Immunolocalization of 91 kDa Ascitic Protein in the Human Placenta

By
Koji KAMI, Noriaki SATO, Toshiyuki MORITA, Isamu ISHIWATA, Masakazu ISHIKAWA and Mitsuo NAKAI

Department of Human Morphology and Physiology, Tokiwa University School of Human Sciences, 1-430-1 Miwa, Mito 310; Ishiwata Obstetrics and Gynaecology Hospital, 1-4-21 Kami-Mito, Mito 310; Department of Obstetrics and Gynaecology, Kitasato University School of Medicine, 1-15-1 Kitasato, Sagamihara 228, Japan

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Summary: A 91 kDa protein, sharing antigenicity with a pregnancy-associated antigen, was isolated from the ascitic fluid of a patient suffered from the gynaecological malignant neoplasms. It was revealed in the human placenta by immunocytochemical techniques, and found to be localized in the stroma, including a fibrous network, fibroblasts and the Hofbauer cells of the terminal and stem villi in the first trimester. Immunoreactive staining was also found in macrophage-like cells, presumable the precursor of the Hofbauer cell, in the stem villi of placenta. It is assumed that macrophage-like cells containing the protein would be originated from the extraembryonic mesoderm or blood vessels, and then migrated to the stroma of terminal villi. During the second trimester, the protein was observed in numerous vesicles among the syncytial trophoblasts. The density of these immunoreactive vesicles and the cytoplasm of Hofbauer cells was rapidly increased with gestational stages, whereas it was gradually reduced towards the late-second trimester. The highest labellings occurred at around 17 weeks or later. On the other hand, at term, the protein was mainly observed in numerous villous blood vessels including endothelial cells. In the light of these results, a hypothesis concerning mechanisms of transplacental permeability of the protein by the placental barrier is discussed further.

The morphogenesis of both foetuses and neoplasms is accompanied by the ontogenetically-programmed appearance and disappearance of several proteins including enzymes. These proteins have been shown to be related to embryonic developments and oncopathological statuses.

On the other hand, in clinical evidences, the placentalogical light of these proteins is also focussed on characteristics of the phenemena in relation to pregnancies and malignancies. So far, a numerous placental protein; such as hCG-A and B, hPL, PAPP-A and B, PP1, SP1, CA 125 and Ca 19—9, were involved in oncodevelopmental aspects at cellular and molecular levels. However, they seemed to be limited to the clinical usefulness. In recent years, a novel 94 kDa pregnancy-associated protein isolated from the placenta has been extensively studied from a variety of aspects. There is a body of evidence showing that an alpha2-protein of the placenta exists to be identical with the protein obtained from malignant ascitic fluids of a patient with the advanced ovarian cancer, whereas a molecular size is found to be 91 k daltons by SDS-PAGE procedures. Moreover, it is assumed that the protein would play important roles in both tumour-associated statuses and pregnancy-associated evidences.

To date, however, nothing is known about the microscopical distribution of the protein in the human placenta and neoplasm. The aim of this study was to address questions of where 91 kDa ascitic protein is localized in the placenta using an immunocytochemical technique. A part of this study was presented in the 9th International Congress of Histochemistry and Cytochemistry, at Maastricht, The Netherlands.

Materials and Methods

Chemicals
Biotin-labelled goat anti-rabbit IgG, avidin-biotin-horseradish peroxidase (HRP) complex and avidin-
Preparation of 91 kDa ascitic protein and its antibody

Ascitic fluid used was obtained from a patient with a malignant ovarian and uterine tumour (mixed mesodermal tumour) at the time of paracentesis, and then centrifuged at 5,000 rpm for 30 min at 4°C. The supernatant was kept at -20°C until use.

An ectopic developmental antigen that appears in malignant ascitic fluid was separated by a new procedure involving DEAE-Sepharose chromatography as described in the previous paper. In brief, pooled ascitic fluid was treated as follows: (1) immunoglobulins were removed using polyethylene glycol 4,000 fractionation; (2) the peak, on DEAE-Sepharose chromatography at pH 6.6, showing the highest specific reactivity to the “First-Generation” antiserum was collected; (3) the protein was separated by a preparative-polyacrylamide gel electrophoresis (PAGE); (4) the antibody against the protein was generated in rabbits; (5) the antibody was applied to a column of normal human plasma (NHP)-coupled Sepharose 4B, then, the “Second-Generation” antibody was obtained within the unbound fractions; (6) a molecular size of the protein was confirmed to be 91 kDa by SDS-PAGE; (7) the monospecificity of the antibody was found using a SDS-PAGE-’SRID’ technique.

In this study, the antibody obtained was used for revealing immunocytochemical characterization of the protein in the human placenta.

Tissue preparation

Thirty-eight placentas were obtained from healthy women at between the 7th and 42nd week of gestation (14 at 7–10 weeks; 15 at 11–20 weeks; 9 at 37–42 weeks at term). Small pieces (1 mm x 3 mm) of villous tissue were excised and fixed by immersion at 4°C for 4–6 h in Zamboni’s fixative with a minor modification, and then processed through graded changes of ethanol and xylene, and embedded in paraffin wax.

Immunocytochemical procedures

All immunocytochemical procedures were carried out at room temperature. Deparaffinized sections kept in 0.01 M phosphate-buffered saline (PBS), pH 7.3, were subjected to the following protocol: (1) enzymatic digestion with 0.1% (w/v) pepsin in 0.05 M Tris-HCl buffer, pH 7.6, containing 0.1% (w/v) CaCl₂, 30 min at 37°C; (2) 3 washes with PBS, 5 min each; (3) blocking of endogenous peroxidase activity with 5 mM sodium metaperiodate solution, 10 min; (4) 3 washes with PBS, 5 min each; (5) blocking free-aldehyde groups with 3 mM sodium borohydride solution, 10 min; (6) 3 washes with PBS, 5 min each; (7) pre-treatment with highly diluted (1:2,500) normal goat serum, 30 min; (8) incubation with the specific antibody diluted at 1:400 to 1:2,000 with PBSA (PBS containing 0.5% bovine serum albumin), 2–4 h; (9) 3 washes with PBS, 10 min each; (10) incubation with biotin-labelled goat anti-rabbit IgG, 30 min; (11) 3 washes with PBS, 10 min each; (12) incubation with avidin-biotin-HRP complex, 30 min; (13) 3 washes with PBS, 10 min each.

In the final step, sections were incubated for 2–5 min in 0.05 M Tris-HCl buffer, pH 7.4, containing 0.02% (w/v) 3,3'-diaminobenzidine tetrahydrochloride and 0.001% (v/v) hydrogen peroxide, and counterstained with 0.1% (w/v) Methyl Green in barbitone buffer at pH 4.0 for 1 h or with Mayer's Haematoxylin in 10 min.

Immunofluorescence antibody procedures followed the above-described protocol except that avidin-biotin-HRP complex was substituted with avidin-biotin-FITC complex.

In order to confirm the specificity of labellings, two concurrent control experiments were performed: (a) preimmune rabbit serum used in place of the specific antibody, and (b) avidin-biotin-HRP complex or avidin-biotin-FITC complex alone, omitting biotin-labelled goat anti-rabbit IgG to identify any non-specific adsorption of the complex.

Results

Human placentae subjected are divided into three groups by their gestational periods; the first (7–10 weeks), second (11–20 weeks) and third (term) trimesters. Figures 1–3 and Table 1 are representative of immunomicroscopical stainings most commonly seen in human placental tissues and their summarized conclusions, respectively. In the control experiments, a specific staining was never observed when a preimmune rabbit serum was used. Neither endogenous peroxidase activity nor autofluorescence was detected.

In the first trimester, the affinity-purified antibody frequently exhibited specific immunostainings to be in close association with stroma of the secondary and tertiary villi; fibrous networks, fibroblasts and the Hofbauer cells (Figs. 1-A ~ 1-C). The distribution of immunoreactive terminal villi was not uniform; some were positive whereas the rests were negative. Labellings in villous stroma were completely disappeared during the second trimester. A few immunogenic dot-structures were also simultaneously localized near apical regions of syncytiotrophoblasts.
The population of these structures gradually increased as gestation progressed. In contrast, immunoreactive evidences were recognized in macrophage-like cells of the stroma in the secondary and tertiary stem chorionic villi (Fig. 1-D). So far, however, the immunostaining was never seen in the umbilical cord.

In the second trimester, immunostainings were demonstrated within a numerous vesicle located in the apical part of syncytiotrophoblasts, although no evidence was seen in the cytotrophoblast (Figs. 2-A — 2-E). The content of immunoreactive vesicles rapidly increased with gestational stages, reaching a maximum around 17 weeks, and then gradually reduced towards a term. The foetal blood vessel also showed positive reactions during the later-half of the second trimester; the lumen of blood vessels and endothelial cells (Figs. 2-C and 2-D). Occasionally, around the latter half of the second trimester, a small number of dot-structures which presented by an immunofluorescence technique were associated with the basement membrane (Fig. 2-F).

At term, the major localization of the protein was shown in the foetal blood vessels including endothelial cells (Figs. 3-A ~ 3-D). Intensity of the immunostaining was distinctly higher than that obtained in the second trimester. In contrast, a few dot-structures demonstrated by immunostainings were only just detected in epithelial folds faced to the maternal space of flattened syncytiotrophoblasts.

**Discussion**

A novel pregnancy-associated protein with a molecular size of 94 k daltons is originally separated from the human placenta at term. Subsequently, the "First-Generation" antibody against the protein reveals a existence of the antigenic molecular mass in the sera of pregnant women and ascitic fluids of patients with advanced ovarian cancer using immunological techniques. In recent years, an ectopic ascitic protein is purified by a new procedure using DEAE-Sepharose chromatography at pH 6.6, and its molecular size is determined to be about 91 k daltons. Furthermore, the "Second-Generation" antibody shows to be useful to semi-quantitative measurements; a higher serum concentration in pregnant women than that estimated in regular-menstruating women, or much higher levels in ascitic fluids of patients with malignant diseases than that of the health, respectively.

The aim of this study was to determine an immunocytochemical characterization of the ectopic antigen in the human placenta using the same antibody against 91 kDa protein. As a result, there are marked differences on the localization of the protein in parallel with gestational stages.

In the first trimester, the antigenic localization was detected to be predominantly limited in the foetal stromal tissue elements of terminal villi including fibroblasts and the Hofbauer cells. In early stage of placentation, it is well known that the basic tissue-skeleton in the stroma of chorionic villi should be derived from the extraembryonic mesodermal cells. The present results are supplemented by the finding that immunoreactive macrophage-like cell populations in the stem villi can be detected. Although the definite cellular classification of these mesodermal cells is still unanswered, it is reasonable to
suppose that the presumable precursor cells should originate from the connective tissue or blood vessels of the stem villi, migrate to the fibrous network of terminal villi and, in this special microenvironment of placentation in the 1st trimester, differentiate into the participant cells which could supply physiological materials programmed.

During the second trimester, the present data clearly demonstrates that a numerous small vesicle with antigenicity were determined in the apical part of syncytiotrophoblasts. The Hofbauer cell represents as an immunoreactive cell population. However, these vesicles of trophoblastic layers and cytoplasmic contents of the Hofbauer cell rapidly reduced by the end of the second trimester. On the contrary, during the course of placentation, the protein is simultaneously characterized by increased accumulation in villous capillaries. Occasionally, immunostainings of dot-structures localized along the basement membrane could be detected. It is postulated that a dynamic permeable transportation of the protein should happen across the membrane until the physiological maturation of placental barrier. This is supported by facts of a high level of the protein found in the maternal blood during the second trimester.\(^{23,24,26}\)

At term, the protein is predominantly observed in the villous blood vessels including endothelial cells. This may be reflected that transplacental permeability of the protein to the maternal blood was resisted by the placental barrier in the later-half period of second trimester.\(^2,23\) It is worth mentioning as a hypothesis that 91 kDa ascitic protein would be a strong candidate of the marker for characterizing the maturity of villous stroma and placental barrier.

So far, however, little is known of the genuine source of this protein. One would expect the protein should be derived from the extraembryonic mesodermal cells of the stem villi during early-placentation stages. However, further detailed studies are needed in order to obtain precise data in relation to the immunoreactive body which was detected in the foeto-placental circulation at term. Studies with an ultramicroscopical characterization using immunocytochemical techniques are now in progress.

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References

9) Kami K and Stoward PJ. Secretory pathway of villotrogenesis in the liver of the cockerel as revealed by immuno-gold and computer-assisted digitization techniques. Histochem J 1991; 23:259–266.


Explanation of Figures

Plate I

Fig. 1. Immunocytochemical staining of the villi in the first-trimester. A: Fibrous stroma (s). ×300. Bar = 50 μm. Negative control (c). ×220. Age; 8 weeks and 6 days. Methyl Green counterstain. B: Fibrous stroma (s). Age; 12 weeks. ×300. Bar = 50 μm. C: Hofbauer cells (H), fibroblasts (f), fibrous stroma (s). Age; 12 weeks. ×1,200. Mayer’s Haematoxylin counterstain. Bar = 10 μm. D: Macrophage-like cells (m) of the tertiary stem villi. Fibrous stroma (s). Age; 7 weeks and 3 days. ×300. Methyl Green counterstain. Bar = 50 μm.
Fig. 2. Immunocytochemical staining of the villi in the second-trimester. A: Vesicular stainings in the syncytial trophoblasts (st). Langhan's cells (L), fibrous stroma (s). x1,200. Bar = 10\,\mu m; Negative control (c). Age: 17 weeks. x220. Methyl Green counterstain. B: Vesicular fluorescence in the syncytial trophoblasts (st). Age: 17 weeks. x600. Bar = 20\,\mu m. C: Syncytial trophoblasts (st), foetal blood vessels (v). Age: 17 weeks. x600. Methyl Green counterstain. Bar = 20\,\mu m. D: Syncytial trophoblasts (st), foetal blood vessels (v). Age: 19 weeks and 6 days. x600. Bar = 20\,\mu m. E: Hofbauer cells (H), syncytial trophoblasts (st), Langhan's cell (L). Age: 17 weeks. x1,200. Bar = 10\,\mu m. F: Basement membrane (arrowheads). Age: 17 weeks. x1,200. Bar = 10\,\mu m.
Plate III

Fig. 3. Immunocytochemical staining of the villi at term. A: Foetal blood vessels including endothelial cells (e). Age: 37 weeks and 1 day. x300. Mayer's Haematoxylin counterstain. Bar = 50 μm; Negative control (c). Age: 39 weeks. x220. Methyl Green counterstain. B: Foetal endothelial cells (e) of the tertiary stem villi (sv) and terminal villi (tv). Age: 37 weeks and 1 day. x600. Bar = 20 μm. C: Foetal endothelial cells (e). Age: 37 weeks and day. x600. Methyl Green counterstain. Bar = 20 μm. D: Foetal endothelial cells (e) of the terminal villi. Age: 37 weeks and 1 day. x600. Bar = 20 μm.