INHIBITION OF ADRENOCORTICAL ACTIVITY BY DEXAMETHASONE IN NEWBORN PIGLETS

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Abstract


Dexamethasone was given orally to half of the piglets of two litters in one 0.25 mg dose within one hour of birth and to half of the piglets of other two litters in two 0.25 mg doses on the 3rd day after birth. The remaining animals in each litter served as controls. All the piglets were exsanguinated 12 hours after treatment. The concentration of 17-hydroxycorticosteroids (17-OHCS) in the plasma of newborn and three-day-old experimental piglets was reduced 64 and 36 per cent, respectively, as against the controls. The capacity of adrenal slices to produce 17-OHCS in vitro on stimulation with corticotrophin was suppressed in the two groups by 20 and 16 per cent. The treatment with dexamethasone also affected ascorbic acid concentration in the adrenal, produced a non-significant increase in blood glucose, reduced ascorbic acid concentration in the blood plasma and liver and had no effect on adrenal mass and on the circulating eosinophile granulocyte level. The demonstration of negative feedback and the decrease of endogenous corticosteroids confirmed functional maturity of the pituitary-adrenal system in newborn piglets. The foetal-maternal relations are analysed from the viewpoint of comparative physiology and the mechanisms controlling corticosteroid level in piglets during the perinatal period are discussed.

Corticosteroids, ACTH, feedback, development, pig.

Investigations of the pathogenesis of splayleg of newborn pigs have shown that the degenerative changes of skeletal muscle are morphologically similar to those seen in corticosteroid myopathy (Zelen and Jirasek 1979). A hypothesis suggested the involvement of a hormonal imbalance, an increased production of corticosteroids in foetuses as a result of stressors to which sows are exposed in the last third of pregnancy (Tuček et al. 1980). The recent prevalence of this congenital disease has drawn attention to the question of physiological preconditions for the development of foetal hyperadrenocorticism, the clinical form of which has not been described (Cleveland and 1970).

Some indication of foetal hyperadrenocorticism can be seen in the observation that the concentration of 17-hydroxycorticosteroids (17-OHCS) in newborn pigs was several times higher on the first day after birth than in older resting pigs (Dvořák 1967). Although few data are available on the endocrine changes in pigs during their foetal development, the observations on adrenal steroids suggest a physiological rise of adrenocortical function of foetal pigs during the last 15 to 20 days of gestation (Dvořák 1972, 1973a; Fevre 1975).

The level of corticosteroids in the blood plasma of sows at farrowing time is substantially lower than in newborn pigs (Fevre 1975; Brenner and Gürler 1977). In man the reverse is the case: the level of cortisol in the umbilical plasma is only 20 to 30 per cent of that of the mothers who have their cortisol concentration several times increased at parturition (Kaupe et al. 1972). Examination of foetal monkeys a few days before birth showed that 58 per cent of their cortisol was of maternal origin (Kittenger 1974). However, there is no evidence of transplacental transfer of maternal adrenocorticotropic hormone (ACTH) in man (Cleveland and 1970) or other mammals. In rats the extirpation of the autonomous hypothalamus-pituitary-adrenocortical axis was described in the last days of foetal development (Corbier and Roffi 1978). Nevertheless, lack or excess of
Table 1

Body, adrenal and liver mass, circulating eosinophile granulocyte count and glucose and ascorbic acid concentration in the organs of piglets after treatment with dexamethasone.

<table>
<thead>
<tr>
<th></th>
<th>1-day-old piglets</th>
<th>3-day-old piglets</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>experimental</td>
<td>control</td>
</tr>
<tr>
<td>No. piglets</td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td>Dose of dexamethasone (mg)</td>
<td></td>
<td></td>
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<tr>
<td>0.25</td>
<td>0.25</td>
<td></td>
</tr>
<tr>
<td>Body mass (kg)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>at treatment</td>
<td>1.29±0.16</td>
<td>1.38±0.18</td>
</tr>
<tr>
<td>at sacrifice</td>
<td>1.42±0.19</td>
<td>1.50±0.19</td>
</tr>
<tr>
<td>Adrenal mass</td>
<td></td>
<td></td>
</tr>
<tr>
<td>actual (mg)</td>
<td>202±45</td>
<td>215±40</td>
</tr>
<tr>
<td>relative (mg/kg)</td>
<td>143±27</td>
<td>148±15</td>
</tr>
<tr>
<td>Liver mass</td>
<td></td>
<td></td>
</tr>
<tr>
<td>actual (g)</td>
<td>32±6</td>
<td>33±4</td>
</tr>
<tr>
<td>relative (g/kg)</td>
<td>23±1</td>
<td>23±1</td>
</tr>
<tr>
<td>Eosinophile granulocytes (10^6/1)</td>
<td>20±20</td>
<td>14±27</td>
</tr>
<tr>
<td>Blood glucose (mmol/l)</td>
<td>7.86±2.75</td>
<td>6.38±2.20</td>
</tr>
<tr>
<td>Ascorbic acid concentration</td>
<td></td>
<td></td>
</tr>
<tr>
<td>of blood plasma (umol/l)</td>
<td>226±56</td>
<td>237±28</td>
</tr>
<tr>
<td>of adrenal (umol/g)</td>
<td>4.43±0.45</td>
<td>4.70±0.56</td>
</tr>
<tr>
<td>of liver (umol/g)</td>
<td></td>
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</tr>
</tbody>
</table>

b  Difference between the groups significant (P < 0.05)
maternal corticosteroids in rats in late gestation affects the secretion of ACTH by the foetal pituitary (D'Angel c a n c e l s the pituitary and adrenal response to or less able to inhibit ACTH secretion and thus to cause a reduction in corticosteroids in variously

The negative feedback of corticosteroids and ACTH lies in that an excess of corticosteroids inhibits the secretion of ACTH. In the cortisol servomechanism a decrease in free cortisol, on the other hand, results in increased secretion of ACTH whereby cortisol level is maintained within a relatively narrow range unless increased secretion of adrenocortical hormones is included under stressful conditions by hypothalamic ACTH-releasing factor. As far as is known, all steroids with corticoid activity are more or less able to inhibit ACTH secretion and thus to cause a reduction in corticosteroid secretion through feedback action. Cortisol is the major inhibitor in those mammals in which it is the main glucocorticoid (G a u n t e t al. 1965). A remarkably high potency in this respect is exhibited by synthetic dexamethasone which is used in human medicine in variously modified tests for examination of the syndromes of adrenocortical hyperfunction. In pigs of 20 kg in body mass pretreatment with dexamethasone at 4 or 8 mg cancels the pituitary and adrenocortical response to some stimuli (D o n a l d e t al. 1968; C o o k e t al. 1974).

The object of the present study was to confirm or to exclude the possibility of the endogeneous production of corticosteroids being suppressed after their exogeneous supply.

Materials and Methods

Large White piglets of 4 litters were employed. Half of the animals in each litter were left untreated to serve as controls. Experimental piglets of two litters (7 animals) were treated with Dexamethasone SP60A (16alpha-fluoro-16alpha-methylprednisolone) orally in a dose of 0.25 mg within one hour of birth when all piglets were weighed at 7 p.m. Experimental piglets of the other two litters (8 animals) were treated with two oral doses of the same drug on the 3rd day after birth at 7 a.m. and 7 p.m. All the experimental and control animals were exsanguinated by decapitation on the following day at 7 a.m. The characteristics of the groups are given in Table 1.

Determination of actual and relative adrenal and liver mass, eosinophile granulocyte count and glucose concentration in the blood, 17-hydroxycorticosteroid (17-OHCS) concentration in the plasma and ascorbic acid concentration in the plasma, adrenal and liver. The adrenals were also used for determination of the production of 17-OHCS by adrenal slices in vitro on incubation with ACTH. The methods have been described in previous publications (D vo ř á k 1972, 1973b, 1974). The results are tabulated as arithmetic means ± standard deviations and shown graphically as means ± standard errors of the means. The significance of the differences of the means was assessed by Student's t-test.

Results

The administration of dexamethasone to newborn and three-day-old piglets had no distinct effect on either their body mass or eosinophile granulocyte count or actual and relative adrenal mass as determined at 12 and 24 hours of the experiment, respectively. Blood glucose concentration was non-significantly higher and ascorbic acid concentration in the plasma and liver were non-significantly lower in the experimental animals than in the controls (Table 1). The animals treated on the third day after birth had a significantly (P < 0.05) higher ascorbic acid concentration in the adrenal and a higher relative liver mass as against the controls.

The most prominent changes occurred in the direct criteria of adrenocortical activity. The administration of dexamethasone reduced significantly (P < 0.01) the concentration of 17-OHCS in the blood plasma and distinctly, though not significantly, the production of 17-OHCS by adrenal slices in vitro on incubation with ACTH (Fig. 1). It is noteworthy that the suppressive effect made itself felt more in newborn piglets: their plasma 17-OHCS level was reduced 64 per cent, whereas that of three-day-old piglets only 36 per cent as against the controls. Similarly, the production of 17-OHCS by adrenal tissue in vitro was depressed 20 per cent in newborn piglets and 16 per cent in three-day-old animals as against the controls.
Concentration of 17-hydroxycorticosteroids (17-OHCS) in the blood plasma and their production in vitro by adrenal slices after stimulation with ACTH in dexamethasone-treated (open bars) and control (hatched bars) piglets aged 1 and 3 days.

**Discussion**

The concentration of 17-OHCS in the plasma of control piglets was in keeping with the high values recorded in the first days after birth (Dvořák 1967, 1972) and with its progressive decrease. In the present case it was 33 per cent lower at 3 days of age than the first day. Concurrently, a somewhat less pronounced decrease was recorded in the production of 17-OHCS by adrenal tissue in vitro. The lower values in the experimental piglets than in the controls show quite convincingly that dexamethasone reduced the production of adrenocortical hormones and their release into the blood stream. Further evidence of it can be seen in the ascorbic acid level which decreases in the adrenals and rises in the blood and possibly also in the liver when enhanced adrenocortical function is induced with exogenous ACTH or by stress on starvation (Dvořák 1973b, 1974). The dose of dexamethasone was apparently not too large and did not act long, for one-day-old piglets did not develop enlargement of the liver which was found in three-day-old piglets treated with two doses or after stimulation of the adrenal cortex for two days (Dvořák 1974). There was only a slight difference between the experimental and control piglets in the count of circulating eosinophile granulocytes. This, together with the decreased level of plasma 17-OHCS in the experimental animals, suggests that dexamethasone which is quick to induce eosinopenia (Blenkinsopp and Blenkinsopp 1967) ceased practically to act 12 hours after administration. However, its waning effect was still perceptible, giving rise to a mild hyperglycaemia. The most important observation is the finding that the dose of synthetic glucocorticoid employed had a suppressive effect on the mechanisms controlling cortisol secretion.

The high values of 17-OHCS in the blood of newborn piglets indicate a certain degree of hypercortisolism as compared with the older animals. A similar state is found in human medicine in patients with Cushing's disease where, however, hypercorticalism is manifested clinically, the hyperfunction of the cortex being responsible for impairment of homeostasis. In piglets this state is, no doubt, physiological and corticosteroid hypersecretion can be regarded as an adaptive process. A high degree of maturity of the
hypothalamus-pituitary-adrenocortical axis in response to stressors has been demonstrated (Dvořák 1973b). Considering their response to suppressive action of dexamethasone, the piglets aged one and three days can be regarded as qualitatively mature also from the viewpoint of the feedback control of the hypothalamus-pituitary-adrenocortical system.

The existence of negative feedback was demonstrated experimentally also in neonatal infants (Kuno et al. 1972). However, their plasma corticosteroid level differs from that found in piglets in that it decreases within 1 to 2 days of birth to the level found in older children. This is obviously the result of suppressive action of cortisol that is transferred from maternal to foetal blood during labour (Tervillä et al. 1969) and reduces cortical secretion in the newborn. There is also evidence in rats that the feedback mechanism is functional during foetal and early postnatal life (Schaap et al. 1965; Klepac and Milković 1979) so that perhaps even endogenous hormones may inhibit stress response on the neonatal pituitary. Large doses of dexamethasone may suppress the increase in corticosteroid level in a state of stress (Donald et al. 1968; Sirett and Gibbs 1969), but dexamethasone does not affect adrenal response to exogenous ACTH (Hart et al. 1969). The sensitivity to ACTH after such treatment may be even enhanced (Bransome 1968).

In the present study this was apparently not the case in the production activity of adrenal slices in vitro.

In the light of the contemporary knowledge and of the observations reported here it seems reasonable to explain the state of adrenocortical activity in neonatal piglets as follows: High 17-OHCS levels found in the blood plasma of piglets in the days of postnatal life are not dependent on the dam, but they are the product of endogenous corticosteroid biosynthesis. Adrenocortical hyperfunction is being prepared in about the last 20 days of intra-uterine development and apparently plays a role in the initiation of parturition. It is conditioned by hyperplasia of the cortex and by a high degree of secretion of foetal ACTH. Feedback mechanisms of the hypothalamus-pituitary-adrenocortical system are functional and appear particularly sensitive in newborn piglets. Treatment with dexamethasone can suppress adrenocortical secretion. The high basal level of endogenous cortisol in the blood apparently suppresses ACTH release more moderately so that except in the first days after birth the subsequent decrease in resting 17-OHCS concentration of the blood is relatively slow. This situation is no obstacle to the response of the hypothalamus-pituitary-adrenocortical system to experimental stressors. When the life of a neonatal pig is endangered in the first days after birth, the high basal plasma 17-OHCS levels may be further increased considerably (Dvořák 1973b).

With regard to the situation during the last third of intra-uterine development of pigs, evaluation is not possible in the absence of species-specific data on the development of foetal-maternal hormonal relations and feedback mechanism. In stressful situations, mainly of acute character, to which sows are exposed, maternal glucocorticoids can apparently pass through the placenta by passive diffusion as is the case with the other steroids (Hill and Longo 1980). However, when exposed to protracted and repeated action of stressors, sows cannot be expected to develop a long-term rise in adrenocortical hormone level (Aberle et al. 1976) in consequence of the adaptation of the organism to environmental conditions. The situation may be complicated by potential differences in the metabolism of foetal and maternal steroids and by the influences of maternal nutrition on hormonal regulation of foetal development and on foetal glucocorticoid receptors (Mulay et al. 1980). Provided that the feedback mechanisms of foetal pigs are functional, which can reasonably be assumed to be so at the end of their intra-uterine life, then a passively induced rise in their corticosteroid concentration would suppress their own production and this would not result in a permanently increased corticosteroid level unless a prolonged exogenous supply of glucocorticoids was involved. This view is supported by the finding of reduced actual and relative adrenal mass of both newborn and ten-day-old piglets from sows treated with dexamethasone for three days during advanced pregnancy (Hühn and Kinpel 1979), since high plasma corticoid level in growing piglets is generally associated with high relative adrenal mass.

Inhibice adrenokortikální aktivity novorozených selat dexamethasonem

Dexamethason v perorální dávce 0,25 mg byl podán polovině selat dvou vrhů během jedné hodiny po narození, polovině selat dalších dvou vrhů dvakrát 3. den života. Za 12 hodin byla všechna vykrvena. U pokusných novorozených selat byla ve srovnání s kontrolami
snížena koncentrace plazmatických 17-hydroxykortikosteroidů (17-OHCS) o 64 %, u tří- denních o 36 %. Schopnost produkovat 17-OHCS in vitro při stimulaci s ACTH byla potla- čena u prvních o 20 %, u druhých o 16 %. Ošetření dexamethasonem ovlivnilo koncentra- ci kyseliny ascorbórové v nadledvíně, neprůzráčné zvýšilo hladinu krevní glukózy a sníž- lo koncentraci kyseliny ascorbórové v krevní plasmě i v játřech, nepůsobilo na hmotnost nadledvis ani nemělo důsledek na počet cirkulujících eosinofilních granulocytů. Prokázá- ná negativní zpětná vazba se snížením endogenních kortikosteroidů potvrzuje funkční zralost systému hypofýza - nadledviny u novorozených selat. Je diskutováno o feto- -maternitních vztazích z hlediska komparativní fyziologie a o mechanismech kontrolujících hladinu kortikosteroidů v perinatalním údobí vývoje selat.

Торжжение адренокortикальной активности новорожденных поросят дексаметазоном

Дексаметазон пероральной дозой 0,25 mg давали половине поросят двух племенных гнезд в течение одного часа после опороса, половине поросят двух последующих гнезд два раза на 3 сутки жизни. Все они были через 12 часов обезврежены. По сравнению с контрольными группами подопытных поросят была понижена концентрация плазматических 17-оксихидроксикортикостероидов (17-OHCS) на 64 %, у поросят в возрасте 3 суток - на 36 %. Способность продукции 17-OHCS в пробирке при стимуляции с ACTH была у первых живот- ных подавлена на 20 %, у второй группы - на 16 %. Подача дексаметазона оказала влияние на концентрации аскорбиновой кислоты в надпочечной железе, несущественно увеличивая уровень глюкозы в крови и понижая концентрации аскорбиновой кислоты в кровяной плазме и печени. Дексаметазон не оказывал влияния на массу надпочечных желез и на численность циркулирующих eosinofilních granulocyte. Установления отрицательные обратные связи с понижением endogenních kortikosteroidů подтверждает функциональную зрелость системы гипофиз - надпочечные железы у новорожденных поросят.

Обсуждается вопрос плодово-материнских отношений с точки зрения сравни- тельной физиологии и механизмов, контролирующих уровень кортикостероидов в перинатальный период развития поросят.

References