Adrenal Cortical Insufficiency—a Life Threatening Illness With Multiple Etiologies

Marcus Quinkler, Felix Beuschlein, Stefanie Hahner, Gesine Meyer, Christof Schöfl, Günter K. Stalla

SUMMARY

Background: The clinical signs of adrenal cortical insufficiency (incidence, ca. 25 per million per year; prevalence, ca. 400 per million) are nonspecific, and misdiagnoses are therefore common. Glucocorticoid substitution therapy has been in use for 50 years but is not a wholly adequate treatment. Our understanding of this disease remains incomplete in many ways.

Methods: We selectively searched the Medline database for publications on adrenal cortical insufficiency, with particular attention to studies from the year 2000 onward (search terms: “adrenal insufficiency” or “Addison’s disease” or “hypopituitarism”).

Results: Hydrocortisone substitution therapy is often given in doses of 10–25 mg/day, timed according to the circadian rhythm. Gastrointestinal and other, febrile infections account for 30–50% of life-threatening adrenocortical crises. Such crises affect 8 of 100 persons with adrenal cortical insufficiency per year and must be treated by the immediate administration of glucocorticoids and fluids. When persons with adrenal cortical insufficiency are acutely ill or are otherwise under unusual stress, they may need additional amounts of hydrocortisone, often in the range of 5–10 mg but occasionally as high as 200 mg. The sustained administration of excessive amounts of steroid can shorten patients’ lives by several years. Inappropriate substitution therapy can cause other major medical conditions, such as metabolic syndrome and osteoporosis.

Conclusion: Important measures for the prevention of adrenocortical crises include improved care by treating physicians, education of patients and their families, the provision of emergency identifying documents, and the prescription of glucocorticoid emergency kits.

Cite this as:

Primary adrenal cortical insufficiency, known as adrenal insufficiency (AI) or Addison’s disease, is rare with a prevalence of approximately 100/1 million/year (1, 2). The incidence of primary AI is approximately 5/1 million/year and has been rising in recent years (1–3) (e1). Autoimmune-mediated adrenalinosis accounts for over 80% of cases in industrialized countries (2). Most patients are young to middle-aged, with more females than males affected. However, this disease affects patients of all ages and in patients under 30 years there is no sex disparity (e2).

Irreversible damage to the adrenal cortex leads to insufficient production of glucocorticoids, mineralocorticoids, and androgens. Over the course of their illness, nearly 60% of patients with autoimmune-mediated AI will be diagnosed with further autoimmune diseases as part of a polyglandular autoimmune syndrome (Table 1).

The secondary form of adrenal cortical insufficiency is caused by a dysfunction at the level of the pituitary (incidence: 20/1 million/year; prevalence 400/1 million). The main causes are the displacement of corticotrophic cells by pituitary macroadenomas or traumatic injury (Table 1) (4, e3). A deficiency of adrenocorticotrophic hormone (ACTH) blocks the stimulation of cortisol production. The adrenal cortex atrophies, and cortisol secretion dwindles. Due to the variety of causes, there is no peak incidence according to age or sex.

By far, the most common cause of adrenal cortical insufficiency is pharmacotherapy with synthetic glucocorticoids (0.5 to 2% of the population) (5). Such therapy can lead to suppression of the hypothalamic-pituitary-adrenal axis (HPA axis) with atrophy of the corticotrophic cells of the pituitary gland and the adrenal cortex (tertiary adrenal cortical insufficiency). High-dose steroid therapy (20–30 mg prednisolone equivalent) can lead to suppression of the regulatory cycle after just a few days (6, e4). In general, the risk of adrenal cortical insufficiency increases with the dose and duration of therapy. Depot preparations and evening administration of higher glucocorticoid doses also increase the risk. Adrenocorticosuppression is difficult to predict in individual cases, hence all patients, even those receiving low-dose glucocorticoid therapy, must be generally considered at risk for the development of adrenal cortical insufficiency (7).
<table>
<thead>
<tr>
<th>Causes</th>
<th>Notes</th>
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<tbody>
<tr>
<td><strong>Primary adrenal cortical insufficiency (AI)</strong></td>
<td></td>
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<tr>
<td>Isolated autoimmune adrenalitis</td>
<td>Autoimmune adrenalitis = most common cause of primary AI in Western countries (&gt;80%), of which 30-40% as isolated disease, 21-Hydroxylase antibody often positive</td>
</tr>
<tr>
<td>Polyglandular autoimmune syndrome Type 1</td>
<td>Hypoparathyroidism, chronic mucocutaneous candidiasis, other autoimmune diseases, lymphomas (rare), mutation in the AIRE gene, autosomal recessive</td>
</tr>
<tr>
<td>Polyglandular autoimmune syndrome Type 2</td>
<td>Hypo/hyperthyroid, premature ovarian failure, vitiligo, type 1 diabetes mellitus, pernicious anemia, association with HLA-DR3 (approximately 60% of patients with autoimmune adrenalitis)</td>
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<tr>
<td>Infections</td>
<td>Tuberculosis (most common cause in developing countries), CMV, HIV, Mycosis (e.g. histoplasmosis)</td>
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<td>Bilateral adrenal hemorrhage</td>
<td>Meningococcal sepsis, primary antiphospholipid syndrome, septic shock</td>
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<td>Extensive adrenal metastases</td>
<td>e.g. renal, lung, breast, gastric, or colon carcinomas, lymphoma</td>
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<td>Bilateral adrenalectomy</td>
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<td>Drugs</td>
<td>e.g. mitotane, etomidate, ketoconazole, fluconazole can cause AI. Rifampicin, phenytoin, barbiturate, carbamazepine accelerate cortisol metabolism</td>
</tr>
<tr>
<td>Adrenogenital syndrome (AGS)</td>
<td>Congenital enzyme defect of steroid biosynthesis (21a-hydroxylase [85%], 11 b-hydroxylase, and others), autosomal recessive, salt deficiency (75%), virilization in girls</td>
</tr>
<tr>
<td>Adrenoleukodystrophy</td>
<td>Neurological disturbances (frequent), hypogonadism, X-linked recessive trait, mutation in the X-ALD gene, accumulation of long chain fatty acids (&gt;C24)</td>
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<tr>
<td>Familial glucocorticoid resistance</td>
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<tr>
<td>Familial glucocorticoid deficiency</td>
<td>Genetic ACTH insensitivity, type 1-3 FGD</td>
</tr>
<tr>
<td>Congenital adrenal hypoplasia</td>
<td>Hypogonadotropic hypogonadism, X-linked mutation in the DAX-1 gene</td>
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<tr>
<td>Triple A syndrome</td>
<td>Achalasia, alacrima, neurological disturbances, autosomal recessive mutation in the triple-A gene</td>
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<td><strong>Secondary adrenal cortical insufficiency</strong></td>
<td></td>
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<tr>
<td>Tumors of the pituitary and hypothalamic regions</td>
<td>E.g., pituitary adenoma, Rathke’s cyst, craniopharyngioma, meningioma, metastases</td>
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<tr>
<td>Pituitary / hypothalamic surgery</td>
<td>—</td>
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<td>Radiation to the pituitary and hypothalamic regions</td>
<td>—</td>
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<tr>
<td>Pituitary infarction / Sheehan syndrome</td>
<td>—</td>
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<tr>
<td>Autoimmune hypophysitis</td>
<td>Lymphocytic, IgG4 associated, drug associated (e.g. ipilimumab, tremelimumab), xanthomatous</td>
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<td>Granulomatous disease</td>
<td>Sarcoidosis, histiocytosis X, Wegener’s granulomatosis</td>
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<tr>
<td>Infections</td>
<td>Abcess, tuberculous meningitis</td>
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<tr>
<td>Traumatic brain injury</td>
<td>—</td>
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<tr>
<td>Genetic causes</td>
<td>E.g. mutations in PROP-1, LHX-4, HESX1, TPIT, POMC genes</td>
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<tr>
<td>Isolated ACTH deficiency</td>
<td>Autoimmune, mutations in PC-1, POMC, or TPIT genes</td>
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<td><strong>Tertiary adrenal cortical Insufficiency</strong></td>
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<td>Chronic glucocorticoid therapy</td>
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<td>Endogenous Cushing’s disease</td>
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<tr>
<td>Isolated CRH deficiency</td>
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CRH, corticotropin releasing hormone; CMV, cytomegalovirus; HIV, human immunodeficiency virus; ACTH, adrenocorticotropic hormone
The current article aims to provide new insight into the management of patients with adrenal cortical insufficiency. A selective literature search of Medline was performed with special focus on recent studies published since the year 2000 (search words: "adrenal insufficiency" or "Addison’s disease" or "hypopituitarism"). The search yielded 451 articles, and relevant articles were selected.

Clinical information and diagnosis
The typical symptoms of adrenal cortical insufficiency are presented in Table 2 (8). In cases of secondary adrenal cortical insufficiency, depending on the underlying condition, there is often complete hypopituitarism with additional signs of growth, sexual, or thyroid hormone dysfunction.

The diagnosis of adrenal cortical insufficiency is made by the combination of low morning blood cortisol level (<100 nmol/L; 3.6 µg/dL) and/or an insufficient cortisol increase to less than 500 nmol/L (<18 µg/dL) after intravenous administration of 250 µg ACTH1–24. The insulin tolerance test (ITT) is considered the gold standard (cortisol increase >500 nmol/L is considered normal); however, its implementation is more arduous. In terms of using basal hormone levels to make a diagnosis, ACTH levels are elevated in primary adrenal cortical insufficiency while levels are low or low-normal in secondary AI (1). The majority of patients with autoimmune-mediated primary adrenal insufficiency show evidence of 21-hydroxylase antibodies in the serum (e5); however, this test can be foregone since a positive result has no therapeutic consequence.

In terms of imaging, the gold standard to diagnose secondary adrenal cortical insufficiency is magnetic resonance imaging of the pituitary and hypothalamus regions with contrast in 2 mm cuts. In addition, an ophthalmologic investigation to rule out chiasmal syndrome is advisable in cases of macroadenoma (>1 cm).

Problems can emerge after long-term pharmacotherapy with glucocorticoids or after treatment for endogenous hypercortisolism, even in cases of apparently sufficient endogenous cortisol production or glucocorticoid replacement. Symptoms mimic those of adrenal cortical insufficiency and are referred to as steroid withdrawal syndrome (e6) (Box 1). The etiology remains unclear; however, relative glucocorticoid resistance mediated through a reduction of glucocorticoid receptors is suspected. The evaluation of endogenous cortisol production is only indicated when test results would have therapeutic consequences, e.g., before discontinuation of therapy or prior to planned surgery. To avoid distorted results, the clinician should ensure a sufficient time interval between testing and the last glucocorticoid administration (at least 4 to 5 plasma half-lives of the glucocorticoid administered, i.e., with hydrocortisone at least 18, and better 24 hours), as well as a sufficiently reduced daily dose (<15 to 20 mg hydrocortisone or <3 to 4 mg prednisolone per day). The tests administered are those listed above.

Hormone replacement therapy
Glucocorticoids
Academic research in the area of glucocorticoid replacement demonstrates a low level of evidence (placebo-controlled trials are impossible because patients die without replacement). However, a great deal of evidence is available from clinical practice.

Physiological cortisol profile and the effects of cortisol—Glucocorticoids have pleomorphic effects on metabolism, including an increase in blood glucose, protein catabolism, activation of bony metabolism with a net reduction of bone mass, and immunomodulatory effects. Acute events such as physical or emotional stress, inflammatory diseases, or injuries can rapidly multiply cortisol levels. Basal secretion is subject to a circadian rhythm, with peak levels in the early morning and a secretion nadir around midnight. These physiologic regulatory mechanisms (9) complicate both the interpretation of basal cortisol values for diagnostic endocrinology and also replacement therapy, which must ideally be adapted to individual circumstances as well as the time of day (10).

History and glucocorticoids used to date for the treatment of adrenal cortical insufficiency—Although Thomas Addison characterized the clinical picture of his namesake disease or primary adrenal cortical insufficiency already in 1855 (e7), it was not until 1936 with the discovery of cortisol by Kendall, Wintersteiner, and Reichenstein and the subsequent chemical synthesis of the hormone that the foundations for replacement therapy were laid.

Current glucocorticoid replacement therapy, dosage, and clinical monitoring—The standard daily replacement dose is between 10 and 25 mg of conventional hydrocortisone (=cortisol). To reproduce the circadian rhythm it is divided into two or three separate doses with approximately two thirds of the entire dose given in the morning (e.g., 10–5–5 0 or 15–5–0 0 mg). Patients with secondary adrenal cortical insufficiency often need slightly lower doses than patients with primary AI. Replacement therapy with prednisolone is also an option. Because of its more extended and powerful action, prednisolone is taken in a single morning dose of 3 to 5 mg, as the biological potency is some six times that of hydrocortisone (e8). The dosing of glucocorticoids as hormone replacement therapy cannot be monitored through hormonal or biochemical parameters. The foundations of therapy supervision are based instead on specific patient history (quality of life, capabilities, inabilities, recurrence and frequency of adrenal crises) and clinical parameters (weight monitoring, development of Cushingoid symptoms, bone density measurements) (4, 10).

Adjustment for stress, illness, and surgery—When adrenal cortical insufficiency is present, the rapid cortisol increase in response to illness and stress occurring physiologically fails to take place. Thus, the replacement dose must be temporarily increased by 10 to 25 mg. The following recommendations are empirically documented: additional intake
of 5 to 10 mg hydrocortisone for prolonged physical activity or increased psychological stress (10). In cases of infection, fever, or minor surgical procedures, the daily dose should be increased to 30 to 75 mg (twice or three times the customary dose) (4, 10). For surgery, childbirth, or intensive treatment, the recommended dose is 100 to 200 mg per day, and for sepsis, 200 to 300 mg per day. During the third trimester of pregnancy, the daily dose should be increased to 25 to 35 mg (11). The dose should also be increased in cases of manifest hyperthyroidism (1). In septic ICU patients with previously healthy adrenals, a hydrocortisone dose (initial 100 mg, followed by 10 mg/hr for at least 7 days) is recommended on the assumption of a relative cortisol deficiency only in cases of septic shock refractory to volume or catecholamine administration (12) (e9).

Interaction with other medications/therapies: By affecting the key enzyme of cortisol metabolism, CYP3A4, various medications and foods can influence hydrocortisone effects by augmenting (e.g., ritonavir, diltiazem, fluoxetine, grapefruit, licorice) or diminishing (e.g., antiepileptics, barbiturates, rifampicin, exenatide) its effect (13) (e10). Hydrocortisone can also decrease the anticoagulation effects of coumarin derivatives as well as increase blood levels of cyclosporine (13). In patients with type 1 diabetes mellitus, it is important to account for the effects of cortisol on glucose metabolism. To avoid nocturnal hypoglycemia, low-dose administration of hydrocortisone in the evening can be useful (14).

Mineralocorticoid therapy—Mineralocorticoid replacement is needed on a regular basis only in patients with primary adrenal cortical insufficiency. Replacement is with fludrocortisone, given in a single dose between 0.05 and 0.1 mg. Treatment is monitored using blood pressure (target: normal values), electrolytes (target: normalized serum Na and K values), and renin concentration (target: upper normal range) measurement. A reduction of the fludrocortisone dose in cases of hypertension, and an increase in dose during pregnancy and extremely hot weather should be considered (1, 10).

Therapy with dihydroepiandrosterone (DHEA)—In both primary and secondary adrenal cortical insufficiency there is a deficiency of DHEA. DHEA acts indirectly through bioconversion into androgens, and directly with DHEA-mediated neurosteroid and immunomodulatory effects (e11). Clinical studies have shown a positive influence of DHEA on mood, sexuality, and health-related quality of life; however, meta-analysis attributes this only moderate value (15). An oral morning dose of 25 to 50 mg DHEA in women with adrenal cortical insufficiency brings serum androgen levels up into the normal female range (16, 17). Positive effects and clinical signs, e.g., regrowth of secondary hair or improved skin moisture, should only be expected after months of therapy and show high inter-individual variation. Treatment monitoring consists of determining the serum DHEA S-levels and serum androgens. DHEA administration must be decided on an individual basis, and succeeds only when despite optimal adjustment of glucocorticoid replacement a persistent marked limitation of general health or libido is present. The patient should also be informed that this is an off-licence therapy, which is not reimbursed by statutory health insurance companies in Germany.

### Shortcomings of current therapy

#### Morbidity

**Adrenal crisis**—Acute adrenal crisis is a life-threatening condition, which will affect approximately one in 13 patients over the course of each year (18) (for causes, cf. Box 1). Despite established glucocorticoid replacement therapy, adrenal crisis remains one of the most common causes of death among patients with chronic adrenal cortical insufficiency (19). The symptoms of adrenal crisis include fatigue, nausea and vomiting, and hypotension and are thus non-specific, which can lead to misdiagnosis. However, the prognosis of acute adrenal crisis is critically dependent on rapid parenteral administration of hydrocortisone (or other glucocorticoids) by the attending physician.
**Osteoporosis**—High glucocorticoid replacement doses over 25 to 30 mg hydrocortisone per day provoke bony changes, i.e. osteoporosis (20). As doses are lowered (<25 mg hydrocortisone per day), so are the case numbers of osteoporosis (21). Synthetic steroids such as prednisolone, because of their higher potency (e8), appear to have more substantial effects on bone (21).

**Metabolic cardiovascular risk factors**—The physiologic circadian rhythm of cortisol affects fluctuations in glucose tolerance at various times of the day. In secondary adrenal cortical insufficiency, the dose amount of daily glucocorticoid replacement correlates with increased body mass index, high cholesterol and triglyceride levels, and an increased prevalence of diabetes mellitus (22, 23). Hydrocortisone administration after 5 pm induces greater insulin resistance than morning administration (9).

**Quality of Life**—In patients with adrenal cortical insufficiency, quality of life is significantly restricted compared to that of the healthy general population (24). However, this seems not to depend on the type of glucocorticoid taken or the frequency of hydrocortisone dosing (25, 26). Instead, it appears that dose quantity and the non-physiological timing in administration of the glucocorticoid are responsible (27).

**Mortality**

Inadequate steroid adjustment in stressful situations as well as chronic over-dosing lead to a 1.5 to 2-fold increase in mortality (reduced life expectancy in females approximately three years and in males approximately 11 years [19]). Leading causes of death are adrenal crisis (at least 25%), infections, cardiovascular disease, and malignancy (28, 29). There is particular risk for patients diagnosed under the age of 40 and for patients with concomitant type 1 diabetes mellitus (19, 28).

Patients with secondary adrenal cortical insufficiency also show an increased mortality rate, which appears to be produced by cardiovascular events (30–32). Hydrocortisone replacement doses >25 mg in particular lead to increased mortality (33).

**Improvements in current therapy**

**Prevention of adrenal crises**

Not administering corticosteroids in a crisis situation, e.g., out of concern for potential immunosuppression in cases of infection, is medical malpractice. The most important element of crisis prevention and management is the intensive and continuing instruction of patients and the people surrounding them.

Treatment for acute adrenal crisis is immediate replacement of glucocorticoids (100 mg hydrocortisone IV followed by another infusion of 100 to 200 mg hydrocortisone over 24 hours) and fluids. Depending on the triggering event, additional treatment appropriate to the situation, e.g., antibiotic therapy, is also necessary. Immediate initiation of therapy may under no circumstances be delayed while waiting for laboratory results.

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**BOX 1**

Common triggering factors for adrenal crisis and frequency by percent*  
- Gastrointestinal infection (22–33%)
- Other febrile infections (17–24%)
- Surgery (7–16%)
- Intense physical activity (7–8%)
- Psychological stress (4–6%)

*according to (18)

**BOX 2**

Strategies to prevent adrenal crisis*  
- Emergency identification card
- Continuing education of patient and family  
  - Dose adjustment in stressful situations / discussion of typical stress situations (fever, trauma, surgery)  
  - Vomiting and diarrhea as urgent indications for parenteral glucocorticoid administration  
  - Symptoms of acute adrenal insufficiency
- Prescribing a hydrocortisone “emergency kit” (e.g., 100 mg hydrocortisone 21-hemisuccinate as ampoules and glucocorticoid suppositories, e.g., 100 mg prednisolone suppositories)
- Instruction in self-injecting hydrocortisone

See also www.endokrinologie.net/krankheiten-glukokortikoide.php (German website), the section „Nebenniere, Steroide und Hypertonie“ (adrenal, steroids, and hypertension) of the German Society for Endocrinology (DGE)
which emergency instructions and contact details of an experienced endocrinologist are listed (Box 2). A European emergency card is now available (eFigure 2). Each patient should also be equipped with an “emergency kit,” and receive appropriate instructions regarding its use (Box 2) (10, 35).

New medications
Although none of the available glucocorticoid preparations can fully mimic the circadian rhythm of cortisol, there are some new developments.

Delayed release preparations
Hydrocortisone with modified release (5 and 20 mg tablets)—This new form of hydrocortisone replacement has been approved in several European countries (including Germany) since the end of 2012 for the treatment of adrenal cortical insufficiency in adults. It consists of an outer shell with rapid-release hydrocortisone surrounding a core with a delayed-release preparation. It is taken in a single dose daily in the morning. In the randomized controlled licensing study for approval, use of the extended-release form significantly improved quality of life, blood pressure, and metabolic profiles (36).

Prednisone with modified Release (1, 2, and 5 mg tablets)—This delayed-release prednisone tablet is taken evenings around 10 pm and begins to work around 3 am. The preparation is approved for patients with rheumatoid arthritis. In a small, open-label study, this tablet produced improvements in morning fatigue and complaints compared to conventional preparations of prednisolone taken at 8 am by patients with adrenal cortical insufficiency (37).

Hydrocortisone with delayed release—This delayed-action hydrocortisone tablet is also used in the evening and acts in the early morning hours (38, 39); however, it is still in development.

Hydrocortisone pump therapy—The continuous administration of hydrocortisone by a pump significantly improved quality of life for seven patients in a pilot study. A larger study is currently planned in Norway. Through pump therapy, hydrocortisone administration closely approximates the physiological profile (27, 40).

Conflict of interest statement:
All involved authors are participating in a register study sponsored by Viropharma Inc.

Prof. Quinkler works as a consultant for Viropharma Inc. He received reimbursement of conference participation fees, travel and accommodation costs, as well as fees for the preparation of scientific meetings from Viropharma.

Prof. Beuschlein received reimbursement of travel and accommodation costs from Viropharma, and he received fees for the preparation of scientific meetings for Viropharma.

Dr. Hahner works as a consultant for Viropharma Inc. She received reimbursement for travel and accommodation costs, as well as fees for the preparation of scientific meetings for Viropharma.

Dr. Meyer received reimbursement for conference, travel, and accommodation costs from Lilly and Ipsen.

Prof. Schönli received fees for consulting from Viropharma.

Prof. Stalle declares that aside from participating in the sponsored register study, there are no conflicts of interest.

KEY MESSAGES

- Adrenocortical insufficiency is a rare but life-threatening disease with various causes.
- Hydrocortisone is the first choice for glucocorticoid replacement, in which a rigid treatment schedule should be rejected for a flexible day-to-day modification (e.g., ±5 mg hydrocortisone).
- Complications occur because of replacement doses set too low (adrenal crises) or too high (metabolic syndrome, osteoporosis). Monitoring of therapy is performed primarily according to clinical criteria.
- Infectious diseases are the main risk for the development of adrenal crisis, and they must be treated early and vigorously. In cases of diarrhea and vomiting, immediate parenteral administration of 100 mg hydrocortisone is necessary.
- Continuing instruction of patients and relatives is essential. Patients should be supplied with an emergency identification card and an emergency kit.

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REFERENCES


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WICHTIGE ÄRZTLICHE INFORMATION

DIESER PATIENT BRAUCHT TÄGliche STEROID-ERSATZTHERAPIE
Im Falle einer schweren Erkrankung, Unfall, Erbrechen oder Durchfall, müssen sofort Hydrocortison 100mg oder ein anderes Glucocorticoid iv/im und physiologische Kochsalzinfusionen verabreicht werden, um eine lebensbedrohliche Nebennieren-Krise zu vermeiden.
Für weitere Infos: www.endokrinologie.net/krankheiten-glukokortikoide.php

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Für weitere Infos: www.endokrinologie.net/krankheiten-glukokortikoide.php

IMPORTANTE MEDICAL INFORMATION

THIS PATIENT NEEDS DAILY STEROID REPLACEMENT THERAPY
In case of serious illness, trauma, vomiting or diarrhoea, Hydrocortisone 100mg iv/im or equivalent glucocorticoid doses and iv saline infusion must be administered without delay to avoid life-threatening adrenal crisis.
For further info see: www.endokrinologie.net/krankheiten-glukokortikoide.php