described in this report and the nomenclature and characteristics of each of the reagents employed is summarized in Figure 11. The carbohydrate groups in these hCG isoforms as well as the percent nicking were analyzed in an earlier study (26) and are directly relevant for defining the nature of these new antibodies in this report.

10

Two antibodies designated B151 and B152 were selected by the use of radiolabeled hCG isoforms, chorioCG C5 and pregnancy hCG CR127. Each displayed preferential binding to C5 as compared to CR 127 since this was the selection criterion. However, upon performing liquid phase immunoassays and calculating affinity constants, it was clear that these two antibodies were very different in specificity (Figure 8). It was found that antibody B151 had one order of magnitude higher affinity both for C5, which is nicked and hyperglycosylated choriocarcinoma hCG, and for CR127 hCGn (813) as compared to CR 127 hCG or nick-free CR 127(814) (see Figure 11 for reagent descriptions). B151 was clearly an antibody with a strong preference for binding to various forms of nicked hCG. Antibody B152 was different in that although it displayed one order of magnitude preference for C5 hCG over CR127 hCG, it recognized the nicked and non-nicked forms of CR 127 hCG, hCG derived from normal pregnancies, to an equal extent.

30 Liquid Phase Assays

Figure 8 shows potency comparisons of liquid phase immunoassays of both B151 and B152 antibodies comparing competitors: 1. standard CR 127 pregnancy hCG (which has a 20% content of nicked hCG); 2. C5 chorio CG (100% nicked and hyperglycosylated); 3.813, nicked CG made from CR 127 by

35

purification, and 4.814, non-nicked hCG derived from CR 127. The labeled ligand was C5 chorio CG. It is apparent that B151 (Fig 8A) shows a preference for nicked forms of hCG. C5 chorio-CG or 813 hCGn bind with similar affinities. The 5 slightly lower potency of 813 hCGn may be ascribed to its 20% contamination with non-nicked hCG. B152 only shows a preference to C5, the hyperglycosylated chorio CG (Fig 8B). 813 hCGn is no more potent a competitor than nick-free 814 hCG.

10

#### Immunometric Two Site Assays

A variety of two site antibody formats were tested. Figure 9 displays these results. It is apparent that B151 cannot bind simultaneously with antibodies (designated by us a site 15 IV) (27) to the beta subunit and beta subunit core (B201 and B204) nor with antibodies directed towards the determinant which exists in heterodimeric hCG as represented by antibody B109 (site III, to which A109 also belongs) (27). In contrast, a general beta antibody which binds to the most 20 common and potent hCG antigenic site previously designated by us as site II (B108 or B207) binds well simultaneously with both B151 and B152 antibodies. B152 binds simultaneously to all antibodies tested except for those to the beta COOH-terminal region (CTP) (28) in contrast to B151 25 which binds well to CTP antibodies. B151 may represent a newly revealed hCG epitope which only exists on nicked hCG as reported in this manuscript.

Using B152 as capture and B207 or B108 as detection antibody, 30 produces an assay which measures all normal pregnancy forms of hCG (both intact and nicked and beta subunit) to a similar extent but prefers binding to the form of hCG or beta subunit from its immunogen, C5. As predicted earlier from the liquid phase studies, this assay does not prefer nicked forms of hCG 35 but hyperglycosylated forms of hCG such as C5. B152 and CTP 104 as well as several other monoclonal antibodies to the

COOH-terminal region of HCG beta cannot bind simultaneously to C5, implying that this region is part of or very close to the epitope of B152. These data taken together with the apparent B152 preference for hyperglycosylated hCG, implies  
5 that the carbohydrate of the CTP region may be part of the B152 epitope. Further studies of the behavior of B152 in two-site assays confirm this hypothesis as detailed below.

Characteristics of the B152-B207-I<sup>125</sup> radioimmunometric two-  
10 site assay

In order to better understand what this assay is measuring, we compared the relative binding potencies of a series of isoforms of hCG shown in figure 9 (also see methods): 1. C5, choriocarcinoma hCG 2. 814, non-nicked hCG. 3. 813, nicked  
15 hCG (80% nicked). 4. M4 mole-derived hCG, 98% nicked hCG with negligible hyperglycosylation. 5. M1A hCG, non-nicked and not significantly hyperglycosylated but missing 80% of the hCG beta COOH-terminus. The B152 two-site assay prefers to bind  
20 both 813 and 814, nicked and non-nicked hCG of normal pregnancy. This confirms that B152 does not display significant preference for the nicked form of hCG but rather for the form with carbohydrate differences. This is also confirmed by the potency of M4 which is also 100% nicked as  
25 is C5 but is not hyperglycosylated and displays a potency similar to CR127 hCG whether nicked or non-nicked. M1A is the least potent ligand and is the only one missing most of its beta COOH-terminal peptide confirming the role of this region in the B152 epitope.

30 In order to further explore the nature of the B152 binding site, a commercially available peroxidase-labeled general hCG $\beta$  antibody (4001) was employed as a detection antibody in a two-site enzyme immunometric system. Eight different hCG forms were evaluated in this system illustrated in Figure 14.  
35 Results are analyzed in terms of relative immunopotency (based on the slope of the regression line) in Figure 10.

Linear regression correlation analysis was performed to compare the relationship of the immunopotencies of preparations 814, C5, M4, C7 and P8 one at a time with the carbohydrate differences as well as nicking differences among the 5 heterodimeric isoforms of hCG. The correlation results for each comparison are as follows: 1. Tetrasaccharide O-linked core:  $R^2=0.9147$   $P=0.0108$ , significant; 2. Triantennary branched moieties N-linked on  $\beta$ :  $R^2=0.8853$   $P=0.0171$ , significant; 3. Sialic acid O-linked:  $R^2=0.3062$   $P=0.3332$ , not significant; 4. Sialic acid N-linked on  $\beta$ :  $R^2=0.2289$   $P=0.4149$ , not significant; 5. Percent nicking in  $\beta$  subunit:  $R^2=0.0984$   $P=0.6072$ , not significant.

To summarize this analysis, the immunopotency of hCG isoforms measured with B152 as a capture antibody in a two site assay correlates best with hyperglycosylated core moieties of both O and N-linked amino acids but not with sialic acid residues not with the degree of nicking of the  $\beta$  subunit.

20

### Discussion

The various isoforms of hCG have received increasing attention during recent years for their potential diagnostic value in problem pregnancies such as Down syndrome, preeclampsia, trophoblastic diseases and early pregnancy loss. Nicked hCG at high concentrations has been associated with trophoblastic disease and other abnormal states of hCG production. Although nicked hCG has been measured by a variety of qualitative techniques such as immunoblotting, as well as by direct isolation and sequence analysis of such hCG isoforms from urine, it is only recently that some investigators have been measuring nicked forms of hCG by a variety of subtractive immunoassays or non-specific hCG assays with the addition of scavenger antibodies since direct, relatively specific antibodies were not available

(22;29). We had earlier shown that our commonly used antibody, B109, to heterodimeric hCG reacted poorly to nicked forms of hCG and this antibody was employed by other groups in subtractive methods in attempts to quantify the content of nicked hCG in blood and urine(22;29).

In this report, we described development of two antibodies produced using choriocarcinoma-derived nicked hCG as immunogen. This led to a direct assay for nicked hCG. By employing a nicked, hyperglycosylated form of hCG from a single choriocarcinoma patient, we have developed two antibodies with distinct specificities. One antibody (B151) prefers binding to nicked forms of hCG, regardless of the origin of the hCG molecule (normal pregnancy or choriocarcinoma). Nick-free CR127 hCG has the lowest affinity to this antibody as would be expected for a nick-directed antibody. While B151 was directed towards an epitope dependent upon peptide bond cleavages in beta loop 2, B152 binding was not affected by peptide bond cleavages within this loop. Since the main difference between C5 hCG and CR 127 hCGn was the hyperglycosylation of the former, it was inferred that B152 was chiefly a carbohydrate-directed antibody. When B151 is used as capture antibody and virtually any general beta antibody as detection antibody in a two site assay, little cross-reaction with hLH is observed. B151 cannot bind simultaneously with antibodies that are directed chiefly to the hCG beta core region (site IV such as B201 and B204) nor can the antibody bind with site III antibodies directed to heterodimeric hCG such as B109 or A109 but it can bind at the same time as antibodies to the beta COOH-region. B151 may represent a new hCG epitope revealed after nicking of beta loop 2. The creation of new epitopes by nicking of the hCG molecule has been reported by others (30).

The second antibody, B152, preferentially binds hCG with the type of carbohydrate modifications prevalent in choriocarcinoma CG C5 regardless of the state of the nicking of the polypeptide chain. This was shown by the measurement of the relative immunopotencies of several well characterized hCG isoforms as compared to their contents of N and O-linked carbohydrate moieties, sialic acid, and percentages of nicking. A significant correlation of B152 binding to hyperglycosylated hCG isoforms but not to those with sialic acid or nicking differences was demonstrated. Antibody B152 appears to have at least partial specificity towards the hCG beta COOH-terminal region. This is shown by failure of a monoclonal antibody to bind simultaneously with beta COOH terminal antibodies such as CTP 104. In studies of B152 as a two site assay, it was shown that hCG isoform MIA which is missing most of its COOH-terminal region binds poorly to B152.

Each antibody has a different application in accordance with its specificity. Up to the present time, only indirect assays were available to quantify nicked hCG, such as subtractive assays or assays with scavenger antibodies added. Development of B151 permitted formulation of direct assays for nicked hCG which have been applied in studies of early pregnancy (25,31). These measurements may have diagnostic applications for certain cancers (18-19) and in the detection of Down Syndrome (22). Application of the B152 antibody resulted in development of assays which can detect differences in the carbohydrate portion of hCG. Major potential applications of this antibody include detection of Down Syndrome (23) and recognition of pregnancies destined for early pregnancy loss (25,32) This antibody is unusual since it is rare for carbohydrate discriminating antibodies to be developed to hCG. The only earlier such development was antibodies to the hCG beta COOH-terminal region by the use of the isolated peptide bound to carrier (28,33). It is

of interest that at least part of the epitope of B152 is also directed towards the beta COOH-terminal region. As reported earlier, the C5 choriocarcinoma form of hCG was the only such hCG displaying 100% hexasaccharide structure on its O-Serine 5 linked carbohydrate moieties in the beta COOH-terminal region. This may have resulted in production of this rare antibody. A second choriocarcinoma derived hCG isoform, C7, displayed 69% of this same O-linked core hexasaccharide structure and proved to be of similar potency as C5 with the 10 B152 epitope.

In conclusion, we have produced two novel monoclonal antibodies each with potential clinical utility: B151 which can be used to measure nicked forms of hCG with better 15 specificity than any antibody reported. B152 which is a unique antibody to a choriocarcinoma form of hCG and can discriminate hyperglycosylated from standard pregnancy hCG.

### **Materials and Methods**

#### 20 Hormones

Nick-free hCG (814) and nicked hCG (813) were prepared from pooled urine standard hCG (batch CR 127) by hydrophobic chromatography, as described previously (14). C8 hCG was purified from an individual with normal pregnancy, M1 and M4 25 hCG were both purified from individuals with gestational trophoblastic disease (hydatidiform mole), and C5 and C7 hCG from individual with malignant trophoblastic disease (choriocarcinoma), as described elsewhere (16,26). The N-terminal peptide sequences of the separated  $\alpha$ - and  $\beta$ - 30 subunits of CR127; P8, M1, M4, C5 and C7 hCG and complete N- and O-linked oligosaccharide structures have recently been published (26). Two peaks were observed by Sephacryl S100 HR chromatography during purification of hCG preparation M1. The peak eluting in the position of standard hCG (M1), and 35 the peak eluting later (M1A) were purified separately. M1

and M4 hCG were both purified from individuals with hydatidiform mole. They were isolated as described earlier (16).

#### 5 Immunogens

Choriocarcinoma hCG designated C5 was isolated from a single patient as described earlier (16) and its complete carbohydrate analysis has recently appeared (26).

#### 10 Immunization of Mice

Mice were immunized intraperitoneal with choriocarcinoma hCG preparation C5 diluted in saline (600 µg/mouse) and mixed 1:1 with Freund's adjuvant. After 3 consecutive immunizations (50 µg/mouse) at 3-4 week intervals, animals were retested for 3 months and then given an intraperitoneal booster immunization (34). Ten days after the last booster, the sera from mice were tested for binding (in liquid phase radioimmunoassays) to both iodinated CR 127 hCG and to iodinated C5 hCG.

20

#### Liquid Phase Assays

Liquid phase assays were performed using a solution of 80µl 0.3M PBS with .02% sodium azide, 50 µl tracer, 20µl normal mouse serum, 50µl 1% horse serum free of gamma globulin for 25 titrations ( when competition studies were performed, 50µl of competitor solution were substituted to generate dose/response or Scatchard curves), 100µl of diluted sera followed by incubation 1 hr at 37C and then overnight at 4C. Next day 100µl rabbit anti-mouse sera was added, incubated 30 10 min at 37C, then 1 hr at ambient, then centrifuged, supernatant aspirated, and pellets counted. The mouse whose antisera had the greatest discrimination between binding of radiolabeled C5 and radiolabeled CR 127 hCG was sacrificed and their spleen's were used for hybridoma production. The

methods and materials used for the fusion work were described earlier (27;34).

#### Characterization of Antibodies

5 The cloned monoclonal antibodies, B151 and B152 were each studied in the liquid phase competitive radioimmunoassays using C5 as radiolabeled ligand and C5 and CR127 hCG as competitors. The method is as described under liquid phase assay method. Affinity constants were calculated by  
10 Scatchard plots as described earlier (35-37).

#### Two Site Assays

Two site assay testing was conducted at Yale University (enzyme immunometric) and Columbia University  
15 (radioimmunometric) as described earlier (24,37). Briefly, Immulon microtiter wells are coated with capture antibody, at a titer determined to provide the best combination of sensitivity and range. The plates were then washed and then blocked with 1% BSA in PES. After a further wash standards,  
20 clinical samples and controls were added to the coated wells (200 $\mu$ L/well). Plates are incubated, then samples are removed and plates washed. Labeled antibody (either radiolabeled or peroxidase labeled) is then added, and plates incubated further. The response variable was generated by a gamma  
25 counter (Packard Instruments Cobra) or by absorption spectrophotometry as appropriate. A cubic spline curve is generated for standard values, and sample values read from this curve. Regression lines and all graphs were created using Sigmaplot 4.01 from SPSS software, Chicago, Ill.  
30 Linear regression analysis of immunopotency) as compared to each of the carbohydrate differences and nicking differences of the hCG isoforms was accomplished in Instat 1998, GraphPad Software, Inc., San Diego California USA.

### Third Series of Experiments

The data plotted in Figures 15, 16 and 17 were derived from IVF patients who had collected first morning void urines daily, commencing with the day of embryo transfer. These 5 subjects had participated in the "donor egg" variant of IVF therapy; i.e., the recipients did not receive exogenous hCG.

A total of 22 cycles were evaluated. In 10 of these cycles, no hCG was detected, indicating that the blastocyst failed to implant in the endometrium. There were five early pregnancy losses, in which a pregnancy was established but failed to carry to term. There were three normal pregnancies with multiple (2-3) fetuses and four singleton pregnancies.

15 The multiple fetus pregnancies produced higher quantities of hCG than did the singleton pregnancies, as would be expected. However, in both categories of successful pregnancies, there was clearly an inversion of the B152/B109 isoform ratio, while in the failing pregnancies, this ratio remains 20 elevated, indicating that the relative persistence of B152 hCG isoform and B109 isn't the predominate isoform.

Ongoing statistical analysis is particularly directed at determination of minimum constellation of determinations 25 necessary to develop a clinically useful discriminant function for the detection of problem pregnancies at a point where useful discriminant intervention (quite possibly the administration of exogenous hCG) would be effective.

30 Loss of patient #10 (Figure 17D) best exemplifies this; her pregnancy was not lost until after sample collection had ceased. And on the basis of hCG concentration, one could not predict a trouble pregnancy - but with the comparison of ratio with concentrations would predict the problem when 35 other indicators appear to be normal.

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**What is claimed is:**

1. A method of predicting pregnancy outcome in a subject by determining the amount of an early pregnancy associated molecular isoform of hCG in a sample comprising:
- 5
- (a) contacting a sample with an antibody which specifically binds to the early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG;
- 10
- (b) measuring the amount of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample; and
- 15
- (c) comparing the amount early pregnancy associated molecular isoform of hCG in the sample determined in step (b) with either (i) the amount determined for temporally matched, normal pregnant subject(s) or (ii) the amount determined for non-pregnant subject(s), wherein the relative absence of the early pregnancy associated molecular isoform of hCG in the sample indicates a negative outcome of pregnancy for the subject.
- 20
- 25
- 30 2. A method of predicting the likelihood of a negative pregnancy outcome in a female subject comprising:
- (a) contacting a sample from the subject with a capture antibody which specifically binds to an early pregnancy associated molecular isoform of hCG under conditions permitting formation of a
- 35

complex between the antibody and the early pregnancy associated molecular isoform of hCG;

- 5 (b) contacting any complex formed in step (a) with a labelled detection antibody under conditions permitting binding to the complex the capture antibody and the hCG isoform;
- 10 (c) measuring the amount of labeled detection antibody bound to the complex so as to thereby determine the amount of the early pregnancy associated molecular isoform of hCG in the sample; and
- 15 (d) comparing the amount early pregnancy associated molecular isoform of hCG in the sample determined in step (b) with the amount determined for a normal pregnant subject, wherein the relative absence of the early pregnancy associated molecular isoform of hCG in  
20 the sample indicates a negative outcome of pregnancy for the subject.
3. The method of claim 1, step (a) further comprising a  
25 second antibody which specifically binds to hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG.
- 30 4. The method of claim 3, wherein the capture antibody is B152.
5. The method of claim 4, wherein the detection antibody  
35 is B207.

6. A method of predicting the likelihood of a negative pregnancy outcome in a female subject comprising:
- 5 (a) contacting a sample from the subject with a capture antibody which specifically binds to an early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG;
- 10 (b) measuring the amount of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample; and
- 15 (c) comparing the amount measured in step (b) with the amount determined by contacting the same sample with a second capture antibody which specifically binds to intact non-nicked hCG without substantially cross-reacting with
- 20 said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG and a second antibody, wherein a high ratio of amounts determined for said first capture antibody relative to the second capture
- 25 antibody indicates a positive outcome of pregnancy for the subject, a low ratio indicates a negative outcome of pregnancy for the subject.
- 30
7. The method according to claim 5, wherein the second capture antibody is B109 and the second antibody is B108.
- 35 8. The method of claim 1, step (c) comprising comparing the amount early pregnancy associated molecular

- 5 isoform of hCG in the sample determined in step (b) with either (i) the amount determined for temporally matched, normal pregnant subject(s) or (ii) the amount determined for non-pregnant subject(s), wherein amounts of the early pregnancy associated molecular isoform of hCG in the sample similar to amounts of early pregnancy associated molecular isoform of hCG in temporally matched pregnant samples indicates a positive outcome, amounts of early pregnancy associated molecular isoform of hCG in the sample similar to amounts of early pregnancy associated molecular isoform of hCG.
- 10
9. The method of claim 1, wherein the sample is a urinary sample or a blood sample.
- 15
10. The method of claim 1, wherein the sample is an aggregate sample taken from at least one day.
11. The method of claim 1, wherein the antibody is labeled with a detectable marker.
- 20
12. The method of claim 11, wherein the detectable marker is a radioactive isotope, enzyme, dye, magnetic bead, or biotin.
- 25
13. The method of claim 12, wherein the radioactive isotope is  $I^{125}$ .
14. A method of predicting pregnancy outcome in a subject by determining the amount of an early pregnancy associated molecular isoform of hCG in a sample comprising:
- 30
- (a) contacting a capturing antibody which specifically binds to the early pregnancy associated molecular isoform of hCG with a solid
- 35

matrix under conditions permitting binding of the antibody with the solid matrix;

- 5 (b) contacting the bound matrix with the sample under conditions permitting binding of the antigen present in the sample with the capturing antibody;
- 10 (c) separating the bound matrix and the sample;
- (d) contacting the separated bound matrix with a detecting antibody which specifically binds to hCG under conditions permitting binding of antibody and antigen in the sample;
- 15 (e) measuring the amount of bound antibody on the bound matrix, thereby determining the amount of early pregnancy associated molecular isoform of hCG in the sample;
- 20 (f) comparing the amount early pregnancy associated molecular isoform of hCG in the sample determined in step (e) with either (i) the amount determined for temporally matched, normal pregnant subject(s) or (ii) the amount determined for non-pregnant subject(s), wherein amounts of the early pregnancy associated molecular isoform of hCG in the sample similar to amounts of early pregnancy associated molecular isoform of hCG in temporally matched pregnant samples indicates a positive outcome, amounts of early pregnancy associated molecular isoform of hCG in the sample similar to amounts of early pregnancy associated molecular isoform of hCG in the non-pregnant samples indicates a negative outcome of pregnancy for the subject.
- 25
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- 35

15. The method of claim 14, further comprising:  
(a) removing of the sample from the matrix; and  
(b) washing the bound matrix with an appropriate  
5 buffer.
16. The method of claim 14, wherein the capturing  
antibody is B152.
- 10 17. The method of claim 14, wherein the detecting  
antibody is B207.
18. The method of claim 14, step (a) further comprising  
a second capturing antibody which specifically binds  
15 to intact non-nicked hCG without substantially cross-  
reacting with said antibody under conditions  
permitting formation of a complex between the  
antibody and the early pregnancy associated molecular  
isoform of hCG.
- 20 19. The method according to claim 18, wherein the second  
capturing antibody is B109.
20. The method of claim 14, step (d) further comprising  
25 a second detecting antibody which specifically binds  
to hCG without substantially cross-reacting with  
said antibody under conditions permitting formation  
of a complex between the antibody and the early  
pregnancy associated molecular isoform of hCG.
- 30 21. The method of claim 14, step (f) comprising comparing  
the amount of the early pregnancy associated  
molecular isoform of hCG determined in step (e) for  
said antibody with the amount determined in step (b)  
35 for the second antibody, wherein a high ratio of  
amounts determined for said antibody relative to the

second antibody indicates a positive outcome of pregnancy for the subject, a low ratio indicates a negative outcome of pregnancy for the subject.

- 5 22. The method of claim 14, wherein the sample is a urinary sample or a blood sample.
23. The method of claim 14, wherein the sample is an aggregate sample taken from at least one day.
- 10 24. The method of claim 14, wherein the antibody is labeled with a detectable marker.
- 15 25. The method of claim 24, wherein the detectable marker is a radioactive isotope, enzyme, dye, magnetic bead, or biotin.
26. The method of claim 25, wherein the radioactive isotope is I<sup>125</sup>.
- 20 27. A method for determining the amount of early pregnancy associated molecular isoforms of in a sample comprising:
- 25 (a) contacting the sample with an antibody which specifically binds to an early pregnancy associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG; and
- 30 (b) determining the amount of complexes formed thereby determining the amount of early pregnancy associated molecular isoform of hCG in the sample.

28. The method of claim 27, wherein the antibody specifically binds a region of the early pregnancy associated molecular isoform of hCG comprising a carbohydrate moiety.
29. The method of claim 27, wherein the antibody is produced by the hybridoma cell line accorded ATCC Accession No.HB-12467 .
30. The method of claim 27, wherein the antibody is B152.
31. A diagnostic kit for determining the amount of early pregnancy associated hCG is a sample comprising:
- (a) An antibody which specifically binds to an early pregnancy associated molecular isoform; and
  - (b) a solid matrix to which the antibody is bound; and
  - (c) reagents permitting the formation of a complex between the antibody and a sample.
32. The diagnostic kit of claim 31, wherein the antibody is B108 as detection antibody and B109 as capture antibody.
33. The diagnostic kit of claim 31, wherein the antibody is B207 as detection antibody and B152 as capture antibody.
34. The diagnostic kit of claim 31, 32 or 33 further comprising control sample(s) normal pregnant sample(s).

35. The diagnostic kit of claim 31, 32, or 33 wherein the antibody is labeled with a detectable marker.
36. The diagnostic kit of claim 35, wherein the detectable marker is a radioactive isotope, enzyme, magnetic bead, dye or biotin.
37. The diagnostic kit of claim 36, wherein the radioactive isotope is I<sup>125</sup>.
38. An antibody which specifically binds to an early pregnancy associated molecular isoform of human chorionic gonadotropin.
39. The antibody of claim 38, wherein the antibody specifically binds to a region of the early pregnancy associated molecular isoform of human chorionic gonadotropin comprising both a carbohydrate moiety and protein part.
40. The monoclonal antibody of claim 38 designated B152.
41. A hybridoma cell accorded ATCC Accession No.HB-12467, producing the monoclonal antibody of claim 40.
42. The early pregnancy associated isoform of hCG of claim 1.
43. The early pregnancy associated isoform of hCG recognized by the monoclonal antibody of claim 40.
44. A method for detecting non-trophoblast malignancy in a sample comprising:
- (a) contacting a sample with an antibody which specifically binds to the early pregnancy

associated molecular isoform of hCG under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG;

5

(b) contacting the sample with a second detection antibody which specifically binds to intact non-nicked hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG,

10

(c) measuring the amount of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample; and

15

(d) comparing the amount of early pregnancy associated molecular isoform of hCG in the sample determined in step (b) with the amount of early pregnancy associated molecular isoform of hCG in the sample determined in step (c), wherein a positive detection of early pregnancy associated molecular isoform detected in step (b) and a relative absence of the early pregnancy associated molecular isoform of hCG

20

25

detected in step (c) indicates the presence of non-trophoblast malignancy in the sample.

30

45. The method of claim 44, wherein the antibody is B152.

35 46. The method of claim 44, wherein the complex formed in step (a) is detected by a detection antibody.

47. The method of claim 46, wherein the detection antibody is B207.
48. The method of claim 44, wherein the second antibody  
5 is B109.
49. The method of claim 44, wherein the complex formed in step (b) is detected by a detection antibody.
- 10 50. The method of claim 49, wherein the detection antibody is B108.
51. The method of claim 44, wherein the non-trophoblast malignancy is ovarian malignancy or prostate malignancy or some other non-trophoblast malignancy.  
15
52. The method of claim 44, wherein the sample is a urinary sample or a blood sample.
- 20 53. A method for detecting gestational trophoblast disease in a sample from a subject comprising:
- (a) contacting a sample with an antibody which specifically binds to the early pregnancy associated molecular isoform of hCG under  
25 conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform of hCG;
- (b) contacting the sample with a second antibody  
30 which specifically binds to intact non-nicked hCG without substantially cross-reacting with said antibody under conditions permitting formation of a complex between the antibody and the early pregnancy associated molecular isoform  
35 of hCG;

- 5 (c) measuring the amount of complexes formed, thereby determining the amount of the early pregnancy associated molecular isoform of hCG in the sample due to binding with the first antibody, and late pregnancy associated molecular isoform of hCG in the sample due to binding with the second antibody;
- 10 (d) determining the ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated molecular isoform of hCG in the subject; and
- 15 (e) comparing the ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated molecular isoform of hCG in the sample determined in step (c) over time, wherein a continuing high ratio of early pregnancy associated molecular isoform of hCG to late pregnancy associated molecular isoform of hCG in the sample determined in step (c)
- 20 indicates the presence of gestational trophoblast disease in the subject.

54. The method of claim 48, wherein the antibody is B152.
- 25 55. The method of claim 48, wherein the second antibody is B109.
56. The method of claim 48, wherein the gestational trophoblast disease is choriocarcinoma or
- 30 hydatidiform mole.
57. The method of claim 48, wherein the sample is a urinary sample or a blood sample.

100

58. An antibody which specifically binds to an early pregnancy associated molecular isoform of human chorionic gonadotropin.
- 5 59. The antibody of claim 58, wherein the antibody specifically binds to a region of the early pregnancy associated molecular isoform of human chorionic gonadotropin comprising a carbohydrate moiety.
- 10 60. The monoclonal antibody of claim 58 designated B152.
61. A hybridoma cell accorded ATCC Accession No. HB-12467, producing the monoclonal antibody of claim 60.
- 15 62. The antibody of claim 58 labeled with a detectable marker.
63. The antibody of claim 58, wherein the detectable  
20 marker is a detection antibody.
64. The antibody of claim 58, wherein the antibody specifically binds to the nicked form of human gonadotropin.
- 25 65. The antibody of claim 58, wherein the antibody specifically binds an epitope dependent upon peptide bond cleavage in beta loop 2.
- 30 66. The antibody of claim 58, wherein the antibody binds the human gonadotropin at the beta COOH-terminal region.

FIG. 1

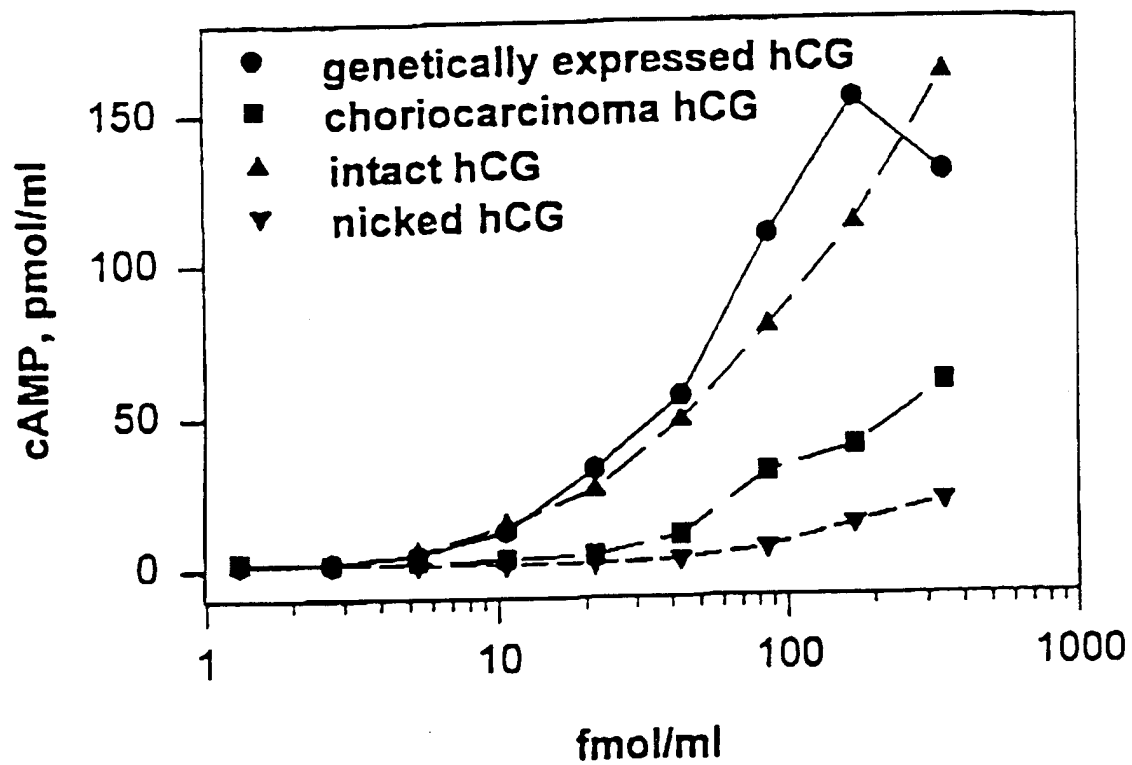


FIG. 2A

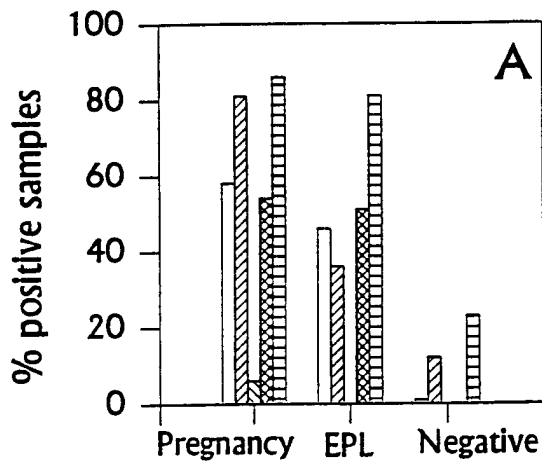
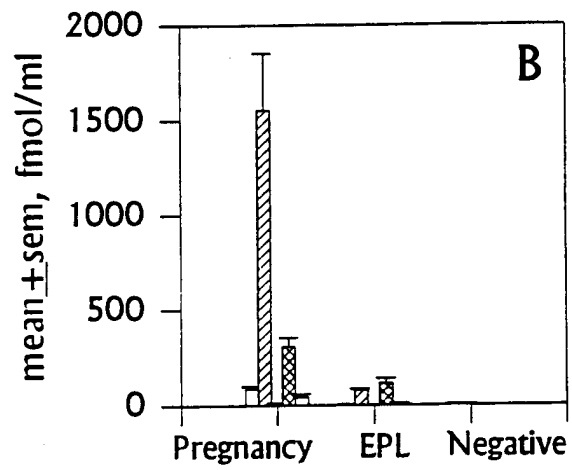
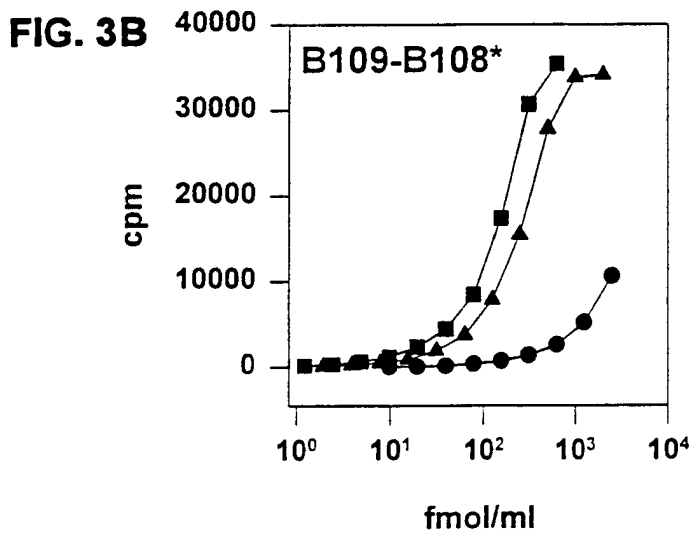
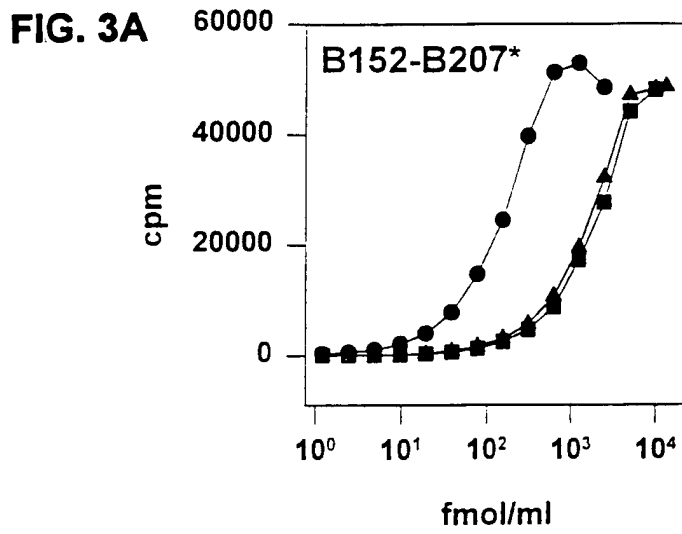


FIG. 2B



- hCG (B109-B108\*)
- hCG (B152-B207\*)
- nhCG (B151-B207\*)
- hCGβ (B201-C104\*)
- hCGβcf (B210-B108\*)

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- choriocarcinoma hCG (C5)
- pregnancy hCG
- ▲— pituitary hCG

FIG. 4

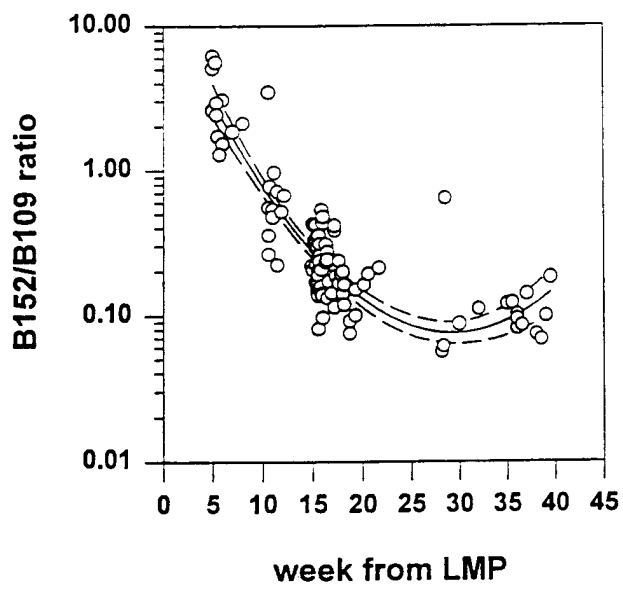
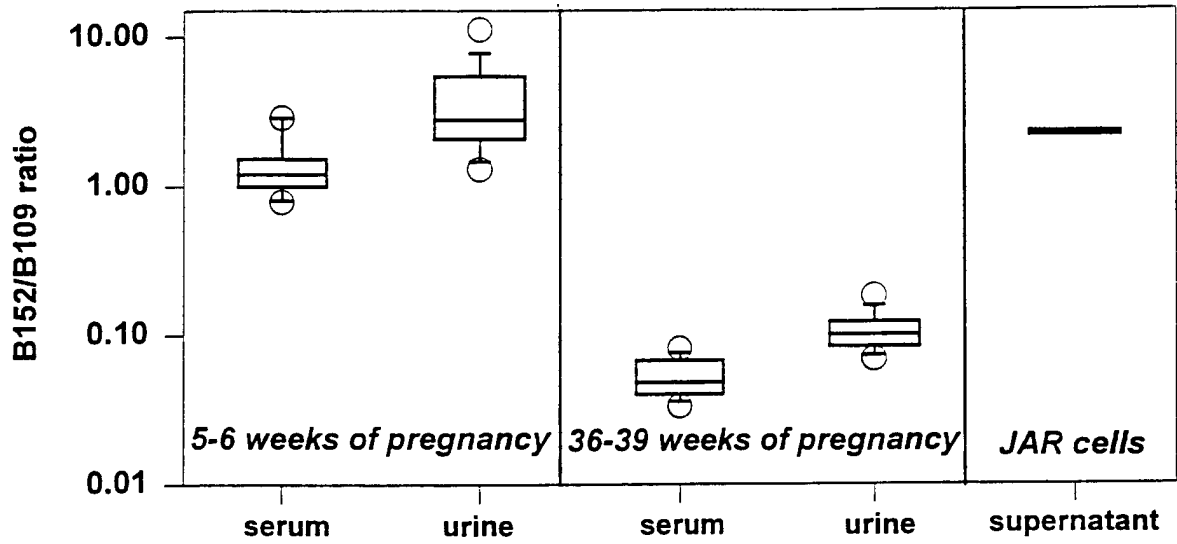
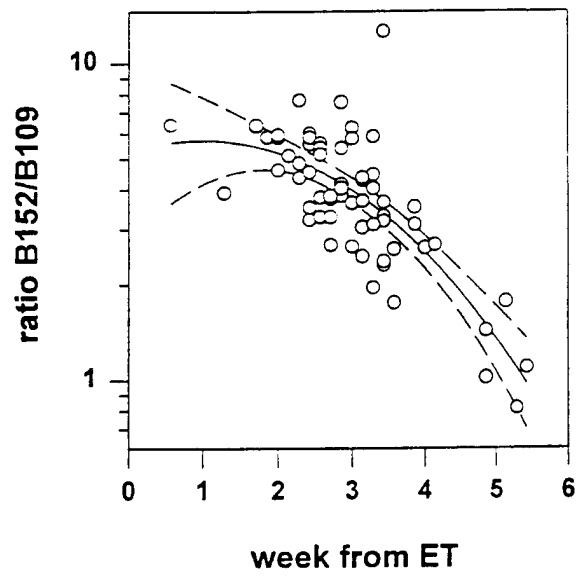


FIG. 5



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FIG. 6



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FIG. 7A

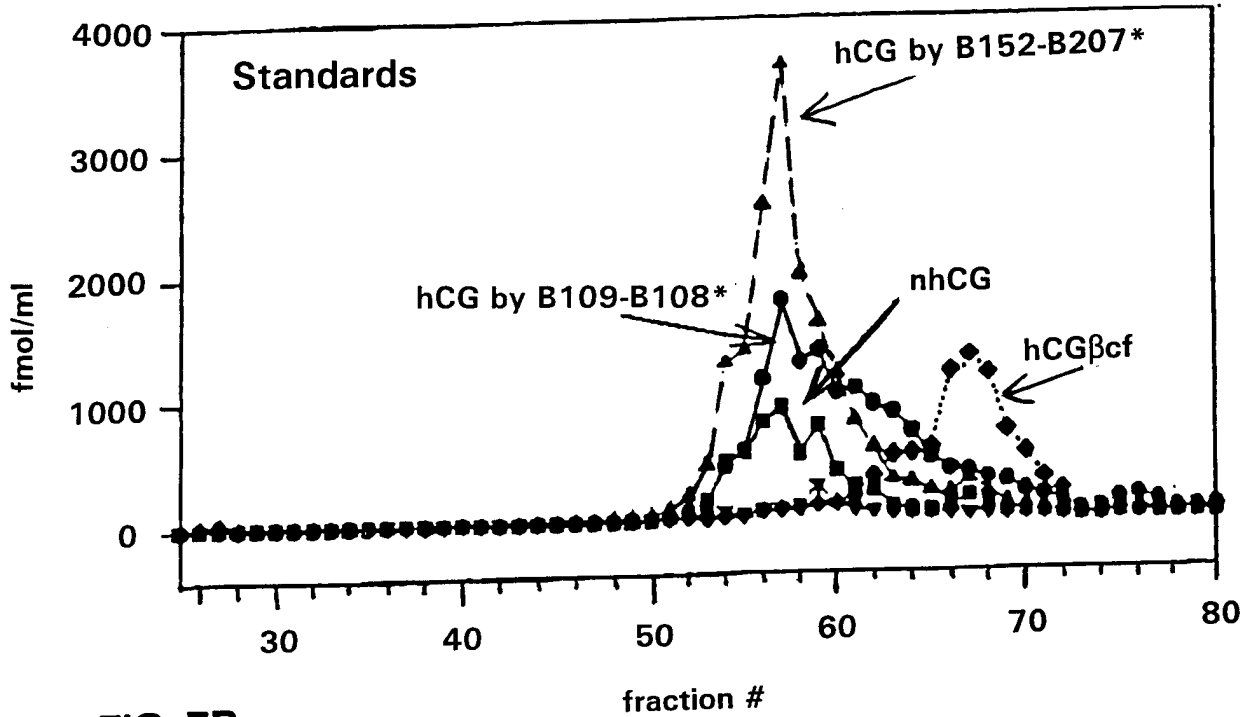
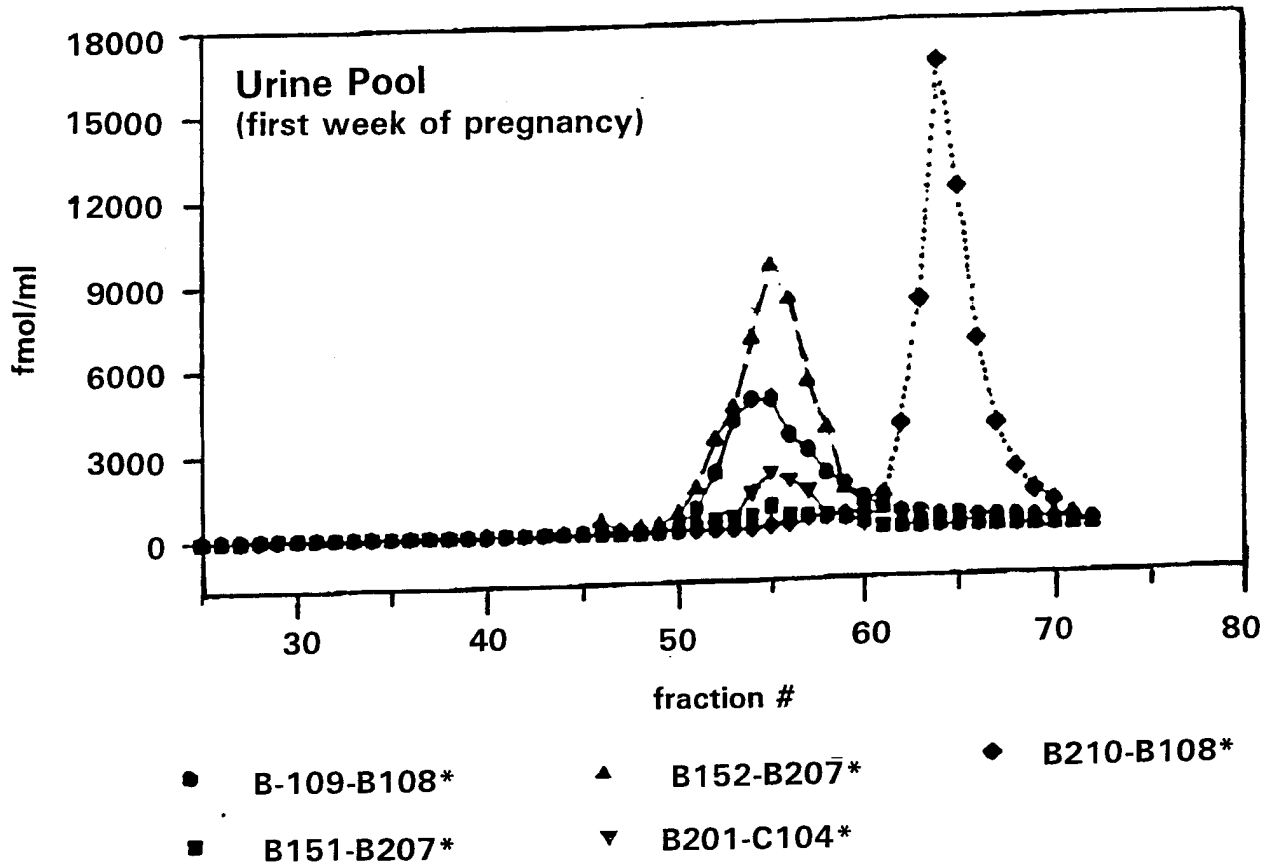


FIG. 7B



SUBSTITUTE SHEET (RULE 26)

FIG. 8A

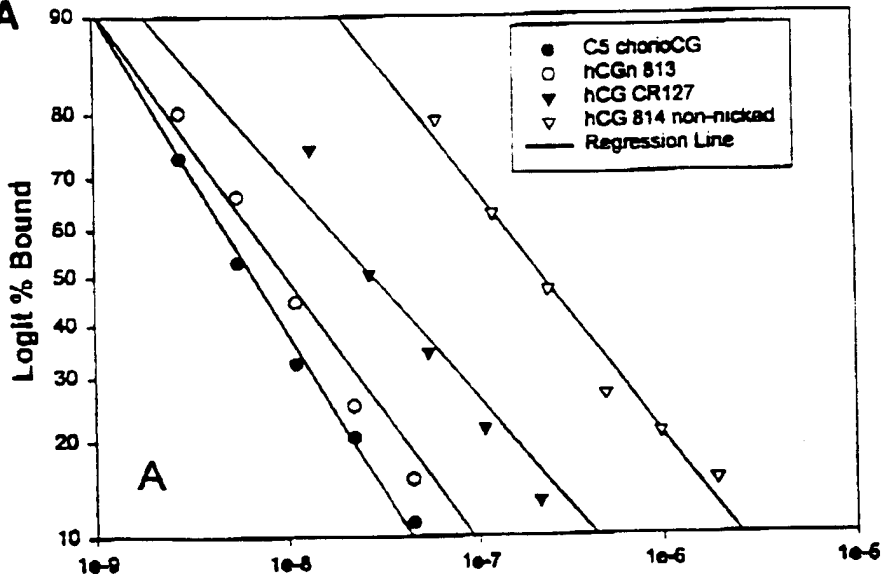
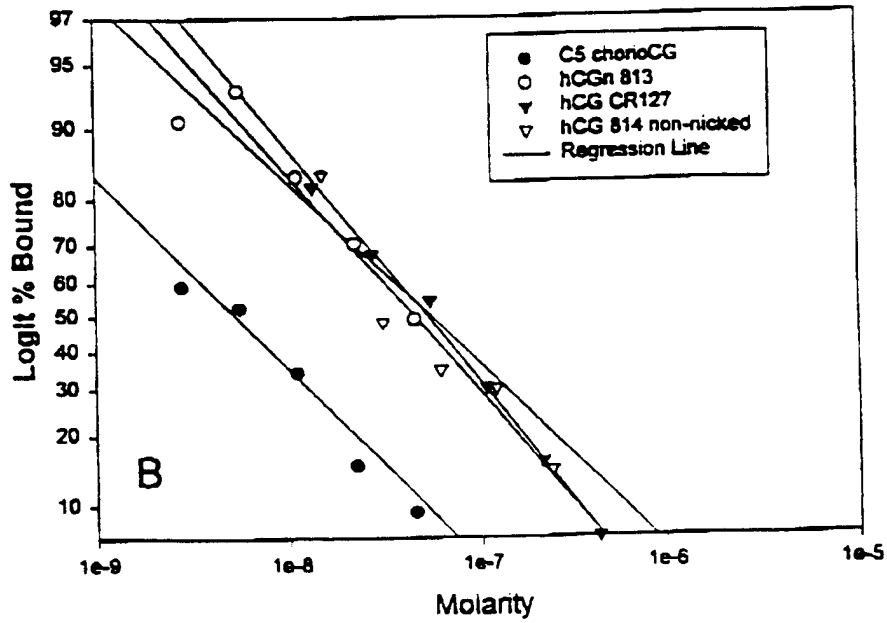


FIG. 8B



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FIG. 9A

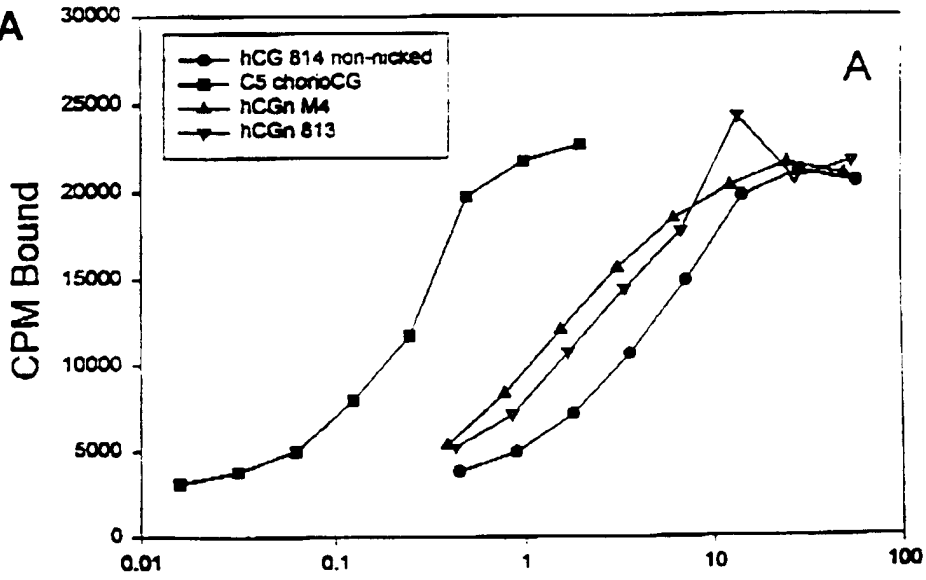
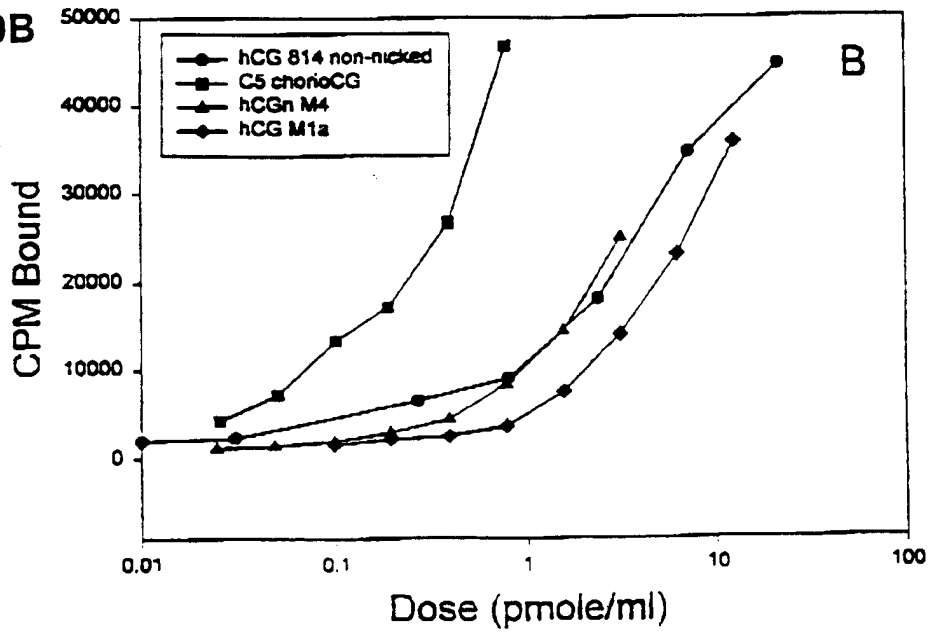
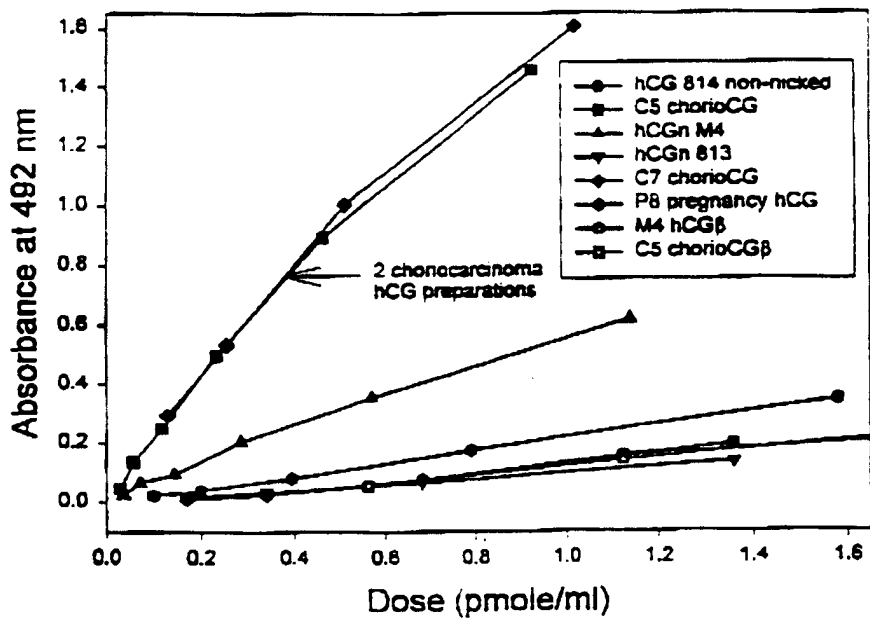


FIG. 9B



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FIG. 10



**FIG. 11**

**Table I. Characteristics of the Reagents Used to Define Antibody Specificity. The peptide and carbohydrate structures of the reagents used were determined earlier (26). The % nicked  $\beta$ -subunit refers to the proportion of molecules with cleavages (missing peptide bonds) in the region  $\beta$ 43 to  $\beta$ 48. The % tetrasaccharide core is the proportion of O-linked oligosaccharides with tetrasaccharide (vs. disaccharide) core structure, and the % sialic acid, is the proportion of O-linked structures with antennae terminated by sialic acid residues. The proportion of triantennary N-linked oligosaccharides on  $\beta$ -subunit is given, as is the corresponding % sialic acid.**

Name	Source	N-Sialic acid <sup>a</sup>	O-Sialic acid <sup>b</sup>	% triantennary N-linked on $\beta$	% tetrasaccharide O-linked core	% $\beta$ nicking
814 hCG	CR 127 hCG <sup>b</sup>	95	66	5	19	19
C5 chorioCG	Choriocarcinoma	95	58	48	100	100
M4 mole CG	Mole pregnancy	120	49	30	20	98
813 hCGn	CR 127 hCG	ND	ND	ND	ND <sup>c</sup>	80
C7 chorioCG	Choriocarcinoma	68	53	48	69	3
P8 hCG	Pregnancy	94	73	21	13	0
M4 mole $\beta$	Mole pregnancy	120	49	30	20	98
C5 chorio $\beta$	C5 chorioCG	95	58	48	100	100
CR 129 $\beta$	CR 129 hCG	96	63	11	17	19
HLH1-1	Pituitary hLH	ND	NA <sup>d</sup>	NA	NA	NA
M1A	Mole pregnancy	98.5	NA	16.5	<15% CTF <sup>e</sup>	24

<sup>a</sup>: % sialic acid residues per sugar chain, N-linked on  $\beta$ .

<sup>b</sup>: % sialic acid residues per sugar chain, O-linked on  $\beta$ .

<sup>c</sup>: The "CR" series of hCG reference preparations were made at Columbia University and were distributed internationally as reference materials for purified hCG. CR 119 is also known as the 3<sup>rd</sup> international immunoassay reference preparation for hCG.

<sup>d</sup>: ND is not done; NA is not applicable to that reagent.

<sup>e</sup>: Less than 15% of the beta COOH-terminal region is present on this preparation.

**FIG. 12**

**Table II: Affinity Constants<sup>a</sup> Determined by Liquid Phase Competition Assays Using C5 as Tracer Ligand**

Antibody	Competitors			
	C5 chorioCG	Nicked hCG CR 127 (813)	Parent CR 127 hCG <sup>b</sup>	Nick-free hCG CR127 (814)
B151	4.4 x 10 <sup>8</sup>	3.8 x 10 <sup>8</sup>	4.2 x 10 <sup>7</sup>	1.3 x 10 <sup>7</sup>
B152	3.5 x 10 <sup>8</sup>	5.4 x 10 <sup>7</sup>	4.7 x 10 <sup>7</sup>	5 x 10 <sup>7</sup>

<sup>a</sup>K<sub>a</sub> as L/M

<sup>b</sup> hCG CR 127 is an NIH-distributed hCG reference preparation produced at Columbia University.

**FIG. 13A**

Table III: Matrices of data for binding characteristics of different pairs of detection antibodies using B151 or B152 as capture antibody.

A. Relative Cross-Reactivities of Two Site Assay Using B151 as Capture Antibody

Ligand	B207 <sup>a,c</sup>	B204 <sup>a</sup>	B201 <sup>a</sup>	B108 <sup>a</sup>	B109 <sup>a</sup>	A109 <sup>a</sup>	CTP104
C5	100%	< <sup>b</sup>	<	100%	<	<	100
813 CR 127 hCGn	100%	<	<	100%	<	<	47%
814 CR127 hCG	12%	<	<	37%	<	<	14%
HCG β	2%	<	<	2%	<	<	<
C5 β	5%	<	<		<	<	<
HCGβ core	<	<	<		<	<	<
HLH	2%	<	<	3%	<	<	<
HLHβ	5%	<	<		<	<	<
HCG α	<	<	<	3%	<	<	<
Maximum binding <sup>b</sup>	50%	0%	0%	13%	0%	0%	83%

<sup>a</sup> labeled detection antibodies

<sup>b</sup> < out of low range detection

<sup>c</sup> this particular assay format was applied in O'Connor et al (25).

**FIG. 13B**

**B. Relative Cross-Reactivities of Two Site Assay Using B152 as Capture Antibody**

Ligand	B207 <sup>a,d</sup>	B204 <sup>a</sup>	B201 <sup>a</sup>	B108 <sup>a</sup>	B109 <sup>a</sup>	A109 <sup>a</sup>	CTPI04 <sup>a</sup>
C5	100%	100%	94%	42%	53%	100%	<
813 CR 127 hCGn	10%	< <sup>b</sup>	<	15%	32%	64%	<
814 CR127 hCG	7%	<	<	30%	100%	26%	<
HCG β	6%	20%	19%	11%	<	<	<
C5 β	190%	100%	100%	100%	<	<	<
HCGβ core	<1%	<	<	<	<	<	<
HLH	<1%	<	<	<	<	<	<
hLHβ	<1%	<	<	<	<	<	<
HCG α	<1%	<	<	<	<	<	<
Maximum binding <sup>b</sup>	64%	2%	44%	80%	2%	14%	25%

The molar quantity of ligand required to produce binding equal to 50% of the maximum binding achieved by C5 was determined. Cross-reactivity shown in this table as a percentage is calculated by dividing the molar quantity of the standard by the molar quantity of the other ligand at 50% maximum binding dose.

<sup>a</sup> labeled detection antibodies

<sup>b</sup> maximum binding represents the total quantity of radiolabeled detection antibody which can bind to the plate in the system described.

<sup>c</sup> < out of low range detection

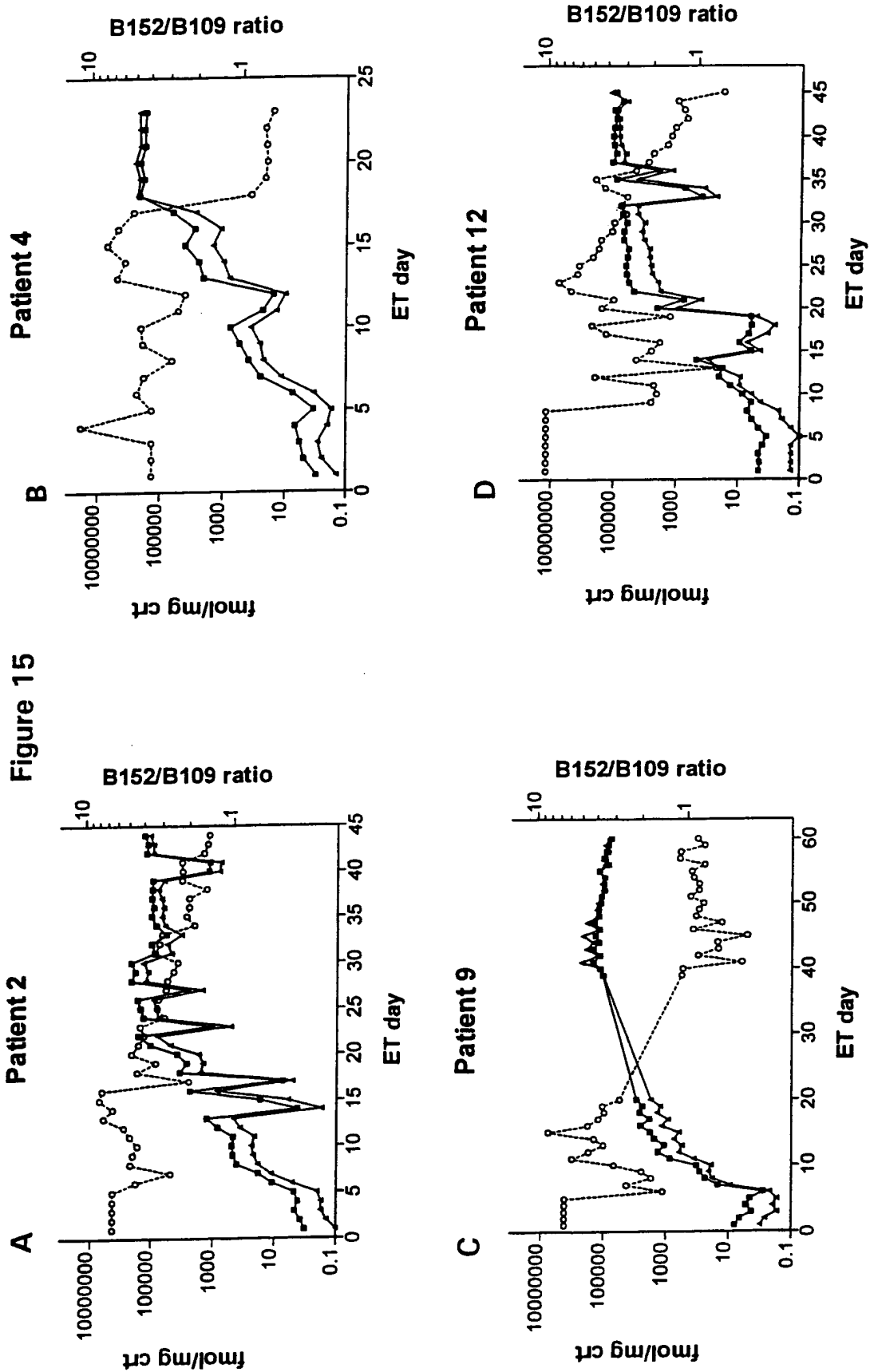
<sup>d</sup> this particular assay format was applied in O'Connor et al (25).

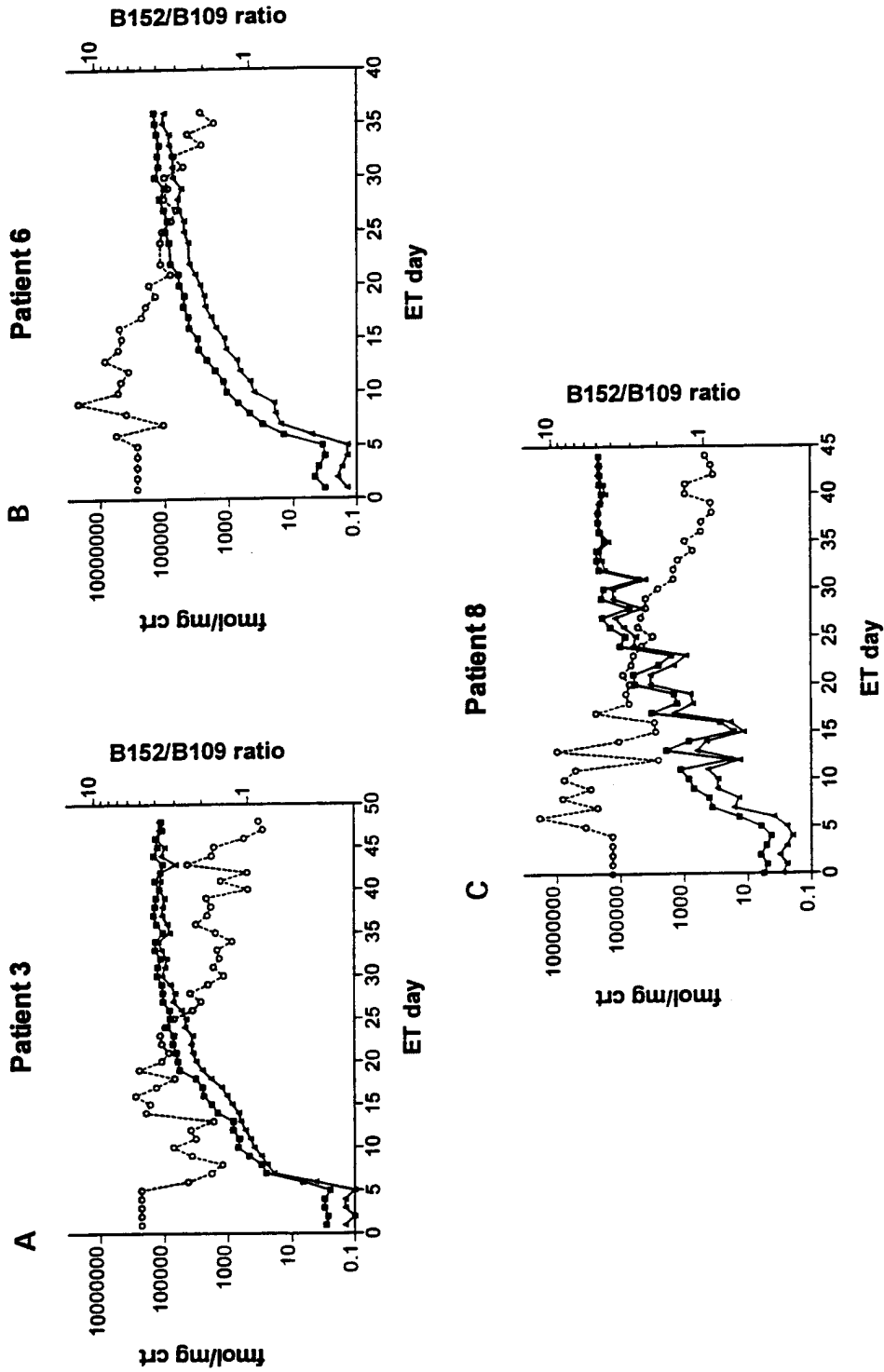
**FIG. 14**

Table IV. Immunoreactivity of antigens in the B152 immunoradiometric assay. The dose-response curves used to provide data for this table are shown in Figure 3. Each curve was fitted with 4-5 points. Slope and coefficient of determination ( $R^2$ ) were determined using a non-linear regression algorithm. Slopes were used as an indicator of antigen potency. Relative potency was estimated as the slope of antigens relative to the slope of C5 Choriocarcinoma hCG (the immunogen)

Reagent	Slope <sup>a</sup>	S.E.	R <sup>2</sup>	Relative Potency
814 hCG	0.1588	0.0098	0.992	10.2%
C5 chorioCG	1.5603	0.1015	0.983	100%
M4 mole hCGn	0.5317	0.0240	0.992	34%
813 hCGn	0.0986	0.0021	0.999	6.3%
C7 chorioCG	1.4515	0.1246	0.985	93.0%
P8 hCG	0.2192	0.0031	0.999	14.0%
M4 mole hCG $\beta$	0.1038	0.0069	0.991	6.67%
C5 chorioCG $\beta$	0.1286	0.0042	0.998	8.24%
CR 129 hCG $\beta$	Only 2 points			
HLH batch I-1	No response			

<sup>a</sup> Slope are from figure 3 as calculated in Sigmaplot 4.01 by linear regression analysis. Units of slope are pmole/ml absorbance at 492nm.

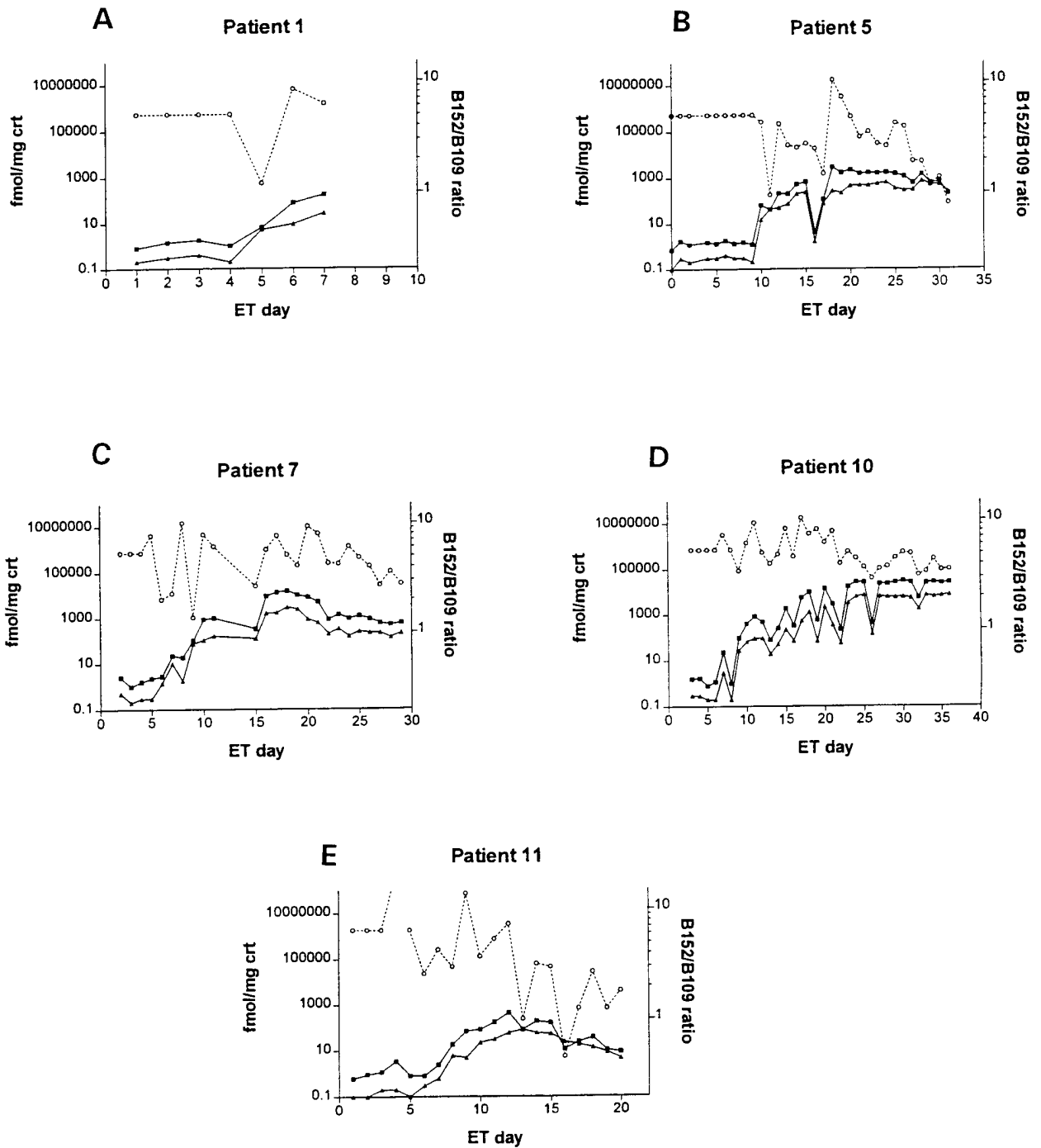




—●— hCG as determined by assay B152-B207\*, fmol/mg creatinine  
—■— hCG as determined by assay B109-B108\*, fmol/mg creatinine  
...○... Ratio=B152-B207\*/B109-B108\*

Figure 16

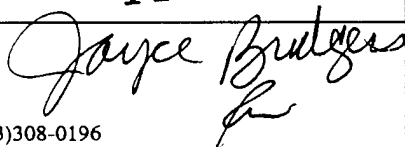
18/18  
Figure 17



—■— hCG as determined by assay B152-B207\*, fmol/mg creatinine  
 —●— hCG as determined by assay B109-B108\*, fmol/mg creatinine  
 - - - ○ - - - Ratio=B152-B207\*/B109-B108\*

## INTERNATIONAL SEARCH REPORT

International application No.  
PCT/US00/13197

<b>A. CLASSIFICATION OF SUBJECT MATTER</b> IPC(7) :Please See Extra Sheet. US CL :Please See Extra Sheet. According to International Patent Classification (IPC) or to both national classification and IPC		
<b>B. FIELDS SEARCHED</b> Minimum documentation searched (classification system followed by classification symbols) U.S. : Please See Extra Sheet.  Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched  Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) EAST and WEST Patent Database, STN-biosis, caplus, haplus, medline.		
<b>C. DOCUMENTS CONSIDERED TO BE RELEVANT</b>		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X,P	WO 99/41584 A2 (THE TRUSTEES OF COLUMBIA UNIVERSITY IN THE CITY OF NEW YORK) 19 August 1999, see full document.	1-66
X	BIRKEN, S. et al. Development and Characterization of Antibodies to a Nicked and Hyperglycosylated Form of hCG from a Choriocarcinoma Patient. Endocrine. April 1999, Vol. 10, No. 2, pages 137-144, see abstract.	1-66
Y	KOVALEVSKAYA, G. et al. Evaluation of Nicked Human Chorionic Gonadotropin Content in Clinical Specimens by a Specific Immunometric Assay. Clinical Chemistry. January 1999, Vol. 45, No. 1, pages 68-77, see abstract.	1-66
<input checked="" type="checkbox"/> Further documents are listed in the continuation of Box C. <input type="checkbox"/> See patent family annex.		
*A*	Special categories of cited documents: document defining the general state of the art which is not considered to be of particular relevance	*T* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
*E*	earlier document published on or after the international filing date	*X* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
*L*	document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	*Y* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
*O*	document referring to an oral disclosure, use, exhibition or other means	*G* document member of the same patent family
*P*	document published prior to the international filing date but later than the priority date claimed	
Date of the actual completion of the international search 21 JUNE 2000	Date of mailing of the international search report 11 AUG 2000	
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. (703) 305-3230	Authorized officer LISA V. COOK Telephone No. (703)308-0196 	

## INTERNATIONAL SEARCH REPORT

International application No.  
PCT/US00/13197

## C (Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	COLE, L. A. et al. The Heterogeneity of Human Chorionic Gonadotropin (hCG). III. The Occurrence and Biological and Immunological Activities of Nicked hCG. <i>Endocrinology</i> . 1991, Vol. 129, No. 3, pages 1559-1567, see full document.	1-37, 42, and 44-57
Y	BERGER, P. et al. Variants of Human Chorionic Gonadotropin from Pregnant Women and Tumor Patients recognized by Monoclonal Antibodies. <i>Journal of Clinical Endocrinology and Metabolism</i> . 1993, Vol. 77, No. 2, pages 347-351, see full document.	1-66
Y	ACEVEDO, H.F. et al. Expression of Membrane-associated Human Chorionic Gonadotropin, Its Subunits, and Fragments by Cultured Human Cancer Cells. <i>Cancer</i> . 01 April 1992, Vol. 69, No. 7, pages 1829-1842, see full document.	1-66
A	HOERMANN R. et al. Molecular Heterogeneity of Human Chorionic Gonadotropin in Serum and Urine From Patients with Trophoblastic Tumors. <i>Clinical Investigation</i> . 1993, Vol. 71, pages 953-960, especially pages 958-959.	1-66

**INTERNATIONAL SEARCH REPORT**

International application No.  
PCT/US00/13197

**A. CLASSIFICATION OF SUBJECT MATTER:**

IPC (7):

C12Q 1/68, 1/70; G01N 21/64, 33/00, 33/53, 33/531, 33/533, 33/543; C07H 21/02, 21/04; C12N 15/00; B05D 1/18; B01J 13/00

**A. CLASSIFICATION OF SUBJECT MATTER:**

US CL :

435/5, 6, 7, 7.21, 7.9, 91.1, 91.2, 94, 183, 270; 436/501, 504, 508, 518, 808, 811; 536/23.1, 24.3, 24.31, 24.32, 25.32; 935/1, 2, 3, 5, 8, 19, 76, 77, 78, 85, 111; 530/300, 350, 387.1, 388.1, 388.2, 391.1, 811, 812, 815;

**B. FIELDS SEARCHED**

Minimum documentation searched

Classification System: U.S.

435/5, 6, 7, 7.21, 7.9, 91.1, 91.2, 94, 183, 270; 436/501, 504, 508, 518, 808, 811; 536/23.1, 24.3, 24.31, 24.32, 25.32; 935/1, 2, 3, 5, 8, 19, 76, 77, 78, 85, 111; 530/300, 350, 387.1, 388.1, 388.2, 391.1, 811, 812, 815;

专利名称(译)	用hCG分析预测受试者怀孕结果的方法		
公开(公告)号	<a href="#">EP1183390A1</a>	公开(公告)日	2002-03-06
申请号	EP2000935949	申请日	2000-05-12
[标]申请(专利权)人(译)	纽约市哥伦比亚大学理事会		
申请(专利权)人(译)	哥伦比亚大学纽约市受托人		
当前申请(专利权)人(译)	哥伦比亚大学纽约市受托人		
[标]发明人	OCONNOR JOHN BIRKEN STEVEN KOVALEVSKAYA GALINA		
发明人	O'CONNOR, JOHN BIRKEN, STEVEN KOVALEVSKAYA, GALINA		
IPC分类号	G01N33/53 C07K16/26 C12N15/02 G01N33/577 G01N33/76 C12Q1/68 C12Q1/70 G01N21/64 G01N33/00 G01N33/531 G01N33/533 G01N33/543 C07H21/02 C07H21/04 C12N15/00 B05D1/18 B01J13/00		
CPC分类号	G01N33/76 C07K16/26 G01N33/57434 G01N33/57449 G01N2333/59 G01N2800/368 Y10S436/814 Y10S436/818		
代理机构(译)	LAWRENCE , JOHN		
优先权	09/311428 1999-05-13 US		
其他公开文献	EP1183390A4 EP1183390B1 EP1183390B8		
外部链接	<a href="#">Espacenet</a>		

#### 摘要(译)

本发明提供了通过测定样品中hCG的早期妊娠相关分子同种型的量来预测受试者的妊娠结果的方法。本发明进一步提供了用于确定样品中人绒毛膜促性腺激素 ( hCG ) 的早孕相关分子同种型的量的方法。本发明还提供了用于确定样品中与早孕相关的hCG的量的诊断试剂盒。本发明另外提供了一种抗体，其特异性结合人绒毛膜促性腺激素的早期妊娠相关分子同种型。最后，本发明提供了用于检测样品中的滋养层细胞或非滋养层恶性肿瘤的方法。