1 Mapping the steroid response to major trauma from injury to recovery: a prospective cohort study

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35 Abstract

36 Context: Survival rates after severe injury are improving, but complication rates and outcomes are variable.

37 Objective: This cohort study addressed the lack of longitudinal data on the steroid response to major trauma and38 during recovery.

39 Design: We undertook a prospective, observational cohort study from time of injury to six months post-injury at
40 a major UK trauma centre and a military rehabilitation unit, studying patients within 24 hours of major trauma
41 (estimated New Injury Severity Score (NISS) >15).

42 **Main outcome measures:** We measured adrenal and gonadal steroids in serum and 24-h urine by mass 43 spectrometry, assessed muscle loss by ultrasound and nitrogen excretion, and recorded clinical outcomes 44 (ventilator days, length of hospital stay, opioid use, incidence of organ dysfunction and sepsis); results were 45 analysed by generalized mixed-effect linear models.

Findings: We screened 996 multiple injured adults, approached 106, and recruited 95 eligible patients; 87 survived. We analysed all male survivors <50 years not treated with steroids (N=60; median age 27 [interquartile range 24-31] years; median NISS 34 [29-44]). Urinary nitrogen excretion and muscle loss peaked after one and six weeks, respectively. Serum testosterone, dehydroepiandrosterone and dehydroepiandrosterone sulfate decreased immediately after trauma and took two, four and more than six months, respectively, to recover; opioid treatment delayed dehydroepiandrosterone recovery in a dose-dependent fashion. Androgens and precursors correlated with SOFA score and probability of sepsis.</p>

53 Conclusion: The catabolic response to severe injury was accompanied by acute and sustained androgen 54 suppression. Whether androgen supplementation improves health outcomes after major trauma requires further 55 investigation.

56 Précis

A cohort study in male survivors of major trauma revealed acute and sustained androgen suppression and protein
catabolism including muscle loss. Serum androgens correlated with probability of sepsis.

59 **INTRODUCTION**

60 Over 5 million people worldwide die each year from serious injury (1), with almost 25% caused by road 61 traffic collisions (RTC) (2). In England alone, there are 5400 trauma deaths and 20,000 severe injuries treated by 62 the National Health Service annually (3). Since 2012, the establishment of 22 trauma centres in England has been 63 accompanied by a 19% improvement in survival odds following injury (4). During this time, the UK also received 64 severely injured military trauma patients from the conflict in Afghanistan (5,6).

Improvements in short-term outcomes have been achieved through early resuscitation and acute care (7), 65 often informed by approaches pioneered on the battlefield. However, improvement in survival is often offset 66 67 during the weeks following acute major trauma by the systemic inflammatory response syndrome (SIRS), which is associated with increased risks of infection, multi-organ dysfunction or failure (MOD/MOF), and death (8.9). 68 Simultaneously, the hypothalamic-pituitary-adrenal axis (HPA) is thought to drive a hypermetabolic and overtly 69 70 catabolic response. Importantly, in this profound catabolic state, patients lose valuable lean muscle and suffer from increased rates of infection and poor wound healing. Moreover, the dynamic nature of this response, especially 71 72 beyond the first few days following injury and during recovery remains poorly described and understood, limiting 73 the evidence base for novel therapeutic interventions. Burn injury also produces an extreme inflammatory and 74 catabolic response after injury, previously targeted by anabolic steroid analogues (10) and beta-blockade (11). The dynamic changes in endogenous glucocorticoids and their influence on adrenal steroid metabolism after severe 75 injury are not well characterized. We know that pro-inflammatory cytokines activate the enzyme 11β-76 77 hydroxysteroid dehydrogenase type (11 β -HSD1) responsible for tissue-specific activation of glucocorticoids 78 through conversion of inactive cortisone to active cortisol (12). However, only scarce data exist on what happens 79 to early sex steroids and their precursors during this catabolic state.

To address these gaps in knowledge, we have undertaken a detailed prospective study of the endocrine and metabolic response to severe injury in military and civilian populations, recruiting patients within 24 hours of major trauma and following up for the six months post-trauma. This was undertaken to identify predictive biomarkers and therapeutic targets as well as to explore the optimal timing for therapeutic interventions that could promote better recovery after severe traumatic injury.

85 MATERIALS AND METHODS

86 Study Design and Protocol

This prospective cohort study was conducted in the Royal Centre for Defense Medicine and the Oueen 87 Elizabeth Hospital Birmingham, a major UK trauma centre and the primary receiving facility for UK military 88 personnel injured abroad. Military and civilian trauma patients with an estimated New Injury Severity Score 89 (NISS) >15 were recruited (13). NISS was used to ensure those with significant extremity trauma but lower ISS 90 were included (14). Patients with significant head injury or pre-injury neoplastic conditions were not eligible. 91 92 None of the patients received etomidate during their treatment. Informed consent was obtained from personal consultees until recovering capacity. The protocol was approved by the NRES Committee South West - Frenchav 93 94 11/SW/0177 and MOD REC 116/Gen/10.

A daily patient review allowed the injured to be clinically phenotyped. Bespoke study management software (Clinical RESearch Tool – CREST) tracked the patient and their clinical data were entered prospectively and used to calculate Acute Physiology and Chronic Health Evaluation II (APACHE II), Sequential Organ Failure Assessment (SOFA), and Simplified Acute Physiology Score (SAPS) (15,16). Sepsis was defined using Bone's criteria of an infection associated with SIRS (17), current at the time of the study.

Details of opioid administration were collected from the electronic health record and prescribing system (PICS) (18) on each of the study patients. The total amount of opioid given during their hospital stay was appropriately weighted and totalled with equivalence to an oral dose of 10mg morphine, adjusting for potency, delivery method and opioid preparation (19).

Blood and urine samples were collected within 24 hours of acute injury and at 3, 5, 10, 14, 21, 28 days and 2, 3, 4 and 6 months during recovery post-trauma. Blood sampling occurred between 0730 and 0900 hrs; serum was then separated and frozen at -80°C for batched analysis. We separately collected morning blood samples from 37 healthy age- and sex-matched controls, to provide a comparator for the steroid data.

108 Assessment of Protein Catabolism

As a surrogate marker for muscle mass, we undertook longitudinal measurements of muscle thickness by 109 110 a well validated method using portable ultrasound as described by Campbell et al. (20,21). Ultrasound measurements were taken from four different muscle sites (biceps brachii, radial forearm, rectus femoris and rectus 111 abdominis) at weekly intervals while in hospital and at 3, 4, 5 and 6 months following discharge. All ultrasound 112 assessments were performed by two trained operators. Measurements were performed three times at each muscle 113 site and the mean of the three measurements was recorded. The dominant arm was favoured for ultrasound 114 115 assessment of muscle mass unless it was missing or unable to be measured where wounds were extensive. This 116 non-invasive method was chosen over others such as creatine (methyl-d3) dilution (D3-creatine) (22) due to the strict dietary requirements for methy-d3 estimation, which was not practical in the context of major trauma. 117 Similarly, we did not undertake MRI measurement of muscle mass due to the risk associated with repeated 118 119 transport of critically ill patients to scanning facilities (23).

Urinary urea excretion was measured and used to estimate Total Urinary Nitrogen (TUN) excretion as
 described by Milner et al. (24) [Estimated Nitrogen Excretion: Urinary Urea Excretion (mmol/l) x 0.028 x 1.25 =
 Total Urinary Nitrogen excretion (g/l)].

123 Steroid Analysis

Serum concentrations of adrenal and gonadal steroids were measured using liquid chromatography-124 125 tandem mass spectrometry (LC-MS/MS) analysis, employing a validated multi-steroid profiling method (25). In 126 brief, serum steroids were extracted via liquid/liquid tert-butyl-methyl-ether (MTBE), evaporated, reconstituted, 127 and analysed by LC-MS/MS for cortisol and cortisone. Serum androgens and androgen precursors (DHEA, 128 androstenedione, testosterone) were measured following oxime derivatization (26,27). Serum DHEA sulfate (DHEAS) was measured following protein precipitation (28,29). Steroid metabolite excretion analysis in 24-h 129 urine samples was carried out by gas chromatography-mass spectrometry (GC-MS) in selected-ion-monitoring 130 131 (SIM) mode, as previously described (30).

Serum concentrations of sex hormone-binding globulin (SHBG) and luteinizing hormone (LH) were
analysed on the Roche Modular System (Roche Diagnostics, Lewes, UK) by two-site sandwich immunoassay
using electrochemiluminesence technology.

135 Statistical Analysis

The raw data were evaluated by analysis of variance (ANOVA). In addition, paired and unpaired Student's
t-test, Chi-Square Analysis and Mann-Whitney tests were used where appropriate.

Generalized linear mixed-effects models (31) were used to examine the change in variables over time. Patients were included in models as random effects to account for repeat measures over time on the same individuals. Time was modelled using restricted cubic splines (32) to allow for flexible relationships (33). Severity scores were modelled as Poisson distributions due to their skewness and non-negative ranges. Plots of predicted average fixed effects with 95% confidence intervals were produced for the first four weeks and first six months post- injury as required. Analyses were conducted in R using libraries lme4, effects, rms and ggplot2.

144 **RESULTS**

145 Patient Recruitment and Clinical Characteristics of the Final Study Cohort

We screened 996 multiply injured adults. The majority of the 889 excluded patients had a NISS ≤ 15 , others had a significant head injury as their major injury component, and two were excluded due to a pre-injury diagnosis of cancer. Of the 102 patients recruited into the study, two withdrew and re-assessment in five revealed an actual NISS ≤ 15 , leaving a study cohort of 95 patients (**Fig. 1A**).

Excluding eight fatalities and seven patients who had received steroid therapy, 80 survivors completed sample collection over 6 months. To minimize confounders, we excluded the small groups of women (n=9) and age-advanced men (n=11), leaving our final study cohort of 60 men <50 years of age (**Fig. 1B**).

A summary of the cohort characteristics is shown in **Fig. 2**. Median age was 27 (interquartile range (IQR) 24-31) years, median NISS was 34 (IQR 29-44), and patient day-1 (=day of major trauma) APACHE II score was 21 (IQR 14-25). Patients remained ventilated on the intensive care unit (ICU) for a median of 9 (IQR 5-16) days. Median length of hospital stay was 36 (IQR 19-56) days. Improvised Explosive Device (IED) (n=33; 55%) and 157 RTC (n=11; 18%) were the most common causes of injury. Twenty-five (42%) patients had at least one septic
158 episode and most occurred in the second week (Fig 2C).

159 Glucocorticoid Biosynthesis and Metabolism After Major Trauma

Serum cortisol concentrations increased slightly after injury, peaking at 408 (IQR 249-511) nmol/L at two 160 weeks (Fig. 3A; Suppl. Fig. 1A) (34). However, concentrations remained within the wide range observed in 161 healthy controls. Serum concentrations of the inactive glucocorticoid metabolite cortisone were lower than normal 162 after injury, and increased slowly over time, but this trend was not significant (p=0.08) (Fig. 3B; Suppl. Fig. 1B) 163 (34). The serum cortisol-to-cortisone ratio, a marker of systemic 11β-HSD activities (Fig. 3C), peaked at two 164 weeks post-injury and returned to normal at around eight weeks. Consistent with these findings, urinary steroid 165 166 metabolite excretion analysis revealed an increase in glucocorticoid metabolite excretion in weeks 2, 4 and 8 after major trauma, alongside changes in steroid metabolite ratios indicative of increased systemic 11B-HSD1 and 167 decreased 11 β -HSD2 activities, as assessed by (5 α -tetrahydrocortisol + tetrahydrocortisol)/tetrahydrocortisone 168 and cortisol- to-cortisone ratio, respectively (Suppl. Fig. 2+3) (34). 169

170 Androgen Biosynthesis and Activation After Major Trauma

171 Serum concentrations of the adrenal androgen precursor dehydroepiandrosterone (DHEA) were very low 172 after injury (p < 0.0001, compared with healthy controls) but recovered to the normal range by three months postinjury (Fig. 3D, Suppl. Fig. 1) (34). In contrast, its sulfate ester, DHEAS, demonstrated sustained suppression; 173 174 median serum DHEAS concentrations did not recover to values within the healthy reference range, even at the end 175 of the 6-month study period (Fig. 3E). Consequently, the serum DHEA-to-DHEAS ratio (Fig. 3F) increased by 176 week 2 compared with controls and failed to return to normal during the 6-month study period. The serum cortisol-177 to-DHEAS ratio (Fig. 3G) increased post-injury, peaking at 2 weeks, followed by a gradual decrease, but without returning to normal by the end of the 6-month study period. 178

Serum concentrations of the androgen precursor androstenedione (**Fig. 3H**) were below the reference range immediately after injury, recovering to the mid reference range at 2 weeks post-injury. Thus, serum androstenedione concentrations recovered much faster than DHEA, suggestive of rapid downstream activation of DHEA to androstenedione. Serum testosterone (**Fig. 3I, Suppl. Fig. 4A+B**) (34) was very low following injury, starting to increase after two weeks, and recovering to the healthy sex- and age-matched reference range approximately eight weeks after injury. This was mirrored by acute suppression of serum LH immediately after injury, followed by recovery to the normal range approximately 2 weeks after injury (**Suppl. Fig. 4C+D**) (34). Serum sex hormone-binding globulin (SHBG) (**Suppl. Fig. 4E+F**) (34) concentrations were subnormal immediately post-injury, but quickly returned to the healthy reference range between injury and day 7.

Consistent with the observed decrease in circulating androgens, 24-h urinary steroid metabolite excretion 189 190 analysis revealed a steep decrease in the major androgen metabolites androsterone and etiocholanolone at 2, 4 and 8 weeks (Suppl. Fig. 4A+B) (34). Similarly, urinary DHEA excretion, representing the sum of unconjugated 191 DHEA and DHEA sulfate, sharply decreased to very low concentrations at 2, 4 and 8 weeks, with a transient 192 193 increase in 16 α -hydroxylation of DHEA at 2 weeks (Suppl. Fig. 4C+D) (34), possibly linked to the systemic decrease in DHEA sulfation (Fig. 3D-F). The overall decrease in androgen production was paralleled by a 194 profound decrease in systemic 5 α -reductase activity (Suppl. Fig. 4E+F) (34), and hence in androgen activation, 195 as 5α -reductase is responsible for converting testosterone to the most potent and rogen 5α -dihydrotestosterone. 196

197

Protein Catabolism After Major Trauma

The 24-hour total urinary nitrogen (TUN) excretion increased immediately after trauma, peaking at 25.0 \pm 16.1 g/day at the end of the first week, returning to below 15.0 g/day by week-4. The mean maximum rate of nitrogen excretion was 33.0 \pm 21.3 g/day (Fig. 4A). The normalization of TUN excretion coincided with the gradual recovery of adrenal and gonadal androgen production (Fig. 4B+C).

The biceps brachii muscle was the most reliable site for ultrasound measurement of muscle thickness; dressings, amputations and other wounds hampered the measurements of the other muscle areas. Changes in biceps brachii muscle thickness followed a U-shaped curve after injury, reaching a nadir at 6 weeks (day-1 after trauma compared with week-6, p=0.001). The mean muscle loss was $22.7\pm12.5\%$ (Fig. 4D). Similar to TUN, muscle thickness recovered alongside gradually increasing adrenal and gonadal androgen production (Fig. 4E+F).

207 Clinical Course of Post-Traumatic Recovery and Serum Androgen Dynamics

The relationship between adrenal and gonadal androgens and the Sequential Organ Failure Score (SOFA) 208 209 and probability of sepsis are illustrated in Fig. 5. During the first four weeks, serum DHEA, DHEAS, and testosterone all correlated with the clinical SOFA score (autocorrelation factor (ACF) = 0.85, 0.90 and -0.79, 210 respectively). The serum concentrations of all three steroids also showed strong associations with the probability 211 212 of sepsis (R=-0.85, 0.85 and -0.97 for serum DHEA, DHEAS and testosterone, respectively). SOFA score and probability of sepsis also correlated strongly with the DHEA: DHEAS ratio (autocorrelation factor (ACF) = -0.94213 214 and -0.96 respectively) and with the serum cortisol-to-DHEAS ratio, negatively for the SOFA score but positively for probability of sepsis (autocorrelation factor (ACF) = -0.81 and 0.89, respectively) (Suppl. Fig. 5) (34). 215

216 Opioid administration and endocrine recovery

To examine whether opioid administration affected endocrine recovery, we modelled the impact of the total cumulative in-patient opioid dose on circulating steroid concentrations during recovery from major trauma. For this purpose, we categorised patients according to cumulative opioid dose. Modelling took into account the differences in ISS, length-of-stay (LOS), ICU LOS, and SOFA score.

The adjusted modelling revealed a dose-dependent impact of opioid treatment, with a higher initial peak of serum cortisol and the cortisol/cortisone ratio in those on higher doses (\geq 3000mg) while those on lower doses had initially lower serum cortisol concentrations but showed better recovery of cortisol and cortisol/cortisone 2 months into the recovery period, with broad interindividual variability in those with high cumulative opioid doses (Fig. 6).

226 Opioid administration showed a pronounced, dose-dependent effect on adrenal and gonadal androgen 227 production, with significantly delayed recovery of serum DHEA and DHEAS in patients on higher opioid doses 228 (p=0.029, p=<0.001 respectively; **Fig. 7**). By contrast, serum testosterone concentrations, which were initially 229 equally suppressed in all cumulative dose groups, showed a much faster recovery in individuals who received 230 higher (\geq 3000mg) total cumulative opioid doses. However, these confidence intervals were large for these model 231 estimates (**Fig. 7**).

232 DISCUSSION

In this study, we have characterized the response of adrenal and gonadal steroids and catabolic metabolism to severe injury, describing the related dynamic changes for six months post-injury. Modelling the data has allowed us to provide a detailed description of the transition from catabolism to anabolism during recovery from severe injury, including investigating the impact of cumulative in-patient opioid dose. Our data are the first to provide detailed adrenal and gonadal steroids beyond the first days after trauma in a large cohort of young patients, with all patients recruited prospectively and steroid analysis carried out by tandem mass spectrometry.

239 As summarized in a recent meta-analysis (35), previous data on serum cortisol after injury are limited to small cohorts derived from elective surgery, rarely followed up for more than two days. In our study, serum 240 241 cortisol quickly returned to normal following slight initial increases after acute trauma. In contrast, serum 242 cortisone remained low for three months post-injury. Our study revealed an initial phase of minor glucocorticoid activation with a transient increase in the serum cortisol-to-cortisone ratio, with changes in urinary glucocorticoid 243 metabolites indicative of increased 11β-HSD1 activity. The cortisol-activating enzyme 11β-HSD1 is the major 244 245 enzyme converting inactive cortisone to cortisol and has been shown to be upregulated systemically and locally in response to inflammation, thereby dampening the inflammatory response (36,37). Skeletal muscle expresses 11β-246 HSD1 (38), Previous studies reported increased 11β-HSD1 activity in an animal model of trauma haemorrhage 247 248 (39). and improved wound healing in mice treated with 11β-HSD1 inhibitors (40). However, human data after 249 trauma are lacking. There is substantial evidence indicating a reduced cortisol clearance in critical illness, due to 250 decreased cortisol inactivation in liver and kidney (41), this mechanism could also be responsible for the slight 251 changes in cortisol and cortisone we observed. This was corroborated by the observed reduction in the urinary 252 cortisol-to-cortisone ratio, which is reflective of 11B-HSD2 activity.

Interestingly, patients on higher opioid doses, showed a higher early peak in cortisol production after trauma, followed by persistently lower circulating cortisol during the recovery period, as compared with patients on lower opioid doses. Previous reports have described suppressive effects of opioids on the HPA axis, though studies in smaller mammals have indicated an acute stimulatory effect of opioid administration on serum cortisol concentrations (42,43). 258 We observed a pronounced and sustained loss of adrenal and gonadal androgen synthesis within the first 24 hours following acute major trauma. The recovery of circulating DHEA and testosterone concentrations took 259 260 two and four months post-injury, respectively, and DHEAS remained pathologically suppressed at the end of the six-month follow-up period. In a mouse model of acute inflammation, sustained suppression of the expression of 261 the DHEA sulfotransferase SULT2A1 and its sulfate donor enzyme, PAPSS2, have been described (44). We 262 reviewed 23 previous studies that measured serum DHEA and DHEAS in critically ill patients (Suppl. Table 1) 263 (45), but most studies followed patients for only a few days and relatively few patients suffered from acute trauma. 264 265 One previous study measured both serum DHEA and DHEAS in 181 patients with septic shock, and 31 patients 266 with acute hip fracture (46). Serum DHEAS was decreased in both groups, while DHEA was increased in sepsis but decreased after trauma. This suggested an inflammation-mediated downregulation of DHEA sulfation after 267 268 trauma, resulting in a dissociation of serum DHEA and DHEAS. In our study, this was also observed, as indicated 269 by a sustained increase in the serum DHEA/DHEAS ratio and persistently low serum DHEAS concentrations.

270 A number of previous studies have described an association of infection and mortality with low circulating 271 DHEAS concentrations and a raised serum cortisol-to-DHEAS ratio in patients with trauma (47-50). In vitro 272 studies have demonstrated that cortisol decreases neutrophil superoxide production, which is counteracted by 273 coincubation with DHEAS (47). Furthermore, we have previously shown that DHEAS, but not DHEA, directly 274 enhances neutrophil superoxide generation; a key mechanism of human bactericidal function via activation of protein kinase C- β , independent of and receptor signalling (51). In the present study, carried out in severely 275 injured men younger than 50 years of age, we observed suppression of both serum DHEA and DHEAS post-injury. 276 277 indicating that the loss of adrenal androgen synthesis is a trauma-related event. Importantly, we showed for the 278 first time that this decrease in circulating adrenal androgen precursors is sustained for several months, and that 279 DHEAS remains low even six months post-injury.

Alongside the decrease in adrenal androgen synthesis, we observed a near complete loss of gonadal testosterone production and pituitary LH secretion immediately after trauma. Both the gradual recovery of adrenal and gonadal androgen production paralleled the decrease in catabolism, as assessed by urinary nitrogen excretion and biceps muscle thickness. The suppression of the hypothalamus-pituitary-gonadal (HPG) axis after severe

injury shown in our study is supported by the literature (52-55). Our prospective, longitudinal data demonstrate 284 that suppression of the HPG axis is of shorter duration than that of the HPA axis. In traumatic brain injury studies, 285 286 a significant proportion of patients go on to develop anterior pituitary dysfunction including secondary 287 hypogonadism (56). However, in our study traumatic brain injury was an exclusion criterion. While limited data from patients with burns and critical illness have suggested a central, hypothalamic-pituitary cause of trauma-288 related hypogonadism (57,58), the evidence prior to our study has been limited. Our data indicated a central cause 289 of suppression to the gonadotrophic axis, with a decrease in both pituitary LH and gonadal testosterone. 290 291 Interestingly, we observed a differential impact of the cumulative opioid dose on adrenal and gonadal androgens, respectively, with a significantly delayed recovery of DHEA and DHEAS, but a trend towards faster recovery of 292 gonadal testosterone synthesis in patients with higher cumulative opioid doses. Previous data on opioid effects on 293 294 adrenal androgen production are very scarce, but our findings with respect to gonadal testosterone biosynthesis 295 contrasted previous studies describing suppressive opioid effects on the HPG axis (42,43).

296 Our study revealed a loss of both adrenal and gonadal androgen production in young and middle-aged 297 men after major trauma. This effect was further enhanced by long-lasting suppression of androgen-activating 298 systemic 5a-reductase activity, as demonstrated by urinary steroid metabolite analysis. Androgens are important in wound healing, erythrocytosis, bone density and muscle mass (59). The catabolic state that occurs following 299 trauma thus presents a significant challenge. The use of androgens to ameliorate catabolism has some precedent. 300 301 as evidenced by the use of the synthetic androgen, oxandrolone, that has some proven benefit in treating burn injury (10). A meta-analysis of 15 Randomised Controlled Trials including 806 burns patients by Li et al, showed 302 significant benefits (P < 0.05) for using oxandrolone, including less net weight loss, lean body mass loss, nitrogen 303 loss, donor-site healing time, and length of stay in the catabolic and rehabilitative phases (60). The use of 304 305 oxandrolone in major trauma was investigated in two intensive care studies but no benefit was demonstrated 306 (61, 62).

The strengths of our study include its prospective nature, narrow age range of the patients, single gender, single site for recruitment and analysis and detailed follow-up over six months as well as the measurement of circulating (and in a smaller cohort also excreted) steroid hormones by state-of-the-art mass spectrometry assays. Analysing a young to middle-aged patient cohort has also reduced the confounding effects of age-related co-morbidities. Another strength is the unique opportunity our study offered for analysis of the opioid effects on endocrine recovery, facilitated by detailed prospective, longitudinal phenotyping with dedicated software.

Our study was limited by the diverse nature of major trauma patients in relation to injury pattern and the 314 315 involvement of military casualties. The timing and number of observations during our study was pragmatic and 316 some statistical comparisons were made using modelled data. While we measured total cortisol and cortisone by tandem mass spectrometry, we did not measure free cortisol or cortisol-binding globulin. We were only able to 317 318 measure urinary steroid excretion in a sub-cohort of patients, as accurate and repeated collection of 24-h urine 319 proved very challenging under ICU conditions. The estimation of nitrogen excretion was pragmatic due to the diverse nature of the patients and we were not able to record nitrogen intake. Ultrasound estimation of muscle 320 321 thickness was performed at four different body sites, but many individuals had limbs missing or extensive wounds that prevented measurements. While imperfect, the longitudinal nature of these measurements allowed us to model 322 323 these changes over time.

In conclusion, in this most detailed and first prospective study of the steroid response to major trauma, we followed the patients from severe injury to six months of recovery, revealing pronounced and sustained decreases in adrenal and gonadal androgen biosynthesis. Recovery of androgen production in the severely injured patients was mirrored by a switch from catabolism to anabolism as reflected by recovery of muscle mass and a decrease in nitrogen loss. Adrenal and gonadal androgens correlated with risk of sepsis. It is tempting to suggest that an anabolic intervention with androgens or androgen precursors could have a beneficial effect on health outcomes during recovery from major trauma. However, this will need to be investigated by future intervention studies.

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- 523 Figure Legends
- **Figure 1.** Consort diagram. (A) recruitment process and (B) subgroup selection for analysis for sixty male
- 525 survivors of severe injury (NISS>15) under 50 years of age who had not been given exogenous steroids were
- 526 analysed.
- 527 Figure 2. Patient Characteristics of the Analysis Cohort. (A) Demographics, (B) Mechanism of Injury and (C)
- the distribution of septic episodes for 60 male survivors from severe injury (NISS>15) under 50 years of age.
- 529 Figure 3. Serum steroids in 60 male survivors of severe injury (NISS>15) under 50 years of age. Serum
- 530 concentrations shown include cortisol (A), cortisone (B), the cortisol-to-cortisone ratio (C), DHEA (D), DHEAS
- 531 (E), the DHEA-to-DHEAS ratio (F), the cortisol-to-DHEAS ratio (G), androstenedione (H), and testosterone (I).
- 532 Data are represented after modelling of the raw data (Suppl. Fig. 1) using a non-linear mixed effects model that

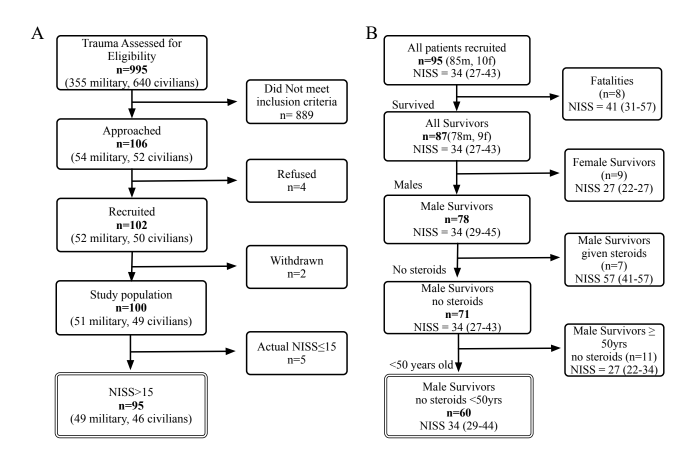
accounts for unbalanced repeated measures using a 4-knot cubic spline. Modelled data are shown as means and
95% confidence intervals.

Figure 4. The relationship between (A) Urinary Nitrogen Excretion or (B) biceps muscle thickness with (B and D) DHEA and (C and F) testosterone, over time for young (<50), severely injured (NISS>15) males who had survived and not been given anabolic steroids. Muscle thickness data was modelled using a mixed effects technique; modelling time as a 6 and 7-knot restricted cubic spline respectively provided the best fit. Data are means and 95% confidence intervals for model-based predicted fixed effects of time are shown.

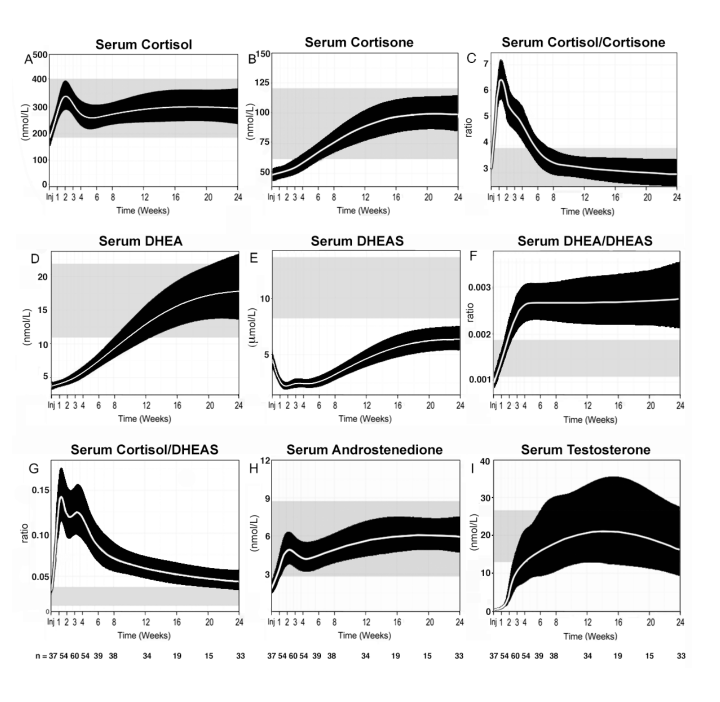
Figure 5. Sequential Organ Failure Assessment (SOFA) score and probability of sepsis in relation to endocrine response. SOFA and sepsis are related serum concentrations of DHEA (Panels A+B), DHEAS (Panels C+D), and testosterone (E+F). Data were modelled using a non-linear mixed effects model that accounts for unbalanced repeated measures using a 4-knot cubic spline. Modelled data are reported as means and 95% confidence intervals.

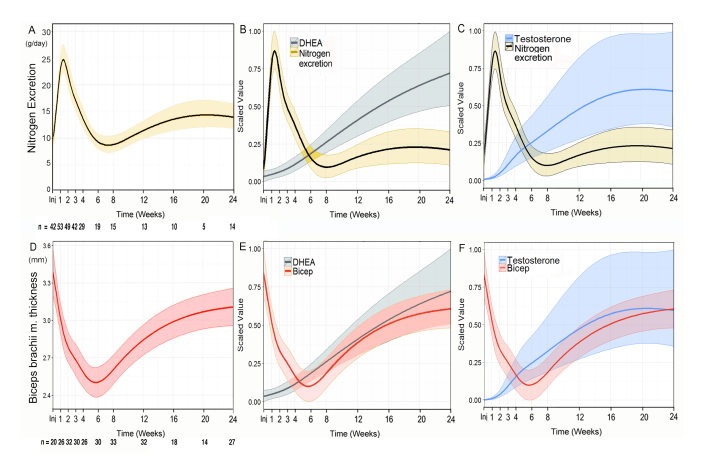
Figure 6. Impact of total inpatient opioid dose on circulating glucocorticoids after major trauma. Serum concentrations are scaled for cortisol, cortisone, the cortisol-to-cortisone ratio and the cortisol-to-DHEAS ratio. Data are represented after modelling of the raw data using a non-linear mixed effects model that accounts for unbalanced repeated measures using a 4-knot cubic spline. Modelled data are shown as means and 95% confidence intervals.

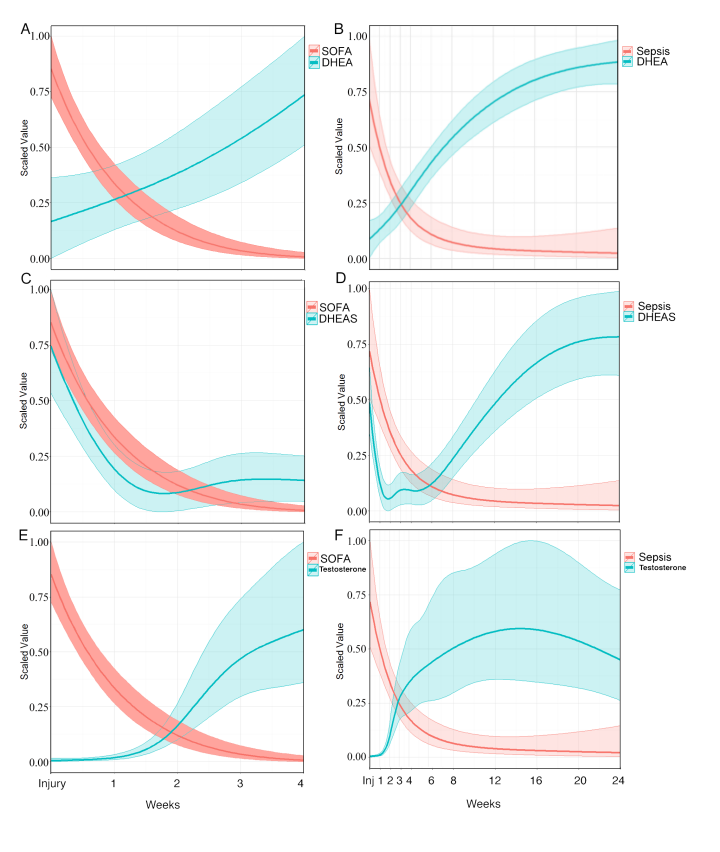
Figure 7. Impact of total inpatient opioid dose on serum androgen and androgen precursors after major trauma. Serum concentrations are scaled for DHEA, DHEAS, the DHEA/ DHEAS ratio, and testosterone. Data are represented after modelling of the raw data using a non-linear mixed effects model that accounts for unbalanced repeated measures using a 4-knot cubic spline. Modelled data are shown as means and 95% confidence intervals.

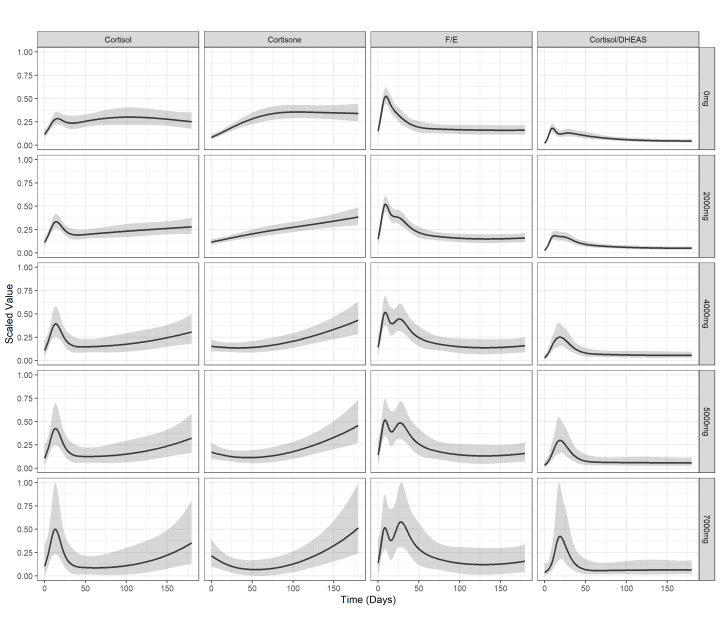


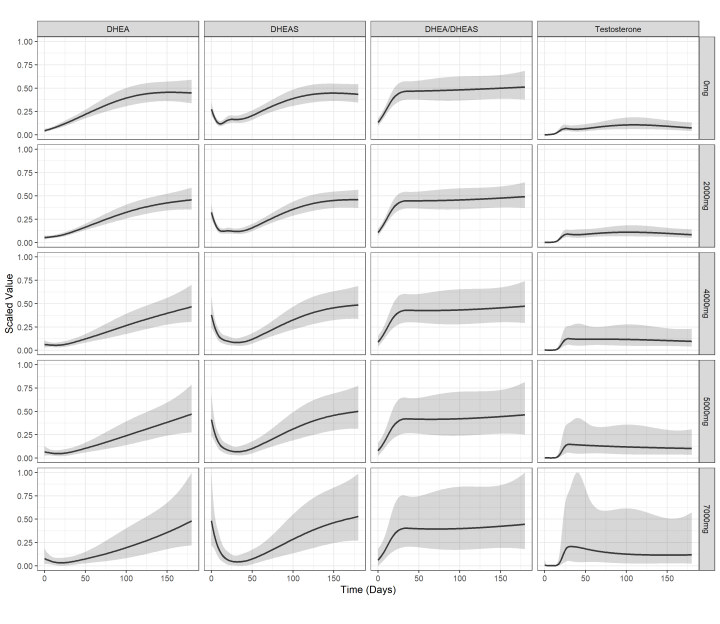
Demographics		B 35-
Patients n=60	Median (IQR)	Ē ³⁰⁻
Age	27 (24-31)	<u>1</u> \$2 25 -
GCS	14 (3-15)	st 25 -
ISS	25 (17-32)	⁶ 15 -
NISS	34 (29-44)	10 -
TRISS	94 (56-98)	5 -
APACHE 2 (Dayl)	21 (14-25)	0 Evaluation BTC CSW Fall Stabilize
SAPS 2 (Dayl)	48 (26-54)	0 Explosion RTC GSW Fall Stabbing Mechanism of Injury
SOFA (Dayl)	9 (6-10)	C
Septic Episodes	1 (0-2)	(Ξ ¹⁵] I
RBCs (Day1), Units	10 (2-17)	sis (
FFP (Day 1), Units	9 (1-15)	æ10- ∎_
Crystalloid/colloid (Day1), L	0 (0-1)	vith
Ventilator Days	9 (5-12)	sti 5 -
ICU Length of Stay, Days	9 (5-16)	Patients with Sepsis (n)
Operative procedures	5 (3-7.5)	
Hospital Length of Stay, Days	36 (19-56)	$0 \frac{1}{48 \text{ hrs} 1 2 3 4 6 8 12 16 20}$ Time (Weeks)

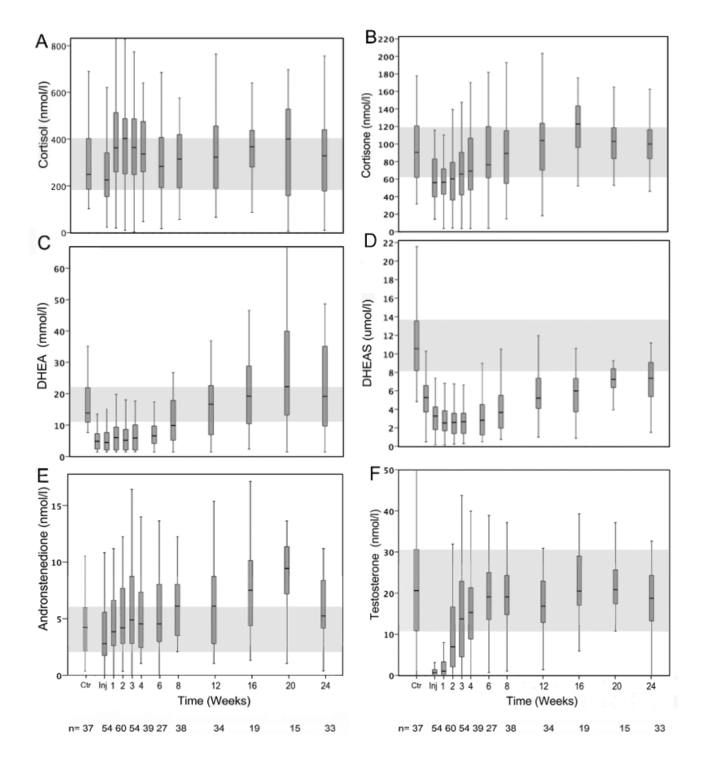


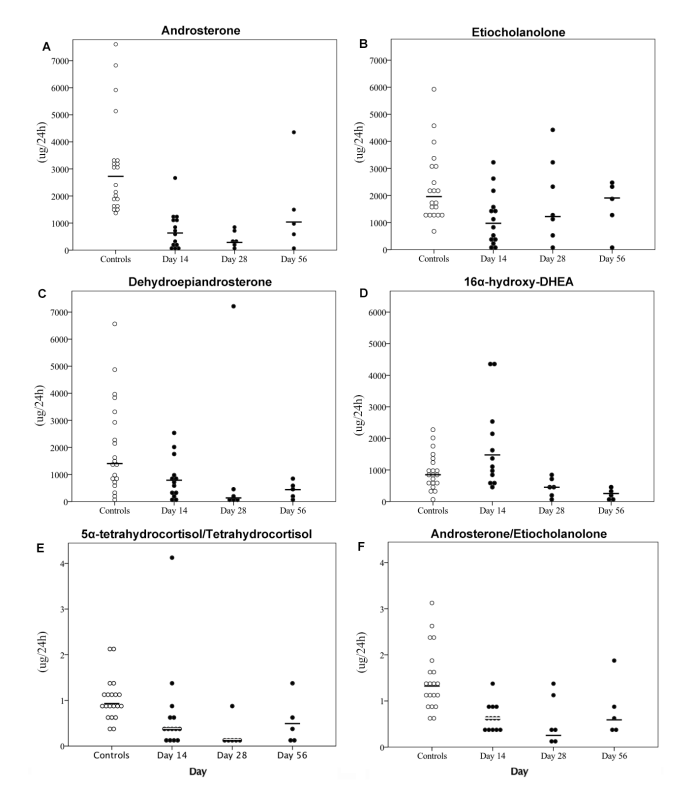


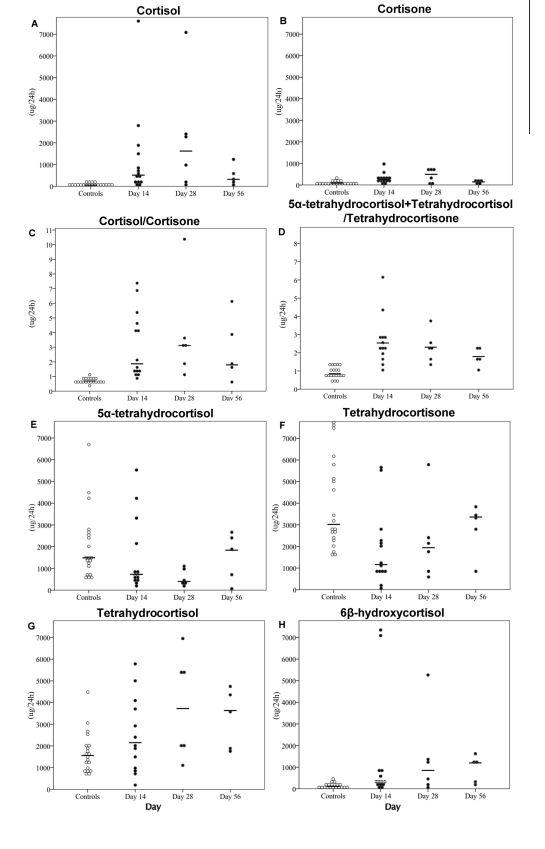


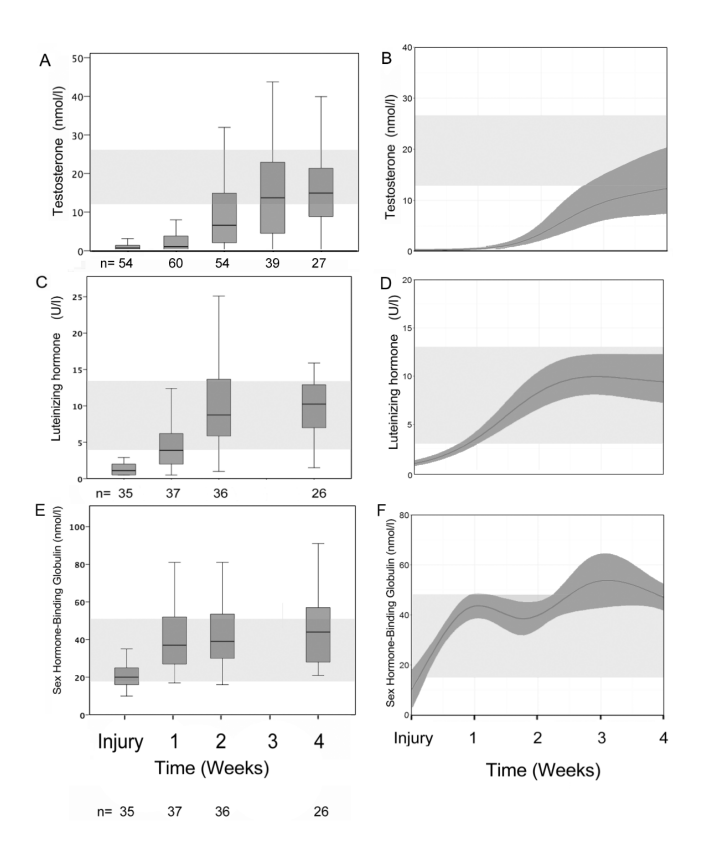


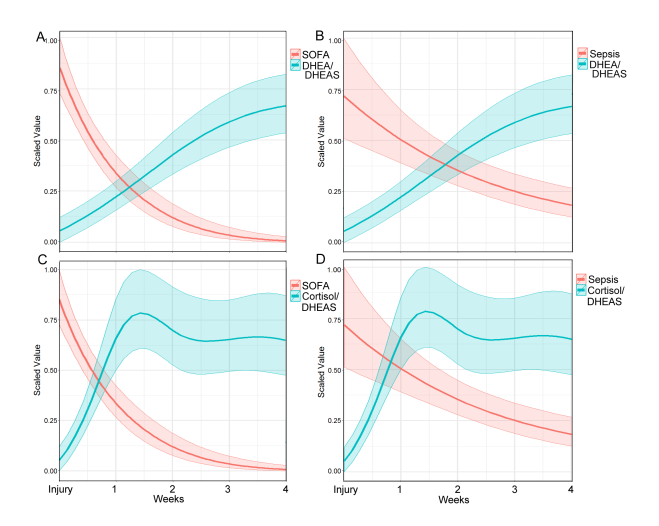












Author	Year	Population	Measurements	Comparison	Outcome
Kolditz (38)	2010	59 patients hospitalized pneumonia	CRP, IL6, TNFa, cortisol baseline and after simulation, ACTH, DHEA, DHEAS on the morning of hospital admission	Clinically stable on day 4 vs clinically unstable on day 4	DHEA/DHEAS ratio was significantly different – higher in stable patients (ratio of 11 vs 7)
Sharshar (39)	2010	103 ventilated ICU patients	IGF, prolactin, TSH, FSH ,LH, estradiol, testosterone, DHEA, DHEAS, cortisol on day 1 when 7 days after wakening	Correlation to SAPSII F vs M	SAPS II assessed at awakening was inversely correlated with plasma levels of DHEA
Klouche(40)	2007	36 ICU patients with critical illness, severe sepsis or septic shock	24h after admission, samples at 0800hrs for cortisol, ACTH, and DHEAS	Elderly vs nonelderly	DHEAS 0.52 nmol/l vs 0.93 in >75 year old vs <75 years old
Chinga-Alayo(41)	2005	113 ICU patients	First hour after admission to ICU – cortisol, thyrotropin, t3, t4, cortisol, prolactin, GH, DHEA	88 Survivors vs 25 non-survivors	No difference in DHEA (86 vs 73 µg/dl)
Butcher(42)	2005	35 elderly and 9 young patients with hip fracture undergoing surgery	serum adrenal stress hormones DHEAS and cortisol immediately following injury	Young vs elderly	DHEAS was higher young trauma group (6.29 \pm 2.41 μ M) compared with elderly hip fracture patients (1.82 \pm 1.58 μ M).
Beishuizen(43)	2002	30 patients with septic shock, 8 with multiple trauma, and 40 controls	serial measurements of DHEAS , cortisol, TNFa, IL-6, and ACTH over 14 days or until death.	Septic shock vs multiple trauma vs controls, Survivors vs non-survivors	DHEAS was lower in septic shock than trauma, both lower control. DHEAS -ve correlation with age, IL-6 and APACHE II scores in both patient groups. DHEAS lower in non-survivors.
Sharshar(44)	2011	102 patients in ICU who regained consciousness after 7 days ventilation	IGF-1, prolactin, TSH, FSH, LH, estradiol, progesterone, test-, DHEA, DHEAS, cortisol 1 st day awake	Survivors vs non- survivors	DHEA and DHEAS were different in men but not in women (lower in non-survivors).
Van den Berghe(45)	2002	33 men long critical illness and age 50, BMI - controls. RCT 5 days of GHRP-2, GHRP2 + TRH infusion or pulsatile	IGF-I, IGFBPs, thyroid hormones, gonadal and adrenal steroids, proinflammatory, metabolic and inflammation markers daily.	Comparison between different therapies	Neither of the foregoing adrenal steroids was altered by any of the study drugs. Replacement requires replacement of somatotrophic, thyrotrophic and gonadotrophic axes to reduce catabolism.
Spratt(46)	1993	postmenopausal women; 20 with acute critical illness vs 110 healthy controls	Day 1-5 DHEA, DHEAS, androstenedione, testosterone, estrogen, gonadotropins, cortisol	Patients vs controls	Admission levels of DHEA and DHEAS were not elevated in patients. Serum DHEA decreased by day 5 in with gonadotropins
Gottschlich(47)	2009	40 patients age 3–18 %TBSA 50.1 were randomly assigned to zolpidem or haloperidol.	2 week study; epinephrine, norepi-, GH, melatonin, DHEA, serotonin, cortisol -0600hrs each study day.	Therapy 1 vs 2	Both drugs were associated with increased DHEA levels (P < .03); no other hormones were affected by medication .

Suppl. Table 1. Longitudinal Studies in critically ill patients analysing DHEA and DHEAS.

Arlt (48)	2006	cross-sectional study consisting of 181 patients with septic shock, 31 patients with acute trauma, and 60 healthy controls.	cortisol, DHEA, and DHEAS were measured before and 60 min after ACTH stimulation.	Septic shock vs acute trauma vs healthy controls	DHEAS lower in septic shock & trauma compared to controls, DHEA was increased in sepsis, decreased in trauma. Sepsis; cortisol and DHEA not increased after ACTH. Most severely ill higher cortisol:DHEA. Cortisol:DHEA increased non-survivors septic shock.
Mueller (49)	2014	179 patients hospitalized with community acquired pneumonia.	DHEA, DHEAS, cortisol	Correlation to PSI score	Correlation between PSI score and DHEAS, cortisol/DHEA, cortisol/DHEAS and DHEA/DHEAS. In age, gender adjusted analysis, DHEA but not DHEAS associated with all-cause mortality.
Ven den Berghe(50)	1995	20 critically ill polytrauma receiving dopamine twere studied to evaluate dopamine withdrawal	DHEAS, prolactin, cortisol during dopamine infusion	Dopamine withdrawal vs continued dopamine	Withdrawal of dopamine 25% increase DHEAS within 24hrs, not DHEAS levels when dopa- continued 2x nights. Prolactin undetectable dopa- infused, increased after 24 hrs of withdrawal.
Dimopoulow (51)	2007	203 severely ill -trauma(93), medical(57), or surgical (53).	24h of admission ICU AM sample to measure cortisol, ACTH and DHEAS.	149 survivors and 54 nonsurvivors	Nonsurvivors had a lower incremental rise in DHEAS (1065 vs. 1642 ng/ml) than survivors.
Osorio(52)	2002	38 patients scheduled for cholecystectomy	DHEAS, ACTH, cortisol, hGH, IGF-1, and IGFBP-3 preop, and then 2 and 7 days after surgery.	Preop to day 2 to day 7 after surgery	Reduction in DHEAS on days 2 and 7 after surgery versus the preop values .
Almoosa(53)	2014	30 men ventilated for >24 hrs for acute respiratory failure.	Blood samples on ICU day 1 and day 3, serum testosterone.	Day 1 and 3 levels, compared to reference	DHEAS levels normal. Total and free testosterone correlated inversely ventilator days and ICU LOS.
Dolecek (54)	2006	29 polytraumas and 28 burned patients (evaluated by their Burn Index, BI) were followed	Bone markers , iPTH, calcium, inorg PO4- , 250H vit. D3, testosterone, DHT, free test- , cortisol, 17β estradiol, DHEAS, TNF α , cytokines.	days 1-7-14-28, of the burned at 1-7-14-28- 56, as well as after 6 and 12 months.	All androgens (T, DHT, FT) decreased significantly in the males, DHEAS decreased in male and females.
Dossett(55)	2008	991 injured patients remaining in the ICU for at least 48 hours	Sex hormones (estradiol, progesterone, testosterone, prolactin, and DHEAS)	Survivors vs non- survivors	Estradiol elevated in nonsurvivors . Estradiol most severely injured had highest. Progesterone, test-, DHEA-S were higher in nonsurvivors; ability to accurately predict death lower than estradiol .
Folan(56)	2001	191 men and women in ICU	DHEA, DHEA-S, and cortisol within 24 hrs of admission and compared with admission APACHE II scores.	APACHE 2 scores and Surgical(SICU) and Medical ICU	The correlations between APACHE II scores and DHEA data for women in the ICU correlation between APACHE II and DHEAS women in SICU.
llias (57)	2007	83 men and 11 women with multiple trauma	TSH, ffT4, T3, ACTH, prolactin, cortisol and DHEAS.	Survivors vs non- survivors	ACTH and DHEAS higher survivors. APACHE II and Cortisol post-Synacthen, DHEAS, TSH*age assessed survival/non-survival better than APACHE II, SOFA or IS scores alone.
Brorsson(58	2014	50 trauma patients admitted to a level-1-trauma centre	Serum & saliva cortisol from injury to five days after trauma. CBG, DHEA & DHEAS twice 5 days after trauma.	Temporal change	A significant decrease over time was observed in DHEA, and DHEAS.
Bergquist (59)	2016	16 adult male patients with burn injury (14.5–72%TBSA)	Plasma cortisol, cortisone, corticosterone, 11-deoxycortisol, DHEA, androstenedione, test-, pregnenolone, , estrone and estradiol and progesterone.	Changes 0, 1, 3, 7, 14, 21 days post burn	Burn injury alters endogenous steroid biosynthesis, reduce testosterone and DHEA levels to 3 weeks and elevated estrone concentrations post injury. Concentrations of glucocorticoids, progestagens and androgen precursors correlated positively TBSA.

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