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Increased NOS coupling by the metabolite tetrahydrobiopterin (BH4) reduces preeclampsia/IUGR consequences

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ABSTRACT
Preeclampsia (PE) is a high-prevalence pregnancy disease characterized by placental insufficiency, gestational hypertension, and proteinuria. Overexpression of the A isoform of the STOX1 transcription factor (STOX1A) recapitulates PE in mice, and STOX1A overexpressing trophoblasts recapitulate PE patients hallmarks in terms of gene expression and pathophysiology. STOX1 overexpression induces nitroso-redox imbalance and mitochondrial hyper-activation. Here, by a thorough analysis on cell models, we show that STOX1 overexpression in trophoblasts alters inducible nitric oxide synthase (iNOS), nitric oxide (NO) content, the nitroso-redox balance, the antioxidant defense, and mitochondrial function. This is accompanied by specific alterations of the Krebs cycle leading to reduced L-malate content. By increasing NOS coupling using the metabolite tetrahydrobiopterin (BH4) we restore this multi-step pathway in vitro. Moving in vivo on two different rodent models (STOX1 mice and RUPP rats, alike early onset and late onset preeclampsia, respectively), we show by transcriptomics that BH4 directly reverts STOX1-deregulated gene expression including glutathione metabolism, oxidative phosphorylation, cholesterol metabolism, inflammation, lipoprotein metabolism and platelet activation, successfully treating placental hypoxygenation, gestational hypertension, proteinuria and heart hypertrophy. In the RUPP rats we show that the major fetal issue of preeclampsia, Intra Uterine Growth Restriction (IUGR), is efficiently corrected. Our work posits on solid bases BH4 as a novel potential therapy for preeclampsia.

1. Introduction
Preeclampsia (PE) is a major hypertensive complication of human pregnancy characterized by a generalized endothelial dysfunction, which is likely triggered by defective endovascular trophoblast invasion of maternal spiral arteries. This condition leads to gestational hypertension associated with proteinuria from mid-gestation, which are classical hallmarks of PE [1]. This syndrome affects around 5% of pregnant women worldwide, and is a leading cause of maternal mortality. Clinically it is classical divided into two presentations, early and late onset (EOPE and LOPE), according to the moment when the symptoms appear, before or after 34 weeks of amenorrhea. Moreover,
PE is associated with a predisposition to strokes, chronic arterial hypertension, heart and renal dysfunction, and increased risk of vascular dementia later in life [2,3].

Placental hypoxia, oxidative stress due to reactive oxygen species (ROS) and related lipid peroxidation are well-described features of preeclampsia [4–6]. Although metabolic profiles of placentas from early-onset PE show downregulation of the glutathione metabolism (i.e. antioxidant defense), to date it has not been stated whether oxidative stress is a cause or a consequence of the disease [7]. Since three decades, the reactive nitrogen species (RNS) nitric oxide (NO) is reported as an endothelium protector able to alleviate preeclampsia [8]. Nitric oxide is a physiological signaling molecule with multiple effects including on DNA synthesis, inflammation, cardiac function, and hypertension [9].

Uregulation of a specific NO synthase (NOS), i.e. inducible NOS (iNOS), linked to hypertension was observed in placentas of PE patients, leading to the suggestion of NOS inhibitors for clinical treatment of PE [10,11]. Therefore, BH4 has been proposed for the treatment of pulmonary hypertension of the newborn, as well as several neurodegenerative diseases [13–16]. Also, the addition of BH4 to human placental homogenates including from preeclamptic pregnancies stimulates NOS activity, supporting the use of BH4 in PE [17].

Hypertension is also caused by the decrease in NO and i-malate, which results from the insufficiency of fumarase (or fumarate hydratase, FH), a key enzyme in the tricarboxylic acid cycle (TCA), or Krebs cycle, that catalyzes the reversible hydration of fumarate to i-malate and is further converted to i-arginine, the substrate of NO [18]. In a rat model of salt-sensitive hypertension due to fumarase insufficiency, i-malate supplementation increased i-arginine and NO levels, and attenuated hypertension [18]. Interestingly, although i-arginine has been shown effective for preventing PE in high-risk pregnancy, the fumarase-related i-malate pathway has not been investigated in PE [19].

The winged-helix transcription factor Storkhead box1 (STOX1) is involved in the genetic forms of preeclampsia [20]. This gene encodes two major isoforms, STOX1A and STOX1B [20]. Although both isoforms can bind the STOX1 DNA binding sites STRE1 and STRE2 (for STOX-responsive element 1 and 2) that we have recently identified, only STOX1A contains a transactivating domain, and specifically deregulates gene expression linked to the nitroso-redox (RNS-redox/ROS) balance [21,22]. STOX1A overexpression confers PE-like gene-expression to trophoblastic cell lines [23] and PE symptoms to pregnant mice carrying transgenic embryos [24] by deregulating more than 2000 endothelial genes [25]. STOX1A overexpression rat induces nitroso-redox imbalance and mitochondrial hyper-activation both in vitro and in vivo [26]. STOX1A overexpressing mice recapitulate long-term cardiovascular consequences such as cardiac stress with heart hypertrophy and fibrosis, consequences of human PE [27]. Overexpression of STOX1A and STOX1B has opposite effects on trophoblast function, in particular trophoblast fusion and syncytialization (acceleration and inhibition, respectively). Therefore, STOX1A/STOX1B imbalance affects gene expression possibly leading to placental abnormalities and related preeclampsia [22]. Finally, STOX1A is abundantly expressed in the brain and its expression is correlated with the severity of late-onset Alzheimer’s disease (LOAD), suggesting that STOX1A controls a conserved pathway shared between placenta and brain [28].

Here, we questioned whether STOX1 overexpression affects the fumarase-related i-malate pathway, and by this way induces iNOS, which in turn alters NO levels, with subsequent nitroso-redox imbalance, metabolic dysregulation, and finally PE-symptoms. Our data reveal that STOX1A and STOX1B overexpression oppositely affect i-malate level and fumarase activity, and also alter iNOS and NO levels, the nitroso-redox balance, the antioxidant defense and bioenergetic metabolism. Treatment with either the NOS coupling agent BH4 or the NOS inhibitor L-NAME showed increase of i-malate with the former and decrease with the latter in PE trophoblasts, possibly explining the pro-preeclamptic effect of L-NAME. In vivo, we used two models of preeclampsia, the STOX1 mice [24] and the RUPP rat model, mimicking early and late onset preeclampsia, respectively. Transcriptomic analysis of mice placentas in vivo showed that BH4 rescues a high proportion of STOX1-deregulated genes including those involved in glutathione metabolism, oxidative phosphorylation, cholesterol metabolism, inflammation, lipoprotein metabolism and platelet activation. Finally, with BH4 we successfully treated gestational hypertension, proteinuria and heart hypertrophy in PE pregnant mice, demonstrating that BH4 is a potential therapeutic drug for preeclampsia. In the RUPP rat, the treatment corrected efficiently the foeto-placental phenotype.

2. Results

2.1. STOX1 regulates the hypoxia marker HIF1A and the fumarase-related i-malate pathway

We investigated STOX1 overexpression in JEG-3 choriocarcinoma trophoblasts that have been stably transfected with: i) the empty pCMX vector containing the CMV promoter (BD3 cell line), or ii) the pCMX-STOX1A vector that overexpresses STOX1A (AA6 cell line), or iii) the pCMX-STOX1B vector that overexpress STOX1B (B10 cell line) (Fig. 1A). Following a defective invasion of the trophoblast cells in the maternal spiral arteries, the maternal vessels keep their contractile arterial vasoreactive functions; it is currently admitted that this defect leads to rapid local variations in oxygen pressure (hypoxia or alternance of hypoxia-hyperoxia) at the materno-fetal interface and eventually to placental dysfunction and to preeclampsia. We simulated hypoxia in these cellular models through stabilization of the hypoxia-inducible factor 1α (HIF1α) via CoCl_{2}, a well-known hypoxia mimetic agent [29] (Fig. 1A). RT-qPCR confirmed STOX1A overexpression in AA6 cells (Fig. S1A), and STOX1B overexpression in B10 cells (Fig. S1B). STOX1A overexpressions did not affect STOX1B levels and vice-versa, and CoCl_{2} treatment further increased STOX1A and STOX1B expression (in AA6 and B10 cells, respectively) (Figs. S1A–B). STOX1A/B overexpression (in the presence and in the absence of CoCl_{2}) was confirmed at the protein level by immunofluorescence using a polyclonal antibody that recognizes both STOX1 isoforms (Figs. S1C–D). CoCl_{2} treatment did not further increase the levels of STOX1A protein, possibly indicating an altered turnover of STOX1A under these conditions.

STOX1A overexpression (AA6 cells) reduced, whereas STOX1B overexpression (B10 cells) increased, the levels of the stabilized (processed) HIF1A form [30,31] compared to control BD3 cells, as shown by Western blots (Fig. 1B). These data suggest that STOX1B overexpression mimics a hypoxic condition, whereas STOX1A overexpression does not promote the expression and activation of HIF1A. CoCl_{2} treatment increased both HIF1A forms in control BD3, confirming its hypoxia mimetic function, and to a lesser extent in AA6 cells, whereas it failed to do so in B10 cells (Fig. 1B). Altogether, the results indicate that STOX1 overexpression reduces the expression and stabilization of HIF1A and thereby the HIF1A-driven response in hypoxia.

Under hypoxia, the expression, processing, and stabilization of
HIF1A are induced through canonical oxygen-dependent regulation and fumarate production are possibly connected since L-HIF1A are induced through canonical oxygen-dependent regulation. In our experimental paradigm, CoCl₂ treatment decreased the levels of fumarate (Fig. 1C–D) in STOX1A (AA6) cells compared to all other conditions, and STOX1A (AA6) cells displayed reduced levels of i-malate (that is produced from fumarate) even in the absence of CoCl₂ treatment (Fig. 1E). i-malate is a crucial metabolite involved in NOS function and hypertension. To note, the levels of i-malate were slightly higher than control in STOX1B (B10) cells, and remained higher also upon CoCl₂ treatment. Thus, under chemical hypoxia not only STOX1 overexpression poorly activated HIF1A, but STOX1A overexpression also reduced the levels of fumarate and i-malate, whereas these factors were moderately affected by STOX1B overexpression.

HIF1A stabilization is modulated by the activity of fumarase. The fumarase specific activity (Fig. 1F), consisting of total fumarase activity (measured with a colorimetric assay) (Fig. S1H) normalized to the amount of enzyme (assessed by immunofluorescence) (Fig. S1 E,F), was similar in STOX1A (AA6) and control cells (BD3), while it was lower in STOX1B (B10) cells. To note, the levels of fumarase assayed by IF were confirmed by a second test, fumarase quantitative assay based on sandwich ELISA assay (Fig. S1G). CoCl₂ treatment increased the fumarase specific activity by about two-fold in control BD3 and STOX1B (B10) cells, thereby restoring control levels in B10 cells, but reduced this activity in STOX1A (AA6) cells (Fig. S1H). Thus, under chemical hypoxia low levels of i-malate in STOX1A overexpressing cells were correlated with low levels of both the substrate (fumarate) and the enzyme responsible for its conversion (fumarase). Upon the same principle, higher levels of i-malate in STOX1B overexpressing cells and controls under chemical hypoxia were correlated with high levels of fumarate and fumarase activity.

### 2.2. STOX1 regulates iNOS and NO levels, the nitroso-redox balance and the antioxidant defense

The regulator of the hypoxia response HIF1A, the levels of fumarate, i-malate, and the fumarase activity are affected by ROS (insufficient fumarase activity increases H₂O₂ levels [34–37]) as well as by the NO metabolism, including iNOS [10,11,18,37,38]. iNOS (NOS2) levels measured by immunofluorescence (IF) were higher in STOX1A (AA6) and STOX1B (B10) compared to control (BD3) cells (Figs. S2A and 2A), and were reduced in all cells upon CoCl₂ treatment. Accordingly, NO levels detected by the diaminofluorescein-2 diacetate (DAF-2DA) probe [39], were higher in STOX1A (AA6) and STOX1B (B10) compared to BD3 cells (Fig. 2B). The use of this technique, while possibly influenced by dehydroascorbic acid and ascorbic acid, may suffer some interferences [40], nevertheless this approach is the reference for innumerable studies for more than 20 years [41–44]. Similarly, upon CoCl₂ treatment, NO was reduced in all cells except B10, suggesting that under hypoxia STOX1B overexpression is able to maintain high levels of NO despite reduced amount of iNOS (Fig. 2B).

Generation of the RNS NO, the ROS O₂ (superoxide) and subsequent formation of the RNS ONOO⁻ (peroxynitrite) are hallmarks of NOS uncoupling [9,13]. Accumulation of iNOS in STOX1A (AA6) and STOX1B (B10) cells was not associated with over-generation of O₂, detected with the dihydroethidum (DHE) probe [41], and ONOO⁻ detected with the dihydrorhodamine 123 (DHR123) probe [42], compared to control (BD3) cells (Fig. 2C–D). No changes were observed upon CoCl₂ treatment, except reduced ONOO⁻ in controls, and to some extent in STOX1B (B10), compared to the corresponding untreated cells.
intensity accumulation but no superoxide and peroxynitrite accumulation, suggesting altered antioxidant defence in STOX1 cells or poor NOS showing altering the nitroso-redox balance [26], induces iNOS and NO (Fig. 2D). Altogether, STOX1 overexpression, which we previously

Fig. 2. NO metabolism and antioxidant defense upon STOX1 overexpression and hypoxia. (A) Quantification of inducible NOS (iNOS; NOS2) fluorescence intensity per cell (images in Fig. S2A). Assessment of (B) Nitric oxide (NO) using the DAF-2 DA probe, (C) the ROS anion superoxide (O$_{2}^{-}$) using the DHE probe, and (D) the ROS peroxynitrite (ONOO$^{-}$) using the DHR123 probe, expressed as percentage of the untreated control BD3 cell line. (E) Total SOD activity (measured as inhibition of the activity of SOD, percentage). (F) Catalase activity expressed in nM/min/ml. Immunofluorescence, n = 60 cells from three independent experiments, mean ± SD. Dosages (NO, O$_{2}^{-}$, peroxynitrite), n = 3–6. Enzyme activities (SOD, catalase), n = 6 for untreated BD3/A6/B10 in NO and O$_{2}^{-}$ assays, n = 3 independent experiments for all other experiments, mean ± SD, *p ≤ 0.05 **p < 0.01 ***p < 0.001 based on one-way ANOVA versus the untreated control (red stars) or the corresponding untreated condition (blue stars). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

(Fig. 2D). Altogether, STOX1 overexpression, which we previously showed altering the nitroso-redox balance [26], induces iNOS and NO accumulation but no superoxide and peroxynitrite accumulation, suggesting altered antioxidant defence in STOX1 cells or poor NOS uncoupling.

Peroxynitrite and NO accumulation have been associated with low superoxide dismutase (SOD) activity [45]. Therefore, we tested SOD that metabolizes O$_{2}^{-}$ into the ROS hydrogen peroxide (H$_{2}$O$_{2}$) [46–48] and is the first line of defence against oxidative stress. The levels of mitochondrial SOD2 (MnSOD), tested by IF, were not affected by STOX1 overexpression nor by CoCl$_{2}$ treatment (Figs. S2B–C), and total SOD activity was not altered beyond a 5% range (Fig. 2E). However, upon CoCl$_{2}$ treatment SOD activity decreased in STOX1B cells (Fig. 2E). The product of SOD activity, the ROS H$_{2}$O$_{2}$, detected with the 2',7'-dichlorodihydrofluorescein diacetate (DCF-DA) probe [42], was essentially not affected, and increased upon STOX1B overexpression in normoxic conditions, in parallel with a mild increase of SOD activity (Fig. S2D). Hence, we reasoned that catalase activity may have been altered instead. Catalase is the antioxidant enzyme that metabolizes H$_{2}$O$_{2}$ into H$_{2}$O and O$_{2}$ [48,49] and also catalytically scavenges peroxynitrite. Catalase can be inactivated by NO accumulation and by single

2.3. STOX1 regulates the bioenergetic metabolism

We then assessed whether the observed alterations were associated with mitochondrial stress in STOX1 overexpressing cells. Indeed, insufficient fumarase activity increases H$_{2}$O$_{2}$ levels by altering the expression of mitochondrial OXPHOS complexes [34–37]. Decreased fumarase and increased H$_{2}$O$_{2}$ levels were indeed observed upon STOX1B overexpression in normoxia (see above, Fig. S1H and Fig. S2D). Moreover, HIF1A stabilization is associated with severe damage of the mitochondrial oxidative phosphorylation (OXPHOS) complexes [32]. The amount of the mitochondrial ROS superoxide anion (mO$_{2}^{-}$) [52], detected by MitoSOX Red staining, was slightly higher in STOX1B (B10) than STOX1A (AA6) and control (BD3) cells, and returned back to control levels under CoCl$_{2}$ treatment (Fig. 3A and B). Intact mitochondria are classically detected by the mitochondrial marker citrate synthase (CS) that catalyzes the condensation of oxaloacetate with the glycolytic end-product acetyl-coenzyme A (acetyl-CoA) to produce citrate and CoA in the Krebs cycle [53–56] (Fig. 3C). Both STOX1A (AA6) and STOX1B (B10) reduced CS activity (~37% and ~54%, respectively) compared to control (BD3) cells, suggesting a reduction of active mitochondria (Fig. 3D). Conversely, CoCl$_{2}$ treatment strongly activated CS activity in STOX1 overexpressing cells beyond the levels of controls, suggesting an increase of active mitochondria in these cells (Fig. 3D). However, the levels of the CS substrate acetyl-CoA were lower in STOX1B (B10), and upon CoCl$_{2}$ treatment also in STOX1A (AA6), compared to control (BD3) cells (Fig. 3E), indicating that CS activity in STOX1 overexpressing cells is possibly limited by the amount of its reactants, and therefore may not correspond to reduced mitochondrial activity.

To meet bioenergetic demands, ATP is produced by glycolysis as well
Fig. 3. Mitochondrial ROS and mitochondrial bioenergetic upon STOX1 overexpression and hypoxia. (A) 3D reconstructions of cells stained with MitoSOX (red) to detect mitochondrial ROS (superoxide anion, O$_2^-$), counterstained with Hoechst (blue). Scale bar 10 μm, and (B) Quantification of MitoSOX fluorescence intensity per cell. (C) Schematic representation of Krebs cycle including the reaction catalyzed by citrate synthase from oxaloacetate (substrate) to citrate (product). (D) Citrate synthase activity expressed in mU/mg of protein. (E) Acetyl-CoA expressed in nmol/g protein. (F) Total ATP expressed as percentage of the untreated control BD3 cell line with non-OXPHOS and OXPHOS fractions. Immunofluorescence, n = 60 cells from three independent experiments, mean ± SD. Dosage (ATP, Acetyl-CoA), enzyme activity (citrate synthase), n = 3 independent experiments, mean ± SD, *p ≤ 0.05 **p ≤ 0.01 ***p ≤ 0.001; based on one-way ANOVA versus the untreated control (red stars) or the corresponding untreated condition (blue stars). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
as mitochondrial OXPHOS \[55,56\]. Total ATP levels increased by 2.5-fold in STOX1A (AA6) and 5.5-fold in STOX1B (B10) compared to control (BD3) cells (Fig. 3F), indicating that either their mitochondria generate more ATP or, alternatively, that this ATP is largely produced by glycolysis. Compatible with the second hypothesis, the fraction of ATP in the presence of the OXPHOS inhibitor oligomycin (i.e. the OXPHOS fraction) remained low in all samples (less than 15%) and ATP changes were mostly not dependent on OXPHOS (Fig. 3F). Glycolytic ATP appeared also responsible for the inverted situation upon CoCl₂ treatment, with control and STOX1A (AA6) overexpressing cells displaying high, and STOX1B (B10) overexpressing cells low, ATP levels (Fig. 3F). Nevertheless, CoCl₂ treatment strongly enhanced the OXPHOS fraction in STOX1A (AA6) cells. Thus, in hypoxia mimetic condition STOX1A overexpression restored, at least in part, mitochondrial respiration and activated CS activity without affecting the ROS O₂⁻.

### 2.4. The BH4 metabolite controls STOX1 effects through iNOS regulation

The nitric oxide signalling pathway modulates mitochondrial respiration in particular in hypoxia. Moreover, iNOS expression is involved in metabolic syndromes and chronic metabolic inflammation \[57,58\]. We reasoned that increasing the iNOS coupling via the BH4 cofactor \[14,17\] or, alternatively, inhibiting iNOS with L-NAME \[10,11,59\] are potential therapeutic strategies to counteract dysfunction due to STOX1 overexpression in preclampsia. Either BH4 or i-NOS treatment maintained STOX1 isoforms accumulation at the mRNA level (Figs. S3A–B), albeit slightly reduced the STOX1 protein, compared to untreated cells (Figs. S3C–D). Either treatment reduced iNOS in STOX1A and STOX1B to the levels of untreated cells (Fig. 4A; S3E), and dramatically reduced the ROS H₂O₂ in all conditions (Fig. S3F). Either treatment also reduced the high mitochondrial ROS in STOX1B (B10) cells (Figs. 4B and S3G).

The RNS peroxynitrite was reduced in STOX1A and control cells by

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Fig. 4. The BH4 metabolite counteracts STOX1 effects through iNOS regulation and rescue of L-malate levels. (A) Quantification of iNOS/NOS2 fluorescence intensity per cell. (B) Quantification of mitochondrial ROS fluorescence intensity per cell. (C) Percentage of peroxynitrite RNS relative to the untreated control BD3 cell line. (D) Fumarase specific activity (ratio of fumarase activity/fumarase protein content) expressed as percent of untreated control BD3 cell line. Levels of L-malate (E) and fumarate (F) expressed in nmol/mg of protein. Data of untreated cells are from the corresponding panels in Fig. 2A (iNOS), Fig. 3B (mitochondrial ROS), Fig. 2D (RNS), Fig. 1F (fumarase specific activity), Fig. 1D (fumarate), Fig. 1E (L-malate). Immunofluorescence, n = 60 cells from three independent experiments, mean ± SD. Dosages (peroxynitrite, fumarate, malate) and enzyme activity (fumarase), n = 3 independent experiments, mean ± SD. *p ≤ 0.05 **p ≤ 0.01 ***p ≤ 0.001 based on one-way ANOVA versus the untreated control (red stars) or the corresponding untreated condition (pink or purple stars). (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
either treatment, but not in STOX1 cells with \( L \)-NAME (Fig. 4C). Thus, the two treatments appear equivalent in rescuing key ROS/RNS alterations due to STOX1 overexpression, with the exception of \( L \)-NAME that failed to reduce RNS in STOX1 (B10) cells.

These two treatments have then been assessed for regulation of the antioxidant response. The total antioxidant SOD activity and the mitochondrial component of this activity (SOD2) were not remarkably different in STOX1 overexpressing compared to control cells, and either BH4 or \( L \)-NAME had little effect on these parameters, except slightly reduced SOD activity and SOD2 levels in STOX1 (B10) cells upon BH4 treatment (Figs. S4A-C). Conversely, the activity of the other major antioxidant, catalase, which was elevated in STOX1 cells, was largely reverted by either treatment, and in particular BH4, despite roughly comparable levels of the protein in all cell lines (Figs. S4D-F).

We then assessed whether these treatments affected the bioenergetic metabolism. Either treatment increased CS activity in all cells, essentially rescuing low CS levels in STOX1 cells, except BH4 in STOX1A cells (Fig. S5A). In addition, both treatments strongly decreased the central bioenergetic metabolite acetyl-CoA levels in all cell types, and more severely upon STOX1A overexpression (Fig. S5B). Finally, BH4 slightly increased and \( L \)-NAME dramatically reduced ATP levels in STOX1A overexpressing cells, whereas \( L \)-NAME strongly increased ATP levels in STOX1B overexpressing cells, this being essentially due to the non OXPHOS-related ATP production (Figs. S5C-D). Despite both treatments increased the fraction of mitochondrial ATP when STOX1A and STOX1B are overexpressed, in particular STOX1A cells treated with BH4 and to a larger extent STOX1B cells treated with \( L \)-NAME (Fig. S5C), in absolute values the contribution of mitochondrial ATP remained quite modest as most ATP was not produced through OXPHOS. These observations suggest that in all cells, acetyl-CoA does not systematically feed the Krebs cycle to produce ATP, despite BH4 or \( L \)-NAME treatments tend to some extent to tilt the balance toward the consumption of acetyl-CoA and mitochondrial ATP production.

Finally, we assessed whether the fumarate and \( L \)-malate pathways are affected by iNOS upon treatment with one or the other molecule. The level of the enzyme fumarase, measured by immunofluorescence (Figs. S6A-B) and sandwich ELISA assay (Fig. S6C), was comparable in untreated cells, and upon BH4 as well as \( L \)-NAME treatment it increased in STOX1B cells whereas was essentially not affected in control and STOX1A cells. Despite minor variations in the amount of enzyme, both treatments strongly reduced the fumarase activity in control and STOX1B cells, contrary to STOX1A cells where the opposite effect was observed, and more strikingly upon BH4 treatment (Fig. 4D, Fig. S6D). Thus, BH4 and \( L \)-NAME resulted in high levels but poor activity of fumarase in STOX1B cells, raising the possibility that in these cells the enzyme was produced to compensate for lack of activity. Conversely, either treatments strongly increased the fumarase activity in STOX1A cells (despite unchanged amount of the enzyme). The high fumarase activity correlated with increased levels of its product \( L \)-malate in STOX1A cells treated with BH4. Conversely, \( L \)-NAME treatment decreased the levels of \( L \)-malate in these cells. Despite minor differences, control and STOX1 expressing cells, with or without BH4 and \( L \)-NAME displayed comparable levels of \( L \)-malate (Fig. 4E), indicating that variations in the level of this substrate are not responsible for the amounts of the downstream product (\( L \)-malate). In summary, BH4 and \( L \)-NAME have globally comparable effect on control and STOX1 overexpressing cells, showing that iNOS modulation affects the fumarate/\( L \)-malate pathway in normal and NO overexpressing cells, but only upon \( L \)-NAME treatment (with BH4) rescued the levels of \( L \)-malate in STOX1A overexpressing cells, whereas \( L \)-NAME not only failed to do so but actually reduced the levels of fumarate in these cells (Fig. 4F).

### 2.5. BH4 restores STOX1-deregulated placental gene expression in vivo

While the mode of action of BH4 in vitro as a NOS coupling agent is well described [14,17], the actual impact of this metabolite in the placenta is not known. We questioned whether in vivo BH4 treatment, provided in drinking water, can restore STOX1-deregulated gene expression in placenta of wild-type (WT) female mice crossed with STOX1A transgenic male mice (heterozygous \( Tg \)-STOX42), a genetic design leading to a preeclamptic phenotype in the mothers (Fig. 5A) [21, 24, 27]. Mice placentas at 16.5 days post-coitus (late gestation) were analyzed at the whole transcriptome level. Total RNAs from pools of three placenta from different mice in four conditions were used (control (Ctrl), Ctrl + BH4, STOX1 (preeclamptic-mimic) mice, and STOX1+BH4) and hybridized to Clarionm5 microarrays.

Out of 22,206 transcripts analyzed, 2476 were modified (ANOVA F test <0.05). The PCA clustering was able to successfully separate the four conditions (Fig. 8B). Webgestalt ontology analysis was used to characterize the BH4 effects by comparison of the STOX1 and STOX1+BH4 conditions, as shown by comparison of KEGG pathways [60]. Among the highest scored pathways in the STOX1 versus STOX1+BH4 condition, genes in “Oxidative Phosphorylation” and “Glutathione metabolism” were up-regulated, and genes in “Platelet activation” and “Cholesterol metabolism” were down-regulated (Fig. S5C). The GSEA analyses of these pathways are represented in Fig. 5D, showing normalized enrichment scores above 2.

To analyze in more detail the effects of BH4 and STOX1 on various genes, we identified a high proportion of genes that were deregulated by STOX1 and brought back to basal level by BH4 treatment. Overall, 565 genes were modified more than 2-fold between CTLs and STOX1 placentas, and amongst those, 84 (~15%) were brought back to ~30% of their gene expression levels in the control placentas (Fig. 5E). These 84 genes were submitted to network analysis using STRING, and were successfully clustered, revealing several clusters (Interferon-regulated genes, Inflammation regulators, Lipoprotein metabolism) (Fig. S7). In conclusion, these in vivo transcriptomic results prove that BH4 acts by compensating alterations induced by STOX1 at the gene level. Some of these genes are highly relevant in the context of NO physiology, such as Nostin (NO Synthase Trafickier), a factor that colocalizes with eNOS, and redistributes subcellularly the synthase [61].

In preeclampsia, endothelial cells play a major role since the disease is characterized by a global endothelial dysfunction. Therefore, analyzing the expression data under the angle of endothelial cells is particularly relevant. We submitted our bulk microarray data to the deconvolution algorithm dTangle to characterize the cell types, their proportion and their gene expression profile [62]. To deconvolute these data we needed dTangle with an available single cell experiment conducted on mice placenta [59]. The algorithm used this reference to identify specific cell markers and the proportion of the different cell types (trophoblasts, endothelial cells, decidual stromal cells, mesenchymal fetal cells and blood cells) (Fig. 6A). Using the microarray data, we used deconvolution algorithms from the signature of the major placental gene categories known in mice [63]. Using the 10 prominent genes of five categories (trophoblast, endothelial cells, decidual stromal cells, mesenchymal fetal cells and blood cells) on our data, we detected easily the five major gene categories without significant changes in our dataset from the Marsh and Blelloch paper used as a reference [63].

Using these genes we could evaluate the proportion of the different cell types, where endothelial cells represented about 14% of the cells, trophoblasts 45%, and among these, 84 (~15%) were brought back to ~30% of their gene expression levels in the control placentas (Fig. 5E). These 84 genes were modified more than 2-fold between CTLs and STOX1 placentas, and amongst those, 84 (~15%) were brought back to ~30% of their gene expression levels in the control placentas (Fig. 5E). These 84 genes were submitted to network analysis using STRING, and were successfully clustered, revealing several clusters (Interferon-regulated genes, Inflammation regulators, Lipoprotein metabolism) (Fig. S7). In conclusion, these in vivo transcriptomic results prove that BH4 acts by compensating alterations induced by STOX1 at the gene level. Some of these genes are highly relevant in the context of NO physiology, such as Nostin (NO Synthase Trafickier), a factor that colocalizes with eNOS, and redistributes subcellularly the synthase [61].
Fig. 5. In vivo STOX1 model of preeclampsia and BH4 effect on placenta transcriptomics. (A) In vivo model of preeclampsia and preclinical trial using BH4 supplementation in healthy and preeclamptic pregnant mice. (B) Principal Component Analysis of gene expression in control and STOX1-overexpressing placentas at 16.5 days post-coitus, with or without BH4 treatment. (C) Enrichment of gene expression of KEGG pathways in STOX1 placentas with or without BH4 treatment. (D) Enrichment of specific pathways related to placental function in this comparison. (E) BH4 rescues the alteration of gene expression induced by STOX1 overexpression in the placenta (threshold chosen: genes with more than two fold changes, and corrected to <30% of the value in control samples).
Fig. 6. Cell proportions in the placenta are altered by BH4 treatment, and major endothelial genes are deregulated. (A) Using deconvolution, we were able to dissociate the different major cell types present in the mouse placenta. The proportion of cells was altered by BH4 for trophoblasts and for decidual stromal cells, while the other cell types did not differ quantitatively. (B) In the endothelial cells, several genes were deregulated. The column ‘Specificity’ refers to the endothelial specificity, with *** for genes expressed exclusively in the endothelium, while ** and * refers to genes that are expressed in other tissues, albeit predominantly in the endothelial cells.
2.6. BH4 successfully prevents major preeclampsia symptoms in mice in vivo

We then validated whether BH4 treatment in drinking water is beneficial against gestational hypertension, proteinuria, and heart hypertrophy, three major preeclampsia symptoms that are recapitulated in the STOX1 mice model [1,21,24,27]. For the first PE symptom, daily monitoring of the blood pressure (BP) by assessment of systolic arterial pressure throughout gestation [24] showed that preeclamptic pregnant mice developed a gestational hypertension from 5.5 days post-coitus (early-gestation) compared to WT pregnant mice (Fig. 7A). BH4 treatment successfully prevented gestational hypertension in preeclamptic pregnant mice and did not change BP in WT pregnant mice compared to the corresponding untreated pregnant mice (Fig. 7A). Analysis of BP by three periods corresponding to early-, mid- and late-gestation showed that BH4 treatment successfully prevented gestational hypertension in preeclamptic pregnant mice throughout these gestational periods (Fig. S9). In WT pregnant mice BH4 treatment slightly decreased BP during first period (early gestation) whereas slightly increased BP in the second (mid-gestation) period (Fig. S9).

For the second PE symptom, preeclamptic pregnant mice displayed proteinuria evaluated as the ratio between albumin and creatinine (ACR ratio) in the urine [24] compared to WT pregnant mice. BH4 treatment successfully prevented proteinuria in preeclamptic pregnant mice without affecting this ratio in WT pregnant mice (Fig. 7B).

The third PE symptom, the cardiac hypertrophy evaluated as the ratio of heart weight over tibia length [27,64] strongly increased in preeclamptic compared to WT pregnant mice (Fig. 7C). BH4 treatment successfully prevented cardiac hypertrophy in preeclamptic mice, and slightly increased the ratio in treated WT pregnant mice (Fig. 7C). Several echographic parameters were measured at 16.5 days of gestation on mice carrying transgenic embryos with or without BH4 treatment (Fig. 7D). Consistently with the actual measures after sacrifice, the heart weight estimated by in vivo echography was reduced to normal level by BH4, and the interventricular septum thickness (a predictor of human coronary disease [65]) was reduced. Several Doppler parameters were corrected by the treatment (increased myocardial performance, increased outflow from the right ventricle and decreased blood velocity for the two sides of the renal arteries).

Furthermore, at term, the placentas were weighed and revealed a decreased placental weight in mice carrying transgenic embryos and this was corrected by BH4 treatment at 18 days post coitus (dpc) (Fig. 7E). There was no reduction in fetal size in this experiment but this is consistent with previous observations showing that in the STOX1 model there is an efficient in utero catch-up growth of the fetus at the end of gestation. The intrauterine growth restriction (IUGR) phenotype detectable clearly at 16.5 dpc, diminishes strongly after 17.5 days [66]. Altogether, this pre-clinical study validated that BH4 treatment throughout gestation successfully prevented the major preeclampsia symptoms.

2.7. BH4 successfully prevents IUGR in the surgical RUPP model of late preeclampsia in rats in vivo

The Reduced Uterine Placental Perfusion (RUPP) model was used to validate the BH4 action on PE symptoms using an independent animal model (Fig. 8A). This surgical procedure is performed at 14 dpc characterizing a clinical phenotype more consistent to late onset preeclampsia (LOPE) rather than to early onset preeclampsia (EOPE) like the STOX1 transgenic mouse. Maternal blood pressure was increased in RUPP animals at 19 days post-coitum. The LOPE status of the RUPP animals was also illustrated by the fact that the heart weight was not significantly modified, despite reported impaired systolic function in this model by Doppler ultrasound echocardiography [67]. The BH4 did not correct the increased blood pressure in this LOPE model. In the RUPP dams, there was an expected decrease in fetal weight (from 2.34 ± 0.29 g to 2.22 ± 0.25 g). BH4 treatment at 3.5 mg/kg largely compensates this (average weight 2.54 ± 0.3 g), while the high dose (7 mg/kg) appeared deleterious, Fig. 8C. These observations were consistent with placental weight that was partly corrected with BH4 (Fig. 8D), but no differences were found for placental efficiency (Fig. 8E). Also, consistent with these observations was the effect of acetylsalizylic acid to assess endothelium-dependent vasorelaxation in isolated uterine artery rings that was brought back to normal in the BH4-treated females (Fig. 8F).

There was no effect detectable in the aorta using the same ex-vivo procedure suggesting that the BH4 treatment modulates hemodynamics, at least partially, and specifically in the utero-placental circulation.

3. Discussion

There is now a wealth of literature stressing the importance of nitrosative stress in preeclampsia [68,69]. At the center of nitrosative stress are the NOS enzymes, that need coupling with the tetrahydrobipterin (BH4) cofactor to produce NO, an essential element of vascular health. Therefore insufficient BH4 concentrations were envisaged long ago as causative in PE [70], as mentioned in a recent review [71]. Consistently, specific mutations of eNOS were associated with an increased risk of PE [72]. BH4 production was shown to be negligible in the mature placenta, suggesting that it should originate from other sources, such as the fetus, and will possibly being limited in pathological pregnancies [73]. A first very elegant preclinical study using BH4 as a treatment in PE was published in 2007 [74]. In this rat model, PE-like symptoms were induced by deoxycortisone acetate (DOCA) injection, and BH4 addition (sepiapterin) was shown to induce relaxation in mesenteric arteries in organ chambers, ex vivo. This descriptive paper nevertheless did not address the fine molecular modification triggered by the treatment, especially in vivo. In the present study, we aimed at a better understanding of BH4 function by cell biology experiments in vitro, and by transcriptome analysis of the placenta in the STOX1 mouse model of EOPE [24], treated or not with BH4. To determine whether any benefits with BH4 are specific for EOPE, we utilized a classical model of LOPE by applying BH4 therapy to the RUPP rat model.

The preeclampsia-associated transcription factor STOX1 is a key regulator of the oxygen-dependent nitroso-redox balance and mitochondrial metabolism [26]. Placental hypoxia associated with oxidative/nitrosative stress is a major feature of preeclampsia [4–6], and STOX1A/STOX1B imbalance has been shown to deregulate nitroso-redox balance and trophoblast function at gene expression level [4–6,22]. Here, we show that the preeclamptic condition generated by STOX1A overexpression affects multiple linked gene pathways in trophoblasts; in particular it deregulates the HIF1α-driven response in hypoxia, which affects the fumarase-related α-malate pathway, induces iNOS thereby altering NO levels with subsequent nitroso-redox imbalance, metabolic dysregulation, and finally PE-symptoms (Fig. 9).

Importantly, we show that under chemical hypoxia only STOX1A overexpression reduces the fumarate/fumarase/α-malate pathway, whereas STOX1B overexpression generates the opposite effect, possibly explaining the specificity of STOX1A overexpression in conferring PE-like gene-expression to trophoblastic cell lines [23] and PE symptoms to pregnant mice. Treatment with the NOS coupling agent BH4 (but not the NOS inhibitor L-NAME) increased α-malate in PE trophoblasts, restoring control conditions. Transcriptomic analysis of mice placentas in vivo confirmed that BH4 rescues a high proportion of STOX1-deregulated genes including those involved in glutathione metabolism, oxidative phosphorylation, cholesterol metabolism, inflammation, lipoprotein metabolism, and platelet activation. Finally, with BH4 treatment we successfully corrected gestational hypertension, proteinuria, placental weight and heart hypertrophy in PE pregnant mice as well as Intra-Uterine Growth Restriction and placental dysfunction in rats, demonstrating that BH4 is a potential therapeutic drug for preeclampsia/IUGR.

Oxyn-independent alteration of HIF1α stabilization upon STOX1...
overexpression indicated that non-canonical regulation of HIF1α through the fumarase pathway was affected, in agreement with the notion that the activity of this key Krebs cycle enzyme and its related oncometabolites fumarate and α-malate can deregulate the HIF1α status [31–33]. In support of this model, gestational hypertension is a major PE symptom, and in a rat model of salt-sensitive hypertension due to fumarase insufficiencies, α-malate supplementation attenuated hypertension through the modulation of NOS and NO metabolism [18]. Associated with alteration of fumarase, and the related fumurate and α-malate metabolites, we indeed showed that both STOX1 isoforms strongly affect the NO metabolism, in particular increasing the iNOS content but decreasing the iNOS coupling as could be also in placentas of PE patients [10,11]. These findings strengthen and help clarify the link between STOX1 overexpression and gestational hypertension in preeclampsia.

The specificity of STOX1A in PE was supported by our findings that STOX1A/STOX1B isoforms oppositely deregulate the stabilization of the hypoxia marker HIF1α, consistently with the opposite effects of STOX1A and STOX1B overexpression shown on trophoblast fusion [22]. In particular, we observed that STOX1A overexpression decreases α-malate and dysregulates the antioxidant defense with dramatically low levels of catalase in hypoxia-mimetic conditions. Our findings are consistent with the hypothesis that a STOX1-dependent vicious circle of iNOS uncoupling, leading to nitroso-redox imbalance and mitochondrial stress, is a major deregulated pathway in PE. In agreement with this picture, in hypoxia iNOS and NO levels were reduced in STOX1A overexpressing cells, whereas STOX1B overexpressing cells were able to maintain high levels of NO despite reduced amount of the NO synthase (iNOS). Related to iNOS alterations, both STOX1 isoforms strongly affected the nitroso-redox balance including peroxynitrite management, and the antioxidant defense remarkably increasing catalase activity. However, switch of catalase activity from over-activation to control levels in hypoxia mimetic condition, and without relevant changes in the enzyme content, was observed only upon STOX1A overexpression, further linking this factor with altered response upon hypoxia.

Both STOX1 isoforms strongly reduced citrate synthase (CS) activity, the first enzymatic and pacemaker step of the mitochondrial Krebs cycle, and CS activity was over-activated in hypoxia mimetic conditions, correlated with the activation of mitochondrial respiration. Despite these changes, the global increase of total ATP levels upon STOX1 overexpression appeared essentially due to anaerobic glycolysis. Again, an opposite effect was observed in hypoxia-mimetic conditions upon overexpression of the different STOX1 isoforms, with STOX1A furtherly enhancing total and mitochondrial ATP, whereas STOX1B increased mitochondrial ATP while reducing total ATP levels. These findings indicate that STOX1A overexpression induces a syndrome of preeclampsia of metabolic origin that is exacerbated in hypoxia.

Our findings place iNOS at a central position in PE-related symptoms and reveal iNOS as a major target for PE treatment. However, iNOS coupling (treatment with BH4) rather than iNOS inhibition (treatment with the non-specific NOS inhibitor L-NAME) appears effective in restoring altered control, non-preeclamptic values. In this context, alteration of the α-malate pathway (fumarate/fumarase/α-malate) and the bioenergetic metabolism upon L-NAME helps explaining why this treatment triggers PE symptoms including hypertension in treated rats and promotes, rather than cures, preeclamptic symptoms [8,75]. Moreover, BH4 (but not L-NAME, at least in STOX1B cells) regulated nitroso-redox stress through reduction of peroxynitrite. Further, BH4 increased ATP availability, whereas L-NAME dramatically reduced ATP levels in STOX1A overexpressing cells. Interestingly, a study suggested that NO produced by endothelial NOS (eNOS) is not relevant for the L-NAME effect, suggesting that L-NAME rather acts through iNOS modulation [76]. In our paradigm, however, iNOS inhibition did not restore key altered parameters. Conversely, modulating iNOS (via iNOS coupling) and mitochondrial stress with BH4 restored this multi-step pathway in vitro, also showing that the levels of α-malate and fumarate activity directly depend on NOS coupling and possibly mitochondrial stress.

Nevertheless, inhibition of NOS by L-NAME and increasing NOS coupling with the BH4 cofactor share common effects in vitro, which include a reduction of STOX1 protein accumulation, reduction of iNOS and the central metabolite acetyl-CoA, and stimulation of mitochondrial OXPHOS.

In our first pre-clinical model, we demonstrated that BH4 directly restores STOX1-deregulated gene expression in placentas including glutathione metabolism, oxidative phosphorylation, cholesterol metabolism, inflammation, lipoprotein metabolism and platelet activation, key pathways involved in preeclampsia [7,77,78]. Our results correlate with the metabolic profiles of women preeclamptic placentas, indicating that glutathione metabolism is one of the key pathways to target in preeclampsia [7]. We show that BH4 is able to target this pathway. To note, we previously performed a transcriptomic analysis to understand the beneficial effects of low-dose aspirin treatment in this mouse model of preeclampsia, and aspirin is now clearly demonstrated to be efficient against preeclampsia [24,79,80]. The aspirin effects were exclusively achieved by alterations of placental gene expression that were disconnected from STOX1 effects, and massively down-regulated genes involved in coagulation, regulated by the HNF1β transcription factor [80]. By contrast, here we show that BH4 directly acts by rescuing STOX1-modulated gene expression. This suggest that BH4 is a more powerful and effective treatment than aspirin, and also that the combination of the two drugs could be in the future the most efficient direction towards a complete protection against preeclampsia symptoms. In the RUPP rats, we showed that the IUGR phenotype is restored by BH4, showing that in this LOPE model, extremely different from the STOX1 mouse model of EOPE, the therapy has beneficial effects as well.

Very recently, Chuaiphichai and coworkers demonstrated that a mouse model deficient for generating BH4, the Gch1 knockout mice presents clearly with the preeclampsia syndrome features, and induces alterations of the uteroplacental vascular development essential for regulating the blood pressure during pregnancy [81]. These concordant findings strengthen our observations, since different causes of preeclampsia symptoms posit BH4 as an essential molecule in gestation.

In the STOX1 mice, gestational hypertension, proteinuria and heart hypertrophy are all prevented by BH4 therapy in our mouse model that also mimics the long-term cardiovascular complications associated with preeclampsia in women [3,27]. Given the long term cardiovascular effects of PE, and its association with a predisposition to at least strokes, chronic arterial hypertension, heart and renal dysfunction, this novel STOX1A/STOX1B-linked multi-step pathway and the associated BH4
therapy could be of interest also for these complications. Interestingly, STOX1A accumulates in the brain of late-onset Alzheimer’s Disease (AD) patients [28] and induces Tau phosphorylation in AD [82]. If the same pathway controlled by STOX1 in preeclampsia is present in Alzheimer’s disease, the rescue of STOX1A-dependent altered parameters by BH4 could be beneficial for such disease.

In addition to utilizing the STOX1 model, we strengthened our results implicating BH4 as a beneficial therapy to reduce maternal symptoms of EOPE by including the rat RUPP model that recapitulates LOPE. In this case, we could not induce a maternal amelioration of the blood pressure, while the foetal and placental weight were restored by the BH4 treatment.

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Fig. 8. *In vivo* evidence of BH4 treatment benefit on IUGR in the RUPP rat model. (A) Structure of the uterine vascularization of the rat uterus. In red are the silver clips that restrict blood flow to 100 μM in the ovarian arteries and to 230 μM in the abdominal aorta. (B, C, D) Mean arterial pressure (MAP), fetal and placental weights analyzed in the RUPP rats. Placental efficiency (D) is calculated by dividing the body weight by the placental weight for every feto-placental unit. (E) Isolated uterine artery function was evaluated using cumulative doses of Acetylcholine (Ach) via wire myography. The area under the curve (AUC from the curve obtained as relaxation against doses of Ach) was considered. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)
4. Conclusions

In summary, STOX1 accumulation in a preeclampsia model impaired iNOS function and the related NO metabolism, which in turn deregulated a multi-step pathway including the nitroso-redox balance with peroxynitrite management, the antioxidant defence, the Krebs cycle including citrate synthase and fumarase activities, the fumarase-related metabolites L-malate and fumarate, and bioenergetic metabolism with acetyl-CoA, mitochondrial OXPHOS, and increased glycolysis to meet the ATP demand, altogether associated with dysregulation of hypoxia HIF1A stabilization. BH4 treatment increases iNOS coupling thereby reverting this STOX1A-dependent multi-step pathway in vitro, rescuing STOX1-deregulated gene expression in the placenta, and treating the major symptoms of preeclampsia in vivo. In an experimental animal model independent of STOX1, the RUPP rat, we showed that the treatment is efficient to correct the IUGR effects of this surgical procedure. Our discoveries suggest that BH4 is a promising therapy to cure adverse maternal-fetal symptoms associated to preeclampsia.

5. Material & methods

5.1. Animals experiments (breeding, treatment and ethics)

The mouse work (female mice) was performed under the local regulations and ethic committees: Animal Care Committee of the Paris Descartes University (agreement no. 02731.02). Placentas were collected at 16.5 days post coïtum and placed in TriZol prior to RNA extraction \[24,66\]. BH4 (dose estimated through normal water consumption at 10 mg/kg/day) or pure water was given in the drinking water for two mice per cage by 15 ml frozen and changed twice a day during all the gestational period following the detection of pregnancy by ultrasonography between days E6.5 or E7.5, this dose having been chosen from previously published studies \[83\]. In more detail, the mice were breed in trios from Friday night to Monday morning with one male mouse (either WT or Tg42 transgenic–heterozygous-) and two WT female mice at 2 months of age, all in the FVB/N genetic background. Overall, they were 9 WT mice without treatment, 7 with BH4, 15 mice carrying transgenic embryos without BH4, and 12 treated with BH4.

To include the RUPP model, studies in rats were conducted in
received from Charles River Laboratories (Wilmington, MA, USA) be accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals with protocols approved by the Institutional Animal Care and Use Committee at the University of Mississippi Medical Center (agreement no. 1511). Timed-pregnant SAS Sprague-Dawley rats were obtained from Charles River Laboratories (Wilmington, MA, USA) between gestational day 10 and 11 and placed on Envigo 8640 diet. On day 14, the RUPP procedure was performed by first placing rats under isoflurane anesthesia (Butler Schein Animal Health, Dublin, OH, USA). An abdominal incision was made and a silver clip (0.203 mm, internal diameter) placed around the sub-renal abdominal aorta above the uterine arteries along with a clip (0.1 mm, internal diameter) on each branch of the ovarian arteries (Fig. 8). The control group consisted of a Sham surgery on gestational day 14, without clip placement. BH4 was dosed in rats from day 13–18 of gestation. Total number of rats was 13 for Sham 13, 15 for RUPP, 6 for RUPP + BH4 3.5 mg/kg/day and 3 for RUPP + BH4 7 mg/kg/day. BH4 was reconstituted in dionized H₂O and frozen as aliquots to administer 2X daily (mornings and evenings) at a final dose of 3.5 or 7 mg/kg body weight given orally by mixing into 0.5 mg peanut butter as vehicle. On gestational day 18, rats were implanted with carotid catheters under isoflurane anesthesia for assessment of conscious blood pressure on day 19 followed by anesthetizing and collecting tissues to weigh fetuses and placentas for calculating pregnancy biometrics. Also, aortas and uterine arteries were isolated and prepared to evaluate endothelium-dependent vasorelaxation in response to increasing concentrations of Acetylcholine. We have previously published specifics regarding these assays for blood pressure and vascular function in pregnant rats [84,85].

5.2. Systolic arterial pressure measurements

Blood pressure was measured by tail-cuff plethysmography using the CODA8 (four channels) device (EMKA Technologies). After five days to one week training, female mice were placed in restrainers on a warming platform (37 °C). The cuff releases progressively the pressure of the cuff and the apparatus records the volume variation of the tail, algorithmically determining the systolic and diastolic pressure, with at least 8 satisfactory measurements per day, taken at the same hour of the day. Systolic and diastolic BP always displayed similar curve profiles in all groups analyzed, therefore only systolic BP is shown.

5.3. Assessment of albumin/creatinine ratio (ACR) in urine

Urine was collected by soft abdominal pressure on the female mice to induce urination, and the samples were stored at −80 °C until use. However, urine samples were difficult to collect from all females at every time point, and therefore results were pooled within each group for each gestational period (early, mid and late). ACR was measured by ELISA using the Albuwell M kit (#1011, Ethos Biosciences) following the accurately the specifications of the manufacturer. For ACR ratio, data are from n = 7 for WTxWT, n = 5 for WTxWT + BH4, n = 10 for WTxTgSTOX42 and n = 9 for WTxTgSTOX42+BH4.

5.4. Heart weight and tibia length measurements

At days 16.5, a part of the mice in each group were sacrificed. The hearts were collected, the blood removed by pressure on an absorbing paper, and weighted. The posterior right tibia was dissected free and measured, to caculate the heart weight to tibia length ratio. Data are from data from n = 6 for WTxWT, n = 6 for WTxWT + BH4, n = 4 for WTxTgSTOX42 and n = 6 for WTxTgSTOX42+BH4.

5.5. Placental transcriptome

Total RNAs from pools of three placenta from different mice in the four conditions were used (Ctrl, Ctrl + BH4, STOX1 (preeclamptic-mimic) mice, and STOX1+BH4), and hybridized to ClarionM mice microarrays (#902931, Thermofisher Scientific), that interrogate more than 22,000 annotated genes. The microarray data was deposited in the EBI database for open accessibility to all the interested researchers. The analyses were carried out using the TAC tool from Affymetrix and the online tool WebGestalt [60] which makes it possible to carry out straightforwardly Gene Set Enrichment Analyses and confront them to Gene Ontology, Molecular Processes or KEGG databases. STRING protein networks analysis was performed using https://string-db.org/cgi/network.pl. Complete transcriptomic data are available under the EBI accession number M-TAB-9314.

5.6. Cell culture and treatments

The cell line overexpressing STOX1A (AA6, overexpressing STOX1A) was described by Rigourd et al., 2008 [23], as well as the control (BDS). In the present study we developed another cell line overexpressing STOX1B (B10). For this, we used plasmids encompassing the STOX1B isoform of STOX1, under the control of the CMV promoter. The cells (cultivated in DMEM GlutaMax (GIBCO-Life technologies) + penicillin/streptomycin and 10% heat-decomplemented Fetal Calf Serum, GIBCO-Life technologies), were transfected in the same time with a plasmid encoding a resistance gene to Geneticin G-418 in a 1:20 proportion (for a 60 cm² plate, 10 μg of STOX1 expression plasmid and 0.5 μg of resistance plasmid) to ascertain that the two plasmids entered the cells following transfection with lipofectamin 2000 (GIBCO-Life technologies) in standard conditions. Selection was carried out ~3 weeks at 500 μg/ml G418 (GIBCO-Life technologies). Resistant clones were isolated and cultivated separately, amplified and maintained in DMEM with Geneticin. The expression of STOX1B was tested by a specific RT-qPCR.

Cells were treated with 0.2 mM cobalt chloride for 24 h to mimic hypoxia-induced HIF1A stabilization, with or without 100 μM BH4 (T4425, Sigma Aldrich) or for 24h with 100 μM t-NAME (N5751, Sigma Aldrich).

5.7. Total RNA extraction and RT-qPCR

Total RNA was isolated from cells using the RNAeasy kit (Qiagen), then reverse-transcribed with SuperScriptIV Reverse transcriptase (Invitrogen). Real-time quantitative PCR was performed using PowerUp Sybr Green PCR Master Mix (ThermoFisher) and the rate of dye incorporation was monitored using the StepOne Plus RealTime PCR system (Applied Biosystems). Three biological replicates were used for each condition. Data were analyzed by StepOne Plus RT PCR software v2.1 and Microsoft Excel. Three biological replicates were used for each condition. Data were analyzed by StepOne Plus RT PCR software v2.1 and Microsoft Excel. Real-time PCR Ct values were analyzed using the 2 ΔΔCt method to calculate the fold expression. qPCR Primers used to detect STOX1 isoforms mRNA are: STOX1A forward-GCTCTTTGTGCCCTCGACAT, reverse-ATTCTCCACGGACACAGATG and STOX1B forward-ATGCCAGTCTCCATGACAT, reverse-TGCTTTGCA-CATTTTGCAAGA. The primers for STOX1B are expected to amplify a 100bp fragment on STOX1B transcript and not a 2265 bp fragment on STOX1A, which does not occur in RT-qPCR experiments, with elongation time limited to 10 s per cycle.

6. Immunofluorescence, immunoblot, reagents, and antibodies

Cells plated on glass slides were fixed with 2% (wt/vol) paraformaldehyde (PFA) and permeabilized with 0.5% Triton X-100. The glass slides were incubated in blocking buffer [BSA 5% (wt/vol) in PBS] overnight at 4 ºC then 1 h at room temperature with the primary antibody, and finally with the secondary antibody and 10 μg/ml Hoechst 33,342 for 1 h at room temperature. STOX1A/8 (HPA037845), HIF1A (SAB2101039), MsNod/SOD2 (polyclonal rabbit-Cy3, S1450), Catalase (SAB450383), DAB20A (4-Diaminofluorescein Diacatect, D2813), DCF-DA (‘2’,‘7’-Dichlorodihydrofluorescein diacatect, D6883), DHE
7. Detection of nitric oxide, the ROS O$_2^-$ and H$_2$O$_2$, the RNS ONOO$^-$ and mitochondrial ROS

Fluorescent molecules were used to measure NO (DAF2DA), ROS anion superoxide O$_2^-$ (DHE), ROS hydrogen peroxide H$_2$O$_2$ (DCF-DA) and RNS peroxynitrite ONOO$^-$ (DHR123) in freshly extracted total proteins. This was done in standard culture conditions, by spectrofluorometry and using microplate reader TECAN Infinite 200. Mitochondrial ROS (anion superoxide O$_2^-$) were detected using MitoSOX Red after 1h incubation at 37 °C, and fluorescence intensity was quantified after 3D-confocal microscopy acquisition and reconstruction. Experiments were done in triplicate.

7.1. Fumarate, 1-malate and acetyl-coA content measurements

The fumarate content was measured with the Fumarate Assay Kit (Colorimetric assay, MAK060-1 KT, Sigma-Aldrich), the 1-malate content with the Malate Assay Kit (Colorimetric assay, MAK067-1 KT, Sigma-Aldrich), and the Acetyl-CoA content with the Acetyl-Coenzyme A Assay Kit (Fluorometric assay, MAK039-1 KT, Sigma-Aldrich) following the manufacturer’s instructions. Tests were performed with freshly extracted total proteins. Experiments were done in triplicate.

8. Total ATP measurement and OXPHOS contribution

For ATP levels, cells were treated with 10 μM oligomycin for 1 h (for glycolytic ATP) or untreated (for total ATP levels); then, ATP levels were assessed with the CellTiter-Glo Luminescent assay (Promega), according to supplier instructions.

8.1. Citrate synthase activity, fumarase activity, SOD activity and catalase activity

Citrate synthase activity was measured with the Citrate Synthase Activity Colorimetric Assay kit (K318-100, BioVision). In the same plate, Fumarase levels were tested with sandwich ELISA assay before total and specific fumarase activities with the Fumarase Specific Activity Assay Kit (Colorimetric assay, AB110043, ABCAM), Total SOD activity was assessed with the Superoxide Dismutase Activity Colorimetric Assay kit (AB65354, ABCAM) that measures the inhibition activity of xantine oxidase by SOD, and catalase activity with the Catalase Assay kit (AB83464, ABCAM) following the manufacturer’s instructions. Tests were performed on freshly extracted total proteins. Experiments were done in triplicate.

8.2. Three-dimensional confocal acquisition, reconstruction, and quantification

For STOXA1/B, iNOS/NOS2, Fumarase, MnSOD/SOD2, Catalase and MitoSOX Red imaging, confocal acquisitions were made using a spinning-disc Yokogawa CellVoyager CV1000 with a 40X air objective (Biolmagerie Photonique UTechS PBI, Institut Pasteur, Paris). Optical slices were taken each 1 μm interval along the z axis covering the whole depth of the cell, at resolution of 1.024/1.024 pixels. Each image was 3D-reconstructed using Maximum Intensity Projection (MIP). Scale bar = 10 μm. Fluorescence quantification was done using a single-imaging frame collection and FIJI ImageJ software (postacquisition analysis). For each condition, 60 cells were analyzed from three independent experiments.

8.3. Protein extraction and immunoblot

Cells were lysed with lysis buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1% Triton X-100, 0.1% SDS, 2 mM DTT, and protease inhibitor mixture). Lysed cells were not centrifuged, and the whole extract was collected. The protein content was determined with the Bradford reagent (Sigma-Aldrich), and 20 μg of protein were loaded for SDS/PAGE. After blotting, Trans-blot Turbo nitrocellulose membranes (Bio-rad) were probed with primary antibodies, then with HRP-conjugated secondary antibodies from Thermo Fisher Scientific.
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