Migratory trophoblast cells express a newly identified member of the prolactin gene family

Dustin O Wiemers*, Rupasri Ain*, Shigeki Ohboshi and Michael J Soares

Institute of Maternal-Fetal Biology and Departments of Molecular & Integrative Physiology and Obstetrics & Gynecology, University of Kansas Medical Center, Kansas City, KS 66160, USA

(Requests for offprints should be addressed to M J Soares, Department of Molecular and Integrative Physiology, University of Kansas Medical Center, Kansas City, Kansas 66160, USA; Email: msoares@kumc.edu)

*D O Wiemers and R Ain contributed equally to this work.

Abstract

Rodents possess an expanded prolactin (PRL) family of genes. These genes encode for a family of structurally related hormones/cytokines that are expressed most prominently in the anterior pituitary, uterus and placenta. In this study, we have identified a new member of the rat PRL family through a search of the National Center for Biotechnology Information expressed sequence database. The cDNA was sequenced and its corresponding mRNA characterized. On the basis of existing nomenclature, the rat cDNA was termed PRL-like protein-N (PLP-N). PLP-N has structural features indicative of its inclusion in the PRL family and is most closely related to PRL-like protein-F (PLP-F) and proliferin related protein (PLF-RP). A survey of PLP-N mRNA expression by Northern analysis indicated that PLP-N showed extensive expression in the metrial gland and minimal expression in the chorioallantoic placenta or other tissues. Expression of PLP-N mRNA was restricted to migratory trophoblast cells. Junctional zone trophoblast cells isolated from day 13 of gestation placenta differentiated in vitro and exhibited a capacity for PLP-N expression. In summary, we have discovered a new member of the PRL family that is prominently expressed in migratory trophoblast cells residing in the metrial gland.

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Introduction

The prolactin (PRL) family has undergone species-specific expansions, most notably in rodents, and especially in the mouse and rat. This expansion arose from gene duplication events and represents a specialized adaptation to pregnancy (Soares & Linzer 2001). In the rat, the PRL family consists of at least 19 members, including PRL, placental lactogens (PLs), PRL-related proteins (PRPs), PRL-like proteins (PLPs), and proliferin-related protein (PLF-RP). Genes encoding the PRL family are situated on chromosome 17 of the rat genome (Cooke et al. 1986, Deb et al. 1991c, Roby et al. 1993, Cohick et al. 1996, Dai et al. 1996a,b; Shah et al. 1998). Members of the PRL family are most prominently expressed during pregnancy in the anterior pituitary, uterus or trophoblast cells of the placenta (Dai et al. 2002, see Soares & Linzer 2001 for a review). Expression is precisely orchestrated in tissue- and gestationally specific patterns for each member of the PRL family (Dai et al. 2002, see Soares & Linzer 2001 for a review). Two compartments of the chorioallantoic placenta (junctional and labyrinth zones) and three trophoblast cell lineages contribute to the elaboration of the hormones. The junctional zone of the chorioallantoic placenta is situated at the maternal interface and contains trophoblast giant cells and spongiotrophoblast cells. Both trophoblast giant cells and spongiotrophoblast cells contribute to the synthesis of PRL family hormones. Hormone production begins at the ontogeny of junctional zone development and continues for the duration of pregnancy (Dai et al. 2002). During the last week of gestation, a new placental source arises. The labyrinth zone of the chorioallantoic placenta is located at the fetal interface and contains a population of trophoblast giant cells, which produce a unique subset of PRL family hormones (Campbell et al. 1989, Dai et al. 2000, 2002). The temporal- and spatial-specific patterns of hormone production probably have an effect on hormone delivery and the biology of pregnancy. Members of the PRL family target and influence a wide range of maternal targets, including the reproductive tract, liver and other classic PRL targets (Lin & Linzer 1999, Müller et al. 1999, Bittorf et al. 2000, Wang et al. 2000, Ain et al. 2002, Yu-Lee 2002) and vasculature (Jackson et al. 1994, Corbacho et al. 2002). Some members act via classical mechanisms involving the
PRL receptor-signaling pathway, whereas others influence their targets via mechanisms that are at present not well understood.

In this report, we have identified a new member of the rat PRL family through a search of the National Center for Biotechnology Information (NCBI) Expressed Sequence Tag database (EST). We have named this new member ‘PRL-like protein-N’ (PLP-N). Most interestingly, PLP-N is uniquely expressed in migratory trophoblast cells situated outside the chorioallantoic placenta. These PLP-N-positive trophoblast cells penetrate and envelop specialized blood vessels within the metrial gland, a mesometrial compartment of the uterus providing the blood supply to the chorioallantoic placenta.

Materials and Methods

Animals and tissue preparation

Holtzman rats were obtained from Harlan Sprague Dawley Inc. (Indianapolis, IN, USA). Both non-pregnant and pregnant rats were utilized. To obtain timed pregnancies, females were caged overnight with fertile males. The presence of sperm in the vaginal smear was designated day 0 of pregnancy. Tissues including uterus, metrial gland and placental tissues were dissected from pregnant animals. Briefly, embryos and placentas with their encapsulating decidua tissue were removed from the uterus. Further dissections were performed with the aid of a dissecting microscope (×10–20 magnification). The overlying decidua tissue and the underlying yolk sac/umbilical insertion were removed by means of fine forceps and irisectomy scissors. The chorioallantoic placenta was separated into two components: junctional zone and labyrinth zone. The junctional zone was identified by its pale appearance, from the absence of fetal blood, and separated from the labyrinth zone, a richly vascularized tissue, by means of fine forceps and 23-gauge needles. Metrial glands were obtained via dissection of mesometrial uterine tissue immediately adjacent to the conceptus. The metrial gland is a heterogeneous tissue comprised of blood vessels, immune cells, trophoblast cells and fibroblasts (Selye & McKeown 1935, Pijnenborg 1981, Peel 1989). All tissues were snap-frozen in liquid nitrogen for northern blot analysis. For in situ hybridization, uterine and placental tissues were removed intact without dissection and frozen in dry-ice-cooled heptane. All tissue samples were stored at −80°C until required for use. Tissue samples were collected from a minimum of three different rats for each data point. The University of Kansas Medical Center Animal Care and Use Committee approved all procedures for handling and experimentation with rodents.

Cell culture models

Two trophoblast cell culture systems were examined for their ability to express PLP-N. The Rcho-1 trophoblast cell line was derived from a rat choriocarcinoma and is capable of differentiating along the trophoblast giant cell lineage (Faria & Soares 1991, Peters et al. 2000). Cells were routinely plated at a density of 2–2.5 × 10^5 cells per 75 cm^2 flask in RPMI-1640 culture medium supplemented with 20% fetal bovine serum (FBS), 50 µM 2-mercaptopethanol, 1 mM sodium pyruvate, 100 U/ml penicillin, and 100 µg/ml streptomycin at the initiation of each experiment. Cells were transferred to differentiation medium (NCTC-135 culture medium supplemented with 10% horse serum, 50 µM 2-mercaptoethanol, 1 mM sodium pyruvate, 100 U/ml penicillin and 100 µg/ml streptomycin) on day 3 of culture and fed every other day with fresh culture medium. Cultures were terminated after 1, 4, 8 and 12 days of culture and RNA extracted for northern blot analysis. During 12-day culture, cells proceed from a proliferative to a differentiated state (Hamlin et al. 1994, Peters et al. 2000).

Primary junctional zone cell cultures were established from dissected day 13 rat chorioallantoic placentas as previously described (Lu et al. 1994). Tissues were cut into small pieces with iris scissors and dissociated with Dispase (48 mg/ml) and DNase I (80 U/ml) for 30 min at 37°C with continuous shaking. The isolated cells were then resuspended in RPMI-1640 culture medium supplemented with 1 mM sodium pyruvate, 50 µM β-mercaptoethanol, 100 U/ml penicillin, 100 µg/ml streptomycin and 10% FBS and filtered through a nylon mesh (70 µm). The cell suspension was washed and then plated at a density of 2–2.5 × 10^6 cells per 25 cm^2 flask in the supplemented RPMI-1640 culture medium. The purity of the cultures was >96% trophoblast. The cells were cultured for 7 days with daily changes of fresh culture medium. At the end of the culture, cells were harvested and RNA extracted for northern blot analysis.

Characterization of the PLP-N cDNA

The NCBI EST database was searched using the translated Basic Local Alignment Search Tool (BLAST) analysis (Altschul et al. 1997) with amino acid sequences for members of the PRL family as queries. Three ESTs possessing homology to rat PLF-RP were identified (UI-R-CXO-bws-e-04–0-UI, UI-R-CX0-bxi-h-07–0-UI and UI-R-CXOs-ccv-a-03–0-UI). The three ESTs showed 99% sequence identity among each other. The UI-R-CXO-bws-e-04–0-UI cDNA was obtained from the University of Iowa Rat Gene Discovery Program and Research Genetics (Huntsville, AL, USA) and used for further analysis. Both strands of the cDNA were sequenced by the Biotechnology Support Facility of the University of Iowa Rat Gene Discovery Program and UI-R-CXO-bws-e-04–0-UI cDNA was obtained from the National Center for Biotechnology Information (NCBI) Expressed Sequence Tag database (EST). We have named this new member ‘PRL-like protein-N’ (PLP-N). The location of the signal peptide for PLP-N was determined with the aid of SignalP analysis (Altschul et al. 1997). For in situ hybridization, uterine and placental tissues were removed intact without dissection and frozen in dry-ice-cooled heptane. All tissue samples were stored at −80°C until required for use. Tissue samples were collected from a minimum of three different rats for each data point. The University of Kansas Medical Center Animal Care and Use Committee approved all procedures for handling and experimentation with rodents.

Cell culture models

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determined by homology and the SignalP software program (version 2.0.b2, Nielsen et al. 1997). The rat PLP-N gene was identified by BLAST analysis (Altschul et al. 1990) of the NCBI rat genome database (http://www.ncbi.nlm.nih.gov/genome/seq/RnBlast.html) with the rat PLP-N cDNA nucleotide sequence.

Analysis of mRNA expression

The expression of rat PLP-N mRNAs in the rat was assessed by biochemical and cytochemical procedures.

Northern blot analysis

Northern blots were performed as previously described (Faria et al. 1990, Orwig et al. 1997b). RNA was extracted from tissues using TR.izol (Chomczynski & Sacchi 1987). Total RNA (20 µg) was separated on 1% formaldehyde–agarose gels and transferred to nylon membranes. Blots were probed with [32P]-labeled cDNAs for PLP-N, PLP-C or PL-II (Dai et al. 2002). Glyceraldehyde-3-phosphate dehydrogenase (G3PDH) cDNA was used as an internal reference to ensure integrity of the RNA samples.

In situ hybridization

PRL family mRNAs were detected in placental tissues using non-radioactive in situ hybridization as previously described (Braissant & Wahl 1998). A plasmid containing the PLP-N cDNA was linearized and used as a template for the synthesis of sense and anti-sense digoxigenin-labeled riboprobes according to the manufacturer’s instructions (Roche Molecular Biochemicals, Indianapolis, IN, USA). Tissue sections were air dried and fixed in ice-cold 4% paraformaldehyde in PBS for 15 min. Pre-hybridization was carried out in a humidified chamber at 50 °C in 5 × standard saline citrate (SSC), 50% deionized formamide, 1 × Denhardt’s reagent, 10% dextran sulfate and salmon sperm DNA (100 µg/ml). Hybridizations were performed in the same incubation conditions overnight. Slides were washed in 23 × SSC at room temperature for 30 min, followed by treatment with RNase-A (100 ng/ml) and additional washes with 23 × SSC for 30 min at room temperature, followed by washes with 2 × SSC for 1 h at 65 °C and 0.1 × SSC for 1 h at 65 °C. Tissue sections were blocked for 30 min and incubated with alkaline phosphatase-conjugated anti-digoxigenin antibody (1:500) in blocking buffer (Roche Molecular Biochemicals) for 2 h at room temperature. Slides were then washed and detection was performed using nitro blue tetrazolium (250 µg/ml) and 5-bromo-4-chloro-3-indolyl-phosphate (225 µg/ml; Roche Molecular Biochemicals).

Immunocytochemistry

Immunocytochemical analyses were used for the purpose of identifying trophoblast cells. All analyses were performed on 10 µm tissue sections prepared with the aid of
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identity of rat PLP-N with other members of the rat PRL family ranged from 26% to 39%. PLP-I, PLP-M, PLF-RP and PLP-F exhibited the highest level of overall amino acid sequence identity with the newly discovered rat PLP-N (Fig. 1B). The predicted signal peptides of PLP-I, PLP-M and PLP-N were very similar, whereas PLF-RP, PLP-F and PLP-N showed more extensive homology in their mature proteins (Fig. 1B). This latter characteristic was most notable between amino acids 41–52 of the mature PLP-N protein sequence. This region is homologous with the aromatic domains found in PLF-RP and PLP-F (Sahgal et al. 2000) and with amino acids encoded by exon 3 from members of the PLP-C subfamily (Dai et al. 1996b, 1998, Orwig et al. 1997d). Furthermore PLF-F, PLP-RP and PLP-N were classified into the same subfamily by analysis with the CLUSTAL W software program (Fig. 2). In the mouse, this subfamily includes PLP-N, PLP-F, PLF-RP and two additional PRL family members not yet discovered in the rat, PLP-E and PLP-O (Wiemers et al. 2003). The rat PLP-N gene was identified by BLAST analysis (Altschul et al. 1990) to be located on BAC clone CH230–190E4 (GenBank Accession No. Ac133698). This BAC clone is situated on chromosome 17 within the rat PRL family locus. Rat PLP-N possesses a 6-exon–5-intron gene structure similar to that reported for mouse PLP-N (Wiemers et al. 2003). Nucleotide sequences for the rat PLP-N cDNA showed 100% identity with the corresponding exon sequences of the rat PLP-N gene.

**PLP-N northern analyses**

The site(s) of PLP-N expression was assessed by northern blot analyses. A survey of several rat tissues (brain, heart, kidney, liver, muscle, intestine, lung, spleen, testis, ovary, uterus and uteroplacental compartment) indicated that PLP-N mRNA expression was restricted to the uteroplacental compartment (Fig. 3). A more systematic examination of expression in the uteroplacental compartment was performed. PLP-N mRNA levels were estimated in the junctional and labyrinth zones of the chorionic trophoblast placenta and in the metrial gland (Fig. 4). PLP-N mRNA was low or undetectable in all placental tissues (Fig. 4). In contrast, PL-II mRNA was strongly expressed in both the junctional and labyrinth zones of the chorionic trophoblast placenta, consistent with earlier observations (Deb et al. 1991a, Dai et al. 2002). We further examined PLP-N and PL-II expression in cells of the trophoblast cell lineage. The Rcho-1 trophoblast cell line is a useful model for...
investigating the trophoblast giant cell lineage (Faria & Soares 1991, Peters et al. 2000). We investigated PLP-N and PL-II mRNA levels in proliferating and differentiating Rcho-1 trophoblast cells. PL-II mRNA was abundantly expressed in the differentiating trophoblast cells, whereas PLP-N mRNA was not detectable (Fig. 4). The metrial gland harvested from day 20 of gestation was a rich source of PLP-N mRNA (Fig. 4). Metrial gland PLP-N expression was initiated between day 14 and 16 and increased as gestation progressed (Fig. 5).

In some of the PLP-N northern blots a higher molecular weight hybridizing band was detected. The identity of the larger hybridizing band was not determined. The larger hybridizing band could be an incompletely or alternatively processed PLP-N transcript.

Collectively, the northern blot experiments indicated that the metrial gland was a prominent source of PLP-N; and the chorioallantoic placenta and the trophoblast giant cell lineage were not major contributors to PLP-N synthesis. However, these experiments did not lead to the identification of the cellular source of PLP-N.

In situ localization of PLP-N

The major source of PLP-N, the metrial gland, contains a mixture of cell types, including immune cells, stromal...
cells, smooth muscle cells, endothelial cells and trophoblast cells (Pijnenborg et al. 1981, Peel 1989, Stewart & Mitchell 1992). In situ hybridization analyses were performed with sense and antisense probes in order to determine the cellular site of PLP-N synthesis. Sense probes did not provide detectable hybridization in any of the tissues investigated. Antisense probes localized PLP-N mRNA to migratory trophoblast cells. At day 14 of gestation, endovascular trophoblast cells possessed PLP-N mRNA (Fig. 6B,D). As gestation progressed, extraplacental migratory trophoblast cells positive for PLP-N mRNA were abundant. Both endovascular and interstitial migratory trophoblasts situated throughout the metrial gland expressed PLP-N mRNA (Figs 6B,D, 7B–D). Red arrowheads in Figs 6 and 7 are directed toward endovascular trophoblast cells and red arrows point towards interstitial trophoblast cells. Cells expressing cytokeratin and PLP-N possess an overlapping spatial distribution within the metrial gland (Fig. 7A). Cytokeratin immunoreactivity has previously been used as a means of identifying cells of the trophoblast lineage within the uteroplacental compartment (Hunt & Soares 1988, Kruse et al. 1999).

In vitro differentiation of PLP-N producing cells
There is a spatial continuum of trophoblast cells from the junctional zone of the chorioallantoic placenta to the metrial gland. This observation suggests that the junctional zone contains a population of cells capable of differentiating into the trophoblast cells. Consequently, primary cell cultures were established from the junctional zone of the gestation day 13 chorioallantoic placenta according to procedures previously described by members of our laboratory (Lu et al. 1994). The junctional zone trophoblast cells were cultured for 7 days and harvested for RNA analysis. As PLP-N mRNA is restricted to migratory trophoblast cells, we used PLP-N mRNA as a means of identifying migratory trophoblast cells. PLP-N mRNA expression was minimal in gestation day 13 or day 18 junctional zone dissected tissue, but was readily detected in the junctional zone cells after 7 days of culture (Fig. 8). The cultured gestation day 13 junctional zone cells also expressed PLP-C mRNA, a member of the PRL family and marker of spongiotrophoblast cell development (Deb et al. 1991b,c, Lu et al. 1994). PLP-C mRNA was not expressed in the metrial gland (Fig. 8). Thus junctional zone trophoblast cells showed the capacity to differentiate into both spongiotrophoblast cells and migratory trophoblast cells of the metrial gland.

Discussion
To date, 20 PRL-related genes have been identified in the rat. Each possesses the prototypical PRL structural backbone. In this report, evidence is presented for the discovery of a new member of the rat PRL family through mining of the NCBI EST database. On the basis of existing nomenclature, the new member was named PRL-like protein-N (PLP-N). Structurally, PLP-N exhibits significant similarities with other members of the PRL family. It is uniquely expressed in a population of migratory trophoblast cells that populate the metrial gland during the last trimester of pregnancy.

The PRL family represents a species-specific expansion of genes involved in adaptations to pregnancy (Mouse Genome Sequencing Consortium 2002). Although there are some notable differences, the PRL family genes of the mouse and rat, including PLP-N, exhibit striking similarities. A mouse ortholog for rat PLP-N has been recently described (Wiemers et al. 2003). Structures of the mouse and rat PLP-N gene, mRNA and proteins are very similar. Tissue expression patterns for mouse and rat PLP-N show some differences. In the mouse, PLP-N is expressed in trophoblast cells of the chorioallantoic placenta and in migratory trophoblast cells, whereas in the rat, PLP-N is only expressed in migratory trophoblast cells (present study). The significance of these two distinct patterns of expression is unknown. The human PRL family consists of a single member, PRL, whereas other species such as the cow possesses a multi-member PRL family with an independent pattern of expansion (Schuler & Kesler 1992, Soares & Linzer 2001). Orthogs for PLP-N have not been identified in the human, the cow or any species other than the mouse and rat.

PLP-N shares structural similarities with other members of the PRL family. These include an extensive amino acid
sequence identity within the predicted signal peptides of PLP-I and PLP-M. The mature PLP-N protein demonstrates considerable homology to PLF-RP and PLP-F, including conservation of the aromatic domain found in PLF-RP, PLP-F and members of the PLP-C subfamily (Sahgal et al. 2000). This domain is enriched in aromatic amino acids and is encoded by a short exon situated between exons II and III of the prototypical PRL gene structure, as is found in several members of the PLP-C subfamily (Dai et al. 1996b, 1998, Orwig et al. 1997a) and in mouse PLP-N (Wiemers et al. 2003). The biological significance of the aromatic domain is not known. Some insights about the biology of PLP-N may be gained from research on the physiology of PLF-RP and PLP-F: PLF-RP targets endothelial cells, where it regulates angiogenesis (Jackson et al. 1994, Bengtson & Linzer 2000, Regulier et al. 2001), whereas PLP-F modulates hematopoiesis (Zhou et al. 2002). Whether PLP-N influences endothelial or hematopoietic cells remains to be determined.

Migratory trophoblast cells synthesize PLP-N. These cells exist within the chorioallantoic placenta at midgestation and leave the chorioallantoic placenta during the last week of gestation. At midgestation, migratory trophoblasts are primarily situated within blood vessels, as best typified by endovascular trophoblast cells lining the central placental artery (Bridgman 1949, Pijnenborg et al. 1981, Zybina & Zybina 2000, Zybina et al. 2000). As gestation advances, endovascular trophoblast cells are located within mesometrial uterine blood vessels located outside the
chorioallantoic placenta. In addition, beginning between days 14 and 15 of gestation in the rat, interstitial trophoblast cells systematically leave the junctional zone of the chorioallantoic placenta (Bridgman 1949, Pijnenborg et al. 1981). They penetrate through the mesometrial decidua and colonize the mesometrial triangle, forming intimate relationships with the uterine vasculature. This mesometrial structure is referred to as the metrial gland (Selye & McKeown 1935). The metrial gland represents the major site of blood flowing to the placenta and, ultimately, to the fetus. The biological activities of PLP-N are not known. On the basis of the localization of this cytokine within migratory trophoblast cells of the metrial gland, we postulate that it may target cells within the metrial gland, including myometrial or vascular smooth muscle cells, immune cells, or uterine stromal cells during the last trimester of gestation. PLP-N is uniquely expressed only in endovascular and interstitial migratory trophoblast cells, and thus represents a useful biological marker for these cell types (present study). These migratory trophoblast cells appear to arise from the junctional zone of the chorioallantoic placenta (present study). The factor(s) underlying trophoblast cell ‘decisions’ to leave or remain within the rat chorioallantoic placenta are not known. Human trophoblast cell migration is regulated by an interplay of locally acting cytokines and growth factors, including

Figure 7 Identification of migratory PLP-N-expressing trophoblast cells. Conceptuses were isolated at day 16 of gestation and 10 µm cryosections were prepared. (A) Cytokeratin immunolocalization within the mesometrial conceptus. (B–D) In situ detection of PLP-N mRNA expression. PLP-N cDNA was used as template for the synthesis of digoxigenin-labeled antisense RNA probes. (C) Section corresponding to the boxed area shown in (B). (D) Section corresponding to the boxed area shown in (C). Original magnifications: (A), (B) × 20; (C) × 45; (D) × 200. Red arrowheads indicate the location of endovascular trophoblast cells and red arrows show the location of interstitial trophoblast cells. Please note that sense probes did not provide detectable hybridization in any of the tissues investigated.
colony stimulating factor-I, epidermal growth factor and insulin-like growth factor-I (Bass et al. 1994, Hamilton et al. 1998, Aplin et al. 2000, Nasu et al. 2000). We propose that trophoblast cell PLP-N expression may be an effective tool for monitoring the differentiation of the trophoblast cell invasive phenotype.

In summary, a new member of the PRL family, PLP-N, has been discovered. PLP-N expression uniquely identifies migratory trophoblast cells, including those localized to the metrial gland.

Acknowledgements

We would like to thank Davin Watne and Jared T Soares for assistance with the preparation of some of the figures.

Figure 8 PRL family gene expression in junctional zone trophoblast cell cultures. Total RNA was isolated from gestation day 13 junctional zone tissue (D13 JZ), gestation day 18 junctional zone cells cultured for 7 days (JZ cells), gestation day 18 metrial gland (D18 MG). Relative levels of PLP-N and PLP-C were determined by northern blot analysis. G3PDH was used as a control to demonstrate loading accuracy and the integrity of the RNA.

Funding

This work was supported by grants from the National Institutes of Health (HD020676, HD029797, HD033994, HD037123, HD039878). Rupasri Ain was supported by a postdoctoral fellowship from the American Heart Association.

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Received in final form 30 July 2003
Accepted 4 August 2003
Made available online as an Accepted Preprint 7 August 2003