Original Article

Extracellular ATP induces albuminuria in pregnant rats

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Abstract

Background. As circulating plasma ATP concentrations are increased in pre-eclampsia, we tested whether increased plasma ATP is able to induce albuminuria during pregnancy.

Methods. Pregnant (day 14) and non-pregnant rats were infused with ATP (3000 µg/kg bw) via a permanent jugular vein cannula. Albuminuria was determined, and blood samples were taken for leukocyte counts, plasma ATP and plasma haemopexin activity. At Day 20 of pregnancy, rats were sacrificed, fetuses and placentas weighed and kidney and placental tissue were snap frozen for immunohistology.

Results. ATP infusion induced albuminuria exclusively in pregnant rats, together with increased neutrophil counts, decreased staining for glomerular sialoglycoproteins and CD39 expression, significant intraglomerular monocyte infiltration and increased glomerular intracellular adhesion molecule-1 (ICAM-1) expression. Plasma haemopexin activity was increased in saline-infused pregnant rats as compared to non-pregnant rats but was inhibited in pregnant ATP-infused rats (to non-pregnant levels). At the end of pregnancy (Day 20), increased plasma ATP level was exclusively seen in ATP-infused pregnant rats. In pregnant rats as compared with non-pregnant rats, we found decreased expression of glomerular AT1 receptors, which was increased after ATP infusion exclusively in pregnant animals.

Conclusion. The present study shows that ATP infusion induced albuminuria exclusively in pregnant rats. Why extracellular ATP showed this pro-inflammatory response exclusively in the pregnant condition is unclear but is probably related with relatively enhanced non-specific immunity and inflammatory reactions characteristic for the pregnant condition.

Keywords: albuminuria; extracellular ATP; haemopexin activity; inflammation; pregnancy

Introduction

Haemopexin (Hx), which belongs to the family of acute phase proteins, shows significant protease activity [1]. Accordingly, active Hx is able to induce proteinuria in rats after intra-renal infusion of Hx in saline [2]. Since proteinuria during pregnancy is a hallmark of pre-eclampsia (PE), a common complication of pregnancy, we tested whether Hx activity was increased during pre-eclampsia. In contrast to our expectation, plasma Hx activity appeared to be significantly lower in patients with pre-eclampsia as compared to normal pregnant control individuals of the same gestational age [3]. Thus, in pre-eclampsia, proteinuria appeared not to be associated with increased plasma Hx activity.

However, in plasma samples from healthy pregnant women, we observed enhanced plasma Hx activity [3] as compared with non-pregnant women. We have shown that Hx activity can be inhibited with serine protease inhibitors or nucleotides like extracellular ATP [4,5]. Indeed, the decreased Hx activity during pre-eclampsia appeared to be due to an increased titre of plasma ATP observed in patients with pre-eclampsia [3]. In vitro treatment of plasma samples from pre-eclamptic patients with ATP-hydrolyzing phosphatases, like apyrase or alkaline phosphatase, leads to reactivation of the Hx activity in these plasma samples [3]. This is in line with previous data, showing that Hx activity can be inhibited by ATP in vitro and subsequently reactivated by apyrase or by endothelial cells expressing ecto-apyrase (CD39) [6]. The possibility of reactivating inactivated Hx by alkaline phosphatase could be the base of a future therapeutic approach, since alkaline phosphatase is a naturally occurring enzyme, the concentration of which is increased in pregnancy and decreased in pre-eclampsia [7]. Interestingly, alkaline phosphatase is presently used in phase II clinical trials for other disorders (http://clinicaltrials.gov/ct2/show/related/NCT00727324).

The observation that pre-eclampsia is associated with increased plasma ATP concentrations has led to the hypothesis that extracellular ATP per sé may be toxic, particularly
during pregnancy. Therefore, we now conducted experiments in which pregnant and non-pregnant rats were infused with ATP (3000 µg/kg BW) during 1h at Day 14 of pregnancy. Urinary protein excretion, glomerular inflammation and expression of glomerular angiotensin II receptor 1 (AT-1R) as well as various inflammatory parameters of the circulation were evaluated.

Materials and methods

Animals
All animal experiments were approved by the Animal Care Ethics Committee of our University. Female Wistar outbred rats (about 200 g) were kept in a temperature- and light-controlled room (lights on from 7:30 AM till 7:30 PM) with free access to food and water. Until selection for experiments, vaginal smears were taken daily, and rats were rendered pregnant by housing them on pro-oestrus with fertile males for one night.

When spermatozoa were detected in the smear the next day, this day was designated as Day 0 of pregnancy. In all rats, a cannula was inserted into the right jugular vein under fluothane/oxygen anaesthesia according to the method of Steffens [8]. This cannula allows stress-free blood sampling and infusions.

Experimental design
Pregnant rats were infused with 3000 µg/kg bw ATP in 2.0 ml saline (n = 10) or with 2.0 ml saline alone (n = 10) on Day 14 of pregnancy. Non-pregnant rats were infused with 3000µg/kg bw ATP in 2.0 ml saline (n = 8) or saline alone (n = 8) on di-oestrus. For determination of albumin excretion, rats were placed in metabolic cages for 24 h 2 days before (Day 12), immediately after (Day 15), 3 days after (Day 17) and 6 days after the infusion (Day 20). At this time, none of the rats had delivered or started parturition. At the same days, blood samples (0.4 ml in EDTA) for white blood cell counts, plasma ATP concentrations and plasma Hx activity were taken from the jugular vein cannula. Six days after the infusion (i.e. Day 20 in pregnant rats, which is almost the end of pregnancy in the rat, delivery is at Day 21), rats were sacrificed by bleeding under fluothane/oxygen anaesthesia. Fetuses and placentas were collected and weighed, and kidney and placenta tissue samples were snap frozen and prepared for immunohistology.

Measurement of urinary albumin
Rats were placed in metabolic cages for 24 h from 11:00 AM until 11:00 AM the next day. The volume of urine samples was measured, and urinary albumin levels were determined using rocket electrophoresis with rabbit-anti-rat albumin antiserum (Nordic Immunological, Tilburg, The Netherlands) as described before [9].

Leukocyte counts and differential leukocyte counts
Blood leukocytes were counted using a microcellcounter (model Sysmex F800, Toa Medical Electronics, Kobe, Japan). A smear was made from each blood sample and stained according to standard methods with the May–Grunwald–Giemsa method. After evaluation of 500 cells per smear, relative and absolute numbers of neutrophils, lymphocytes and monocytes were calculated according to standard methods.

Evaluation of plasma Hx activity
Protease activity of plasma Hx was evaluated by the ‘glomerular ECM stripping assay’ as described previously [2,5,10]. This in vitro assay is based on the potential impairment by Hx of glomerular extracellular matrix molecules (ECM), such as expression of sialoglycoproteins or CD39 expression. Acetone-fixed cryostat normal rat kidney tissue is incubated with either (1:8 diluted plasma samples (100 µl)section for 60 min; plasma from 3 and 6 days after the infusion) from pregnant rats infused with saline, pregnant rats infused with ATP; non-pregnant rats infused with saline and non-pregnant rats infused with ATP. Following incubation, sections were washed and stained for glomerular ECMs (i.e. sialoglyco-
Results

Urinary albumin excretion

Figure 1 shows that ATP infusion increased 24-h albumin excretion, peaking at Day 17, exclusively in pregnant rats. No significant alterations after ATP infusion in non-pregnant rats were observed; no effect of saline infusion was observed.

Colloidal iron staining

To demonstrate that loss of glomerular charge selectivity can play a role in the albuminuria observed, we stained kidney sections for glomerular sialoglycoproteins, reflecting glomerular anionic sites. Mean arbitrary scores for glomerular sialoglycoproteins are shown in Figure 2a. It can be observed that the amount of reaction product is significantly decreased in pregnant ATP-infused rats as compared with pregnant saline-infused rats. No difference of glomerular anionic sites between saline-infused pregnant and saline-infused non-pregnant rats could be observed; nor did we observe an effect on glomerular anionic sites after ATP infusion with respect to this staining in non-pregnant rats. Representative glomeruli of control pregnant rats (infused with saline) (left panel) and pregnant rats infused with ATP (right panel) are shown in Figure 2b. These photomicrographs show a clear loss of stainability for sialoglycoproteins of the capillary walls and the mesangium in a glomerulus of an ATP-infused pregnant rat as compared with a glomerulus of a saline-infused pregnant rat.
crographs show strong staining along the capillary walls and the mesangial area of the glomerulus of a pregnant rat infused with saline (left photomicrograph) and a clear loss of stainability for sialoglycoproteins of the capillary walls and the mesangium in a glomerulus of an ATP-infused pregnant rat (right photomicrograph).

**Glomerular inflammation**

To evaluate whether the loss of glomerular sialoglycoproteins reflecting that affected charge selectively is induced by an inflammatory response and/or oxidative stress, we evaluated glomerular inflammation by staining kidney sections for intraglomerular influx of monocytes, neutrophils and lymphocytes as well as for the expression of ICAM-1, indicating endothelial cell activation. As a parameter for the presence of oxidative stress, we measured the expression of CD39, an ecto-enzyme which is extremely sensitive to oxidative stress [13].

It can be seen that the mean number of intraglomerular monocytes increased after ATP infusion in pregnant rats as compared to non-treated pregnant rats (Figure 3). We found no effect of ATP infusion on glomerular monocyte number in non-pregnant rats. At this interval after infusion, i.e. 6 days, we only found a single neutrophil or lymphocyte in the glomeruli of the kidneys in the various groups of rats; no differences between the four groups of rats were observed for neutrophil or lymphocyte numbers (results not shown).

Since we found increased numbers of monocytes but not of neutrophils or lymphocytes in the glomeruli of pregnant rats following ATP infusion, we evaluated whether there was a relationship between the number of infiltrated glomerular monocytes and the amount of albumin excreted in the urine (Figure 4). In Figure 4, the regression line represents a linear relationship between the number of ED-1 positive cells per glomerulus at Day 20 vs the 24-h urinary albumin excretion at the same day in pregnant ATP-treated rats ($R^2 = 0.68$; slope 492.4, significantly different from zero, $P < 0.05$). No relationship was found between intraglomerular ED-1 positive cells and 24-h albumin excretion in pregnant saline-treated or non-pregnant ATP- or saline-treated rats.

Upregulation of endothelial ICAM-1 indicates pro-inflammatory activation of these cells. Figure 5 shows increased glomerular ICAM-1 expression in pregnant ATP-infused rats as compared with saline-infused pregnant rats. No effect of ATP infusion was seen upon glomerular ICAM-1 expression in non-pregnant rats. Figure 5b shows representative glomeruli of pregnant rats, infused with saline (left panel) or ATP (right panel) stained for ICAM-1. ICAM-1 expression can be seen in the glomerular tuft in pregnant rats following ATP infusion, indicating activation of endothelial cells but not in pregnant rats following saline infusion.

Thus decreased expression of CD39 may reflect endothelial injury induced by oxygen free radicals released during a local inflammatory response. It can be seen that
glomerular CD39 expression is significantly decreased in pregnant ATP-treated rats as compared with saline-infused pregnant rats (Figure 6a), indicating oxidant injury within glomeruli of pregnant ATP-infused rats. No significant alterations of glomerular CD39 expression were observed after infusion of ATP into non-pregnant rats as compared with saline-infused non-pregnant rats. Figure 6b shows representative glomeruli from pregnant rats stained for CD39 expression. The pregnant saline-infused rat shows a linear staining pattern along the capillary walls (left panel). After infusion of ATP in pregnant rats, significant loss of CD39 throughout the glomerular tuft can be seen (right panel) as compared with glomeruli of pregnant saline-infused rats.

**ATP-induced inflammatory cells in the circulation**

To evaluate the potential systemic nature of the ATP-induced inflammatory response, we studied the number of circulating total leukocytes, neutrophils, monocytes and lymphocytes in ATP-infused pregnant rats vs control animals. The numbers of peripheral leukocytes counted at various intervals after ATP or saline infusion in pregnant and non-pregnant rats are shown in Figure 7. After infusion of ATP in pregnant rats, a significant increase of the mean leukocyte number was seen 1 and 3 days after ATP infusion, while the mean leukocyte number was decreased 6 days after ATP infusion as compared with the mean leukocyte number before infusion. In normal pregnant rats, the mean leukocyte number was also significantly decreased 6 days after the infusion of saline, i.e. at the end of pregnancy. No significant effect was observed regarding mean leukocyte number in non-pregnant rats infused with ATP or saline. As can be seen in the upper right panel of Figure 7, the mean neutrophil number was increased 1, 3 and 5 days after ATP infusion in pregnant rats as compared with the pre-infusion level. Mean monocyte number was decreased in pregnant rats after ATP infusion at Day 20 as compared with the pre-infusion level. Mean lymphocyte number increased 3 days after infusion and decreased 6 days after infusion of ATP as compared with pre-infusion levels (Day 12). No effect of ATP or saline infusion was seen on mean levels of leukocytes, neutrophils, lymphocytes or monocytes in non-pregnant rats.

**Plasma Hx activity and plasma ATP titre**

Next to its pro-inflammatory activity, ATP may act through inactivation of Hx, an effect which has been demonstrated in vitro [4,5]. We therefore measured plasma Hx activity and plasma ATP concentrations in the experimental and control groups. As can be seen from Figure 8, mean plasma Hx activity was increased in saline-infused pregnant rats as compared to saline-infused non-pregnant rats. On the other hand, mean plasma Hx activity is significantly
lower in ATP-treated pregnant rats as compared to saline-infused pregnant rats. Figure 9 shows that 6 days after ATP infusion, mean plasma ATP level was significantly increased in pregnant ATP-treated rats as compared to saline-infused pregnant rats and ATP-treated non-pregnant rats. In saline-infused non-pregnant rats, plasma ATP concentrations were not affected by the infusion.

Glomerular expression of the AT-1R

As activated Hx is able to decrease the expression of the AT-1R upon various cells in vitro [14], it was expected that the AT-1R was diminished in the tissue of control pregnant rats with increased Hx activity but not in ATP-infused rats in which circulating Hx was inhibited. Therefore, we stained kidney sections of the experimental and control rats for the presence of the AT-1R. Figure 10 shows representative micrographs of glomeruli immunohistologically stained for AT-1R expression. In non-pregnant control animals (Figure 10c), positive staining for AT-1R in a mesangial pattern can be observed. In line with our expectations, it can be observed that the glomerular expression of AT-1R is decreased in normal pregnant rats (with circulating active Hx) as compared with control non-pregnant rats, with inactivated Hx (Figure 10a and c). Pregnant rats, in contrast to non-pregnant animals, showed enhanced expression of AT-1R after infusion of ATP as compared to saline infusion (Figure 10b and d).

Bodyweight of fetuses

Finally, we studied the effect of ATP infusion on the weight of fetuses and placentas. Figure 11 shows that fetal weight was slightly but significantly decreased in pregnant rats infused with ATP as compared to saline-infused pregnant rats. No significant effect was seen on placental weight. There were no differences in mean numbers of fetuses between the ATP and saline-infused rats, and we found no fetal resorptions or fetal death in either group (results not shown).

Placental expression of CD39 and CD73

Although placental weight is not affected by ATP infusion, the fact that fetal weight is decreased suggests that placental function is impaired after ATP infusion. As affected placental function may be caused by ischaemia, we evaluated placental expression of CD39 and as well as CD73 activity (Figure 12). Decreased CD39 expression together with increased CD73 activity has been recognized as a hallmark of ischaemia [13,15]. Photomicrographs of representative placental tissue stained for CD39 or CD73 are shown in Figure 12a–f, while semi-quantitative evaluation of the staining intensity, using an arbitrary scale, is shown in Figure 12g. ATP-infused rats showed decreased placental CD39 expression as compared to control rats. It can be seen that CD39 expression is decreased in the maternal vasculature (V) of ATP-treated rats vs saline-infused rats.
(D vs C), while also in the microvasculature of the labyrinth (L), expression of CD39 is decreased in ATP-infused pregnant rats vs control pregnant rats (B vs A). Panels E and F show representative micrographs of enhanced placental activity of CD73 in a pregnant ATP-infused rat (panel F) vs a pregnant saline-infused rat (panel E). In contrast to saline-infused pregnant animals, ATP-treated pregnant rats showed increased CD73 activity in giant trophoblast cells (GT) and between the decidua (DB) and the giant trophoblast cell layer (GT). As can be seen in Figure 12g, the CD73 activity was significantly increased in placental sections of ATP-infused pregnant rats as compared with saline-infused pregnant rats (Figure 12g).

Discussion
As we previously observed enhanced plasma ATP levels in subjects with PE as compared with healthy pregnant women of the same gestational age [3], the notion emerged that extracellular ATP may be a potentially toxic molecule for the pregnant rat. The aim of the present study was to sub-
To instantiate this hypothesis. Therefore, pregnant rats infused with ATP on Day 14 of pregnancy were compared with control pregnant rats infused with saline and with non-pregnant rats which also were given a single infusion of ATP or saline. Since at Day 14 of pregnancy the placenta of the rat has fully developed, we selected this gestational age for the ATP infusion. Slow infusion was carried out via a permanent jugular vein cannula in conscious conditions, which is preferable over a single injection in view of the short half-life of extracellular ATP in vivo [16]. It appeared that exclusively in pregnant rats, urinary albumin excretion associated with decreased loss of glomerular anionic sites occurred after ATP infusion. Next to systemic

![Fig. 9](image_url)

**Fig. 9.** Mean (± SEM) plasma ATP concentrations in saline-infused pregnant rats (black bars), pregnant rats infused with ATP (open bars), saline-infused non-pregnant rats (dotted bars) and non-pregnant rats infused with ATP (striped bars). *P < 0.05, Mann–Whitney U-test, significantly different from saline-infused pregnant rats.

![Fig. 10](image_url)

**Fig. 10.** Photomicrographs of rat glomeruli after immunostaining for the AT-1R. In non-pregnant saline-infused animals (c), positive staining for AT-1R in a mesangial pattern can be observed. Saline-infused pregnant rats (with active Hx in their circulation) show almost negative staining at Day20 of pregnancy (a). Pregnant rats, 6days after infusion of ATP, with inhibited Hx in their circulation, clearly show a positive mesangial staining pattern of expression of the AT-1R (b). Positive staining is also shown in non-pregnant rats 6days after infusion of ATP in non-pregnant rats (d).

![Fig. 11](image_url)

**Fig. 11.** Fetal and placental weight in saline-infused pregnant rats (black bars) and in ATP-infused pregnant rats (open bars). *P < 0.05, Mann–Whitney U-test, significantly different from saline-infused pregnant rats.
as well as glomerular inflammation, also placental ischaemia was observed in ATP-infused pregnant rats vs saline-infused pregnant rats. Finally, the drop of mean fetal weight observed in ATP-treated pregnant rats may be considered a consequence of the pro-inflammatory and ischaemic events occurring in these animals.

Since systemic inflammation seems to be elicited by ATP infusion in pregnant rats as reflected by increased numbers of circulating inflammatory cells and lymphocytosis, we feel that albuminuria may be due to this inflammatory response. This notion is supported by the observations that glomerular ICAM-1 expression is upregulated exclusively in pregnant ATP-infused rats and that the mean intraglomerular influx of ED-1 positive monocytes correlates with the albumin excretion. The absence of significant enhancement of monocytes in the circulation in ATP-treated pregnant rats, in contrast to neutrophils and lymphocytes, is in line with the influx of these cells into the glomeruli of the

Fig. 12. (a–f) Photomicrographs of expression of CD39 and CD73 in the placenta of pregnant rats treated with ATP (b, d and f) and saline-infused pregnant rats (a, c and e) (Day 20 of pregnancy) using an antibody against CD39 and a standard enzyme histochemical method to visualize the activity of CD73. Expression of CD39 can be found in the decidual region (c and d) and in the labyrinthal (L) region (a and b). It is clearly shown that decreased reaction product is visible in the maternal vasculature of ATP-treated rats vs control rats (d vs c), while also the microvasculature of the labyrinth showed decreased staining in ATP-infused rats vs control pregnant rats (b vs a). ATP-treated pregnant rats show increased CD73 activity on giant trophoblast cells (GT) as well as between the decidua and the giant trophoblast cell layer as compared with control rats (f vs e). DB, decidua basalis; GT, giant trophoblast cells; ST, spongiotrophoblast cell layer; V, maternal vein; L, labyrinth; MT, mesometrial triangle. (g) Semi-quantitative expression in arbitrary units of placental CD39 and placental CD73 activity in saline-infused pregnant rats (black bars) and in ATP-infused pregnant rats (open bars). A significant decrease of CD39 expression and a significant increase of CD73 can be seen in ATP-infused vs control pregnant rats. *P < 0.05, Mann–Whitney U-test, significantly different from saline-infused rats.
kidney. Indeed, sequestration of monocytes into the kidney is not unlikely, since similar types of sequestration are observed in human renal disease [17]. The glomerular infiltration of monocytes may lead to enhanced glomerular permeability for albumin. The observation of decreased colloidal iron staining in pregnant ATP-treated rats as compared with the other groups of rats supports the notion that loss of glomerular anionic sites promotes impairment of glomerular perm selectivity leading to albuminuria [18,19]. The question whether ATP exerts a direct toxic effect upon the kidney or rather induces a systemic response affecting the kidney remains to be investigated. However, taking the data together, we feel that during the short circulation time of ATP, a predominant systemic inflammatory response is induced in the present model, possibly through stimulation of purinergic P2x or P2y receptors occurring on both neutrophils and monocytes [20]. Although no massive proteinuria is observed in the present model, it cannot be excluded completely that protein overload may contribute to glomerular damage and recruitment of inflammatory cells [21].

Also, the loss of glomerular CD39 expression supports the concept that oxidant stress is associated with the local inflammatory activity in the glomerulus due to the presence of monocytes. The loss of CD39 expression, which are extracellular matrix molecules extremely sensitive to toxic oxygen metabolites [13], occurring in association with increased glomerular permeability has been demonstrated in other experimental models in the rat, including adriamycin nephrosis [22–24]. Loss of vascular CD39 expression as reflected in kidneys as well as in placentas of ATP-infused pregnant rats may be in line with enhanced plasma ATP in these rats observed at Day 20. Thus, due to the decreased vascular CD39 expression, the normal hydrolysis of extracellular ATP by vascular endothelium is diminished, resulting in enhanced plasma ATP levels at the end of pregnancy in these animals. Moreover, the staining pattern, i.e. decreased CD39 staining concomitantly with enhanced CD73 staining, is characteristic for ischaemia in the placenta [13,15], which may promote enhanced plasma ATP at Day 20 of pregnancy. The pathogenesis of the placental ischaemia in the present model remains to be investigated. It may be speculated, however, that the infused ATP binds to P2Y receptors on the trophoblast, which may induce superoxide anion production [25,26]. Moreover, as the trophoblast surface is coated with CD39, it is likely that degradation products of ATP, like ADP, may contribute to local microthrombus formation in the placental vasculature, promoting local ischaemic injury [27].

Like in the human situation [3], plasma Hx activity is increased during normal pregnancy in the rat as compared to non-pregnant rats. In the ATP-infused pregnant rat, plasma Hx activity was not enhanced as compared with non-pregnant rats. Since extracellular ATP is a potent inhibitor of Hx activity, this might be ascribed to the infused ATP into the circulation of these rats. However, significant increase of plasma ATP could only be detected at Day 20 of pregnancy, whereas diminished Hx activity in these animals occurred before Day 20 as well. The reason for this discrepancy is obscure and needs further experimentation. It is conceivable, however, that the infused ATP binds to circulating Hx molecules inhibiting its activity, whereas the enhanced plasma ATP level detected at Day 20 originates from the ischaemic placenta of these rats shown at the end of pregnancy. Experimental confirmation of significantly less placental ischaemia earlier in pregnancy of ATP-infused rats might help to explain the lack of ATP peaking before Day 20. Be this as it may, it is clear that a single ATP infusion is apparently able to induce a systemic inflammatory reaction (and placental ischaemia at the end of pregnancy) in pregnant rats.

The question as to the potential function of enhanced Hx activity in the healthy pregnant rat emerges. As suggested for the human condition [3], the increase in plasma Hx during normal pregnancy in the rat may be necessary for vascular expansion. We have recently shown that activated Hx is able to shed the AT-1R from various cell types, including endothelial cells and monocytes [14]. It is conceivable that during normal rat pregnancy, increased Hx activity is responsible for AT-1R downregulation, which may contribute to a decreased responsiveness for angiotensin II. In the rat, similar to humans, non-responsiveness for angiotensin II, associated with healthy pregnancy, is necessary for expansion of the vascular bed and appropriate placental perfusion [28]. In the present study, we observed decreased expression of the AT-1R in the glomerular microvasculature of normal pregnant rats as compared to non-pregnant rats. This is in line with loss of the AT-1R observed in normal human pregnancy [29]. In ATP-treated pregnant rats lacking enhanced Hx activity, the AT-1R expression seemed to be increased as compared to normal pregnant rats. Thus, lack of active Hx, induced by ATP infusion, leaves the AT-1R expression unaffected, as reflected by increased glomerular expression of the AT-1R in ATP-infused pregnant animals.

It seems likely that the increased AT-1R expression in ATP-infused pregnant rats is also involved in the albuminuria seen in these rats, since angiotensin II increases intraglomerular pressure and impairs the size-selective function of the glomerular filtration barrier to macromolecules, such as plasma proteins [30]. Also via non-haemodynamic effects, angiotensin II receptor may be involved in proteinuria, since binding of angiotensin II to the AT-1R may result in oxygen-free radical production and release of cytokines and adhesion molecules [31]. Therefore, the upregulation of this receptor in the glomeruli after ATP infusion in pregnant rats may promote the glomerular inflammation seen in these rats. Further analysis of vascular AT-1R in relation to albuminuria and glomerular inflammation in this experimental model is in progress.

In summary, in the present communication, we demonstrate that ATP is toxic for pregnant rats. Infusion of ATP induces albuminuria, systemic inflammation, decreased plasma Hx activity and increased expression of AT-1R. The reason why extracellular ATP shows this pro-inflammatory response exclusively in the pregnant condition is unclear at this moment. It may, however, be related with the relatively enhanced level of non-specific immune activity and inflammatory reactions characteristic for the pregnant condition in humans [32–34] and in rats [12]. This is in line with previous data showing that infusion
ATP induces albuminuria in pregnant rats of a single ultra-low dose of the pro-inflammatory lipopolysaccharide (LPS) resulted in a relatively strong and persistent inflammatory response in pregnant rats in contrast to non-pregnant rats showing little response [12,35].

Supplementary data

Supplementary data is available online at http://ndt.oxfordjournals.org.

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Conflict of interest statement. None declared.

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