

Technical note

Proteomic profiling reveals sub proteomes of the human placenta



Wendy E. Heywood^b, Rhian-Lauren Preece^a, Jeremy Pryce^a, Jenny Hallqvist^b,
Robert Clayton^b, Alex Virasami^a, Kevin Mills^b, Neil J. Sebire^{a,*}

^a Histopathology Department, Great Ormond Street Hospital for Children, Great Ormond Street, London, WC1N 3JH, UK

^b Centre for Translational Omics, UCL Great Ormond Street Institute of Child Health, 30 Guilford Street, London, WC1N 1EH UK

ARTICLE INFO

Article history:

Received 14 August 2017

Received in revised form

25 September 2017

Accepted 28 September 2017

Keywords:

Proteomics

Placenta

Chorion

Decidua

Trophoblast

ABSTRACT

Proteomic characterisation of the placenta has largely been focused on effect of disease, anatomical features or specific cell types. We describe an unbiased proteomic mapping analysis to investigate how the placental proteome changes throughout the organ. A transverse slice of a human placenta was sectioned into 1 × 1 cm samples. Sections were analysed using label free proteomics. Analysis revealed two distinct sub-proteomes that did not have anatomical significance. One had a muscular proteome and the other had distinct immunomodulation functions. Chorionic plate enriched proteins highlighted the fetal tissues high energy requirements whilst mechanisms of the decidua observed included modulation of cortisone levels.

© 2017 The Authors. Published by Elsevier Ltd. This is an open access article under the CC BY-NC-ND license (<http://creativecommons.org/licenses/by-nc-nd/4.0/>).

1. Introduction

There is growing interest regarding mechanisms by which placental function/dysfunction can affect short and long-term outcomes in fetal and postnatal life [1]. Variation in gene expression [2] and the transcriptome [3] of the human placenta have been described revealing its diverse molecular variation. However, to date little has been published in relation to detailed documentation of regional variation or anatomical mapping of the placental proteome. The human proteome atlas is able to describe the placental proteome through transcriptomic analysis and immunohistochemistry in regards to known placental cell types [4] but not in relation to regional or anatomical variation. Placental proteome studies have largely been conducted in respect to identifying changes in disease such as preeclampsia and fetal growth restriction [5,6]. Many studies have used limited samples from poorly defined sites within the placenta, such as a ‘cotyledon’. However, the tissue proteome of an organ can differ markedly according to sub-anatomical regions and functional variation, which can markedly affect methodologies for how proteomic studies should be performed. This study has investigated whether there is variation across the placenta and whether sub-proteomes of the human term placenta represent functional differences in relation to sites of the

placenta proteome map.

2. Methods

2.1. Detailed methods are provided in supplementary information

A placenta from a clinically uncomplicated, term normal vaginal delivery was used for this study, donated for research with appropriate consent and ethical approval as part of a larger study examining placental sampling. The placenta was sectioned as shown in Fig. 1B within 24 h of delivery having been initially refrigerated at 4°. Placental protein was run on a 1D PAGE [7,8] and analysed using label free proteomics as described previously [9].

3. Results & discussion

Total analysis identified 1096 quantifiable proteins of >95% confidence. The extreme end sections which form the peripheral margin of the placenta were removed from the analysis; as the proteomic integrity of these end sections was poor indicating this area undergoes significant degradation post-delivery. To identify if there was anatomical clustering of profiles the data were examined using principle components analysis, which shows an overall view of the variation of the placental proteome (Fig. 1A). The PCA resulted in a four component model ($R^2X(\text{cum}) = 0.581$ and $Q^2(\text{cum}) = 0.126$). The first and second component best described

* Corresponding author.

E-mail address: Neil.Sebire@gosh.nhs.uk (N.J. Sebire).

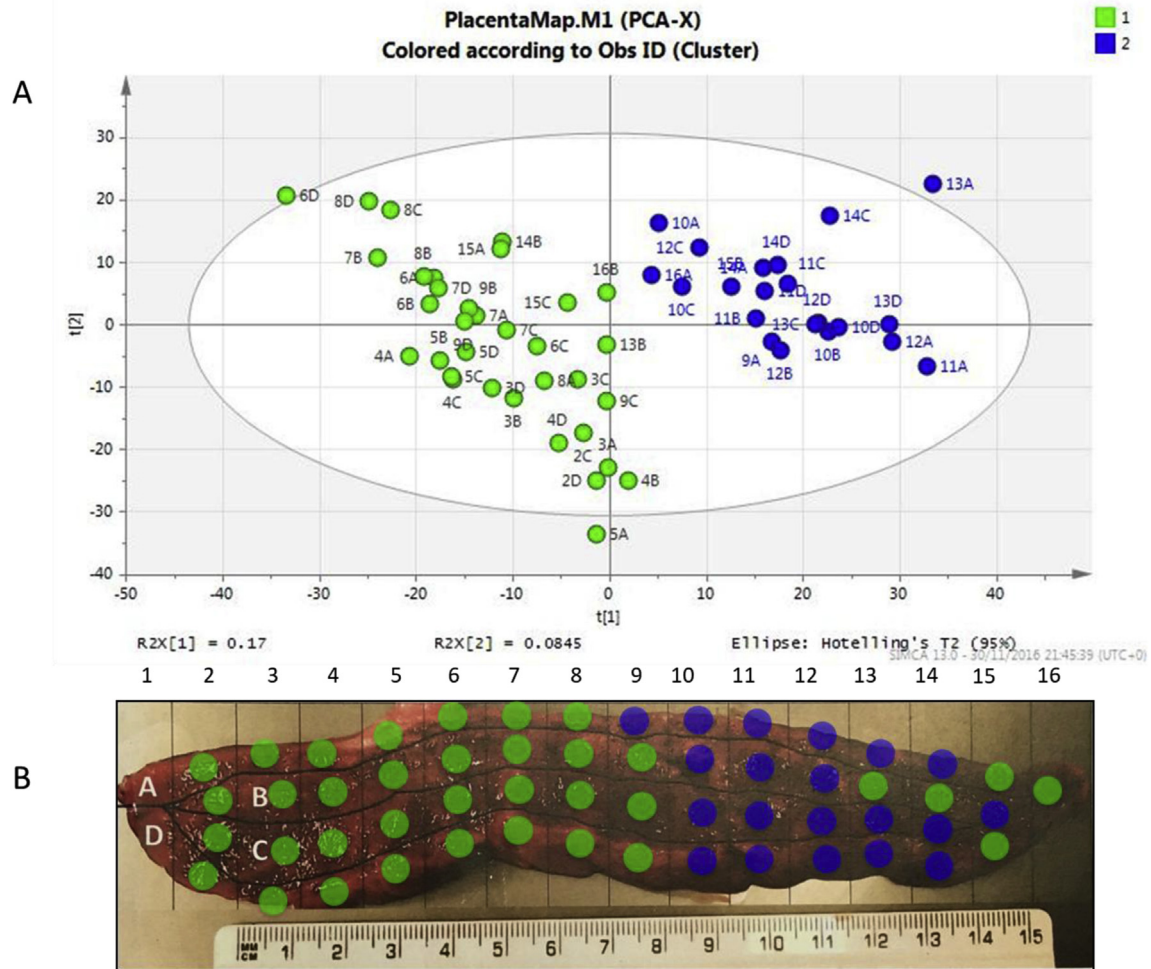


Fig. 1. A Placenta proteome PCA score plot. The score plot for the four component PCA model ($R2X(\text{cum}) = 0.581$ and $Q2(\text{cum}) = 0.126$). The first and second component best described the presence of 2 clusters (colour highlighted) of which first component explains 17% of the variation and component 2 explains 8.5% of the variation. B illustrates the sectioning and depicted subproteome clusters identified in A annotated to the anatomical structure of the placenta.

the separation of two major sub-proteomes of the human placenta.

Gene Ontology analysis of clustered regions demonstrates that cluster 1 is enriched with proteins involved in suppression of the innate immune response, in particular antigen presentation. Pathway analysis (Supplementary data 4) also identifies antigen presentation as a significant pathway. Cluster 2 appears to have enriched proteins involved in exocytosis, muscle hypertrophy and the signalling cascade of SMAD protein import to the nucleus and protein localisation. Pathway analysis also confirms platelet degranulation (or exocytosis) and keratinisation as enriched. The list for cluster 2 also seems to show more extracellular matrix proteins. These analyses suggest presence of sub-proteomes with differing functions.

3.1. Proteins that do not change throughout the placenta

Proteins that show the least variation across the placenta as indicated by a coefficient of variation (CV) of <30 across all sections were proteins involved in integrity of the cytoskeleton and metabolism (supplementary data 2). As expected many other proteins with lower CV values are known housekeeping proteins. These proteins which have uniform expression throughout the placenta could potentially be used for standardising future analyses.

3.2. Comparison by anatomical region; maternal vs fetal

The overall placenta PCA analysis does not show marked clustering according to whether sections are from the maternal and fetal sides of the placenta, indicating that these proteomic differences are small, presumably since the majority of tissue represents parenchymal villi from all sites. To identify which proteins are altered between the maternal and fetal sides rows A and D were compared. Eleven proteins were altered in both clusters for which seven were enriched towards the maternal side and four at the fetal side (supplementary data 5). A selection of interesting differently expressed proteins are shown in a placenta 3D heat map in Fig. 2. These include; Corticosteroid 11-beta-dehydrogenase isozyme 2 which acts to inactivate cortisone which is an essential process in pregnancy. This enzyme is inhibited by liquorice (*Glycyrrhiza uralensis*) and may explain why ingestion of liquorice is not advised during pregnancy [10,11]. Elevated Tryptophan-tRNA ligase, cytoplasmic, may be related to involvement in angiogenesis and shear stress response of the endothelium. Fetal side enriched proteins included the ECM component prolargin, creatine kinase B-type, required for tissues with high energy demands. cAMP-dependant protein kinase type II is involved in signal transduction in particular response to lipid and glucose metabolism and Apolipoprotein AI is a lipid transport protein. Interestingly, a recent proteomic

Download English Version:

<https://daneshyari.com/en/article/5585990>

Download Persian Version:

<https://daneshyari.com/article/5585990>

[Daneshyari.com](https://daneshyari.com)