Role for the thromboxane A₂ receptor β-isofrom in the pathogenesis of intrauterine growth restriction

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Intrauterine growth restriction (IUGR) is a pathology of pregnancy that results in failure of the fetus to reach its genetically determined growth potential. In developed nations the most common cause of IUGR is impaired placentation resulting from poor trophoblast function, which reduces blood flow to the fetoplacental unit, promotes hypoxia and enhances production of bioactive lipids (TXA₂ and isoprostanes) which act through the thromboxane receptor (TP). TP activation has been implicated as a pathogenic factor in pregnancy complications, including IUGR; however, the role of TP isoforms during pregnancy is poorly defined. We have determined that expression of the human-specific isoform of TP (TPβ) is increased in placentae from IUGR pregnancies, compared to healthy pregnancies. Overexpression of TPα enhanced trophoblast proliferation and syncytialisation. Conversely, TPβ attenuated these functions and inhibited migration. Expression of the TPβ transgene in mice resulted in growth restricted pups and placentae with poor syncytialisation and diminished growth characteristics. Together our data indicate that expression of TPα mediates normal placentation; however, TPβ impairs placentation, and promotes the development of IUGR, and represents an underappreciated pathogenic factor in humans.

Intrauterine growth restriction (IUGR) is defined as failure of the fetus to reach its genetically pre-determined growth potential1-2. IUGR can be diagnosed in utero by estimated fetal weight for gestational age less than the 10th percentile, accompanied by Doppler ultrasound detecting decreased, absent or reversed end diastolic flow in the umbilical arteries3-5. IUGR is a leading cause of fetal mortality and morbidity, second only to pre-term delivery4-6; however, IUGR offspring that survive the perinatal period are at increased risk of cardiovascular disease and renal complications in adulthood7. Currently no biomarkers exist that predict pregnancies at risk of developing IUGR nor are there effective preventative treatments. In the absence of effective prophylaxis or treatment the growth restricted fetus is delivered early, often pre-term, which contributes further to the risk of adverse neonatal outcomes8.

IUGR can be classified as either symmetrical or asymmetrical. Symmetrical IUGR results from maternal factors (such as malnutrition and alcohol consumption) and congenital infections. Conversely, asymmetrical growth primarily arises from the presence of a placental pathology3,9. Placentae of growth restricted pregnancies are smaller in weight, have reduced number and diameter of villi and a reduction in the number and area of capillaries within the villi indicating impaired vascularisation and branching angiogenesis10-12. They also exhibit greater rates of infarction, fetal vessel thrombosis and chronic villitis than the placenta of normal pregnancies13. These placental pathologies are precipitated by morphological and biochemical abnormalities at the cellular level. IUGR

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placental weight in rodent models associated with IUGR30,31, suggesting a strong causal role for TXA2 in disease.

Hypoxia and ultimately oxidative stress10.

Peroxydation of the membrane lipid arachidonic acid results in the formation of F-series isoprostanes, which are markers of oxidative stress13,18. However, isoprostanes are also biologically active and exert their effects through activation of the thromboxane (TXA2) receptor (TP)17,19. Bone fide TXA2 generation is also perturbed in IUGR with the TXA2:prostacyclin ratio increased in maternal serum and perfusates of the placental circulation from IUGR pregnancies compared to those with “normal” outcomes20,21. TXA2 overproduction by placentae and platelets is prevalent in human pregnancies complicated by cigarette smoking22, diabetes mellitus23 and alcohol consumption24 which are associated with IUGR. Furthermore, infusion of TXA2, analogues (such as U46619 and STA31) into rodents produces a TXA2-like syndrome characterised by a 10–15% reduction in pup weight, which persisted until 28 and 77 days post-partum for male and female offspring, respectively25–27. The pups from these pregnancies exhibit a pattern of growth restriction characteristic of asymmetrical IUGR with reduced body/liver weight with sparing of brain changes in weight or energy metabolites25,26,28,29. Moreover, TP antagonists block the vasoconstriction and normalised pup and placental weight in rodent models associated with IUGR20,31, suggesting a strong causal role for TXA2 in disease.

The role of the TP is well established in a number of diseases; however, very little information exists on the role of the specific isoforms during pregnancy. The G-protein coupled TP has two isoforms in humans, TPα and TPβ, derived from alternate splicing of a single gene32,33. Both isoforms are identical through the first 328 amino acids but differ in the length of the cytoplasmic C-terminus (with the tails 15 and 79 amino acids for TPα and TPβ, respectively). The cytoplasmic tail is a poor discriminator of G-protein coupling34,35 promoting complacency over the differential role of these two isoforms in human disease. Our study sought to identify the complement of TP isoforms expressed in placentae from normal, healthy and IUGR affected pregnancies and to understand the differential regulation of trophoblast function by TPα and TPβ. We have identified enhanced TPβ protein expression in placentae from IUGR pregnancies compared to those from normal pregnancies. Moreover, we demonstrate that enhanced TPβ expression impairs trophoblast function promoting placental insufficiency that is characteristic of IUGR. Pup weight and placental density in the TPβ-transgenic mice are significantly reduced and trophoblasts in these placentae exhibit impaired proliferation and syncytialisation consistent with IUGR in humans. In conclusion, we propose that dysregulation of TP isoform expression in the placenta is a significant cause of spontaneous IUGR in human pregnancy, which is amenable to targeted therapy to prevent both the short and long term mortality and morbidity associated with this condition.

Results

TPβ expression is increased in the syncytiotrophoblast layer of placentae from growth restricted pregnancies. The distribution and expression of TP isoforms in placental tissue, and whether this changes during IUGR, has not been documented. We assessed TP isoform expression in third trimester placental tissue from “normal” uncomplicated pregnancies and those affected by IUGR (Table 1). As expected, the birth weight of babies from IUGR pregnancies was significantly lighter (P = 0.004) compared with those from control gestationally matched pregnancies (Table 1). Immunoblotting of placental protein extracts revealed a 2.8 fold increase in overall TP expression in IUGR placentae (P < 0.005) (Fig. 1a). This was associated with increased expression of both TPα (1.6 fold; P = 0.017) and TPβ (2.6 fold; P < 0.005) (Fig. 1a) after normalisation to the loading control. When examined by IHC, TPα expression was ubiquitous in control placentae (including the syncytiotrophoblast layer, endothelial cells and villous stroma); however, TPβ expression was essentially absent, with any staining primarily restricted to the endothelium of villous blood vessels (Fig. 1b, arrows). Conversely, TPβ expression was markedly increased in the syncytiotrophoblast layer in IUGR tissues while TPα expression was not different (Fig. 1b). TP isoforms are regulated by alternative splicing. Transcript levels for Total TP and TPα did not change in IUGR; however significantly lower expression of TPβ was evident in IUGR compared to control placental tissues (Fig. 1c). This change in TPβ gene expression also contributed to a change in the ratio of TPα:TPβ (Fig. 1c). These data suggest that neither transcription nor alternate splicing were responsible for the enhanced TPβ protein expression and that protein stabilisation is the most likely explanation.

Table 1. Clinical data on placental samples from “normal” and IUGR affected pregnancies. Clinical characteristics are shown for the control (gestationally matched) and IUGR patients recruited for placental sample collection. Data include maternal age, body mass index (BMI), systolic (SBP) and diastolic (DBP) blood pressure, gestational age at delivery, birth weight and fetal gender. Data are represented as mean (SD) (n = 53 and 22). Independent samples T-test, *P < 0.05, ***P < 0.005.

<table>
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<th>Subgroup</th>
<th>Maternal Age</th>
<th>BMI</th>
<th>SBP (mm/Hg)</th>
<th>DBP (mm/Hg)</th>
<th>Gestational Age at Delivery (weeks)</th>
<th>Birth Weight (g)</th>
<th>Fetal Gender (M/F)</th>
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<td>Control</td>
<td>31.5 (5.0)*</td>
<td>24.9 (6.1)</td>
<td>113.5 (12.4)</td>
<td>67.6 (10.7)*</td>
<td>34.0 (4.4)</td>
<td>2443.0 (989.2)***</td>
<td>67.9%/32.1%</td>
</tr>
<tr>
<td>IUGR</td>
<td>34.1 (3.9)</td>
<td>22.6 (3.2)</td>
<td>119.6 (14.7)</td>
<td>75.2 (10.1)</td>
<td>35.6 (2.9)</td>
<td>1745.0 (632.3)</td>
<td>45.4%/54.5%</td>
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TP isoforms differentially regulate trophoblast proliferation. To explore the individual role of TP in placentation we characterised the effects of TPα and TPβ in the choriocarcinoma cell lines, BeWo and JEG-3, commonly used as in vitro models of placental development. Both BeWo and JEG-3 cells constitutively expressed low levels of TPα but no TPβ (Supplementary Fig. S1). Transfection and stable selection of TPα, TPβ and the "tail-less" deletion mutant (TPΔ328) led to a 5–10 fold increase in receptor expression at the protein level compared to empty vector (GV) transfected cells (Supplementary Fig. S1). Overexpression of TPΔ328 or TPα conferred no growth advantage compared to GV transfected cells or untransfected BeWo (Fig. 2a) or JEG-3 (Fig. 2b) cells. However, activation of TPβ ablated proliferation in BeWo cells (P = 0.014) and reduced growth of JEG-3 cells by 30% (P = 0.001), (Fig. 2a,b, respectively). Thus, TP isoforms show specific and opposite regulation of trophoblast function.
Proliferation is a balance between cell division and cell death. Examination of apoptosis by flow cytometry, either at baseline or under serum-starvation conditions, did not show differences between any of the TP transfected cell lines and the ∅ control (Fig. 2c). Therefore the effects of TP on trophoblast growth must be related to cell cycle progression. Examination of randomly cycling cells indicated expression of TPΔ328 had no effect on BeWo cell cycle progression compared to ∅ transfected or untransfected BeWo cells (Fig. 2d). TPα overexpression elevated the number of cells in S-phase (Fig. 2d) suggesting enhanced G0/G1 exit which was confirmed by reduced ability to arrest cells with L-Mimosine (Supplementary Fig. S2) and enhanced S-phase entry within 24 hours of withdrawal of L-Mimosine (Supplementary Fig. S2). This was accompanied by enhanced phosphorylation of Ser780 and Ser795 in Rb in TPα expressing cells (Fig. 3a), the product of enhanced expression of Cyclin A (3.2 fold), D1 (2.6 fold) and CDK6 (4.1 fold) (Fig. 3b,c). Conversely, TPβ activation promoted accumulation of cells in the G2/M phase (Fig. 2d) suggesting the reduced proliferation was due to an inability to complete cell division. This phase of the cell cycle is driven by Cyclin B-CKD1 complexes, which appear to be normally expressed in TPβ-BeWo cells (Fig. 3b). However, expression of the cell cycle inhibitor proteins p21 (5.2 fold), p27 (3.9 fold) and p53 (2.6 fold) were all increased with TPβ overexpression compared to TPα, TPΔ328, ∅ and untransfected BeWo cells (Fig. 3d; P < 0.05). Therefore the cells were unable to progress through the G2/M phase causing inhibition of TPβ cell proliferation (Fig. 2a).

TPβ expression reduces BeWo syncytialisation. Another essential function for placentation is formation of the fetal-maternal interface, which is lined by multinucleate syncytiotrophoblasts resulting from membrane fusion (syncytialisation). The high expression of TPβ in the syncytiotrophoblasts (Fig. 1b) strongly suggested this function would be regulated by TPβ. Treatment of BeWo cells with the TP agonist I-BOP (200 nM) did not affect basal expression of E-cadherin or syncytin by BeWo cells (Supplementary Fig. S3). Conversely, forskolin treatment (100 μM) reduced E-cadherin and augmented syncytin expression by immunofluorescence (Fig. 4a) and immunoblotting (Fig. 4b) which are both markers of syncytialisation. TPα activation
on BeWo cells accelerated loss of E-cadherin (40% compared to 16% in ∅ cells; Fig. 4a,b) and enhanced syncytin expression (2.5 fold; Fig. 4b) compared to untransfected and ∅ expressing cells (Fig. 4a). TPΔ328 did not augment E-cadherin loss but significantly increased syncytin expression (1.15 fold gain) indicating the effects of TPα on differentiation were mediated by both isoform specific (E-cadherin loss) and common (elevated syncytin expression) pathways (Fig. 4a,b). Conversely, TPβ-BeWo cells retained more E-cadherin membrane staining and higher expression by immunoblotting, with no induction of syncytin expression after forskolin treatment (Fig. 4a,b).

Collectively these data suggest that TPα promotes syncytialisation during placentation while TPβ expression abrogates the process. These findings are consistent with the expression pattern of TP isoforms with high TPα expression in “normal” pregnancies promoting robust placentation while TPβ promotes insufficient placentation associated with pathological pregnancies.

TPβ expression slows trophoblast cell migration. A key step in the pathogenesis of asymmetric growth restriction is the invasion of the uterine wall and remodelling of spiral arteries by cytotrophoblasts. The role of TP isoforms in regulating this process is undocumented. Motility of ∅ and TPΔ328 transfected JEG-3 cells was unchanged, compared to untransfected cells, in the presence of I-BOP (Fig. 5a). TPα-JEG-3 cells had a mild reduction (14%) in migration compared to ∅ controls (P < 0.001), but not compared to TPΔ328-JEG-3 cells (P = 0.160) (Fig. 5a). TPβ overexpression reduced migration (36%) which was significant compared to ∅ (P < 0.001), TPΔ328 (P < 0.001) and TPα (P < 0.001) expressing cells (Fig. 5a).

Migration of cells is a complex process regulated by cytoskeletal dynamics, extracellular matrix (ECM) interactions and planar cell polarity. To determine the mechanism by which TPβ activation inhibited trophoblast migration we examined adhesion of cells to several matrix components. Consistent with the migration data, transfection of JEG-3 cells with ∅, TPΔ328 and TPα did not influence binding of trophoblasts to matrix components (Fig. 5b). Conversely, activation of TPβ enhanced JEG-3 attachment to collagen I and collagen IV based matrices (3 fold and 2.5 fold, respectively) while attachment to fibronectin and vitronectin was unchanged (Fig. 5b). This was significantly different to ∅-JEG-3 (P = 0.025) and TPΔ328-JEG-3 (P = 0.01) cells when binding to collagen I. Thus, the blunted migration in TPβ-JEG-3 cells might be explained by enhanced adhesion to ECM.

To further determine the role of adhesion in the inhibition of trophoblast migration by TPβ we examined focal adhesion formation in these cells. Association of vinculin with integrins denotes focal adhesion complexes actively engaged with matrix binding and is therefore implicated in the control of cell motility36,37. I-BOP
treatment of TPα-JEG-3 cells (Fig. 5c) reduced the levels of vinculin staining (both size and number of focal adhesions) compared to ∅ and TPΔ328 transfected cells (which were no different to untransfected cells) which is indicative of fewer focal adhesions. By comparison, I-BOP treatment of TPβ-JEG-3 cells enhanced focal adhesion size 3 fold compared to TPα and TPΔ328 expressing cells, although focal adhesion number was maintained overall (Fig. 5c). Consistent with the migration data, there was no difference in focal adhesion size and number in TPβ cells at baseline indicating the receptor needed to be activated for these effects. The enhanced focal adhesion size in TPβ-JEG-3 cells is consistent with the greater adhesion to collagen-based matrices and suggests that enhanced adhesion to the matrix is the mechanism by which TPβ activation impedes trophoblast movement.

Focal adhesion kinase (FAK) is centrally involved in the signalling response of integrins, which interact directly with the ECM to transmit signals from the extracellular environment. Consistent with the enhanced size of focal adhesions in TPβ-JEG-3 cells, TPβ activation enhanced FAK phosphorylation at Y397 (the FAK autophosphorylation site) up to 1.8 fold (P = 0.014), Y577 by 2 fold (P ≤ 0.011) and Y861 1.5 fold (P ≤ 0.01) compared to ∅- and TPΔ328-JEG-3 cells (Fig. 5d). Non-significant changes in Y576 and Y925 phosphorylation were also observed in I-BOP treated TPβ-JEG-3 cells compared to the other sub-lines. Phosphorylation of Y577 enhances FAK activation, which would enhance focal adhesion formation, while phosphorylation of Y861 promotes focal adhesion targeting. The enhanced Y577 phosphorylation is indicative of elevated Src activation; however, activation of Src (denoted by enhanced Y418 and loss of Y529 phosphorylation) only occurred in TPα-JEG-3 cells, not TPβ-JEG-3 cells, (Supplementary Fig. S4) indicating the enhanced Y577 phosphorylation is likely the result of another Src family member.
Suppression of placental and fetal growth in TPβ-transgenic mice. Our in vitro data suggest that TPβ activation inhibits normal trophoblast function; however, whether expression of TPβ is a cause or a
consequence of the development of IUGR remained unanswered. We characterised the TPβ-transgenic (TPβ-Tg) mice to determine a causal role for TPβ in the development of IUGR. The pups born to TPβ-Tg dams were on average 26% smaller than pups born to Wt dams (P < 0.001; Fig. 6a,b). In terms of assessed growth potential, 37% of all pups born to TPβ-Tg dams fell below the 10th percentile rank consistent with the human classification of IUGR. There was a 12% reduction in placental weight for pups born to TPβ-Tg dams compared with the pups born to Wt dams (Fig. 6c), although this did not reach significance. Placental diameter was significantly larger in growth restricted pups born to TPβ-Tg dams (P = 0.014; Fig. 6d) resulting in the placentae being less dense (P = 0.029; Fig. 6e) as well as small. These data indicate that pups born to transgenic mice are growth restricted and not just small for gestational age.

The placental pathology of the growth restricted pups born to TPβ-Tg dams was immediately obvious. Assessment of the labyrinth in the placentae from TPβ-Tg dams was 10% smaller than its Wt counterparts (P < 0.001; Fig. 7a,b). To explain the altered labyrinth size we documented the differentiation/syncytialisation and proliferation of trophoblasts in placentae using immunohistochemistry. Consistent with human data, Wt mouse placentae were devoid of E-cadherin staining denoting robust fusion in the syncytiotrophoblast layer (Fig. 7c,d). The area of E-cadherin expression in the labyrinth of growth restricted pups born to TPβ-Tg dams was 5.3 fold higher (Fig. 7c,d) compared to Wt mice. Moreover, the amount of proliferation, as denoted by PCNA staining, was 1.8 fold lower in the labyrinth of placentae from pups born to TPβ-Tg dams versus Wt dams (Fig. 7e,f). Together these changes in cellular function indicate that placentae of growth restricted litters from TPβ-Tg dams have reduced rates of syncytialisation and proliferation. This in vivo data is consistent with the abnormal function of TPβ overexpressing trophoblasts in vitro and clearly implicates enhanced TPβ-expression on the syncytiotrophoblasts of human placentae as causal to the development of IUGR.

Discussion
We report that TP ligands manipulate placentation through direct actions on trophoblasts, in an isoform specific fashion. Stimulation of TPα enhanced trophoblast differentiation and proliferation while TPβ activation prevented these activities and inhibited migration. These data have important potential implications for placentation as progeny of TPβ-Tg mice displayed fetal growth restriction, accompanied by poor trophoblast proliferation and syncytialisation, and reduced placental weight. Most importantly, TPβ expression was enhanced in placentals.
biopsies from growth restricted human pregnancies suggesting isoform switching in the pathogenesis of fetal growth restriction. The cause of idiopathic IUGR, especially asymmetrical IUGR, is unknown. The discovery of the role of the different TP isoforms in placentation, particularly that dysregulation of TPβ is observed in the growth restricted placenta, raises the possibility that this is a significant, yet targetable, pathogenic factor in human pregnancy. The acknowledgement of the role of the different TP isoforms in placentation enriches our perception of the role for TP activation in pregnancy and expands the repertoire of pathophysiological functions ascribed to the receptors.

Our observations add to the dichotomy in the literature over the role of TP isoforms in the regulation of health and disease. TPα is known to be pro-angiogenic, whilst TPβ inhibits angiogenic responses42. Despite these divergent findings both TP isoforms are pro-tumorigenic in epithelial cells in a tissue/organ specific fashion43,44. A previous study documented TP isoform transcripts in primary trophoblasts; however, no studies have documented TP isoform protein expression. This study showed TPα mRNA was ubiquitous and at high levels in primary trophoblasts in vitro but TPβ expression was more variable. However, the outcomes of these pregnancies were not known due to the trophoblasts being isolated from first trimester tissue42. Therefore it is possible that the trophoblasts with high TPβ expression may have been from placentae with some sort of pathology, such as IUGR. Our protein data confirms the robust expression of TPα in trophoblast cells in vitro and in vivo but we do not see the same expression of TPβ in placenta from “normal” pregnancies. TPβ expression in placentae from pathological pregnancies has not been previously documented and it is unknown how the enhanced TPβ expression results in IUGR. The control over TP splicing is likely driven by differential promoter utilization43; however, the lower expression of TPβ mRNA in IUGR placentae suggests that the mechanism is due to post-transcriptional regulation and protein stabilisation. Expression of TPβ, but not TPα, is increased in conditions with elevated free radical production44 due to both increased trafficking of the membrane and protein stabilisation. Such changes are well established in the IUGR placenta and may be the reason for TPβ upregulation in disease.

Previous studies on the pathogenic role of TXA2 in IUGR were performed in small animal model45–47 where TPβ expression is absent. These data suggest that appropriate activation of TPα may promote healthy placentation and angiogenesis; however, overstimulation of TPα may result in ligand-induced receptor desensitization45–47 essentially creating an environment devoid of TP activation. The bona fide IUGR placenta has been reported to produce elevated TXA2 levels45–47 although TXA2 synthase levels have not been documented. Little is known about the relative importance of TP isoforms to placentation in humans or the consequences of TP activation on trophoblasts. Stimulation of primary human villous cytotrophoblasts (from term placenta) with a TXA1 mimetic abrogated differentiation and enhanced apoptosis and p53 expression46. Our data support many of these findings, including the failure to differentiate; however, we did not observe any increase in apoptosis. This may reflect differences between the low concentrations of TP agonist used here (100 nM or lower), which are more reflective of reported (patho)physiological levels49, versus the supra-pathological (10 μM) doses in the original study48. What wasn’t clear from the previous data was the TP isoform responsible for these effects. Our data clearly show that TPβ mediates the adverse effects of TXA2 on placentation, with neither TPα nor TPβ producing the same responses. These data further accentuate the divergent roles for TPα and TPβ as human-specific disease modifiers and emphasize the need for greater understanding of the biology of the individual isoforms. The signalling that perturbs trophoblast function and placentation resides in the divergent C-terminal residues of TPβ. Indeed, we have previously shown that the C-terminus of TPβ is all that is required for the inhibition of angiogenesis49. The 79 residue cytoplasmic tail of TPβ is a poor discriminator of G-protein coupling (TPα and TPβ share >70% of Gα units as second messengers) but has significantly different protein-protein interactions than that of TPα. Recently, protein-protein interactions of the tail of TPβ have been shown to manipulate the behavior of smooth muscle and cancer cells50,51 independent of G-protein coupling. However, both interactions enhance proliferation and migration in the respective cell types and interactions are yet to be identified that inhibit cellular function as described here. We believe that pathways unrelated to G-protein activation, mediated through direct interaction of proteins with the cytoplasmic tail of TPβ, regulate the negative effects of TPβ on placentation and trophoblast migration.

The primary deficit in the original report of the TPβ-Tg mice was a decrease in placental and pup size52; however, reductions in pup and placental weights were not quantified nor was any basis for the growth restriction explored. In the previous study only homozygous transgenic mice showed the IUGR phenotype, which could have resulted from the mixed genetic background (SV129xC57BL/6). Our cohort have been backcrossed to C57BL/6 for a total of 16 generations and still maintain the growth restricted phenotype suggesting that TPβ expression is the cause of the growth restriction. The trophoblast layer of these placentae was poorly formed with smaller labyrinthine zones, most likely a result of the diminished trophoblast proliferation caused by TPβ activation. This is likely to have been compounded by the anti-angiogenic effects of TPβ39, which would have further limited placental growth. The failure to syncytialise would have compromised fetal growth, not only by reducing the transport of nutrients and waste products of metabolism, but because the integration of fetomternal signalling would be compromised in these placentae53. Again, the anti-angiogenic effects of TPβ may also play a role in fetal growth with vascular bed development possibly retarded in the TPβ-Tg pups. Thus, this model represents a unified theory for the origins of idiopathic IUGR in humans and a new target for therapeutic development.

The identification that TXA2 levels are elevated in IUGR, compared to “normal” pregnancies, has resulted in targeting TXA2 generation to provide therapeutic benefit. As in cardiovascular disease, these attempts have primarily focussed on low-dose aspirin (60–150 mg/day) to prevent TXA2 synthesis by platelets. Aspirin crosses the placenta and has a favourable safety profile in pregnancy48,54. Despite this, the outcomes of large scale clinical trials for the use of aspirin to prevent IUGR have been largely negative. Historically, initiation of aspirin therapy after 16 weeks gestation55 has been shown to be less beneficial than early in pregnancy53,54,55. However, whether IUGR is associated with a normotensive or hypertensive pregnancy also changes whether fetal, or maternal and fetal circulations are involved, which changes the response to aspirin therapy41. Moreover, aspirin appears to preferentially inhibit TXA2 production in villous cell types other than the trophoblast53. Our data would suggest that
Figure 7. TPβ expression in mice reduces labyrinthine formation and prevents trophoblast proliferation and differentiation as part of the IUGR phenotype. (a) Representative images of H&E stained sections of placentae from pups born to Wt (n = 89) and TPβ-Tg (n = 48) dams at day 18.5 of gestation. (b) From the H&E stained sections, the size of the labyrinthine zone of placentae from pups (n = 9) born to Wt (□) dams and pups (n = 31) born to TPβ-Tg (■) dams was elucidated using ImageJ software. Representative images (c) and quantification (d) of E-cadherin expression in the labyrinthine zone in placentae from pups (n = 47) born to Wt (□) dams and pups (n = 59) born to TPβ-Tg (■) dams. Representative images (e) and quantification (f) of proliferation (PCNA staining) in the labyrinthine zone in placentae from pups (n = 53) born to Wt (□) dams and pups (n = 59) born to TPβ-Tg (■) dams. All sections stained for E-cadherin and PCNA were imaged at 200x magnification, scale bar: 100 μm. The data are represented as mean ± SEM. Independent samples T-test, *P < 0.05, ****P < 0.001.
all these factors are issues in the effectiveness of aspirin therapy as TPβ is activated not just in response to TXA2, but also isoprostanes[16]. Administration of aspirin to promote appropriate placentation, and negate the effects of TPβ, would only be effective early in pregnancy and may need to target both maternal and fetal circulations, as TPβ is expressed on both fetal and maternal endothelial cells and syncytiotrophoblasts. As such, current therapies for IUGR are insufficient as they do not appropriately target TPβ activation, resulting in persistent disease. Conversely, the use of dual action TP/TXA2 synthase antagonists has not been trialled in IUGR, although these drugs are effective in animal models of IUGR[31], and may represent a better therapeutic option than aspirin.

Collectively our data suggest that the upregulation of TPβ in placentae of IUGR pregnancies initiates a cascade which compromises trophoblast proliferation, invasive potential and differentiation, to produce an environment unable to support placental growth and fetal development. Our data are the first to elucidate the role of the human-specific isofrom of TP, TPβ, as a pathogenic component in the development of IUGR and suggests that targeting TPβ activation, or TP splicing, offers a therapeutic avenue for IUGR, which currently has no effective therapies.

**Materials and Methods**

**Materials.** The TXA2 mimic I-BOP ([(1S-[1α,2α(Z),3β{1E,3S*},4α)]-7-[[3-hydroxy-4-[(4-iopropoxy)-1-butenyl]-7-oxabicyclo[2.2.1]hept-2-yl]-5-heptenoic acid] was from Cayman Chemical. Forskolin and L-Mimosine were from Sigma Aldrich. All cell culture reagents were from Life Technologies, immunohistochemical reagents from DAKO and general chemicals from Sigma-Aldrich unless otherwise stated.

**Tissue collection.** This study was performed in accordance with the guidelines approved by the Northern Sydney Local Health District Human Research Ethics Committee, St Leonards, NSW, Australia and was assigned the site specific assessment number 0912–348M and the Australian national ethics application form number HREC/09/HARBR/165. Collection of human placental tissue was performed with written informed consent obtained prior to delivery. Placentae were collected from non-labouring or minimally labouring women at delivery by lower caesarean section from pregnancies complicated by IUGR or gestationally matched control pregnancies. Patient populations were matched for body mass index (BMI) and smoking, gestational diabetes, pre-existing hypertension and pre-eclampsia were exclusion criteria (Table 1). IUGR was diagnosed prior to delivery by ultrasound, where estimated fetal weight of less than the 10th percentile for gestational age was associated with elevated fetoplacental vascular resistance defined as the Systolic/Diastolic ratio over the 95th percentile or absent or reversed flow in diastole in the umbilical artery, and confirmed by weight at birth. Placental tissue was washed in sterile saline and either snap frozen in liquid nitrogen or fixed in 10% (v/v) neutral buffered formalin.

**Cell culture and transfection with vectors encoding TP isofoms.** Choriocarcinoma cell lines (BeWo and JEG-3) were maintained in RPMI culture medium supplemented with 10% (v/v) fetal calf serum, 2 mM L-glutamine, 10 μg/mL penicillin and 10 U/mL streptomycin. Sub-confluent monolayers were passaged weekly using TrypLE Express. To ensure maximal TP activation, BeWo and JEG-3 cells were treated with 200 nM I-BOP. Syncytialisation of BeWo cells was induced by treating with 100 μM forskolin[56].

The coding sequence for TPa, TPβ, and the deletion mutant (TPΔ298) were cloned into pcDNA3.1. TP receptor constructs were introduced into BeWo and JEG-3 cells using Effectene (Qiagen Pty Ltd.) or Lipofectamine 2000 (Life Technologies) transfection reagents, respectively, following the manufacturers’ instructions. Empty vector (pZ) was used as a control. Stable expression was achieved in BeWo and JEG-3 cells after antibiotic selection using 200 μg/mL and 400 μg/mL Geneticin (G418 sulfate), respectively. Receptor expression was confirmed using semi-quantitative RT-PCR on RNA from stably expressing cells.

**Quantitative gene expression using Droplet Digital PCR.** RNA isolation was performed using TRIzol Reagent according to the manufacturer’s recommendations. Tissue samples were weighed, placed into an M-tube with the appropriate amount of TRIzol Reagent and disaggregated using a gentleMACS Dissociator (Miltenyi Biotec). Cell lines were scraped directly into TRIzol Reagent. RNA quality and concentration were assessed using a NanoDrop 1000 Spectrophotometer (Thermo Fisher Scientific) and subsequently 1 μg RNA was reverse transcribed to cDNA using BioScript Reverse Transcriptase (Bioline). Gene expression was quantitated using the QX200 Droplet Digital PCR (ddPCR) System (Bio-Rad). Reactions were prepared using the QX200 ddPCR EvaGreen Supermix, 50 ng cDNA and the following primers which span the intron-exon boundary responsible for alternative splicing of TP or an internal control gene (HPRT1):

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<th>Primer</th>
<th>Sequence</th>
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<td>TP (Total) Forward</td>
<td>5′-CGGGTTCAAGGGATTCTCG-3′</td>
</tr>
<tr>
<td>TP (Total) Reverse</td>
<td>5′-CTGTTGGAGAGTCTCAAAGGAAG-3′</td>
</tr>
<tr>
<td>TPa Forward</td>
<td>5′-CGACCACCGAGGACAGGAGG-3′</td>
</tr>
<tr>
<td>TPa Forward</td>
<td>5′-CGAGGCTGTGAGGACAGGACAr3′</td>
</tr>
<tr>
<td>TPβ Forward</td>
<td>5′-GAGGCGCTGAGGACAGC-3′</td>
</tr>
<tr>
<td>TPβ Reverse</td>
<td>5′-CGTCACCTGAGGTCAGC-3′</td>
</tr>
<tr>
<td>TPΔ298 Forward</td>
<td>5′-TTCAAGGAAGCACTGTACCCC-3′</td>
</tr>
<tr>
<td>HPRT1 Forward (internal)</td>
<td>5′-GACTCCAGATGTTCTCAACAC-3′</td>
</tr>
<tr>
<td>HPRT1 Reverse</td>
<td>5′-GACTCCAGATGTTCTCAACAC-3′</td>
</tr>
</tbody>
</table>

Droplets were generated from the reactions using QX200 Droplet Generation Oil for EvaGreen on the QX200 Droplet Generator (Bio-Rad) according to the manufacturer’s instructions. The droplets were cycled on a T100 Gradient Thermal Cycler (Bio-Rad) according to the manufacturer’s protocol using an annealing temperature of 60°C. The droplets were analysed using the QX200 Droplet Reader and analysed using QuantaSoft Software (Bio-Rad). Detection of fewer than <15,000 droplets in a sample indicated poor quality and was therefore excluded from the analysis.
Chemokinesis assay. JEG-3 cells were chosen as BeWo cells did not migrate well in these assays. Transfected JEG-3 cells were grown to confluence and motility assessed as previously described\(^9\) in the presence of I-BOP (200 nM) or vehicle control (equivalent volume of ethanol). Media was supplemented with L-Mimosine (400 μM) to negate the effects of proliferation on the outcome of the assay. Photographs were taken at the time of injury (0 hours) and after 24 and 48 hours. T-Scratch software was used to quantify the distance migrated and the migration of I-BOP treated cells was then normalised to vehicle control treated cells.

Adhesion assay. Adhesion of JEG-3 cells to matrix proteins, including collagen I (Life Technologies), collagen IV (Sigma-Aldrich), fibronectin (Life Technologies) and vitronectin (Promega) was assessed as previously described\(^8\). JEG-3 cells were cultured in the presence of 200 nM I-BOP, or vehicle control, for 24 hours prior to the assay. Cells were detached using TrypLE Express, plated into pre-coated 24 well plates at 1 × 10\(^5\) cells/well and incubated for 37 °C for 30 minutes. After washing to remove unattached cells, wells were fixed with 10% (w/v) neutral buffered formalin and adhesion detected by staining with 1% (w/v) methylene blue as previously described\(^9\).

Cell proliferation, apoptosis and cell cycle analysis by flow cytometry. JEG-3 and BeWo cell proliferation was assessed by seeding 2 × 10\(^4\) cells/well into 24 well plates. Cells were cultured in the presence of 200 nM I-BOP, or vehicle control, for up to 7 days. Cells were enumerated by cell counting or staining with 1% (w/v) methylene blue. Cell cycle and apoptosis analyses were performed on randomly cycling, and growth arrested, BeWo cells as previously described\(^8\). DNA content was analysed using a BD FACSCalibur Flow Cytometer and ModFit software. Apoptosis was quantified as the hypodiploid peak in these samples.

SDS PAGE, slot blots and immunoblotting. Protein lysates from transfected BeWo and JEG-3 cell lines were obtained as described\(^9\). For placental tissue, biopsies were weighed, placed into an M-tube with the appropriate amount of lysis buffer containing 1% (v/v) Y30 and disaggregated using a gentleMACS Dissociator (Miltenyi Biotec). For phospho-proteins, lysates were prepared using hot SDS-PAGE buffer as previously described\(^8\). Concentration of protein lysates was estimated using the Pierce BCA Protein Assay Kit (Life Technologies) and 30 μg separated by SDS-PAGE under reducing conditions on 8%-10% (w/v) polyacrylamide gels. After transfer to PVDF membranes, immunoblotting was performed as previously described\(^8\) for: Cyclin A, B1, D1, E, p16, p21, p27, p53 (1:1000, Santa Cruz Biotechnology) and Retinoblastoma Protein (Rb) (pSer\(^780\), pThr\(^795\) and pSer\(^780\)/pThr\(^795\)) (1:1000, Cell Signaling Technology), E-cadherin and synctyin (Santa Cruz Biotechnology), FAK (pY\(^397\), pY\(^397/402\), and pY\(^397/407/411\), 1:2000, Cell Signaling Technology) and Src (pY\(^418\), pY\(^397/413\); 1:2000, Life Technologies). Total protein content was assessed by immunoblotting for GAPDH (Santa Cruz Biotechnology) or unphosphorylated versions of the proteins of interest (FAK/Src/Rb; 1:2000, Santa Cruz Biotechnology and Cell Signaling Technology).

For slot blotting, 100 μg of protein lysate from placental samples was loaded into a manifold (Bio-Rad) and adsorbed onto nitrocellulose membranes under vacuum. Membranes were probed for total TP content (Santa Cruz Biotechnology), TPα, TPβ (using our in-house antibodies)\(^40\) and loading for cellular content was assessed using a Histone H1 antibody (1:1000, Santa Cruz Biotechnology). Protein expression by immunoblotting was detected using the appropriate HRP-conjugated secondary antibodies (1:3000, Dako) in combination with an enhanced chemiluminescence detection system (PerkinElmer) and ImageQuant LAS 4000 (GE Healthcare). Densitometry of protein bands was quantified using ImageJ software.

Immunofluorescent imaging for cell membrane structure and focal adhesion complexes. Cells were plated onto glass coverslips pre-coated with 0.2% (w/v) gelatine and 0.2 μg/mL fibronectin. For syncytialisation assays BeWo cell culture medium was supplemented with 200 nM I-BOP, or with without 100 μM forskolin, and replenished every 24 hours over 3 days. For detection of focal adhesions, JEG-3 cells were exposed to 200 nM I-BOP, or vehicle control, for 24 hours. Immunostaining after fixation in 4% (w/v) paraformaldehyde was performed as previously described\(^9\), which involved incubation with antibodies against E-cadherin (1:500, Santa Cruz Biotechnology) or vinculin (Sigma Aldrich) and detected using an Alexa Fluor 488 conjugated second antibody (1:800; Molecular Probes). Coverslips were mounted onto glass slides and counterstained with DAPI. Slides were imaged using an Olympus DP71 fluorescent microscope at 400x magnification using Olympus DP controller software and then images were overlayed using ImageJ software.

Estimates of placental structure in TPβ-Tg and wild-type mice. The animal component of this study was performed in accordance with the guidelines approved by the Kolling Institute of Medical Research's Animal Ethics Committee, St Leonards, NSW, Australia and was assigned the protocol number 1311–015A. TPβ-transgenic (TPβ-Tg) mice\(^52\) were bred from their original mixed C57Bl6/SV129 background to be congenic on the C57Bl/6 background (>15 generations). Mice were bred as hemizygote crosses with wild-type (Wt) C57Bl/6j mice. Mice were housed within the Kearns facility at the Kolling Institute within individually vented cages under PC2 conditions and had food and water ad libitum. Timed intra-strain mating of Wt and TPβ-Tg mice (10–12 weeks old) were performed using nulliporous female mice. Females with copulation plugs were separated at E0.5 and pregnancies carried until E18.5 when mice were euthanized using inhalation anaesthetic, tissues collected and biometrics on the pups assessed. The cohort of mice included 89 pups born to 14 WT dams and 48 pups born to 7 TPβ-Tg dams. Quantification of labyrinthine area was performed using ImageJ software.

Immunohistochemistry. Formalin fixed, paraffin embedded human and mouse placental tissues were sectioned (4 μm), dewaxed and rehydrated through graded alcohols. Haematoxylin and eosin (H&E) staining was performed to identify placental structure. Immunohistochemistry (IHC) was performed as previously described\(^9\) with antigen retrieval performed at 99 °C for 20 minutes in EnVision Flex retrieval solution (high pH, Dako). Slides were incubated in sequenza racks (Thermo Fisher Scientific) overnight at 4 °C with our primary antibodies that.
against TPα and TPβ (1:400; raised in-house)46, E-cadherin (2 μg/mL; Cell Signaling Technology) and PCNA (1 μg/mL; Santa Cruz Biotechnology) or anti-rabbit IgG isotype control at equivalent concentrations. NovaRed peroxidase HRP substrate kit (Vector Laboratories) was used to visualise the staining following the manufacturer’s instructions. Sections were counterstained using Mayer’s haematoxylin (Merck Millipore) and mounted with Eukitt mounting medium (Grale HDS). All placental sections were imaged using a Nikon ECLIPSE 80i light microscope with a Nikon Digital Sight Control Unit and Nikon Digital Sight DS-5M camera at 200x magnification using NIS-Elements software version 3.

Statistical analyses. - Statistical analyses based on One-way analysis of variance (ANOVA) were performed using SPSS (version 20). LSD or Games-Howell post-hoc tests were used to analyze densitometry readings from western blot analyses and percentage wound closure in scratch assays. Cell proliferation data were assessed using a general linear model analysis as well as linear contrasts within the program R. All other data was analysed using Independent samples T-tests or Mann-Whitney U non-parametric tests using SPSS. All data are displayed as mean ± standard error of the mean (SEM).

References


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Author Contributions
The study and experiments were developed by K.L.P., V.S. and A.W.A. K.L.P., V.S., D.H.U. and A.W.A. performed the experiments. K.L.P., V.S., D.H.U. and A.W.A. analysed the data. K.L.P. and A.W.A. wrote the manuscript. K.L.P., S.M., V.T., A.S., V.C., J.M.M. and A.W.A. assisted with experimental design, discussed results and edited the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

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