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Title: Effect of oxygen on the expression of renin-angiotensin system components in a human trophoblast cell line

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Abstract: During the first trimester, normal placental development occurs in a low oxygen environment that is known to stimulate angiogenesis via upregulation of vascular endothelial growth factor (VEGF). Expression of the placental renin-angiotensin system (RAS) is highest in early pregnancy. While the RAS and oxygen both stimulate angiogenesis, how they interact within the placenta is unknown. We postulated that low oxygen increases expression of the proangiogenic RAS pathway and that this is associated with increased VEGF in a first trimester human trophoblast cell line (HTR-8/SVneo). HTR-8/SVneo cells were cultured in one of three oxygen tensions (1%, 5% and 20%). RAS and VEGF mRNA expression were determined by qPCR. Prorenin, angiotensin converting enzyme (ACE) and VEGF protein levels in the supernatant, as well as prorenin and ACE in cell lysates, were measured using ELISA. Low oxygen significantly increased the expression of both angiotensin II type 1 receptor (AGTR1) and VEGF (both P<0.05). There was a positive correlation between AGTR1 and VEGF expression at low oxygen (r=0.64, P<0.005). Corresponding increases in VEGF protein were observed with low oxygen (P<0.05). Despite no change in ACE1 mRNA expression, ACE levels in the supernatant increased with low oxygen (1% and 5%, P<0.05). Expression of other RAS components did not change. Low oxygen increased AGTR1 and VEGF expression, as well as ACE and VEGF protein levels, suggesting that the proangiogenic RAS pathway is activated. This highlights a potential role for the placental RAS in mediating the proangiogenic effects of low oxygen in placental development.

RESPONSE TO REVIEWERS

Manuscript # PL-15-100105

Title: 'Effect of oxygen on the expression of renin-angiotensin system (RAS) components in a human trophoblast cell line' **Authors:** Sarah J Delforce; Yu Wang; Meg E Van-Aalst; Celine Corbisier de Meaultsart; Brian J Morris; Fiona Broughton-Pipkin; Claire T Roberts; Eugenie R Lumbers; Kirsty G. Pringle

We would like to thank the Editor and reviewers for their generally favorable assessment of our manuscript entitled: 'Effect of oxygen on the expression of renin-angiotensin system (RAS) components in a human trophoblast cell line'. We consider that we have now addressed all of the reviewers' concerns. We have displayed our text corrections in red throughout the manuscript and described them in our response to reviewers (see below). The reviewers' comments are stated first, in italics, followed by our response.

Reviewer #1:

In this study the authors have shown that a low oxygen environment stimulates the expression of AGTR1 and VEGF mRNAs in HTR-8/SVneo cells in vitro. Both AT1R and VEGF are involved in angiogenesis and proliferation, which are key properties of first trimester EVTs. We showed that incubation in a low oxygen environment increased production of ACE and VEGF protein by HTR-8/SVneo cells and increased expression of the Ang II/AT1R pathway. They proposed that the placental RAS is under the influence of many factors including endocrine, molecular and environmental ones. No response required.

General comment: This is a well-written and designed article by experimented team. I have concerns about the reliability of the use of HTR-8/SVneo cells instead primary trophoblasts or other lines (i.e. BeWo, Jeg-3, JAr ...etc). How the authors justify the choice of this line and not another? This justification must be added to the section of discussion.

AUTHORS' RESPONSE: The following passage has now been added to the Materials and methods (page 2, lines 48-55) "HTR-8/SVneo trophoblast cells were chosen for the study as they are a transformed first trimester human extravillous trophoblast cell line. Because of this they are an ideal tool for investigating the effect of low oxygen on the proliferative/angiogenic RAS pathways in placental development and are superior to other cell lines such as choriocarcinoma BeWo cells. We have demonstrated previously that the pro-angiogenic/proliferative pathway of the RAS is expressed in the HTR-8/SVneo cell line [14] as occurs in the first trimester placenta *in vivo* [9], but which is not seen in BeWo cells [14]."

Reviewer #2:

This study by Delforce and colleagues characterize the role of varying oxygen tension on the expression of the renin-angiotensin system in trophoblast cell line HTR-8/SVneo. The main findings of this paper are that the mRNA expression of AGTR1 and VEGF at low oxygen level and the increased VEGF secretion from cells. The methodological approach is sound, although first trimester primary trophoblast cells or explants would have been preferred. The overall conclusion that low oxygen regulates the RAS may be overstated given that only one gene from the RAS is altered (AGTR1) and this was not confirmed at the protein level. This is important since although REN mRNA expression was altered, there were no changes at the protein level. Changes in VEGF expression with oxygen tension are well established and therefore these findings are more confirmatory than novel insights.

AUTHORS' RESPONSE: We agree that the conclusion that low oxygen regulates the RAS was overstated and have revised this section in the Discussion (page 6, lines 164-167): "The study also showed that even small reductions in O_2 have substantial effects on the expression of the AT₁R and pro-angiogenic factors such as VEGF. This was based on our finding that culture in 1% oxygen had a much more powerful effect than culture in 5% oxygen"

Minor comments:

Cell culture: Please provide the passage number(s) of the cell line if possible.

AUTHORS' RESPONSE: Cells were between passages 10-20. This information has now been added to the Materials and methods section of the manuscript (page 2, lines 58-59).

Line 53-54: RNA quality measurements are not appropriate indications of cell viability. Trypan blue exclusion, MTT, apoptosis assays may be required.

AUTHORS' RESPONSE: We have rephrased the text to clarify that RNA quantity, not quality, was used as an indicator of cell viability, page 3 lines 66-69: "RNA quantity was used as an indicator of cell viability and was assessed using the Nanodrop spectrophotometer, no differences in RNA quantity were detected between the treatment groups (data not shown). RNA quality was determined by agarose gel electrophoresis."

Line 73: Please check if this is supposed to be "one microliter" or "one milliliter"

AUTHORS' RESPONSE: We thank the reviewer for noting this error. This sentence was supposed to say "one millilitre" and has now been corrected (page 3, line 89).

Please provide details for regression analysis. Also please provide the R2 value.

AUTHORS' RESPONSE: The following passage has now been added to the Materials and methods (page 4, lines 115-116): "Pearson correlation coefficients were calculated to determine any relationships between the mRNA expression of genes." R² and r values are now present in the figure legends and have been added to the text (page 5, lines 137-138).

Reviewer #3: In this manuscript, the authors describe in vitro experiments demonstrating that low oxygen influence RAS and VEGF in HTR-8/SVneo cells. The authors have shown by different techniques that culture in low oxygen tension (1%) increased AGTR1 and VEGF mRNAs as well as ACE and VEGF proteins expression in trophoblast cells. Data are interesting and add knowledge to the area, however some points should be elucidate before manuscript publication.

AUTHORS' RESPONSE: No response required.

Specific comments:

1- How did the authors define oxygen tension (1%, 5% or 20%) for culture? As the effects on gene and protein expression were observed with very low oxygen tension (1%) it is important to clarify whether this oxygen tension reflects the physiological environment of early gestation.

AUTHORS' RESPONSE: We chose 1% and 5% oxygen tensions to closely reflect the environment experienced by the early gestation trophoblasts within the placental villi as well as the extravillous trophoblasts invading into the endometrium. We have now included a section in the introduction outlining the physiological environment in the first trimester (page 1, lines 7-12): "Rodesch *et al.* [2] established that the oxygen tension in the intervillous space at 8 weeks to be 17.9 mmHg (~2.5%) with a range of 5–30 mmHg (~0.7–4.3%) while the oxygen tension in the endometrium is higher at 39.6 mmHg (~5.7%) with a range of 25–70 mmHg (~3.5–10%). Thus there is an oxygen gradient experienced by the first trimester trophoblasts that can range from ~1% (in the placental villi) to up to 10% in endometrium."

2- Material and Methods- Are VEGF primers also described by Marques et al. (2011)? Did the authors investigate VEGF-A? There are multiple isoforms of VEGF-A. What was the VEGF isoform(s) investigated?

AUTHORS' RESPONSE: The VEGF primers described target VEGF-A and will detect all transcript variants of VEGF-A. VEGF primers were described in Pringle et al. (2011); this reference has been added to the text (page 3, line 81).

3- Material and Methods - Lines 53-54 - There was influence of oxygen tension in cell viability?

AUTHORS' RESPONSE: No influence of oxygen tension on cell viability was observed. This information has now been added to the Materials and methods (page 3, 66-69). Also see response to reviewer 1, above.

4- Material and Methods- Did the authors measure protein content (VEGF, prorenin and ACE) at 24hs of culture?

AUTHORS' RESPONSE: We have measured VEGF protein, as well as prorenin and ACE protein at 24 hours of culture. No effect of oxygen tension on the levels of these proteins was, however, observed. The Materials and methods section has been modified to reflect this (lines 98-99) and these results have now been included in the manuscript:

- Page 5, lines 142-144: "Although measurable levels of *REN* mRNA were found (Fig. 1), prorenin protein was low in both the culture medium and cell lysate of HTR-8/SVneo cells at 24 and 48 h incubation irrespective of the prevailing O₂ (only 48 h data shown; Fig. 3)."
- Page 5, lines 147-148: "No effect of oxygen tension on VEGF protein was observed at 24 h incubation (data not shown)."
- Page 5 lines 153-155: "No effect of oxygen tension on ACE protein in culture medium or cell lysates was observed at 24 h incubation (data not shown)."

5- Did the authors expect to identify ACE2, AGTR2 and MAS1 gene expression in HTR-8/SVneo cells. How to explain the observed data? Could this lack of expression alter expression of other RAS members in HTR-8/SVneo cells.

AUTHORS' RESPONSE: We did not expect to identify *ACE2*, *AGT2R* or *MAS1* gene expression in HTR-8/SVneo cells as we have previously demonstrated that this cell line does not express these mRNAs (Wang et al. 2012). A statement outlining the similarities between these 2 studies has been included in the Results (page 4, lines 126 – 127).

6- Results - Lines 117-118 - What is the physiological meaning of the correlations? What is the molecular correlation between these genes. Manuscript could be improved with more discussion about a hypothesized mechanisms related to the data.

AUTHORS' RESPONSE: The following passage has now been added to the Discussion (page 7, lines 203-209) "We found that *AGT1R* mRNA abundance was highly correlated with levels of both *VEGF* and *ATP6AP2* mRNA. The association between *AGT1R* and *ATP6AP2* mRNA suggests that increased levels of the (pro)renin receptor (ATP6AP2) may stimulate the activation of the placental RAS cascade and in turn increase AT₁R, while the association between *AGT1R* and *VEGF* mRNA suggests that increased expression of VEGF mRNA and protein, and thereby stimulate angiogenesis within the developing placenta."

7- Result - Lines 123 - How much prorenin was expected in the cell cultures? Why the values are low?

AUTHORS' RESPONSE: The values observed in the study were expected and were similar to those obtained in previous studies in which increases in prorenin protein were only observed when HTR-8/SVneo cells were stimulated with cAMP (see Wang *et al.* 2012). This was addressed in the Discussion (page 6, lines 168-176) where we suggest that the expression of prorenin (mRNA and protein) may be influenced by factors other than low oxygen including hormonal activation and/or activation by cAMP: "A low oxygen environment was not associated with increased expression of prorenin mRNA or

its protein. This suggests that a low oxygen tension alone is not responsible for the high levels of *REN* mRNA found in first trimester placentae [10]. It is more likely that increased *REN* mRNA expression in the placenta early in pregnancy is the result of hormonal activation and/or activation by cAMP of *REN* mRNA expression. Human chorionic gonadotropin (hCG) has been recognized as a stimulus for prorenin production by villous placenta [19], as have β -adrenoceptor agonists [19]. We have also shown previously that cAMP increases *REN* mRNA expression and prorenin protein secretion by HTR-8/SVneo cells [14]."

8- Discussion - Please check this statement in lines 141-145.

AUTHORS' RESPONSE: We have adjusted the text because we agree that the effect of $1\% O_2$ was only seen for *AGT1R* mRNA and not the whole RAS cascade. Please also see our response to Reviewer 1, above.

Conflict of Interest Statement Click here to download Conflict of Interest Statement: Conflict of Interest.docx

Conflict of Interest

The authors have no competing interests to declare.

Effect of oxygen on the expression of renin-angiotensin system components in a human trophoblast cell line

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ABSTRACT

During the first trimester, normal placental development occurs in a low oxygen environment that is known to stimulate angiogenesis via upregulation of vascular endothelial growth factor (VEGF). Expression of the placental renin-angiotensin system (RAS) is highest in early pregnancy. While the RAS and oxygen both stimulate angiogenesis, how they interact within the placenta is unknown. We postulated that low oxygen increases expression of the proangiogenic RAS pathway and that this is associated with increased VEGF in a first trimester human trophoblast cell line (HTR-8/SVneo). HTR-8/SVneo cells were cultured in one of three oxygen tensions (1%, 5% and 20%). RAS and VEGF mRNA expression were determined by qPCR. Prorenin, angiotensin converting enzyme (ACE) and VEGF protein levels in the supernatant, as well as prorenin and ACE in cell lysates, were measured using ELISA. Low oxygen significantly increased the expression of both angiotensin II type 1 receptor (AGTR1) and VEGF (both P<0.05). There was a positive correlation between AGTR1 and VEGF expression at low oxygen (r=0.64, P<0.005). Corresponding increases in VEGF protein were observed with low oxygen (P < 0.05). Despite no change in ACE1 mRNA expression, ACE levels in the supernatant increased with low oxygen (1% and 5%, P<0.05). Expression of other RAS components did not change. Low oxygen increased AGTR1 and VEGF expression, as well as ACE and VEGF protein levels, suggesting that the proangiogenic RAS pathway is activated. This highlights a potential role for the placental RAS in mediating the proangiogenic effects of low oxygen in placental development.

Keywords: renin-angiotensin system; pregnancy; placenta; hypoxia; gene expression

Abbreviations

ACE, angiotensin-converting enzyme; ACE1, angiotensin-converting enzyme 1; ACE2, angiotensin-converting enzyme 2; AGT, angiotensinogen; AT1R, angiotensin II type 1 receptor; *AGTR1* mRNA, angiotensin II type 1 receptor mRNA; Ang, angiotensin; ATP6AP2, ATPase, H⁺ transporting, lysosomal accessory protein 2 / (pro)renin receptor; BCA, bicinchoninic acid; EVT, extravillous trophoblast; HIFs, hypoxia inducible factors; iNOS, inducible nitric oxide synthase; RAS, reninangiotensin system; *REN* mRNA; (pro)renin mRNA; VEGF, vascular endothelial growth factor.

Highlights

- The human placenta develops in a low oxygen environment
- The placental renin-angiotensin system (RAS) is highly expressed in early pregnancy
- Culture in 1% O₂ increased AGTR1 mRNA and ACE protein in HTR-8/SVneo cells
- The latter was associated with increased VEGF mRNA and VEGF protein
- Stimulation of the placental RAS by low oxygen may be proliferative and angiogenic

1 Introduction

2 A low oxygen environment during the first trimester of pregnancy is required for 3 optimal placental development that is essential for supplying fetal demand in late 4 gestation. This early gestation low oxygen environment occurs as a result of extravillous 5 trophoblast (EVT) cell proliferation and subsequent invasion of the decidua and its 6 vasculature, initially occluding maternal spiral arterioles from about two weeks after 7 implantation [1]. Rodesch et al. [2] established that the oxygen tension in the 8 intervillous space at 8 weeks is 17.9 mmHg (~2.5%) with a range of 5-30 mmHg 9 (-0.7-4.3%) while the oxygen tension in the endometrium is higher at 39.6 mmHg 10 (~5.7%) with a range of 25–70 mmHg (~3.5–10%). Thus there is an oxygen gradient 11 experienced by first trimester trophoblasts that can range from ~1% (in the placental 12 villi) to up to 10% in the decidua. The low oxygen environment stimulates angiogenesis 13 and vascularization of the placenta. Poor placental development, characterized by 14 insufficient decidual invasion by EVTs, incomplete occlusion of maternal arterioles 15 with inadequate remodeling of maternal spiral arterioles, and early onset of maternal 16 blood flow to the conceptus, ultimately results in poor nutrient and oxygen exchange during the 2^{nd} and 3^{rd} trimesters. These are associated with intrauterine growth 17 restriction and preeclampsia [3, 4]. 18 19 A low oxygen tension stabilizes hypoxia inducible factors (HIFs) which promote 20 angiogenesis and vascularization by activating pro-angiogenic factors such as vascular 21 endothelial growth factor (VEGF), angiopoietins, factors involved in regulation of 22 vascular tone such as inducible nitric oxide synthase (iNOS) and proteins involved in 23 nutrient transfer such as transferrin and glycolytic enzymes [5-8]. 24 Another system that might regulate placental angiogenesis is the renin-angiotensin 25 system (RAS). Tissue RASs have been shown to be involved in the regulation of 26 angiogenesis, as well as cell proliferation and apoptosis [9]. We have shown that mRNA expression of prorenin (REN mRNA), (pro)renin receptor (ATP6AP2 mRNA), 27 angiotensinogen (AGT mRNA), angiotensin (Ang) II type 1 (AT₁R) (AGTR1 mRNA) 28 29 and Ang converting enzyme 2 (ACE2) (ACE2 mRNA) are all very high in early 30 gestation placentae compared with term [10]. In addition, we have shown that the 31 mRNA expression of placental VEGF is correlated with those of REN, ATP6AP2 and 32 AGTR1 mRNAs [10]. Thus the placental RAS is most active during the first trimester

33 and therefore could stimulate angiogenesis, as it does in other tissues. The ocular RAS 34 is stimulated by ischemia. The increased activity of the ocular RAS is associated with a 35 potent angiogenic response mediated via the Ang II/AT₁R pathway [11]. In addition, 36 early renal development requires activation of the RAS by a low oxygen milieu [12]. 37 While a low oxygen environment regulates placental development [13], the extent to 38 which the RAS is essential for normal placental development has not been established. 39 To investigate interactions between a low oxygen milieu and the placental RAS, we 40 examined the effects of low oxygen on the expression of the RAS and VEGF in a first 41 trimester human trophoblast cell line, HTR-8/SVneo, which we have previously shown 42 expresses mRNAs encoding those RAS pathway components that stimulate 43 angiogenesis in the eye and kidney [14, 15].

44

45 Materials and methods

46 Cell culture

47 HTR-8/SVneo cells are an immortalized first trimester trophoblast cell line (a kind 48 gift from Prof. Charles Graham, Queens University, Ontario). HTR-8/SVneo 49 trophoblast cells were chosen for the study as they are a transformed first trimester 50 human extravillous trophoblast cell line. Because of this they are an ideal tool for 51 investigating the effect of low oxygen on the proliferative/angiogenic RAS pathways in 52 placental development and are superior to other cell lines such as choriocarcinoma 53 BeWo cells. We have demonstrated previously that the pro-angiogenic/proliferative 54 pathway of the RAS is expressed in the HTR-8/SVneo cell line [14] as occurs in the 55 first trimester placenta in vivo [9], but which is not seen in BeWo cells [14]. 56 HTR-8/SVneo cells were cultured in RPMI-1640 medium (HyClone), supplemented 57 with 10% fetal bovine serum (SAFC Biosciences), 1 mg/ml antibiotic-antimycotic (Gibco) and 1% L-glutamine in 5% CO₂ in room air at 37°C (cells were between 58 59 passages 10–20). Cells were seeded at a density of 200,000 or 400,000 cells per well for 60 24 h or 48 h incubation, respectively. They were seeded in 6 well plates with 2 ml of 61 incubation medium per well and allowed to settle for 24 h, after which time the medium 62 was changed. Cells were then transferred to sealed oxygen chambers containing either 1%, 5% or 20% O_2 and 5% CO_2 in N_2 and cultured for 24 or 48 h, with chambers 63 64 flushed every 24 h. Cells were harvested and the incubation medium was collected at 24 and 48 h, then snap frozen in liquid nitrogen at -80°C for subsequent protein and

66 mRNA analyses. Three experiments were conducted in triplicate. RNA quantity was

67 used as an indicator of cell viability and was assessed using the Nanodrop

68 spectrophotometer, no differences in RNA quantity were detected between the treatment

- 69 groups (data not shown). RNA quality was determined by agarose gel electrophoresis.
- 70

71 Semi-quantitative real-time reverse transcriptase polymerase chain reaction (qPCR)

72 Total RNA was isolated using the RNeasy mini kit according to the manufacturer's 73 instructions (Qiagen). In addition, we examined the integrity of the total RNA in each 74 sample using gel electrophoresis. RNA samples were DNase treated (Qiagen) and total RNA was spiked with a known amount of Alien RNA (Stratagene), 10^7 copies per ug of 75 76 total RNA, before the RNA was reverse transcribed using a Superscript III RT kit with 77 random hexamers (Invitrogen). The Alien qRT PCR inhibitor alert system serves as a 78 reference for internal standardization [16]. qPCR was performed in an Applied 79 Biosystems 7500 Real Time PCR System using SYBR Green for detection. Each 80 reaction contained 5 µl of SYBR Green PCR master mix (Applied Biosystems), RAS 81 and VEGF primers that we have described previously [10, 17], cDNA reversed 82 transcribed from 10 ng total RNA, and water to 10 µl. Messenger RNA abundance was calculated as described previously, using the $2^{-\Delta\Delta CT}$ method and expressed relative to 83 Alien mRNA and a calibrator sample (a term placental sample collected at elective 84 85 Cesarean section) [10].

86

87 Extraction and quantification of total protein from cells

88 Protein was extracted from cells using a radioimmunoprecipitation assay (RIPA)

89 lysis and extraction buffer. One milliliter of RIPA buffer (50 mM Tris-HCl, 150 mM

90 NaCl, 1 mM EDTA, 1% NP-40, 0.5% sodium deoxycholate, 100 nM sodium

91 orthovanadate and Complete Mini Protease Inhibitor Cocktail tablets (Roche

92 Diagnostics Australia) and 10 µL of 100 nM PMSF were added to each sample.

93 Samples were incubated on ice for 30 min then centrifuged at 13,000 rpm at 4°C for 10

94 min. Supernatants were collected. Protein was quantified using the Pierce BCA Protein

95 assay kit (Life Technologies) according to the manufacturer's instructions.

96

97 Measurement of prorenin, VEGF and ACE by ELISA

98 Prorenin, VEGF and ACE concentrations in culture medium and cell lysates (at both 99 24 and 48 h) were measured using the Human Prorenin ELISA kit (Molecular 100 Innovations), Human VEGF Duoset ELISA kit (R&D systems) and the Human ACE 101 Duoset ELISA kit (R&D Systems), respectively, according to the manufacturers' 102 instructions, as described previously [18]. For prorenin, VEGF and ACE proteins in 103 culture medium and prorenin in cell lysates, all samples were assayed in duplicate on 104 one ELISA plate. Therefore there was no inter-assay variability. For ACE protein in cell 105 lysates, samples were assayed in duplicate over two plates and inter-assay variability 106 was 9.4%. Intra-assay coefficients of variation were 13.1% and 3.2% for prorenin 107 culture medium and cell lysate, 6.8% for VEGF culture medium and 14.2% and 2.9% 108 for ACE culture medium and cell lysate, respectively.

109

110 Data analysis

111 Two-way ANOVA with Tukey's multiple comparisons test was used to determine

112 the effects of differing oxygen tensions on the relative abundance of RAS mRNAs,

113 prorenin, ACE and VEGF protein levels. A Kruskal-Wallis non-parametric test was

114 used to determine the effects of differing oxygen tensions on prorenin concentration in

both the culture medium and cell lysate at 48 h. Pearson correlation coefficients were

116 calculated to determine any relationships between the various mRNA levels. GraphPad

117 Prism (Prism version 6.0) was used for all graphs and correlation analyses and the SPSS

118 statistical package (SPSS for Windows, Release 17.0.0. Chicago) was used for all other

119 analyses. Significance was set at P < 0.05.

120

121 Results

122 Effect of O₂ on RAS and VEGF mRNA expression

123 At all oxygen tensions, HTR-8/SVneo cells expressed genes of the pro-angiogenic

124 pathway of the RAS, i.e., AGT, REN, ATP6AP2, ACE1 and AGTR1, as well as VEGF,

- 125 but expression of genes of the anti-angiogenic RAS pathway, i.e., ACE2, AGT2R, MAS1
- 126 was not detected (Figure 1); this is consistent with previously published data on the
- 127 expression of RAS genes in this cell line [14].

128 Expression of AGT, REN, ATP6AP2 and ACE1 mRNAs was not affected by the 129 prevailing oxygen tension, but AGTR1 and VEGF mRNA expression were significantly 130 higher in HTR-8/SVneo cells cultured in low oxygen (1% O₂) compared with 5% or 131 20% O₂ at 48 h (all P < 0.01). A similar trend was seen in 24 h incubates. This did not, 132 however, reach statistical significance (Fig. 1). 133 There was a significant effect of incubation time on the expression of AGT, REN, 134 ATP6AP2, ACE1 and AGTR1 mRNAs, in that expression of these mRNAs was 135 significantly higher at 48 h compared with 24 h (all P<0.05; Fig. 1). 136 A significant correlation was found between AGTR1 and VEGF mRNA and between AGTR1 and ATP6AP2 mRNA (r=0.6378 and 0.6500 respectively, $R^2 = 0.4067$ and 137 138 0.4733 respectively, both *P*<0.01; Fig. 2). 139 140 Effect of O_2 on the concentration on prorenin and VEGF protein concentration in the 141 culture medium 142 Although measurable levels of REN mRNA were found (Fig. 1), prorenin protein 143 was low in both the culture medium and cell lysate of HTR-8/SVneo cells at 24 and 48 144 h incubation irrespective of the prevailing O_2 (only 48 h data shown; Fig. 3). 145 After 48 h VEGF levels in the culture medium of HTR-8/SVneo cells grown in 1% 146 O₂ were much higher than levels in medium from samples cultured in 5% and 20% O₂ 147 (P < 0.05 and P < 0.01, respectively; Fig. 4). No effect of oxygen tension on VEGF 148 protein was observed at 24 h incubation (data not shown). 149 Effect of O_2 on the concentration of ACE in the culture medium 150 151 After 48 h culture, HTR-8/SVneo cells cultured in 1% and 5% O₂ had significantly 152 higher levels of ACE protein in the supernatant than cells cultured in 20% O_2 (P<0.05; 153 Fig. 5). ACE levels in the cell lysate were not affected by O_2 . No effect of oxygen 154 tension on ACE protein in culture medium or cell lysates was observed at 24 h 155 incubation (data not shown). 156 157 Discussion

158Incubation of primary HTR-8/SVneo trophoblast cells at a low (1%) oxygen tension

159 increased the expression of *AGTR1* and *VEGF* mRNA and stimulated the production of

ACE and VEGF proteins. We propose, therefore, that a low oxygen milieu activates the pro-angiogenic pathway of the RAS by increasing AT_1R and ACE protein levels and that these changes are associated with increases in VEGF, an important angiogenic factor.

164 The study also showed that even small reductions in O_2 have substantial effects on 165 the expression of the AT_1R and pro-angiogenic factors such as VEGF. This was based 166 on our finding that culture in 1% oxygen had a much more powerful effect than culture 167 in 5% oxygen. Thus, increased AGTR1 and VEGF mRNA expression and VEGF protein 168 only occurred at 1% O₂. A low oxygen environment was not associated with increased 169 expression of prorenin mRNA or its protein. This suggests that a low oxygen tension 170 alone is not responsible for the high levels of REN mRNA found in first trimester 171 placentae [10]. It is more likely that increased *REN* mRNA expression in the placenta 172 early in pregnancy is the result of hormonal activation and/or activation by cAMP of 173 *REN* mRNA expression. Human chorionic gonadotropin (hCG) has been recognized as 174 a stimulus for prorenin production by villous placenta [19], as have β -adrenoceptor 175 agonists [19]. We have shown previously that cAMP increases REN mRNA expression 176 and prorenin protein secretion by HTR-8/SVneo cells [14]. Expression of AGT and 177 ATP6AP2 mRNAs were not affected by oxygen. This suggests that their high 178 expression in early gestation placentae [10] is not due to the low oxygen milieu. 179 Although the expression of ACE1 was not stimulated by culturing cells in a low 180 oxygen environment, we did observe increased ACE protein in medium from 181 incubations carried out in low oxygen. Goyal et al. have observed a similar effect [20]. 182 In animal studies they showed that maternal hypoxia was associated with upregulation 183 of placental ACE protein without a corresponding increase in ACE1 mRNA levels [20]. 184 They concluded that this was because hypoxia-regulated miR-27, which regulates ACE 185 translation, was down-regulated [20]. It is possible that a hypoxia-regulated miRNA in 186 the HTR-8/SVneo cell cultures could be similarly affected by oxygen. 187 In a previous publication we reviewed evidence suggesting that the low oxygen 188 milieu of early placental development activated the placental RAS, stimulating 189 angiogenesis and cell proliferation [15]. The RAS pathway most likely to be involved in 190 inducing these effects is the prorenin/AGT/ACE/Ang II/AT₁R pathway because the Ang 191 II/AT₁R blocking drug, candesartan, blocks Ang II-induced endothelial proliferation

192 [21]. Hypoxia inducible factor (HIF-1 α) is essential for normal trophoblast

- 193 differentiation, since both HIF-1 α and AT₁R gene ablation in mice impairs placental
- vascularization [22, 23]. Ang II acting via the AT₁R causes cell proliferation [24].
- 195 We were surprised that a low oxygen milieu had no effect on prorenin expression
- and secretion. There may, however, be a number of factors acting on the placental
- 197 renin/AGT/ACE/Ang II/AT₁R pathway that upregulate different components, so
- 198 ensuring increased placental Ang II production. As mentioned above, hCG and cAMP
- are potent stimuli for placental prorenin release. cAMP also stimulates AGT mRNA
- 200 expression [25]. MicroRNAs (miRNAs) may affect the stability or translation of ACE1
- 201 mRNA as has been observed in animal experiments [20]. AGTR1 mRNA expression in
- 202 HTR8/SVneo cells is, moreover, increased by low oxygen.
- 203 We found that *AGTR1* mRNA abundance was highly correlated with levels of both
- 204 *VEGF* and *ATP6AP2* mRNA. The association between *AGTR1* and *ATP6AP2* mRNA
- suggests that increased levels of the (pro)renin receptor (ATP6AP2) may stimulate the
- activation of the placental RAS cascade and in turn increase AT_1R , while the
- 207 association between AGTR1 and VEGF mRNA suggests that increased AT₁R in first
- 208 trimester placenta may increase expression of VEGF mRNA and protein, and thereby
- 209 stimulate angiogenesis within the developing placenta.
- 210 In conclusion, we have shown that a low oxygen environment stimulates the
- 211 expression of AGTR1 and VEGF mRNAs in HTR-8/SVneo cells in vitro. Both AT₁R
- and VEGF are involved in angiogenesis and proliferation, which are key properties of
- 213 first trimester EVTs. We showed that incubation in a low oxygen environment increased
- 214 production of ACE and VEGF protein by HTR-8/SVneo cells and increased expression
- of the Ang II/AT₁R pathway. We propose that the placental RAS is under the influence
- 216 of a number of endocrine, molecular and environmental factors.
- 217

218 Author Contributions

- 219 E.R. Lumbers and K.G. Pringle were responsible for designing the experiments.
- 220 C.T. Roberts, B.J. Morris and F. Broughton-Pipkin assisted in developing the
- 221 experimental concepts that were the basis of the experiments.
- 222 Y. Wang, S.J. Delforce, M.E. Van-Aalst and C. Corbisier de Meaultsart carried out the
- 223 experiments. Wang and Delforce analysed the data.

- 224 Delforce, Wang, Lumbers, Morris and Pringle interpreted the data and drafted the
- 225 manuscript.
- 226

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- 299

300	LEGENDS TO FIGURES
301	Fig. 1. Effect of O_2 on expression of RAS components in HTR-8/SVneo cells at 24 and
302	48 h. (A) AGTR1 and (B) VEGF mRNA were significantly upregulated by low oxygen
303	(1% O ₂) at 48 h when compared with 5% and 20% O ₂ (*, both $P < 0.01$). (B) VEGF
304	mRNA was significantly increased at 48 h compared with 24 h by 1% O_2 (#, P<0.05).
305	(A) AGTR1, (C) ATP6AP2, (D) REN, (E) ACE1 and (F) AGT mRNA expression
306	significantly increased with incubation time (•, all $P < 0.05$). Data are presented as mean
307	\pm SEM.
308	
309	Fig. 2. The correlations between VEGF and AGTR1 mRNA abundance and between
310	AGTR1 and ATP6AP2 mRNA abundance in HTR-8/SVneo cells cultured at 1% O ₂ . (A)
311	<i>VEGF</i> and <i>AGTR1</i> (r=0.638, R ² =0.407 and P=0.0044); (B) <i>AGTR1 and ATP6AP2</i>
312	$(r=0.650, \mathbb{R}^2=0.473 \text{ and } P=0.0035).$
313	
314	Fig. 3. Levels of prorenin protein in culture medium and cell lysates of HTR-8/SVneo
315	cells cultured in varying oxygen tensions for 48 h. Neither culture medium (A) nor cell
316	lysate (B) prorenin protein levels were affected by oxygen tension. Data are presented
317	as mean \pm SEM.
318	
319	Figure 4. The effect of O_2 on VEGF protein concentration in the culture medium of
320	HTR-8/SVneo cells at 48 h. Culture in 1% O_2 for 48 h significantly increased the
321	concentration of VEGF compared with culture in either 5% or 20% O_2 (*, $P < 0.05$ and
322	$P < 0.01$ respectively). Data are presented as mean \pm SEM.
323	
324	Fig. 5. The effect of O_2 on ACE protein concentration in the culture medium and cell
325	lysate of HTR-8/SVneo cells at 48 h. (A) The concentration of ACE in the culture
326	medium of HTR-8/SVneo cells was significantly increased after 48 h of culture in both
327	1% and 5% O_2 when compared with the concentration measured in medium cultured in
328	20% O ₂ (*, both P <0.05). (B) ACE protein levels in cell lysates were unaffected by
329	oxygen. Data are presented as mean \pm SEM.











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Figure 4



Figure 5

