Gender differences in the long-term effects of perinatal hypoxia on pulmonary circulation in rats

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Submitted 15 November 2002; accepted in final form 9 April 2003

Chronic hypoxia is a well-defined, clinically important cause of pulmonary hypertension. With restoration of normoxia, hypoxic pulmonary hypertension is completely reversible (17). However, if the hypoxic exposure occurs at around the time of birth, the recovery is incomplete, and subtle yet significant alterations of the pulmonary circulation persist well into adulthood. Although characterized variously in different studies, these changes can be summarized as an increased susceptibility to the development of pulmonary hypertension later in life (10, 14, 19, 20, 31, 33).

In 1997, Barer et al. (1) noticed in a preliminary report that the late effects of perinatal hypoxia appear more prominent in female than in male rats. In agreement with this observation, resting pulmonary arterial blood pressure (PAP) and/or pulmonary vascular resistance in adults were slightly but significantly elevated in studies of perinatal hypoxia where pooled male and female rats were studied (10, 19) but not in experiments on males only (14).

The present study was, therefore, designed to test the hypothesis that the long-term effects of perinatal hypoxia on pulmonary circulation differ between genders and that the presence of gonads during maturation plays a prominent role in this variability. The results were reported in a preliminary form (13).

Methods

The project in this study was reviewed and approved by the Animal Studies Committee of the Charles University Second Medical School. All procedures conformed to the European Community and National Institutes of Health guidelines for the use of experimental animals. Wistar rats were purchased from Anlab (Prague, Czech Republic). To analyze the specific, late effects of perinatal hypoxia and gonadectomy on pulmonary circulation, three experiments were performed (1 main and 2 supplementary). Their overall design is schematically illustrated in Fig. 1.

Main experiment: perinatal hypoxia and neonatal gonadectomy. In the main experiment, adult rats were divided into groups based on gender, history of perinatal hypoxia, and gonadal status. This resulted in four male and four female groups in the main experiment: 1) perinatally normoxic with intact gonads; 2) perinatally normoxic gonadectomized as newborns; 3) perinatally hypoxic with intact gonads; and 4) perinatally hypoxic gonadectomized as newborns.

In fact, the groups with the intact gonads consisted of rats that did not undergo any surgery and rats serving as sham surgery controls. Since we did not find any significant differences between sham-operated and nonoperated rats, their data were pooled to enhance clarity by reducing the large number of groups.

Perinatal hypoxia and gonadectomy. The rats were exposed to perinatal hypoxia as previously described (12, 14). Pregnant rats were placed into a normobaric hypoxic chamber (12% O2) 1 wk before the expected date of delivery (term = 3 wk). Newborn pups were kept in the same environment with their mothers for another 5–7 days after birth.
and then kept in room air until maturity (23 wk). Some of the pups were gonadectomized right after removal from the hypoxic chamber in deep ether anesthesia (24). Sham surgery was performed on some of the pups under identical conditions as the gonadectomy.

**Hemodynamic and morphological measurements.** When the rats were adults (24 wk old), they were anesthetized with thiopental (40 mg/kg ip). While they breathed room air spontaneously, their carotid artery was cannulated to measure systemic arterial blood pressure (SAP). Under oscilloscopic monitoring of pressure, the pulmonary artery was then catheterized via the right jugular vein and right ventricle as previously described (14, 16, 23), and PAP was recorded.

Mechanical ventilation with room air was then begun through a tracheostomy at ~60 breaths/min (10 cmH2O peak inspiratory pressure, 0 cmH2O end-expiratory pressure). The chest was opened by sternotomy with extra care taken to minimize bleeding, and an ultrasonic flow probe (2.5-mm SS series with J reflector; Transonic Systems, Ithaca, NY) was placed at the ascending aorta to measure aortic blood flow (T106 flowmeter, Transonic Systems) as an estimate of cardiac output (11, 23). This value relative to body weight is reported as cardiac index (CI). The values obtained with this method are lower than cardiac output in vivo due to the anesthesia and especially the thoracotomy.

The heart was then dissected, and the right and left ventricles and septum were separately weighed (8). Right ventricle to left ventricle plus septum weight ratio (RV/LV+S) was used as an index of right ventricular hypertrophy associated with pulmonary hypertension. The lungs were filled through the trachea with neutral formal solution at a pressure of 12 cmH2O and then placed in the same solution for several days. Lung sections were then cut and stained by the hematoxylin resorcin-fuchsin method. Because the muscular layer of pulmonary vessels is enclosed between two elastic laminae, whereas the nonmuscular vessels only have one lamina, the degree of muscularization of peripheral lung vasculature was determined as previously described (17, 23) by separately counting distal pulmonary vessels (adjacent to alveoli or alveolar ducts, <300 μm in diameter) with and without double elastic lamina. All such vessels in one sagittal section of the left lung of each rat were counted (median 105 vessels). The percentage of the total that is double laminated (%DL) was then calculated (17, 23).

**Supplementary experiment 1: gonadectomy in adulthood.** Because the data in the main experiment showed significant effects of neonatal gonadectomy, a question arose as to whether, for these effects to occur, the gonads must be absent just at the time of the measurement in adulthood or, rather, during the preceding period of development. To address this issue, a supplementary experiment was performed in which male and female rats were exposed to perinatal hypoxia identically as the rats in the main experiment, but the gonadectomy was postponed until maturity (13 wk of age). They were then given 10 wk to recover from the surgery before being measured.

**Supplementary experiment 2: hypoxia in adulthood.** To confirm that the effects observed in the main experiment were due to hypoxia acting specifically at the perinatal period, another supplementary experiment was performed. Rats were exposed to identical hypoxia as in the main experiment (12% O2 for 2 wk), however, not at the perinatal period, but instead after reaching maturity (starting from 12 wk of age). Immediately afterward, some of the rats were gonadectomized. All rats then lived in room air for 10 wk. Only RV/LV+S was determined in this supplementary experiment.

**Statistical analysis.** The data were analyzed using one-factor ANOVA followed by Scheffé’s post hoc test (StatView 5.0.1; SAS Institute, Cary, NC). The Scheffé’s test was chosen.
because of its relative robustness with uneven n between groups. Our groups varied in size because of the pooling of nonoperated and sham-operated controls and because not all measurements were successful in all rats (e.g., excessive bleeding before CI measurement). Statistical significance was assumed when \( P < 0.05 \).

**RESULTS**

Late effects of perinatal hypoxia in gonad-intact males and females. Although adult males were heavier than females (as expected), perinatal history of hypoxia had no effect on body weight in adulthood (Table 1). Perinatal hypoxia had no significant effect on SAP, PAP, CI, or left ventricle plus septum weight in adulthood in either sex (Table 1, Fig. 2, A and D). The relative weight of the right ventricle (RV/LV+S), equal in perinatally normoxic males and females, was significantly increased by the perinatal experience of hypoxia in adult females (Fig. 2B). The degree of peripheral pulmonary vessel muscularization (%DL) was not significantly affected by perinatal hypoxia (Fig. 2C), although in males, the \( P \) value was marginal (0.058).

**Perinatal hypoxia and neonatal gonadectomy.** In perinatally normoxic rats, neonatal gonadectomy had no significant effect on any of the variables measured in adulthood (Table 1, Fig. 2). In perinatally hypoxic males and females, neonatal gonadectomy had no significant effect on body weight, SAP, CI, and left ventricle plus septum weight in adulthood (Table 1, Fig. 2D). Neonatal castration had no effect on PAP and RV/LV+S in adult males with a history of perinatal hypoxia (Fig. 2, A and B). %DL was higher in perinatally hypoxic castrated males compared with perinatally normoxic males with intact gonads (by 85%) or those that had been gonadectomized (Fig. 2C).

**DISCUSSION**

The main new findings of the present study can be summarized in two points. 1) Hypoxia in the perinatal (but not in other) period had a permanent consequence of elevated RV/LV+S in intact females. 2) Ovariectomy at a very young age (but not in adulthood) greatly augmented the effects of perinatal hypoxia in females so that PAP and %DL in adulthood were considerably increased. By contrast, castration of males (neonatal or adult) did not unmask any appreciable long-term effects of perinatal hypoxia on pulmonary circulation except an increased %DL (much less than in females).

The late effects of perinatal hypoxia on pulmonary circulation were first reported in 1990. In a study on male rats, we found that perinatal hypoxia did not cause any significant elevation of PAP, RV/LV+S, or %DL in adulthood (in agreement with our present results), but pulmonary vasoconstrictor reactivity to acute hypoxic challenges was greatly increased by perinatal hypoxia in rats recovering from chronic hypoxia in adulthood (14). We interpreted this finding as indicating that perinatal hypoxia led to increased susceptibility to stimuli, promoting the development of pulmonary hypertension in adulthood. At the same time, Hakim and Mortola (10) reported slightly but significantly elevated vascular resistance in isolated perfused lungs of a mixed group of adult male and female rats that had been exposed to neonatal hypoxia, compared with normoxic controls. Pulmonary vasoconstrictor reactivity to acute hypoxia was also elevated (10).

Although there are discordances in details between our earlier study (14) and that of Hakim and Mortola (10), they are both consistent with an idea that perinatal hypoxia does not cause permanent severe pulmonary hypertension, but that it does increase the susceptibility of adult lung vasculature to insults later in life. This concept was subsequently confirmed several times in various models (2, 15, 19, 20, 33), including a study in humans (31). Thus our present finding of definite, yet relatively weak, effects of perinatal hypoxia on baseline characteristics of pulmonary circula-
tion in intact adult rats is consistent with these previous reports.

In the present experiment, animals with a history of perinatal hypoxia were studied when much older (6 mo) than rats in the previous reports (1.5–4 mo) (2, 10, 14, 15, 19, 20). The fact that the effects of perinatal hypoxia on pulmonary circulation were still evident after a normoxic interval this long suggests that they may be permanent rather than only slowly reversible.

Chronic hypoxic pulmonary hypertension in adulthood is less severe in females than in males (27). This might be, in part, related to less severe acute hypoxic pulmonary vasoconstriction in females (5, 34). It is, therefore, somewhat surprising in our present study that with perinatal hypoxia, the late effects are slightly more prominent in intact females than males. Because the greater resistance of female pulmonary circulation to hypoxia in adulthood is caused mostly by female sex hormones (5, 34), our logical next step was to test the role of gonads in gender variability of the late effects of perinatal hypoxia. We found that early ovariectomy did not eliminate the gender differences, as is the case with hypoxia in adulthood (34). However, on the contrary, early ovariectomy exacerbated them. These data imply two interesting phenomena. One is that ovarial function is protective not only against hypoxic pulmonary hypertension in adulthood but also against the late effects of perinatal hypoxia. The second implication is that without the protective action of the ovaries, pulmonary vasculature of female rats is much more sus-

![Fig. 2. Effects of perinatal hypoxia and neonatal gonadectomy on pulmonary circulation, cardiac index (CI), and right ventricle in adulthood. Intact, rats with intact gonads; Gx, gonadectomized rats; open bars, animals born and raised in normoxia; filled bars, perinatally hypoxic rats. Numbers in parentheses are n of perinatally normoxic per n of perinatally hypoxic animals. Data are means ± SE. A: mean pulmonary arterial blood pressure (PAP) is not significantly affected by perinatal hypoxia in intact males and females but is elevated in perinatally hypoxic females ovariectomized as newborns. B: relative weight of the right ventricle (right ventricle to left ventricle plus septum wet weight ratio, RV/LV+S) is increased by perinatal hypoxia in females but not in males. Left ventricle plus septum weights were unaffected by perinatal hypoxia or neonatal gonadectomy (Table 1). C: muscularization of peripheral pulmonary vessels expressed as percentage of double-laminated (i.e., muscularized) peripheral vessels (%DL) is markedly increased in perinatally hypoxic females ovariectomized as newborns. A similar change in males is much less prominent. D: tendency for lower CI estimate in perinatally hypoxic compared with perinatally normoxic rats did not reach statistical significance. *P < 0.005 differs from all other groups. **P < 0.0001 differs from all other groups. †P < 0.01; ††P < 0.005; †††P < 0.0001. BW, body weight.]

AJP-Lung Cell Mol Physiol • VOL 285 • AUGUST 2003 • www.ajplung.org
ceptible to adverse, long-term effects of perinatal hypoxia. Thus there are marked gender differences in postnatal development of pulmonary circulation that have both gonad-dependent and independent components.

The protective effects of female sex hormones, especially estrogen, on circulation, in general (6, 22, 35), and on pulmonary vessels, in particular (25, 26, 28), have been studied quite extensively. By affecting numerous signaling systems, such as nitric oxide, prostacyclin, endothelin, angiotensin II, and various growth factors, estrogen decreases vascular smooth muscle tone and inhibits vascular wall growth (3, 4, 6, 7, 18, 21, 32). It is likely that one or more of these mechanisms may also be involved in the protective effect of ovaries against long-term effects of perinatal hypoxia on pulmonary circulation. However, despite the attractiveness of estrogen as the agent of protection in this situation, our study does not positively prove its role. Possible involvement of other ovarian hormones cannot be excluded. Hypothetically, changes in gonadotropin hormones secondary to gonadectomy might also play a role. Androgen involvement seems unlikely because its levels in females are markedly lower than in males, where their removal had much less effect.

Because estrogen relaxes the vascular wall and inhibits its growth (6, 22, 35), one likely explanation for the protective effect of the ovaries could have been that it masks the effects of perinatal hypoxia by continually stimulating the synthesis of vasodilators in adulthood, including the time of our measurements. Our data indicate that this is not the case. For the protective effect to occur, the ovaries must be functional during maturation, but not at the time of measurement in adulthood. Thus the ovaries appear to act on the pulmonary vasculature during development to reduce the long-term hypertensive consequences of perinatal hypoxia.

Although the influence of neonatal gonadectomy on the late effects of perinatal hypoxia was much more prominent in female than in male rats, it was not completely absent in males. Namely, the muscularization of peripheral pulmonary vessels was augmented in adult males with a history of perinatal hypoxia if they had been gonadectomized as pups, albeit much less so than in females. If, indeed, the protective effect of the ovaries in females is mediated by estrogen, as speculated above, then the same mechanism might also be in action in males. The much smaller protective effect of the testes compared with the ovaries would correspond to their much smaller estrogen production. Alternatively, an independent, yet smaller protective effect of testosterone cannot be excluded on the basis of our present data.

In adults, the properties of pulmonary (and other) vessels themselves are similar between males and females, and the advantage of a lower susceptibility to hypertension is imparted on the female vasculature by the female sex hormones. The most unusual finding of the present study is, therefore, that without the protective effect of the ovaries, the female pulmonary vessels are not equal to the males in their sensitivity to the late effect of perinatal hypoxia (females are affected much more). The cause of this gender difference is unknown, but it appears that it is an intrinsic property of the affected cells of the vascular wall that is not governed by sex hormones.

The gender differences in the late effects of perinatal hypoxia and gonadectomy are selective to pulmonary circulation: SAP did not differ among the groups regardless of gender, gonadal status, or perinatal history (Tables 1 and 2). This is in agreement with a recent study showing equal SAP in perinatally normoxic and hypoxic rats (29). It is possible that pulmonary selectivity is due to the substantial remodeling of the pul-

### Table 2. Gonadectomy in adulthood does not affect systemic and pulmonary circulation of perinatally hypoxic male and female rats

<table>
<thead>
<tr>
<th>Gonads</th>
<th>Sex</th>
<th>BW, g</th>
<th>SAP, mmHg</th>
<th>PAP, mmHg</th>
<th>CI, ml·min⁻¹·100 g⁻¹ BW</th>
<th>RV/LV + S</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intact</td>
<td>Males</td>
<td>603 ± 12*</td>
<td>127 ± 2</td>
<td>15.6 ± 0.7</td>
<td>7.8 ± 0.4</td>
<td>0.27 ± 0.01</td>
</tr>
<tr>
<td></td>
<td>Females</td>
<td>364 ± 19</td>
<td>123 ± 3</td>
<td>15.9 ± 0.4</td>
<td>6.7 ± 0.6</td>
<td>0.27 ± 0.01</td>
</tr>
<tr>
<td>Gonadectomy (in adulthood)</td>
<td>Males</td>
<td>563 ± 16*</td>
<td>128 ± 3</td>
<td>14.9 ± 0.8</td>
<td>7.2 ± 0.4</td>
<td>0.26 ± 0.01</td>
</tr>
<tr>
<td></td>
<td>Females</td>
<td>392 ± 14</td>
<td>125 ± 3</td>
<td>15.7 ± 0.8</td>
<td>6.0 ± 0.6</td>
<td>0.27 ± 0.02</td>
</tr>
</tbody>
</table>

Data are means ± SE. Parentheses show numbers of rats. All groups were perinatally hypoxic and measured when adult. PAP, mean pulmonary arterial blood pressure; CI, open-chest estimate of cardiac index; RV/LV + S, right ventricle to left ventricle plus septum wet weight ratio. *P < 0.001 males differ from corresponding female group.

### Table 3. Chronic hypoxia in adulthood does not have any delayed effect on relative right ventricle weight

<table>
<thead>
<tr>
<th>Gonads</th>
<th>Sex</th>
<th>Hypoxia in Adulthood</th>
<th>BW, g</th>
<th>RV/LV + S</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intact</td>
<td>Males</td>
<td>No</td>
<td>653 ± 11(7)*</td>
<td>0.29 ± 0.02(7)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>593 ± 17(8)*</td>
<td>0.30 ± 0.02(8)</td>
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</tr>
<tr>
<td>females</td>
<td>No</td>
<td>378 ± 19(4)</td>
<td>0.29 ± 0.01(4)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>363 ± 5(7)</td>
<td>0.29 ± 0.02(7)</td>
<td></td>
</tr>
<tr>
<td>Gonadectomy (in adulthood)</td>
<td>Males</td>
<td>Yes</td>
<td>586 ± 16(8)*</td>
<td>0.30 ± 0.01(8)</td>
</tr>
<tr>
<td></td>
<td>Females</td>
<td>Yes</td>
<td>402 ± 11(7)</td>
<td>0.32 ± 0.03(7)</td>
</tr>
</tbody>
</table>

Data are means ± SE. Parentheses show numbers of rats. All rats were measured 10 wk after a 2-wk exposure to hypoxia in adulthood. *P < 0.0001 males differ from corresponding female group.
monary vasculature that occurs at the time of birth and to the fact that chronic hypoxia causes pulmonary, but not systemic, hypertension. However, the mechanism of acute hypoxic sensing appears to be permanently altered by perinatal hypoxia both in the pulmonary (vasoconstriction) (10, 14) and systemic (vasodilation) (29) vascular beds. Our data do not exclude the possibility that stimuli more relevant to the systemic circulation, acting at the perinatal period, may have late effects similar to those observed in this study with hypoxia and pulmonary circulation.

The prominent vulnerability of female pulmonary circulation to the late consequences of perinatal hypoxia and the protective role of functional ovaries resonate with a clinical experience that the much feared primary pulmonary hypertension is about twice as frequent in women than in men and that it is typically a disease of reproductive age (9, 30). Although there are currently no data to support a mechanistic connection between these observations, it is tempting to speculate that a history of insults during a critical period of development, such as perinatal hypoxia, might combine with female sex hormone alterations to increase the susceptibility to primary pulmonary hypertension. Further examination of this speculation appears warranted.

In conclusion, we found that perinatal hypoxia causes long-term alterations of pulmonary circulation and right heart that are more pronounced in female than in male rats. Chronic hypoxia in adulthood does not have such delayed effects. Gonadectomy at a very young age, but not in adulthood, greatly augments the late pulmonary hypertensive effects of perinatal hypoxia, but only in females. Thus male and female pulmonary vessels inherently differ in sensitivity to perinatal hypoxia. During maturation, ovaries exert a protective influence against the permanent consequences of perinatal hypoxia in lung vessels.

We thank O. Hnilicková and K. Venclíková for skillful technical assistance.

DISCUSSIONS

This study was supported by Grant Agency of the Czech Republic Grants 305/97/0570 and 305/00/1432 and by Czech Ministry of Education Research Project 111300002.

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