EXPERIMENTAL STUDY

Ectopic Agouti protein overexpression increases stimulated corticosterone production without effect on adenylate cyclase activity in mouse adrenal cells

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Abstract

Objective: The antagonism of Agouti protein (AP) and Agouti-related protein on melanocortin receptors suggests an inhibitory role in the regulation of steroidogenesis. However, we have previously demonstrated that ectopic AP overexpression increased restraint-induced corticosterone release and adrenal reactivity to ACTH in mice. A high steroidogenic response to ACTH may be a consequence of a stimulatory AP action on the adenylate cyclase (AC) and/or intracellular steroidogenic enzymes.

The aim of the present study was to estimate the effect of ectopic AP overexpression on the activity of AC and steroidogenic intracellular enzymes.

Methods: ACTH and forskolin were used for AC stimulation, and dibutyryl cAMP and progesterone were used for stimulation of intracellular steroidogenic enzymes in isolated adrenal cells in male C57Bl/6J mice of two Agouti genotypes: Ay/a (ectopic AP overexpression) and a/a (absence of AP in all tissues).

Results: ACTH and forskolin increased cAMP accumulation to the same extent in both Ay/a and a/a mouse adrenal cells (P < 0.001; ANOVA), but resulted in higher corticosterone production in Ay/a mice (P < 0.001 for ACTH and P < 0.01 for forskolin; ANOVA). Dibutyryl cAMP- and progesterone-induced corticosterone production was higher in Ay/a mice than in a/a mice (P < 0.001 for dibutyryl cAMP and P < 0.01 for progesterone; ANOVA).

Conclusions: Ectopic AP overexpression increased stimulated corticosterone production and intracellular steroidogenic enzyme reactivity to cAMP without an effect on AC activity.

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Introduction

The Agouti protein family includes Agouti protein (AP) and Agouti-related protein (AgRP). They have a striking similarity in structure and function (1), although their patterns of distribution in tissues are completely different. AP is a paracrine-signaling factor that is normally expressed only in the skin of rodents, regulating coat color (2). The human homolog of AP is expressed in a wide range of tissues, including adipose tissue, testis, ovary, heart, foreskin, kidney and liver (3). However, its physiological functions in human tissues remain unclear to date. AgRP is expressed primarily in the hypothalamus and adrenal cortex of humans and rodents, and at low levels in testis, lung and kidney of mice (1, 4, 5). Central AgRP is involved in appetite regulation and energy balance, but little is known about AgRP function in adrenals and other peripheral tissues (1).

The most accepted mechanism for AP/AgRP action is a classical competitive antagonism of melanocortin receptors (MCRs) (6, 7). Some findings suggest that AP not only inhibits the binding of agonists, but also alters the interaction of MCRs with intracellular effectors (8). Other studies have demonstrated that both AP and AgRP act as inverse agonists rather than classical competitive antagonists of MCRs (9). Thus, these authors (8, 9) suggest a more complex mechanism by which APs and the melanocortin peptides mediate a physiological response via the MCRs.

Melanocortin peptides (adrenocorticotropin (ACTH) and α-melanocyte-stimulating hormone (αMSH)) are important regulators of the hypothalamic–pituitary–adrenal (HPA) axis. The role of APs in the regulation of HPA function, including steroidogenesis in the adrenal gland, is poorly understood, and findings are very conflicting (10, 11). Melanocortin receptors MC2-R (ACTH receptors), MC3-R, MC4-R and MC5-R have been found in the adrenal cortex (4, 5). The antagonism of APs on MCRs suggests its inhibitory role in the regulation of steroidogenesis. Human recombinant AP was demonstrated to be a potent inhibitor of MC2-R signaling in the adrenal cortical lineage (7). AgRP
was shown to completely abolish αMSH-induced steroid secretion, mediating the inhibitory effect via MC3-R and MC4-R (5, 12). Nevertheless, basal and stress-activated corticosterone production was previously reported to be within normal limits in the Agouti yellow (A⁺/a) obese mice with overexpression of AP (13). Moreover, it has been shown that ectopic AP overexpression increases restraint-induced corticosterone release (11, 13), higher adrenal sensitivity to exogenous ACTH in vivo and an elevated response of the whole adrenal gland to ACTH stimulation in vitro (13).

Apparently the effect of ectopic AP overexpression on adrenal steroidogenesis is not limited only by the inhibition of ACTH receptors. It is possible that AP can modulate post-receptor signal transmission in adrenal cells. AP has been shown to increase the intracellular concentration of Ca²⁺ in myocytes and adipocytes (14, 15). It also increases gene expression in adipose tissue and the β-cells of the pancreas gland (16, 17). High steroidogenic response to ACTH in A⁺/a mice may be a consequence of the stimulatory action of AP on plasma membrane enzyme adenylate cyclase (AC) and/or intracellular steroidogenic enzymes.

The aim of this study was to estimate the effect of ectopic AP overexpression on the activity of AC and steroidogenic cAMP-dependent enzymes in adrenal cells. As a model of AP overexpression we used mice with the autosomal dominant mutation Agouti yellow (A⁺/a) (2). In this report, we have demonstrated that ectopic AP overexpression increased stimulated corticosterone production and intracellular steroidogenic enzyme reactivity to cAMP without an effect on AC activity.

Materials and methods

Materials were obtained from the following sources: corticosterone, ACTH(1-39), dibutyryl cAMP, forskolin, collagenase type 1A, 3-isobutyl-1-methylxanthine (IBMX), bovine serum albumin (BSA, fraction V) from Sigma Chemical Company, St Louis, MO, USA; [³H]corticosterone from Amersham Biosciences, Amersham, Bucks, UK; progesterone from Fluca, Buchs, Switzerland; cAMP RIA kits from Immunotech, Marseille, France.

Animals

The study was undertaken on 3-month-old male C57Bl/6J mice of two Agouti genotypes: A⁺/a with ectopic AP overexpression and a/a with an absence of AP. Mice of a/a and A⁺/a genotypes were siblings obtained from reciprocal crosses A⁺/a × a/a and a/a × A⁺/a. Animals were bred in the vivarium of the Institute of Cytology and Genetics (Siberian Division, Russian Academy of Science, Novosibirsk, Russia). They were housed in groups of five per cage under natural light conditions, at an ambient temperature of 21°C with free access to water and food. Animals were caged separately for 4 days before the experiments. A⁺/a mice weighed 27.1±1.1 g (n = 15) and a/a mice weighed 25.7±0.4 g (n = 13). Because of circadian rhythms, all experiments started at 0900 h. All studies were carried out under the highest standards of humane animal care using the International European ethical standards (86/609-EEC) and Russian national instructions for the care and use of laboratory animals.

Isolation and purification of adrenal cells

Cells were prepared from adrenals by decapitation and collagenase digestion. Briefly, adrenals from five mice (of the same genotype) were cleaned of fat and cut into small pieces. Glands were removed and incubated in Kreb–Ringer bicarbonate buffer (KRB; pH 7.4; 1.5 ml/10 glands) with 4% BSA and 4% collagenase at 37°C in an atmosphere of 95% O₂ and 5% CO₂. After 60 min, the suspension was dispersed by gentle homogenization (repeated pipetting), filtered through nylon gauze and centrifuged at 36 g for 10 min then at 96 g for 10 min at 4°C. The suspension was washed three times with 2.5 ml cold KRB with 0.5% BSA. After the final centrifugation, the cells were resuspended in 2.0 ml KRB with 0.5% BSA and counted with Trypan blue in a hemocytometer chamber. Cells were suspended in KRB with 0.5% BSA to a final concentration of 4×10⁵ cells/ml (to a total of ~14×10⁵ adrenal cells for each experiment) with a viability higher then 90%, as determined using the Trypan blue method.

Incubation of adrenal cells

Alquots (0.2 ml) of adrenal cell suspension (4×10⁵ cells/ml) were incubated in KRB with 0.5% BSA under an atmosphere of 95% O₂ and 5% CO₂ in the presence or absence of different doses of stimulators: ACTH (10⁻¹³, 10⁻¹², 10⁻¹¹, 10⁻¹⁰ M) or forskolin (10⁻⁶, 10⁻⁵, 10⁻⁴ M) or dibutyryl cAMP (10⁻⁵, 10⁻⁴, 10⁻³ M) or progesterone (2 μg/ml). After 2 h of incubation the suspension was centrifuged at 1800 g for 5 min. The cell-free medium was decanted and stored at −20°C until assayed for corticosterone. Samples for the measurement of cAMP accumulation were collected in the same experiments immediately after the samples were decanted for the corticosterone assay. Three experiments were conducted, each in triplicate.

Corticosterone assay

Corticosterone was measured by competitive protein binding using serum from female mice with inner transcrortin, and unlabeled and labeled corticosterone. Intra- and interassay coefficients of variation were less than 5 and 10% respectively.
cAMP assay

In the cAMP experiments, 0.5M IBMX (phosphodiesterase inhibitor) was included in the medium (KRB). After samples were decanted for the corticosterone assay, the cells were resuspended in 0.1 ml aliquots of KRB and cAMP production was terminated by the addition of 0.1 ml perchloric acid (0.2 M). Cells were then disrupted by sonication for 4 min. Samples were precipitated by centrifugation at 180g for 5 min. The cell-free medium was decanted and stored at −20°C until cAMP assay. cAMP accumulations were determined by RIA according to the manufacturer’s specification.

Data analysis

All data are expressed as the means±S.E.M. Statistical analysis was performed using two-way ANOVA with post hoc comparison by Newman–Keuls test. A difference between two means was considered to be statistically significant when P < 0.05.

Results

There were no genotype differences in corticosterone production and cAMP accumulation in untreated adrenal cells in all experiments (Figs 1, 2, 3 and 4).

Effects of ACTH and forskolin on cAMP accumulation in A/y/a and a/a mice

ACTH stimulation: ACTH at doses of 10−13 and 10−12 M did not increase the cAMP level in either A/y/a or a/a mice. ACTH at doses of 10−11 and 10−10 M increased cAMP accumulation to the same extent in both genotypes (P < 0.001; ANOVA) (Fig. 1A).

Forskolin stimulation: Forskolin at a dose of 10−6 M had no effect on the cAMP level and higher doses (10−5 and 10−4 M) increased cAMP accumulation to the same extent in both genotypes (P < 0.001; ANOVA) (Fig. 1B).

Effects of ACTH and forskolin on corticosterone production in A/y/a and a/a mice

ACTH stimulation: All ACTH doses markedly increased corticosterone production in both A/y/a and a/a mice (P < 0.001; ANOVA) (Fig. 2A). The increase was greater in A/y/a mice than that observed in a/a mice (P < 0.001; ANOVA). Corticosterone production achieved a plateau at the 10−12 M ACTH dose in both A/y/a and a/a mice.

Forskolin stimulation: All forskolin doses markedly increased corticosterone production in both A/y/a and a/a mice (P < 0.001; ANOVA) (Fig. 2B). The increase was greater in A/y/a mice than in a/a mice (P < 0.001; ANOVA) with a maximal activation effect at 10−5 M, following a decline at 10−4 M in both genotypes.

Effects of dibutyryl cAMP on corticosterone production in A/y/a and a/a mice

Dibutyryl cAMP at 10−4 and 10−3 M progressively stimulated corticosterone production in both A/y/a and a/a mice (P < 0.001; ANOVA) (Fig. 3). The effect of the Agouti genotype was significant (P < 0.01; ANOVA): at a dose of 10−3 M dibutyryl cAMP, A/y/a mice showed a greater increase of corticosterone production compared with a/a mice (P < 0.05; Newman–Keuls test).

Effects of progesterone on corticosterone production in A/y/a and a/a mice

Progesterone at 2μg/ml stimulated corticosterone production in A/y/a and a/a mice (P < 0.001; ANOVA) (Fig. 4). The effect of the Agouti genotype was significant (P < 0.01; ANOVA): A/y/a mice showed a greater increase of corticosterone production compared with a/a mice (P < 0.001; Newman–Keuls test).
Discussion

We have demonstrated that ACTH-stimulated corticosterone production in adrenal cells was higher in mice with ectopic AP overexpression. This result is consistent with others who have reported that ACTH-induced corticosterone release was increased in vivo (11, 13) and in vitro in mice with ectopic AP overexpression (13). Agouti proteins (AP and AgRP) were shown to be antagonists of MCRs (6, 7). Since AP inhibits hormone action on MCRs, chronic antagonism of AP in A/y/a mice may result in the development of some compensatory mechanisms which intensify ACTH-induced corticosterone production. The mechanisms of the enhanced activation of ACTH-induced steroidogenesis in the adrenals of A/y/a mice are unknown. Harris et al. (11) suggested that in mice with AP overproduction, chronic, partial antagonism of MC2-R led to an increased level of expression of the ACTH receptors in the adrenal cortex. Here we suggested and tested another possible mechanism which involves post-ACTH receptor regulation of membrane AC and/or intracellular steroidogenic enzymes.

We used forskolin to estimate AC activity. Forskolin stimulated steroidogenesis in a dose-dependent manner except for the highest dose of $10^{-4}$ M. This dose of forskolin increased cAMP accumulation but decreased corticosterone production. Apparently, the reduction in corticosterone secretion at this dose of forskolin is independent of AC activation. Forskolin at a dose of $10^{-4}$ M, unlike lower doses, has a direct Ca$^{2+}$ channel-blocking action (18), inhibits glucose transport (19, 20) and can decrease steroidogenic response by this means.

Contrary to ACTH, forskolin acts directly on the AC and increases cAMP level without interaction with
any of cell-surface receptors. We did not show an influence of ectopic AP overexpression on AC activity. Although forskolin-induced cAMP accumulation was the same in A/y/a and a/a mice, the steroidogenic response was greater in A/y/a mice compared with a/a mice. This may be a consequence of the response of high intracellular steroidogenic enzymes to cAMP. Our experiments with exogenous cAMP stimulation confirmed this assumption. The capacity of AP to up-regulate intracellular enzymes has been demonstrated with key intracellular lipid-metabolizing enzymes (21). The experiment with the administration of an excess of progesterone (cAMP-independent activation of steroidogenesis) indicated a higher activity of intracellular steroidogenic enzymes in A/y/a mice. The increased enzyme activity may account for the increased sensitivity of intracellular steroidogenic enzymes to cAMP action in A/y/a mice. A possible explanation for the rise in enzyme activity may be increased enzyme expression levels in mice with AP overexpression. The action of AP on gene expression has been shown in adipose tissue and β-cells of the pancreas gland (16, 17).

AP may act via a modulation of Ca²⁺ concentrations. Calcium plays an important role in the regulation of gene expression and enzyme activity in adrenals (22). AP was shown to increase the intracellular concentration of Ca²⁺ in myocytes and adipocytes (14, 15). It may be assumed that AP also increases intracellular concentrations of Ca²⁺ in the adrenal gland, stimulating steroidogenic enzyme activity.

ACTH-induced cAMP accumulation was also the same in both A/y/a and a/a adrenal cells. These data indicated that higher steroidogenesis in stimulated adrenal cells from the mice with ectopic AP overexpression did not result from changes in AC activity. Moreover, there were no correlations between corticosterone secretion and cAMP production in ACTH-induced adrenal cells. We used a physiological dose range for adrenal cell stimulation. Previously we have found (authors’ unpublished data) that plasma ACTH concentrations in both A/y/a and a/a mice varied from 2 × 10⁻¹⁲ M (basal) to 2 × 10⁻¹⁰ M (stress induced). ACTH at 10⁻¹³ and 10⁻¹² M stimulated steroidogenesis without an increase in cellular cAMP synthesis. Upon the addition of higher concentrations of ACTH (10⁻¹¹ and 10⁻¹⁰ M), cellular cAMP synthesis was significantly increased, but did not lead to a further increase of corticosterone secretion. This phenomenon has also been demonstrated by other authors (23–25). The lack of correlation between corticosterone secretion and cAMP production at the lower ACTH dosage may be due to the binding of cAMP with the cAMP-dependent protein kinase regulatory subunit, which does not allow measurement of the actual increase of cAMP production in cells (26).

On the other hand, ACTH at the lower concentrations may not stimulate steroidogenesis via AC. It has been demonstrated that low ACTH concentrations stimulate steroidogenesis in adrenal cells, increasing Ca²⁺ influx via T-type Ca²⁺ channels (27, 28). The 15-lipoxygenase metabolite(s) of arachidonic acid is also a second messenger for ACTH (25). Higher steroidogenesis in A/y/a mice may be due to a higher response by the adrenal cells to the second messengers other than cAMP. Thus AP may influence AC-dependent and AC-independent ACTH signaling pathways and so affect steroidogenesis.

AP normally is not expressed in adrenals in contrast to AgRP (29). Adrenal AgRP was shown to have an inhibitory paracrine effect on stimulated steroidogenesis acting via MC3-R and MC4-R (4, 5, 12). The pleotropic effect of A/y mutation on energy balance is considered to be due to AP inhibitory action at MC3-R and MC4-R in the hypothalamus (29). If this was the case for the adrenal, overexpression of AP would inhibit adrenal steroidogenesis. We found a contrary effect. Possibly adrenal AP exerts two opposite effects in A/y/a mice: it inhibits steroidogenesis by blocking MC3-R and MC4-R for αMSH action and stimulates steroidogenesis by increasing steroidogenic enzyme activity. The absence of differences between the adrenal function of A/y/a and a/a mice under resting conditions may be due to interference by the opposing tendencies of AP action. The prevalence of one or other effect may depend on the type of adrenal stimulation.

We have here demonstrated, for the first time, that ectopic AP overexpression increased stimulated corticosterone production and intracellular steroidogenic enzyme activity without an effect on AC activity. The exact mechanisms that enhance ACTH-induced steroidogenesis in A/y/a mice remain to be investigated. Understanding the role of APs in the regulation of steroidogenesis in rodents can help to elucidate the physiological function of APs in human tissues.

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