Optimising glucocorticoid replacement in adrenal insufficiency

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Abstract
Several different regimens for steroid replacement have been tried, but head to head studies of different treatments need to be more widely undertaken. Once daily treatment with prednisolone gives the most physiological profile, and studies comparing prednisolone with hydrocortisone are now being undertaken. It is likely that prednisolone 4mg daily will be the optimum treatment used.

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It is well known that steroid replacement with glucocorticoids saves lives in patients with adrenal insufficiency. Excess glucocorticoid exposure, as evidenced by patients with Cushing’s syndrome, can also be fatal. There remains an increased mortality associated with adrenal insufficiency despite glucocorticoid replacement therapy with a standardised mortality ratio of >2. The cause of the increased mortality is yet to be definitively elucidated, but may be due to excess steroid exposure or replacement regimens that are uncoupled from the normal physiological cortisol profile. Remarkably, there is no agreement on how we can optimise replacement therapy for either primary or secondary adrenal insufficiency.

Uncoupling glucocorticoid dose timing from the normal circadian cortisol rhythm can be harmful. Human cells have developed circadian clocks that are kept in synchrony by the hypothalamic-pituitary-adrenal (HPA) axis. Cortisol levels rise at around 04:00 hours, under the influence of the central circadian pacemaker in the hypothalamus driven by ACTH. Misalignment between cortisol levels and clock time is one of the factors contributing to the adverse health outcomes seen in shift workers and problems associated with jet lag.

It is not known how closely we need to mimic the endogenous rhythm of cortisol when we replace glucocorticoids in patients with pituitary or adrenal failure. Normally the HPA axis has both an ultradian and a circadian rhythm. The ultradian rhythm of irregular pulses of activity can only be mimicked with a pump. Mimicking the circadian rhythm accurately would require levels of cortisol to rise before waking, and this can be reproduced either with a pump or with a delayed release oral tablet.

The excess replacement that frequently occurs with twice or thrice daily hydrocortisone is associated with an increase in cardiovascular deaths in these patients, especially with escalating doses, and the late peaks in cortisol might be harmful. A delayed release preparation (Chronocort) taken last thing at night has the theoretical advantage of mimicking the early morning 04:00 hours rise in cortisol, which cannot be achieved with other preparations.

Another strategy available in some countries is the use of dual release hydrocortisone (Plenadren) once daily first thing in the morning. This extremely expensive sustained release preparation fails to replicate the pre-awakening rise in cortisol and we know very little about what happens to blood or tissue levels of cortisol overnight. It relies on continuous absorption of hydrocortisone from the gut which can be variable, a problem that must be guarded against in patients with infectious diarrhoea, when glucocorticoids are crucial. The manufacturers state that Plenadren is commonly associated (>1 in 10 cases) with gastrointestinal side effects.

Comparison of twice-daily cortisone acetate with once-daily slow release hydrocortisone in the DREAM study suggests that a once-daily preparation has fewer side effects than twice-daily cortisone acetate. Lower doses of glucocorticoids have fewer side effects than higher doses. In terms of concentration of glucocorticoids in the blood, once-daily prednisolone has been shown to mimic the circadian rhythm better than any other oral glucocorticoid. A double-blind randomised controlled trial comparing hydrocortisone 15 mg in the morning plus 5 mg hydrocortisone at sunset in patients who were fasting for Ramadan found no difference between this and prednisolone 5 mg at dawn and placebo at sunset.

An open-label study has also found no difference in markers of health such as blood pressure and cholesterol, but found increased satisfaction in patients because they only needed to take medication once daily and found this significantly more convenient (p=0.048).

Prednisolone is rapidly absorbed and has a prolonged cellular effect. Whether this theoretical advantage of prednisolone is clinically useful and results in reduced risk of adrenal crisis is unknown. It is clear that previously used doses of prednisolone have been excessive and 3–4 mg once daily is sufficient replacement in most patients. Low-dose prednisolone has been shown to be superior in reducing androgens, 17-hydroxy progesterone and improving growth velocity in patients with congenital adrenal hyperplasia compared with thrice-daily hydrocortisone.

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From a patient's perspective, infusion pumps are less convenient than tablets. Infusion pumps have been used to mimic the circadian rise in morning cortisol. In addition to normalising morning ACTH levels in patients with primary adrenal failure, improvements in physical functioning and vitality domains of the SF-36 have been noted in open-label studies. In double-blind placebo-controlled conditions, when compared with oral hydrocortisone treatment, the subjective health outcome benefits of pumps are no longer detectable. These studies were performed without any pulsatile or ultradian patterns. Ultradian pulses might have subtle effects on cognition, although long-term benefits have not been proven.

**Conclusion**


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**Conflict of interest**

None

**References**


22. Ramadan fasting in patients with corticosterone deficiency treated either by hydrocortisone or by prednisolone. Available at: https://clinicaltrials.gov/ct2/show/NCT03585829