17α-Ethynyl estradiol-3-sulfate increases survival and hemodynamic functioning in a large animal model of combined traumatic brain injury and hemorrhagic shock: a randomized control trial

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Abstract

**Background:** Traumatic brain injury (TBI) and severe blood loss resulting in hemorrhagic shock (HS) represent leading causes of trauma-induced mortality, especially when co-occurring in pre-hospital settings where standard therapies are not readily available. The primary objective of this study was to determine if 17α-ethynyl estradiol-3-sulfate (EE-3-SO4) increases survival, promotes more rapid cardiovascular recovery, or confers neuroprotection relative to Placebo following TBI + HS.

**Methods:** All methods were approved by required regulatory agencies prior to study initiation. In this fully randomized, blinded preclinical study, eighty (50% females) sexually mature (190.64 ± 21.04 days old; 28.18 ± 2.72 kg) Yucatan swine were used. Sixty-eight animals received a closed-head, accelerative TBI followed by removal of approximately 40% of circulating blood volume. Animals were then intravenously administered EE-3-SO4 formulated in the vehicle at 5.0 mg/mL (dosed at 0.2 mL/kg) or Placebo (0.45% sodium chloride solution) via a continuous pump (0.2 mL/kg over 5 min). Twelve swine were included as uninjured Shams to further characterize model pathology and replicate previous findings. All animals were monitored for up to 5 h in the absence of any other life-saving measures (e.g., mechanical ventilation, fluid resuscitation).

**Results:** A comparison of Placebo-treated relative to Sham animals indicated evidence of acidosis, decreased arterial pressure, increased heart rate, diffuse axonal injury and blood–brain barrier breach. The percentage of animals surviving to 295 min post-injury was significantly higher for the EE-3-SO4 (28/31; 90.3%) relative to Placebo (24/33; 72.7%) cohort. EE-3-SO4 also restored pulse pressure more rapidly post-drug administration, but did not confer any benefits in terms of shock index. Primary blood-based measurements of neuroinflammation and blood brain breach were...
also null, whereas secondary measurements of diffuse axonal injury suggested a more rapid return to baseline for the EE-3-SO₄ group. Survival status was associated with biological sex (female > male), as well as evidence of increased acidosis and neurotrauma independent of EE-3-SO₄ or Placebo administration.

**Conclusions:** EE-3-SO₄ is efficacious in promoting survival and more rapidly restoring cardiovascular homeostasis following polytraumatic injuries in pre-hospital environments (rural and military) in the absence of standard therapies. Poly-therapeutic approaches targeting additional mechanisms (increased hemostasis, oxygen-carrying capacity, etc.) should be considered in future studies.

**Keywords:** Hypovolemia, Brain injuries, Traumatic, Estrogens, Swine, Hemodynamics, Multiple trauma

**Background**

Traumatic brain injury (TBI) and severe blood loss resulting in hemorrhagic shock (HS) individually represent leading causes of trauma-induced mortality and are especially detrimental when combined [1–3]. Hypothermia, acidosis and coagulopathy represent the hallmark complications of HS, ultimately resulting in oxygen debt at the tissue level [4]. Hypovolemia also decreases arterial pressure and increases vasoconstriction, resulting in earlier and more severe cerebral dysautoregulation, reduced blood flow, hypoxia, increased contusion volume and a doubling of mortality rate in concurrent TBI + HS [5, 6]. Although fluid resuscitation is recommended for HS, unless carefully managed it can also exacerbate brain edema and elevate intracranial pressure [7]. Similarly, hyperventilation helps to restore systemic acid–base balance following HS [8], whereas respiratory depression is the most common cause of death in preclinical models of moderate-to-severe TBI [9]. Respiratory complications are also more common in TBI + HS relative to HS alone in humans [10]. Thus, the optimal resuscitation approaches for concurrent TBI + HS remain actively debated [3, 7].

Care is further complicated in remote settings (e.g., injuries occurring in the wilderness, developing countries, or military settings) where resuscitative fluids and blood products are not readily available [11, 12]. Death typically ensues in little more than one hour in the absence of intravenous fluid resuscitation in Class III or IV trauma patients [13]. The family of estrogens (17β-estradiol and 17α-ethinyl estradiol-3-sulfate [EE-3-SO₄]) are naturally occurring steroid hormones that are beneficial in HS [14, 15] and have been shown to be neuroprotective across multiple neural injury models [16, 17]. Specifically, EE-3-SO₄ has been shown to increase short-term survival rate (e.g., 3–6 h) in both rodent [18] and swine [15] models of HS in the absence of typical doses of resuscitation fluids. Proposed mechanisms of action for EE-3-SO₄ include increased cardiac ejection fraction and vasodilation [19–22], increased mitochondrial respiratory complex activity in the myocardium [18, 23], increased cell survival pathways concomitant with decreased cell death pathways, as well as decreased metabolic acidosis and glucose derangement [15, 21]. All of the above effects are dependent on estrogen receptor engagement, where specificity was recently confirmed by estrogen receptor antagonists [20].

Estrogens freely cross the blood–brain barrier (BBB) and have been demonstrated to maintain and regulate the BBB in both humans and rodents [24, 25]. In terms of neuroprotection, seminal work suggests estrogens reduce lesion size and lessen the extent of cell death in the injured brain [17], as well as potentially promoting vascular regeneration following injury [26]. It has been suggested that the estrogen-mediated maintenance of the BBB may also reduce edema after stroke via dampening of the Na–K–Cl cotransporter mechanism [27]. Most pertinent to the current study, EE-3-SO₄ administered one hour following TBI in rodent models resulted in a reduction in intracranial pressure, edema and neuroinflammation while increasing cerebral perfusion pressure and partial pressure of oxygen in brain tissue [17, 28–30], but did not affect markers of diffuse axonal injury [30].

Due to similar homology in terms of hemostatic mechanisms, cardiovascular systems and brain structure (gyrencephalic, similar gray-white matter ratios), swine represent the most commonly utilized species for large animal models of TBI + HS [3]. The majority of these studies have primarily utilized controlled cortical impact or fluid percussion injury, even though closed-head, acceleration injuries represent the most common form of human TBI [31]. A recent study reported acute mortality rates of approximately 88% and 13%, respectively, in an acceleration model of TBI with either 55% or 40% blood loss in the absence of any treatment relative to Shams [32]. In addition to traditional metrics of metabolic derangement associated with HS, results from this study also validated the sensitivity of several blood-based biomarkers for measuring diffuse axonal injury, blood–brain barrier breach and neuroinflammation in swine (glial fibrillary acidic protein [GFAP], neurofilament light chain [NFL], ubiquitin C-terminal hydrolase [UCH-L1], amyloid-beta 40 [Aβ40] and 42 [Aβ42]) that are commonly used in clinical settings [33]. To our knowledge,
there have been no studies examining the efficacy of EE-3-SO$_4$ in a large animal model of TBI + HS.

The current study therefore had two primary aims. The first was to attempt to replicate previous findings of metabolic derangement and neurotrauma in a swine model of closed-head, accelerative TBI + HS (i.e., Placebotreated animals relative to uninjured Shams). The second aim examined the efficacy of EE-3-SO$_4$ to prolong survival in a pre-hospital environment that mimicked more austere levels of care (absence of resuscitative fluids or mechanical ventilation; [32]). Based on previous literature [15, 21], we postulated that EE-3-SO$_4$ would increase survival time and improve hemodynamic functioning, while subsequently decreasing markers of metabolic acidosis, BBB breach and neuroinflammation. Second, we also predicted that there would be a statistically null effect for blood-based and immunohistochemical markers of diffuse axonal injury [30].

Methods
Animal preparation
The methods used in the current study are nearly identical to a previous publication [32] and are therefore only briefly presented here. All animal procedures (see Table 1) were approved by the local Institutional Animal Care and Use Committee (Lovelace Biomedical, FY17-077, FY20-133) and the US Army Medical Research & Development Command Office of Research Protections Animal Care and Use Review Office (DM160115, DM160115.e001) prior to study initiation. The study was conducted in accordance with Animal Research: Reporting In Vivo Experiments 2.0 guidelines [34]. Specifically, eighty sexually mature Yucatan swine (28.18 ± 2.72 kg; 40 females and 40 males; 190.64 ± 21.04 days old at the time of experimental procedures) were obtained from Premier Biosource (formerly S&S Farms, Ramona, California, USA). Animals were screened and vaccinated for common swine diseases by the vendor prior to arrival at the research facility. Upon arrival, animals were examined by a veterinarian and underwent a quarantine (i.e., no experimental procedures) and acclimation period of 7 days, with observation daily prior to the start of experimental procedures. Animals were single-sex group-housed when possible (exceptions made for odd number of animals per sex or behavioral incompatibility) in indoor runs on a 12:12 light/dark cycle. Environmental conditions were maintained between ~16–27 °C and ~30–70% relative humidity. Animals were limit fed (based on age and weight) twice per day and had ad libitum access to water. Animals were randomly assigned in a blocked fashion to the experimental drug (EE-3-SO$_4$ vs. Placebo) and the actual/mock rough ground transport conditions, or to the uninjured Sham cohort. The blocked assignment controlled for biological sex and experimental arm to alleviate concerns about potential time-related effects. All in-life procedures with the exception of rough ground transport, data quality assurance and data scoring were conducted in a blinded fashion, with blind broken immediately prior to conducting the final analyses. Rough ground transport did not significantly affect survival rates ($p > 0.10$) and therefore will be presented in a separate manuscript.

During experimental procedures, animals were maintained under general anesthesia using a combination of isoflurane, midazolam and ketamine (see Additional file 1 for dosing). Femoral arteries were catheterized and flushed every 20 min in conjunction with a single artificial breath. Blood samples were obtained and analyzed using point-of-care devices for primary (glucose, lactate and bicarbonate [HCO$_3$]) and secondary (potential hydrogen [pH], partial CO$_2$ pressure [PCO$_2$], sodium [Na], potassium [K], ionized calcium [iCa]) variables. A Simoa HD-1 Analyzer was used to determine concentrations of NFL, UCH-L1 and GFAP (primary outcome measures), as well as Aβ40 and Aβ42 (secondary outcome measures). Invasive arterial pressure monitoring was used to calculate primary (shock index [SI]; pulse pressure [PP]) and secondary (mean arterial pressure [MAP] and heart rate) hemodynamic variables in 30-s epochs. Focused analyses examined the effects of EE-3-SO$_4$ on hemodynamics at 5 min and 20 min immediately post-administration.

A closed-head TBI was initiated via a pneumatic device targeting a rotation of 250 radians/second in the coronal
plane [32] with a subset of animals monitored for actual head kinematics [35]. Animals were immediately placed in lateral recumbency and subjected to arterial hemorrhage via controlled removal of approximately 40% of estimated total blood volume $161 \pm 48.7$ s after the TBI. Animals were then administered EE-3-SO$_4$ (formulated in the vehicle at $5.0 \text{ mg/mL}$ and dosed at $0.2 \text{ mL/kg}$) or Placebo ($0.45\% \text{ sodium chloride solution}$) as an intravenous administration over $5 \text{ min}$ via a continuous pump ($0.2 \text{ mL/kg}$) and monitored.

Single immunohistochemistry labeling was performed to examine for extravasated serum proteins (Immunoglobulin G; IgG) as markers of blood–brain barrier integrity, axonal pathology (amyloid precursor protein; APP), and upregulation of microglia (ionized calcium-binding adaptor molecule 1; IBA1).

**Statistical plan**

The first series of analyses attempted to replicate previous observations [32] of blood-based biomarker findings of metabolic derangement and neurotrauma in a swine model of closed-head, accelerative TBI+HS (i.e., Placebo vs. Sham). All terminal samples from non-surviving animals were excluded from analyses due to extreme physiological derangement (e.g., $5 \text{ min}$ of apnea) and non-standard data collection times. Generalized linear models (GLM) with appropriate (Gaussian or Gamma) response distributions determined by the model fit or linear mixed-effects (LME) models were utilized for analyses (see Additional file 1). Similar to a previous publication [32], baseline measurements were used as a covariate for all analyses when available.

The second series of analyses examined the efficacy of EE-3-SO$_4$ in promoting survival and improving physiological endpoints (i.e., EE-3-SO$_4$ vs. Placebo). Any animal that did not survive at least $20 \text{ min}$ post-blood loss was excluded from drug-focused analyses. Survival rates between the cohorts were assessed using a Cox proportional-hazards model test. Similar tests (GLM, LME) and methodologies (e.g., baseline as a covariate, terminal samples excluded) were used to investigate differences in physiological markers between EE-3-SO$_4$ and Placebo-treated animals. Any data that were collected with minor variations in protocol were individually reviewed for outlier status (see Additional file 1), with all analyses conducted with and without extreme outliers (results unchanged).

Finally, a series of LME models were fit to determine variables that differed between non-surviving and surviving animals independent of drug assignment. Specifically, surviving animals ($N=16$) were matched to non-surviving animals ($N=16$) based on biological sex, drug assignment and temporal cohort (whenever was possible). The two groups were compared across all primary and secondary variables based on the last successfully acquired timepoint prior to death (timepoint also matched to surviving animals), with matched data eliminated for blood-based biomarkers in the event of death occurring prior to acquisition. Both baseline measurements and acquisition time were entered as additional covariates into the model. The latter controlled for the fact that the temporal course of each biomarker was expected to fluctuate as a function of time post-injury [32]. Due to the exploratory nature of these analyses, individual tests were not corrected for multiple comparisons.

**Results**

**Characterization of model pathology**

No significant (all $p’s > 0.05$) group differences existed at baseline for hemodynamic, point-of-care, or neural biomarkers between Placebo ($N=34$) and uninjured Sham ($N=12$) cohorts. Significant Group × Time interactions were observed for glucose ($F_{4,35.93}=25.45$, $p < 0.001$), lactate ($F_{4,22.21}=6.32$, $p = 0.001$) and HCO$_3$ ($F_{4.23.87}=7.03$, $p = 0.001$) following Bonferroni correction (0.05/3 = 0.017; see Additional file 1: Fig. S1A). Follow-up tests indicated reduced HCO$_3$ (all $p’s < 0.001$; Cohen’s $d = −3.40$ to $−1.68$), increased glucose (all $p’s < 0.01$; $d = 1.38$ to 3.91) and increased lactate (all $p’s < 0.01$; $d = 1.79$ to 2.94) in the Placebo cohort, which all demonstrated evidence of an incomplete recovery trajectory at 295 min post-TBI. Secondary point-of-care (Bonferroni correction at 0.05/5 = 0.01) variables also demonstrated significant Group × Time interactions for Na ($p = 0.001$) and K ($p < 0.001$). Main effects of Group were observed for pH ($p = 0.002$), PCO$_2$ ($p = 0.004$) and iCa ($p < 0.001$), with Time effects presented in Additional file 1.

Significant Group × Time interactions were also observed for the primary hemodynamic variables of PP ($F_{4,34.02}=41.50$, $p < 0.001$) and SI ($F_{4,22.18}=10.72$, $p < 0.001$) following Bonferroni correction (0.05/2 = 0.025). SI was significantly decreased post-TBI for the Placebo relative to Sham cohort ($p = 0.002$, $d = −1.04$), but then increased after blood loss with evidence of an incomplete recovery (all $p’s < 0.001$; $d = 1.56$ to 1.89). In contrast, PP was significantly ($p’s < 0.003$; $d = −5.88$ to $−1.11$) reduced in Placebo animals immediately post-blood loss until 85 min post-TBI, with statistical evidence of full recovery at 145 min post-TBI (see Additional file 1: Fig. S1B). Significant Group × Time interactions (see Additional file 1) were also observed for secondary hemodynamic measurements of HR ($p < 0.001$) and MAP ($p < 0.001$).

Immunohistochemical results indicated significant increases in cortical APP ($\chi^2 = 47.22$, $p < 0.001$, $d = 3.11$), as well as cortical ($\chi^2 = 7.01$, $p = 0.008$, $d = 0.60$) microglial (IBA1) and upregulation of microglia (ionized calcium-binding adaptor molecule 1; IBA1).
analyses based on a priori criteria. There were no statistical differences between remaining EE-3-SO4 (N = 31) and Placebo (N = 33) cohorts on age, body weight or anesthetic time for catheter placement (all p's > 0.05). Comparison of HYGE parameters (Table 2) indicated no significant cohort effects for peak angular velocity, time-to-peak and deceleration time (all p's > 0.05). Similarly, there were no significant differences between the groups in terms of total blood volume removed with the pump, or the amount of time that elapsed between the TBI and the onset of the blood loss procedure (all p's > 0.05). Baseline ionized calcium (iCa) was significantly increased (uncorrected Wald-$\chi^2$ = 4.31, $p = 0.038$) for the EE-3-SO4 (1.41 ± 0.01 mmol/L) relative to Placebo (1.38 ± 0.01 mmol/L) cohort. Otherwise, there were no other significant group differences for all hemodynamic, point-of-care and neural biomarkers between EE-3-SO4 and Placebo groups at baseline (all p's > 0.05; $d = 0.00$ to 0.52).

The Cox proportional-hazards model indicated that the percentage of animals surviving to 295 min post-TBI (i.e., duration of the experiment; Fig. 1) was significantly higher ($\beta = -1.14, Z = -1.70, p_{\text{one-sided}} = 0.044$) for the EE-3-SO4 cohort (28/31; 90.3%) relative to Placebo cohort (24/33; 72.7%). Specifically, there were five deaths in the Placebo cohort relative to 1 death in the EE-3-SO4 cohort by 145 min following TBI, with an almost statistical significance increase in survival rate and time (minutes post-traumatic brain injury [TBI]) for the EE-3-SO4 (solid line) relative to Placebo (dashed line) cohort based on a one-sided Cox proportional-hazards model (Cox PH). The end of the experiment occurred at 295 min post-TBI.

Table 2 Animal characteristics and HYGE parameters

| Measure                  | Placebo (N = 33) | EE-3-SO4 (N = 31) | p       |
|--------------------------|------------------|-------------------|---------|
| Age (days)               | 190.64 ± 22.57   | 186.65 ± 18.72    | 0.433   |
| Weight (kg)              | 28.07 ± 3.22     | 28.12 ± 2.62      | 0.952   |
| reTBV(%)                 | 40.2 ± 1.4       | 40.2 ± 1.5        | 0.970   |
| Peak velocity (rad/s)    | 249.25 ± 6.25    | 248.80 ± 6.35     | 0.777   |
| Decel time (ms)          | 4.07 ± 0.40      | 3.61 ± 0.39       | 0.651   |
| Time to peak (ms)        | 6.15 ± 0.24      | 6.15 ± 0.22       | 0.932   |

Decel time deceleration time, ms millisecond, rad/s radians per second, reTBV removed estimated total blood volume
variables (all $p's > 0.01, d = -0.14$ to $0.23$; Additional file 1: Fig. S2). Results indicated that there were no significant main effects or interactions ($Group \times Time$: see Fig. 2B) between EE-3-SO$_4$ and Placebo cohorts on SI or PP following Bonferroni correction ($0.05/2$ tests; all $p's > 0.025, d = -0.46$ to $-0.15$). Secondary variables of MAP and heart rate were null for all main effects and interactions as well ($d = -0.33$ to $0.17$; Additional file 1: Fig. S3A). Please see Additional file 1 for expected main effects associated with Time.

A drug-focused analysis examining primary hemodynamic measures at 5 min and 20 min immediately

![Box-and-scatter plots](image_url)

**Fig. 2** Box-and-scatter plots depicting primary point-of-care (POC; Panel A) and invasive hemodynamic (Hemo; Panels B and C) markers for EE-3-SO$_4$ (filled circles) and Placebo (unfilled circles) cohorts. Data points are presented at collection times corresponding to Table 1, with POC measurements (glucose, lactate, bicarbonate [HCO$_3$]) occurring at baseline (Base), following drug/Placebo administration (35 m), at hour intervals post-trauma (85 m, 145 m, 205 m), and at the terminal experimental endpoint (Term; ~ 295 min post-traumatic brain injury). Primary hemodynamic markers (shock-index [SI] and pulse-pressure [PP]) were continuously collected over the course of the entire experiment (Panel B), with data points displayed for baseline, immediately post-traumatic brain injury (0 m), immediately post blood loss procedure (25 m), at hour intervals post-trauma (85 m and 145 m), and at the terminal endpoint. Panel C presents a smaller temporal window to capture the predicted rapid effects (5 and 20 min post-drug) of EE-3-SO$_4$ administration (significant Group $\times$ Time interaction). The asterisk denotes significant Group $\times$ Time interaction for PP.
post-drug administration (Fig. 2C) demonstrated a significant Group \( \times \) Time interaction for PP \( (F_{1,62.00} = 6.33, p = 0.014) \). This interaction was driven by a more rapid recovery of PP in the EE-3-SO4 \( (p < 0.001; \) repeated measures \( d = 1.16) \) relative to the Placebo \( (p < 0.001; \) repeated measures \( d = 0.47) \) cohort. No Group effects or interactions were observed for SI values \( (0.05/2 \) tests; all \( p's > 0.025, \) \( d = -0.13) \) or secondary hemodynamic markers of heart rate and MAP \( (0.05/2 \) tests; all \( p's > 0.025, \) \( d = -0.07 \) to \( -0.05) \) cohort.

Immunohistochemical results (Fig. 3A, B) indicated no significant differences for EE-3-SO4 relative to Placebo for either axonal pathology (APP; cortex only) or BBB breach (IgG extravasation) in the cortex or cerebellum following Bonferroni correction \( (0.05/3 \) tests; all \( p's > 0.017, \) \( d = -0.06 \) to \( 0.12) \). Independent \( 2 \times 2 \) (Group \( \times \) Time: 5 min post-injury vs. pre-terminal with skull sensor presence as a covariate) did not demonstrate any Group effects or interactions for primary markers (NFL/UCH-L1/GFAP; Fig. 3C) following Bonferroni correction \( (0.05/3 \) tests; all \( p's > 0.017, \) \( d = -0.14 \) to \( 0.09) \). Secondary blood biomarkers (Aβ40 and Aβ42; Additional file 1: Fig. S3B) indicated a Group \( \times \) Time interaction for Aβ40 \( (F_{1,53.80} = 6.80, p = 0.012) \) that survived Bonferroni correction \( (0.05/2 = 0.025) \), whereas there was no Group effect or interaction for Aβ42 \( (p's > 0.025; \) \( d = -0.25) \). The Aβ40 was characterized by a faster recovery in the EE-3-SO4 \( (p < 0.001; \) repeated measures \( d = 0.54) \) cohort across time relative to the Placebo \( (p < 0.001; \) repeated

![Fig. 3](image-url)
measures \( d = 0.26 \) cohort, partially driven by a higher Aβ40 value immediately post-drug for the EE-3-SO\(_4\) group.

**Secondary analyses**

In terms of overall group composition, results indicated that there was approximately a twofold increase (\( \chi^2 = 8.17, p = 0.004 \)) in the proportion of males in the non-surviving (13/16; 81.3%) versus the surviving group (21/52; 40.4%). There were no differences between the non-surviving and matched surviving samples for other demographics (age and weight) or indices of trauma (peak velocity, time to peak, deceleration time, time between TBI and end of blood loss, and blood loss volume). There were no differences between baseline point-of-care measures, invasive hemodynamic markers, or blood-based biomarkers (all \( p's > 0.05 \)).

All point-of-care (Fig. 4A) and invasive hemodynamic measurement (Fig. 4B) analyses were performed on the last available measurement (prior to moribund criteria being met), and controlled for baseline levels and acquisition time post-injury. Glucose was the sole primary point-of-care variable exhibiting a significant effect, but was higher in surviving relative to non-surviving animals (\( F_{1,20} = 5.76, p = 0.026, d = 0.98 \)), whereas HCO\(_3\) (\( d = 0.80 \)) and lactate (\( d = -0.66 \)) exhibited medium

![Box-and-scatter plots](image)

**Fig. 4** Box-and-scatter plots depicting all primary and significant secondary variables from point-of-care (POC; Panel A), invasive hemodynamic (Hemo; Panel B), and blood protein (Panel C) markers for survival analyses. All data were obtained from the last available, non-terminal timepoint for the non-surviving cohort, and the equivalent time point for each animal’s match (Surviving cohort). Graphed data have been residualized (Rsd.) to account for the effects of initial baseline values and varying measurement acquisition time post-injury. In the case of larger adjustments, this means that negative values are possible. Plots for primary POC measurements (glucose, lactate, bicarbonate [HCO\(_3\)], significant secondary POC measurements (potassium [K], potential hydrogen [pH]), primary hemodynamic measurements (shock-index [SI] and pulse-pressure [PP]), primary blood protein markers (neurofilament light chain [NFL], ubiquitin C-terminal hydrolase [UCH-L1], glial fibrillary acidic protein [GFAP]) and significant secondary protein markers (amyloid-beta 42 [Aβ42]) are displayed. Asterisks denote significant main effects of Survival Status.
to large effect sizes but were non-significant at current sample sizes (both *p*'s > 0.05). Analysis of secondary point-of-care variables indicated significantly lower pH values (*F*$_{1,20}$ = 7.26, *p* = 0.014, *d* = 1.13) and elevated K (*F*$_{1,19}$ = 5.98, *p* = 0.024, *d* = −1.03) for non-surviving animals. No primary or secondary invasive hemodynamic measurement was significant for survival status (all *p*'s > 0.05, *d* = −0.40 to 0.66). NFL values (*F*$_{1,19}$ = 12.72, *p* = 0.002, *d* = −1.46) were significantly higher for non-surviving animals among primary blood protein markers (Fig. 4C). Both GFAP (*d* = −0.72) and UCH-L1 (*d* = −0.74) were not statistically significant, but exhibited medium effect sizes in a similar direction indicating higher pathology (*p*'s > 0.05). Similarly, secondary blood-based biomarkers demonstrated significantly higher levels in non-surviving versus Surviving animals for Aβ42 (*F*$_{1,19}$ = 5.66, *p* = 0.028, *d* = −0.97), with a null Group effect observed for Aβ40 (*p* > 0.05, *d* = −0.60).

**Discussion**

Blood products are not always available in extreme circumstances [1, 36], necessitating the development of novel agents that can both augment the body’s natural response to severe blood loss and mitigate pathological aspects of shock [4]. The current study examined the efficacy of EE-3-SO$_4$ as a treatment for TBI + HS in an austere environment (no mechanical ventilation post-injury, no additional resuscitation fluids, no craniotomy), as frequently occurs in military trauma scenarios and in developing countries [32]. The blocked randomization procedures adequately controlled for all potential major confounders from both demographic variables (non-significant differences in animal age, weight and sex) and experimental (statistically equivalent TBI exposure parameters, pre-injury anesthetic time, blood loss levels, etc.) procedures. Current results replicated previous findings of metabolic derangements, a decrease in MAP in conjunction with increased heart-rate, and both blood-based and immunohistochemical evidence of diffuse axonal injury and blood brain barrier disruption in a large animal model of closed-head accelerative TBI + HS [32].

The administration of EE-3-SO$_4$ increased survival rate, normalized pulse pressure immediately post-drug, and provided preliminary evidence of neuroprotection relative to the Placebo cohort in this fully blinded trial. Previous studies have demonstrated increased survival rates and times for rodent and swine models of HS following intravenous EE-3-SO$_4$ administration [15, 21]. Current findings extend these results to a large animal model of TBI + HS with approximately 40% blood loss, with EE-3-SO$_4$ significantly prolonging survival rate relative to a control cohort (90.3% vs. 72.7%, respectively), albeit at a smaller magnitude relative to previous studies of isolated and severe HS [15]. The mortality rate observed in the Placebo cohort was also roughly commensurate with reported Class III–IV trauma rates [13], providing additional external validity for the closed-head TBI + HS model.

The majority of animals in both groups expired from respiratory distress rather than cardiovascular factors, similar to a previous TBI + HS swine model with 55% blood loss [32]. Acute respiratory failure represents the leading cause of death in preclinical models of isolated TBI [9], complicates clinical care of TBI patients [37], and is more common following TBI + HS relative to HS alone [10]. The current study did not directly quantify the presence of congestion, edema, hemorrhage or microatelectasis in pulmonary tissue as has been done in previous swine models [38], complicating the dissociation of central nervous system involvement in respiratory failure due to TBI. Mechanical ventilation remains the first line of defense for managing acute respiratory distress syndrome in both pre-hospital and hospital setting following complex trauma [39] and is typically used during all phases of preclinical trauma models [3]. However, mechanical ventilation is not available as a treatment option in austere environments to combat respiratory distress, representing a potentially critical factor that should be more carefully considered in future studies for full bench-to-bedside translation.

EE-3-SO$_4$ also more rapidly restored pulse pressure post-administration relative to Placebo, followed by statistically equivalent pressures for the remainder of the experiment. The rapid action of EE-3-SO$_4$ on pulse pressure suggests direct activation of estrogen receptors rather than through genomic signaling [20]. The membrane receptor effects of estrogen include activation of endothelial nitric oxide synthase and the consequent production of nitric oxide, as well as endothelial-independent, rapid mobilization and release of calcium within subcellular compartments leading to increases in Ca$^{2+}$-triggered K$_p$ channel activity [22]. Activation of these receptors collectively results in changes to myocardial contractility and vasodilation of vascular smooth muscle [20]. Over-exuberant vasodilation in the face of severe hypovolemia could be detrimental, but initial vasodilation could also moderate the intense peripheral vasoconstriction seen in TBI + HS, and contribute to the normalization of pulse pressure.

Replication analyses indicated significant post-injury changes in all point-of-care markers of acidosis and other metabolic derangements (glucose, lactate, bicarbonate, etc.), the majority of which did not recover to baseline levels at the end of the 5 h monitoring period. Several of point-of-care markers (pH and potassium)
were significantly more affected in non-surviving relative to surviving animals prior to death, although glucose was unexpectedly higher for surviving animals. Acidosis represents one of the hallmark complications of HS, and non-surviving animals were unable to compensate from a hemodynamic perspective, ultimately resulting in even further increases in oxygen debt at the tissue level [4, 40]. With the exception of glucose, current findings also partially replicate a previous swine model of severe hemorrhage, which reported that increased glucose/potassium/lactate and decreased bicarbonate/MAP were associated with survival [15]. In contrast to previous work in rodents [20], EE-3-SO₄ administration did not significantly affect either point-of-care markers or MAP relative to Placebo, suggesting the need for polytherapeutic approaches to further promote survival and more rapidly restore homeostasis following TBI + HS [7].

Clinical research studies are increasingly using blood-based protein assays to characterize the extent of neurotrauma both in the acute and chronic injury phases of TBI [33]. Previous findings [32] of significant changes in NFL, GFAP and Aβ42 were replicated in our swine models of accelerative TBI + HS, with UCH-L1 and Aβ40 also significant in the current study due to increased statistical power. Several of these blood-based biomarkers demonstrated sensitivity to injury as soon as 35 min post-TBI and were strongly associated with survival, suggesting potential prognostic indications and a portable test for TBI [41]. Similarly, immunohistochemical evidence of diffuse axonal injury (periventricular region only) and blood–brain barrier breach (both periventricular and cerebellar regions) were also present, with previous research suggesting a close coupling between these pathologies [42–44]. In contrast, there were no significant differences between the Placebo and Sham cohorts on an immunohistochemical marker of inflammation (IBA1) following correction for multiple comparisons. The lack a neuroinflammatory response most likely reflects the relatively brief, 5-h post-injury monitoring period employed in the current study, as neuroinflammation has been shown to be present for multiple years post-injury following TBI [45].

Estrogen sulfate has been shown to increase cerebral perfusion pressure, increase partial brain oxygen pressure and decrease intracranial pressure, but not to affect markers of diffuse axonal injury in a previous rodent study [30]. Contrary to our a priori predictions, EE-3-SO₄ showed evidence of normalizing plasma levels of Aβ40 rather than biomarkers traditionally associated with blood brain barrier breach or neuroinflammation. Aβ is a 40–42 amino acid long peptide generated by successive cleavage of amyloid pre-cursor protein by β-secretase followed by γ-secretase [44]. Although Aβ42 is believed to be more toxic, both forms have been shown to be rapidly released post-TBI, persist for weeks to months post-injury, and are typically viewed as potential markers of diffuse axonal injury [46, 47]. Numerous preclinical studies have suggested neuroprotective effects for 17β-estradiol [48], although estradiol is also elevated post-TBI and has been shown to confer an increased risk of death in severe human TBI [49]. However, it remains unknown whether the elevated levels of estradiol post-TBI are due to decreased metabolism (i.e., hydroxylation of estradiol to estrone or increased synthesis due to increased aromatase activity). Although promising, current findings of a more rapid recovery in plasma Aβ40 following EE-3-SO₄ administration require further replication given the lack of efficacy for other female steroidal hormones in clinical TBI trials [50] and current null findings for APP immunohistochemistry.

In the current study, male sex was associated with a nearly twofold increase in mortality rate regardless of drug assignment. There is a rich preclinical literature suggesting that biological sex and associated female endogenous steroidal hormones affect systematic responses to both blunt force and neurotrauma, but with mixed findings in clinical studies [9, 40, 48, 51–53]. Specifically, retrospective clinical studies suggest that female sex may be protective against blunt-force trauma complications such as organ failure and sepsis rather than confer a benefit in terms of mortality [53–55]. Other clinical studies have suggested that only perimenopausal or postmenopausal females demonstrate decreased mortality following isolated moderate to severe TBI [56, 57], whereas pediatric-focused TBI studies indicated increased survival only for post-pubescent females [58, 59]. The latter more closely corresponds to the approximate age of the swine used in the current study and potentially suggests a U shaped relationship between female sex and neuroprotection as a function of age.

There are several limitations to the study that should be noted. First, the current study purposefully did not measure several physiological functions (cerebral perfusion pressure, partial brain oxygen pressure, intracranial pressure, etc.) due to their invasive nature. The study design was intentionally focused on point-of-care and blood based biomarkers that can readily be performed in humans relative to more sophisticated immunohistochemical assays, and our aim to examine a more realistic closed head injury (i.e., intact skull). Blood samples and embedded brain tissue from this study will be made available upon request for additional, secondary analyses. Second, all animals received anesthesia throughout the entire protocol in compliance with the approved ethical framework for this study. Although this is unlikely to have influenced drug-related outcomes due to the
fully randomized and blinded design, it may have artificially inflated mortality rates associated with respiratory depression across both cohorts. The selected anesthetic regimen partially mitigated this confounder through utilization of agents that minimize respiratory depression (i.e., midazolam and ketamine) relative to isoflurane, but in doing so also potentially increased neuroprotective effects [60]. Finally, animals were only monitored for up 5 h post-injury in the current study, which limits the conclusions that can be drawn about more long-term therapeutic effects of EE-3-SO4 or long-term pathophysiologic consequences of the TBI + HS model.

Conclusions
In summary, blood products (whole blood, plasma, etc.) represent the treatment of choice for severe blood loss with or without a concomitant TBI, but are not always available [11, 12]. Current results provide additional support for the efficacy of EE-3-SO4 to promote survival following HS and TBI + HS in austere environments in the absence of fluid resuscitation [15, 20, 21], along with additional salutary effects on hemodynamics. Poly-therapeutic approaches that target additional mechanisms (increased hemostasis, oxygen carrying capacity, etc.) for promoting survival to complement the beneficial effects of EE-3-SO4 should be considered in future studies, along with more in-depth characterization of how EE-3-SO4 potentially mitigates neuronal and pulmonary injury.

Abbreviations
Aβ40: Amyloid beta 40; Aβ42: Amyloid beta 42; APP: Amyloid precursor protein; BBB: Blood–brain barrier; EE-3-SO4: Estradiol-3-sulfate; GFAP: Glial fibrillary acidic protein; GLM: General linear models; HCO3: Bicarbonate; HS: Hemorrhagic shock; IBA1: Ionized calcium‑binding adaptor molecule 1; iCa: Ionized calcium; IgG: Immunoglobulin G; K: Potassium; LME: Linear mixed effects; MAP: Mean arterial pressure; Na: Sodium; NFL: Neurofilament light; PCO2: Partial CO2 pressure; pH: Potential hydrogen; PP: Pulse pressure; SI: Shock index; TBI: Traumatic brain injury; UCH-L1: Ubiquitin C-terminal hydrolase L1

Supplementary Information
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Additional file 1. Supplemental Materials.

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Authors’ contributions
CM, MV, AG, AL, JG, EE, CL, MVG, DS, VS, CRB and SPR analyzed data. AD drafted the tables and figures. AM, DS, VS, AD, RK and IC interpreted the data and drafted the manuscript. AM designed the study protocol and supervised the study. JL, RD, JRL, AM and AD collected data. RK collected and analyzed data. All authors read and approved the final manuscript.

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Availability of data and materials
The datasets used and/or analyzed during the current study are available from the corresponding author on request.

Declarations
Ethics approval and consent to participate
All animal procedures (see Table 1 for summary) were approved by the local Institutional Animal Care and Use Committee (Lovelace Biomedical, FY17-077, FY20-133) and the U.S. Army Medical Research & Development Command Office of Research Protocols Animal Care and Use Review Office (DM160115, DM160115.e001) prior to study initiation.

Consent for publication
Not applicable.

Competing interests
The authors have no conflicts of interest to disclose.

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