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Traumatic injury and exposure to mitochondrialderived damage associated molecular patterns suppresses neutrophil extracellular trap formation

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DOI: 10.3389/fimmu.2019.00685

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Document Version Peer reviewed version

Citation for published version (Harvard): Hazeldine, J, Harrison, P, Dinsdale, R & Lord, J 2019, 'Traumatic injury and exposure to mitochondrial-derived damage associated molecular patterns suppresses neutrophil extracellular trap formation', Frontiers in immunology. https://doi.org/10.3389/fimmu.2019.00685

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1	Traumatic Injury and Exposure to Mitochondrial-derived
2	Damage Associated Molecular Patterns Suppresses Neutrophil
3	Extracellular Trap Formation.
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5	
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17	Short running title: Neutrophil dysfunction post-trauma
18	
19	Keywords: Neutrophils, Neutrophil extracellular traps, Trauma, Mitochondrial-derived
20	DAMPs, Immune suppression.
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Abstract

Major traumatic injury induces significant remodelling of the circulating neutrophil pool and loss of bactericidal function. Although a well-described phenomenon, research to date has only analysed blood samples acquired post-hospital admission, and the mechanisms that initiate compromised neutrophil function post-injury are therefore poorly understood. Here, we analysed pre-hospital blood samples acquired from 62 adult trauma patients (mean age 44 years, range 19-95 years) within 1-hour of injury (mean time to sample 39 minutes, range 13-59 minutes). We found an immediate impairment in neutrophil extracellular trap (NET) generation in response to phorbol 12-myristate 13-acetate (PMA) stimulation, which persisted into the acute post-injury phase (4-72 hours). Reduced NET generation was accompanied by reduced reactive oxygen species production, impaired activation of mitogen-activated protein kinases and a reduction in neutrophil glucose uptake and metabolism to lactate. Pre-treating neutrophils from healthy subjects with mitochondrial-derived damage-associated molecular patterns (mtDAMPs), whose circulating levels were significantly increased in our trauma patients, reduced NET generation. This mtDAMP-induced impairment in NET formation was associated with an N-formyl peptide mediated activation of AMP-activated protein kinase (AMPK), a negative regulator of aerobic glycolysis and NET formation. Indeed, activation of AMPK via treatment with the AMP-mimetic AICAR significantly reduced neutrophil lactate production in response to PMA stimulation, a phenomenon that we also observed for neutrophils pre-treated with mtDAMPs. Furthermore, the impairment in NET generation induced by mtDAMPs was partially ameliorated by pre-treating neutrophils with the AMPK inhibitor compound C. Taken together, our data demonstrate an immediate trauma-induced impairment in neutrophil anti-microbial function and identify mtDAMP release as a potential initiator of acute post-injury neutrophil dysfunction.

76 Introduction

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78 Major injury induces significant phenotypic and functional remodelling of the peripheral 79 neutrophil pool, attributable in part to the emergence into circulation of immature granulocytes (IGs) and highly mature neutrophil subsets (Hampson et al. 2017; Hazeldine et 80 al. 2014; Hazeldine et al. 2017; Leliefeld et al. 2016; Pillay et al. 2012). Alongside changes 81 82 in the surface expression of adhesion molecules and chemokine receptors (Johansson et al. 2011; Visser et al. 2011; Visser et al. 2012), immediate and prolonged impairments in 83 phagocytosis (Hampson et al. 2017; Liao et al. 2013), reactive oxygen species (ROS) 84 production (Hampson et al. 2017; Hazeldine et al. 2017) and chemotaxis (Li et al. 2015; 85 Tarlowe et al. 2003) have been reported for neutrophils isolated from critically-injured 86 patients. Whilst understanding of trauma-induced changes in neutrophil intracellular 87 bactericidal function is well developed, few studies have investigated the impact of injury on 88 the extracellular defensive mechanisms of neutrophils and how soon after injury, any 89 90 compromise occurs.

91

92 Comprised of a DNA backbone decorated with granular and cytosol-derived peptides and enzymes, neutrophil extracellular traps (NETs) are an extracellular anti-microbial defence 93 mechanism deployed to prevent pathogen dissemination (Brinkmann and Zychlinsky 2012). 94 Studies that have examined NET generation post-injury have reported an increase in ex vivo 95 96 NET formation by resting neutrophils (Itagaki et al. 2015; Li et al. 2015), a hyperactivity that may reflect their in vivo exposure to high mobility group box-1 (HMGB-1) and interleukin 97 98 (IL)-33, two NET-inducing alarmins whose circulating concentrations are significantly increased post-injury (Cohen et al. 2009; Tadie et al. 2013; Xu et al. 2017). However, in 99 terms of stimulus-induced NET generation, comparable (Itagaki et al. 2015) or reduced 100 101 (Hampson et al. 2017; Li et al. 2015) NET production in response to stimulation with phorbol 12-myristate 13-acetate (PMA) has been reported post- trauma. Of these studies, only 102 one performed quantitative analysis (Hampson et al. 2017), and neither study that reported a 103 post-injury reduction in NET formation investigated the mechanism(s) responsible (Hampson 104 et al. 2017; Li et al. 2015). 105

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Activation of three non-redundant molecular processes underpin PMA-induced NET 107 generation. Produced by the multi-subunit enzyme nicotinamide adenine dinucleotide 108 phosphate (NADPH) oxidase, ROS generation is essential for the process of chromatin 109 decondensation that precedes NET release (Fuchs et al. 2007; Kirchner et al. 2012). 110 Occuring prior to (Hakkim et al. 2011) or following (Keshari et al. 2013) ROS production, 111 activation of the mitogen activated protein kinases p38 and extracellular signal regulated 112 kinase 1/2 (ERK 1/2), triggers NET formation by inhibiting caspase activation and increasing 113 expression of the pro-survival protein Mcl-1, thus promoting NET production over the 114 induction of apoptosis (Hakkim et al. 2011; Keshari et al. 2013). Finally, glycolysis is a 115 fundamental metabolic requirement for PMA-induced NET formation, with the uptake and 116 breakdown of extracellular glucose a necessity for the process of DNA expulsion (Rodriguez-117 Espinosa et al. 2015). 118

119

Trauma-associated tissue damage results in the release into the circulation of damageassociated molecular patterns (DAMPs), a collection of cytosolic, mitochondrial and nuclearderived proteins and DNA (*Cohen et al. 2009; Kaczmarek et al. 2018; Zhang et al. 2010*).
Whilst renowned for their role in immune activation (*Hazeldine et al. 2015; Zhang et al. 2010*), data are emerging that suggests mitochondrial-derived DAMPs (mtDAMPs), which include N-formylated peptides and mitochondrial DNA (mtDNA), possess immune tolerising

properties. For instance, it has been shown *in vitro* that monocytes pre-exposed to mtDNA (Fernandez-Ruiz et al. 2014) and neutrophils pre-treated with whole mtDAMP preparations (Zhang et al. 2010) exhibit impaired cytokine production and calcium mobilisation respectively upon secondary stimulation. Furthermore, a significant reduction in stimulusinduced ROS production and transmigration was reported for neutrophils pre-exposed to bacterial-derived or synthetic N-formylated peptides (Gabl et al. 2018; Kaczmarek et al. 2018; Li et al. 2015), both of which signal through the same formyl peptide receptor (FPR) as mitochondrial-derived formyl peptides. Based on these observations, the concept of mtDAMP-induced tolerance has been coined and proposed to be a potential mechanistic explanation for the state of peripheral neutrophil dysfunction that develops in the aftermath of major trauma (Kaczmarek et al. 2018; Li et al. 2015).

Here, in a prospective observational study of trauma patients, we have performed for the first time a quantitative assessment of NET production during the pre-hospital, ultra-early (≤ 60 minutes) and acute (4-72 hours) post-injury phases, and assessed the impact that major injury has on the molecular processes and signalling pathways that underpin PMA-induced NET generation. Furthermore, based on the emerging concept of mtDAMP-induced tolerance, we have investigated whether pre-exposing neutrophils isolated from healthy subjects to

144 mtDAMPs *in vitro* results in altered NET generation upon secondary stimulation with PMA 145 and the mechanisms involved.

176 Materials and methods

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178 Study design and setting

180 This manuscript presents data acquired from subjects enrolled into the Brain Biomarkers after 181 Trauma Study, an ongoing prospective longitudinal observational study of adult trauma 182 patients conducted at a single Major Trauma Centre site in the UK (University Hospitals 183 Birmingham NHS Foundation Trust, Birmingham). Ethical approval for the study was 184 granted by the North Wales Research Ethics Committee - West (REC reference: 13/WA/0399, Protocol Number: RG_13-164).

186

Patient enrolment began in the pre-hospital setting, where on a 24/7 basis between March 187 2016 and October 2018, emergency care teams acquired blood samples from adult trauma 188 patients (≥ 18 years) with a suspected injury severity score (ISS) ≥ 8 within 1-hour of injury 189 (defined as the time of phone call to emergency services). In the pre-hospital setting, blood 190 samples were not taken from patients who were deemed unlikely to survive transportation to 191 hospital. Post admission, patients were excluded if they were aged <18 years, if pre-hospital 192 blood samples had been acquired >1 hour post-injury and if clinical assessments confirmed 193 either an ISS <8 or a previous diagnosis of neuro-degenerative disease. No patients received 194 195 blood products in the pre-hospital setting.

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197 Capacity and consent

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Due to the nature of injuries sustained, patients were unlikely to provide informed consent for 199 their participation at the time of study enrolment. Consequently, patient recruitment was 200 201 performed under the guidance of the Mental Health Capacity Act 2005 for research in emergency situations and the Declaration of Helsinki. For patients who lacked capacity, an 202 agreement for study participation was sought from a legal consultee (family member or 203 204 clinician not directly involved in the study), with written consent obtained from the patient once they regained capacity. In instances where the patient did not regain capacity, data were 205 retained in accordance with the agreement of the legal consultee. 206 207

208 Blood sampling

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In the pre-hospital environment, peripheral venous blood samples were acquired during the 210 211 intravenous cannulation of patients or by venepuncture. Once taken, blood tubes were stored at room temperature (RT) until arrival at hospital, where analysis began within 1-hour by a 212 single laboratory researcher on a 24/7 basis. Additional blood samples were acquired 4-12 213 and 48-72 hours post-injury. At all three time points, blood samples were collected into BD 214 Vacutainers[®] (BD Biosciences, Oxford, UK) containing ethylenediaminetetraacetic acid, z-215 serum clotting activator or 1/10 volume of 3.2% trisodium citrate. Full blood counts were 216 performed using a Sysmex XN-1000 haematology analyser (Sysmex UK, Milton Keynes, 217 UK) that measures a white cell differential and IGs, which are defined as promyelocytes, 218 myelocytes and metamyelocytes. The analyser uses fluorescence dyes that label intracellular 219 DNA and RNA, with the intensity of the fluorescence signal directly proportional to the 220 nucleic acid content of the cell. Due to their higher RNA content, IGs are discriminated from 221 mature neutrophils via their stronger fluorescence signal. Daily internal quality control 222 223 measurements (XN check, Sysmex UK) and monthly external quality control samples (UKNEQAS, Watford, UK) ensured instrument performance. 224

Sixty-seven adults (mean age 31 years, range 18-80) served as a cohort of healthy controls (HCs). HCs were volunteers who were not taking any regular medication for a diagnosed illness and did not have an acute episode of infection prior to the time of sampling. The recruitment of HCs was carried out in accordance with the ethical approval granted by the University of Birmingham Research Ethics Committee (Ref: ERN_12-1184) with written informed consent from all subjects. All subjects gave written informed consent in accordance with the Declaration of Helsinki.

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234 Preparation of mtDAMPs and mtDNA

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MtDNA and mtDAMPs were prepared from mitochondria isolated from the K562 tumour cell line (ATCC[®], Teddington, Middlesex, UK) as described previously (*Hazeldine et al.* 2015). MtDNA concentration and protein content within mtDAMPs were determined by spectrophotometry (Nanodrop 2000; Thermo Fisher Scientific, Paisley, UK) and preparations stored at -80°C prior to use.

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242 Neutrophil isolation and treatment

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Neutrophils were isolated by Percoll density gradient centrifugation (Scientific Lab Supplies, 244 Nottingham, UK) with cell purity, which was routinely \geq 99%, determined using a Sysmex 245 XN-1000 haematology analyser. Neutrophils were re-suspended at concentrations of 246 1-10x10⁶ /ml in phenol red free or phenol red containing RPMI-1640 media supplemented 247 248 with 2 mM L-glutamine, 100 U/ml penicillin and 100 µg/ml streptomycin (GPS; Sigma-Aldrich, Dorset, UK), phenol red free or phenol red containing RPMI-1640 media 249 supplemented with GPS and 10% heat-inactivated fetal calf serum (HI-FCS; hereafter 250 251 referred to as complete medium (CM); Sigma-Aldrich), glucose free RPMI-1640 media supplemented with GPS (Gibco, Fisher Scientific UK Ltd, Loughborough, UK), Hank's 252 balanced salt solution (HBSS) supplemented with calcium and magnesium (hereafter referred 253 to as HBSS^{+/+}; Gibco, Life Technologies, Cheshire, UK) or HEPES buffer containing 1 mM 254 Ca^{2+} . 255

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For mtDAMPs and mtDNA experiments, neutrophils were pre-treated for 15 minutes 257 (37°C/5% CO₂) with 40 or 100 µg/ml mtDAMPs or mtDNA prior to secondary stimulation. 258 Prior to inclusion in transmigration, lactate and ROS assays, neutrophils were pelleted, 259 supernatants removed and cells resuspended in specified media. To inhibit FPR-1 signalling, 260 261 neutrophils were treated for 60 minutes (37°C/5% CO₂) with 2.5 µM cyclosporin H (CsH; Abcam, Cambridge, UK) or vehicle control, prior to mtDAMP stimulation. For compound C 262 experiments, neutrophils were treated for 60 minutes with 200 µM compound C (Sigma-263 Aldrich) or vehicle control prior to mtDAMP and PMA treatment. To inhibit calcium-264 calmodulin-dependent protein kinase kinases (CaMKKs), neutrophils were incubated for 60 265 minutes (37°C/5% CO₂) with 2.5 µM STO-609 (Sigma-Aldrich) or vehicle control prior to 266 mtDAMP treatment. To induce AMP-activated protein kinase (AMPK) signalling, 267 neutrophils were treated for 60 minutes with 1 mM AICAR (Sigma-Aldrich) prior to PMA 268 stimulation. 269

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271 Ex vivo NET formation

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Neutrophils (2 x 10^5 in phenol red free or phenol red containing RPMI + GPS or glucose free RPMI-1640 media supplemented with GPS) were stimulated with 25 nM PMA (Sigma-Aldrich) for 3 hours at $37^{\circ}C/5\%$ CO₂. Post-stimulation, supernatants were collected and

centrifuged at 2,200 x g for 10 minutes at 4°C, after which the DNA content of cell-free 276 supernatants was analysed. Briefly, 100 µl aliquots of cell-free supernatant were incubated 277 with 1 µM SYTOX Green dye (Life Technologies) for 10 minutes at RT. Fluorescence was 278 279 measured using a BioTek Synergy 2 fluorometric plate reader (NorthStar Scientific Ltd, Sandy, UK) with excitation and emission set at 485 nm and 528 nm respectively. In our 280 trauma-based studies, DNA quantification was performed using a λ -DNA standard curve 281 282 (Fisher Scientific) with PMA-induced NET generation presented as DNA concentration after subtracting the readings obtained from untreated controls. For mtDAMP experiments, 283 background fluorescence values acquired from SYTOX Green staining of mtDAMPs in the 284 285 absence of neutrophils were subtracted from test readings, with NET production expressed as a fold increase above untreated controls. 286

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288 Visualisation of NETs by fluorescence microscopy

 2×10^5 neutrophils in phenol red free or phenol red containing RPMI + GPS or glucose free 290 RPMI-1640 media supplemented with GPS were seeded onto glass coverslips and incubated 291 for 30 minutes at 37°C/5% CO₂ to allow for cell adherence. Following a 3-hour stimulation 292 with 25 nM PMA (37°C, 5% CO₂), samples were fixed for 30 minutes with 4% 293 paraformaldehyde (37°C, 5% CO₂), washed three times in phosphate buffered saline (PBS) 294 and permeabilised with 0.1% Triton X-100 (Sigma-Aldrich). DNA was then stained with 295 296 1 µM SYTOX Green dye for 5 minutes, after which slides were washed once in PBS, mounted in fluoromount medium and visualized using a LEICA DMI 6000 B microscope 297 298 (LEICA, Milton Keynes, UK) at x20 or x40 objective.

- 299300 **ROS production**
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For *ex vivo* analysis of neutrophils isolated from trauma patients, ROS generation was 302 assessed by lucigenin-amplified chemiluminescence. The effect of mtDAMP pre-treatment 303 on ROS production was examined using luminol-amplified chemiluminescence. In both 304 instances, 100 μ l aliquots of neutrophils (1x10⁶/ml in HBSS^{+/+}) were dispensed into wells of 305 a 96-well white-bottomed flat plate (BD Biosciences), pre-coated with PBS/2% BSA, that 306 contained 25 µl of luminol (pH 7.3; final concentration 100 µM; Sigma-Aldrich) or lucigenin 307 (final concentration 200 μ M; Sigma-Aldrich) and 50 μ l HBSS^{+/+}. Neutrophils were then 308 stimulated with 25 nM PMA or vehicle control, after which ROS generation was assessed at 309 1-minute intervals for 180 minutes using a Berthold Centro LB 960 luminometer (Berthold 310 311 Technologies, Hertfordshire, UK). Experiments were performed in quadruplicate, with ROS 312 production measured as relative light units and calculated as area under the curve (AUC).

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314 Measurement of lactate concentration in cell-free culture supernatants

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Neutrophils $(2x10^{6} \text{ in phenol red free RPMI} + \text{GPS})$ were stimulated for 1, 2 or 3 hours (37⁰C, 5% CO₂) with 25 nM PMA or vehicle control. At each time-point, cell-free supernatants were harvested (800 x g, 5 mins, 4^oC) and samples stored at -80^oC prior to analysis. Lactate concentration in 25 µl aliquots of supernatant was determined using a commercially available lactate assay kit according to manufacturer's instructions (Sigma-Aldrich).

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326 Glucose uptake assay

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Following a 15-minute rest period at $37^{\circ}C/5\%$ CO₂, neutrophils (1x10⁶ in RPMI-1640 media without glucose) were stimulated for 60 minutes ($37^{\circ}C/5\%$ CO₂) with 25 nM PMA or vehicle. With 10 minutes of the stimulation period remaining, the fluorescent glucose analogue 2-N-7-nitrobenzen-20xa-1,3-diazol-4-yl amino-2-deoxyglucose (2-NBDG; Thermo Fisher) at a final concentration of 100 μ M was added. Post-incubation, samples were washed and cells re-suspended in glucose free RPMI in preparation for flow cytometric analysis, which was performed on a CyAn_{ADP}TM bench top cytometer (Dako, Cambridgeshire, UK). 10,000 neutrophils were collected and FL1 mean fluorescence intensity values recorded.

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337 Neutrophil transmigration

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Neutrophils $(1x10^7/ml)$ in HEPES buffer containing 1 mM Ca²⁺ were incubated for 30 339 minutes in a 37^oC water bath with 3 µg/ml calcein-acetoxmethyl ester (calcein-AM, Fisher 340 Scientific), after which cells were pelleted, supernatants removed and neutrophils re-341 suspended at 1×10^{7} /ml in phenol red free CM. A total of 1×10^{6} neutrophils were dispensed 342 into the upper chambers of polycarbonate membrane cell culture inserts with 3 µM pores 343 (Corning, New York, USA) that had been pre-loaded into wells of a 24-well flat bottomed 344 plate (BD Biosciences) containing pre-warmed phenol red free CM and 1 nM LTB₄ (R and D 345 Systems, Abingdon, UK). Following a 90-minute incubation at 37^oC, cell culture inserts were 346 removed and plates read immediately for calcein fluorescence using a BioTek Synergy 2 347 348 fluorometric plate reader with excitation and emission set at 485 nm and 528 nm respectively. Fluorescence intensities were converted into neutrophil numbers via the use of a standard 349 350 curve that was generated from calcein-AM loaded neutrophils that had been incubated 351 alongside the test samples in the conditions described above. The number of neutrophils measured in media in which no chemokine was added was subtracted from the numbers 352 calculated for wells that contained 1 ng/ml LTB₄ in order to determine specific chemokine-353 354 mediated migration.

355

356 Assessment of neutrophil phenotype

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Freshly isolated neutrophils ($1x10^5$ in CM) were stimulated with 100 µg/ml mtDAMPs or 358 vehicle control for 15 minutes at 37°C in a humidified 5% CO₂ atmosphere. Post-treatment, 359 samples were stained on ice for 20 minutes with the following mouse anti-human monoclonal 360 361 antibodies or their concentration-matched isotype controls: 2 µg/ml fluorescein isothiocyanate (FITC)-labelled CD62L (clone DREG56; eBioscience, Hatfield, UK); 1 µg/ml 362 CXCR1-FITC (clone eBIO8F1-1-4; eBioscience); 0.5 µg/ml R-phycoerythrin (PE)-labelled 363 364 CXCR2-PE (clone eBio5E8-C7-F10; eBioscience) or 2.5 µg/ml allophycocyanin (APC)labelled CD11b (clone ICRF44, BioLegend, London, UK). Post incubation, cells were 365 pelleted (250 x g, 5 minutes, 4^{0} C), supernatants discarded and neutrophils washed once in 366 PBS/1%BSA. Following resuspension in PBS, samples were transferred to polypropylene 367 FACS tubes for flow cytometric analysis, which was performed on an AccuriC6TM bench top 368 cytometer (BD Biosciences). 10,000 neutrophils, gated according to their forward scatter 369 370 (FS)/sideward scatter (SS) properties, were acquired for analysis, where receptor expression was measured as median fluorescence intensity (MedFI). 371

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- 376 Cell signalling measurements
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To determine signalling through AMPK and MAPK pathways, cell lysates prepared from 378 $2x10^{6}$ resting neutrophils, $1x10^{6}$ neutrophils stimulated with either 25 nM PMA or 100 µg/ml 379 mtDAMPs for 2-90 minutes $(37^{\circ}C/5\% \text{ CO}_2)$, or $2x10^{6}$ neutrophils stimulated with 100 µg/ml 380 mtDAMPs for 5 minutes following 1-hour pre-treatment with 2.5 µM STO-609 or 2.5 µM 381 382 CsH were separated on 10 or 12% SDS-polyacrylamide gels. Following protein transfer to 383 polyvinylidene difluoride membranes (Bio-Rad, Hertfordshire, UK), blots were probed overnight at 4°C with rabbit anti-human antibodies (Cell Signalling Technology, 384 385 Massachusetts, USA) directed against phosphorylated AMPK (pAMPK), phosphorylated ERK1/2 (pERK1/2), phosphorylated P38 (pP38), lactate dehydrogenase A (LDHA) or 386 pyruvate kinase (PKM2). Post incubation, membranes were washed in tris-buffered saline 387 containing 0.001% tween (TBST) and incubated for 1 hour at RT with a goat anti-rabbit 388 secondary antibody conjugated to horse radish peroxidase (HRP; diluted 1:4000 in TBST; GE 389 Healthcare, Buckinghamshire, UK). HRP activity was detected using enhanced 390 chemiluminescence (Bio-Rad). To confirm equal loading of proteins, blots were probed with 391 antibodies against total ERK 1/2, total P38 (1:1000; Cell Signalling Technology) or β-actin 392 (1:5000, GeneTex, California, USA). Densitometry analysis was performed using Image J 393 software (National Institutes of Health, Bethesda, MD, USA). 394

395

396 Enzyme-linked immunosorbent assays (ELISAs)

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Serum was prepared from blood collected into BD vacutainers containing z-serum clotting activator. Following a 30-minute incubation at RT, blood samples were centrifuged at 1,620 x g for 10 minutes at 4^oC, after which serum was aliquoted and stored at -80^oC until analysed. ELISAs to measure serum concentrations of HMGB-1 (IBL International, Hamburg, Germany), mitochondrial encoded NADH dehydrogenase 6 (ND6; MyBioSource, San Diego, California, USA) and IL-33 (R and D Systems) were performed in accordance with manufacturer's instructions.

405

406 Statistical analyses

407

Statistical analyses were performed using GraphPad Prism[®] software (GraphPad Software 408 Ltd, California, USA). Data distribution was examined using the Kolmogorov-Smirnov or 409 Shapiro-Wilk normality test. For data that followed a normal distribution, paired student T-410 411 tests, a repeated measures ANOVA with Bonferroni multiple comparison post hoc test or a one way ANOVA with Dunnett's multiple comparison post hoc test were performed. For 412 non-normally distributed data, a Wilcoxon matched-pairs signed rank test, a Friedman test 413 414 with Dunn's multiple comparison post hoc test or a Kruskal-Wallis with Dunn's multiple comparison post hoc test was performed. For box and whisker plots, whiskers represent 415 minimum and maximum values. Statistical significance was accepted at $p \le 0.05$. 416

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- 426 **Results**
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428 Patient enrolment and demographics429

1,070 adult trauma patients were screened for study inclusion, with 87 subjects enrolled into
the study (Supplementary Figure 1). Of these, 62 patients with a mean age of 44 years (range
19-95 years) and mean injury severity score of 26 (range 9-57) had their immune function
analysed (Table 1). The mean time of pre-hospital blood sampling was 39 minutes post-injury
(range 13-59 minutes).

435

436 PMA-induced NET production is impaired post-trauma437

438 Compared to neutrophils isolated from HCs, neutrophils acquired from trauma patients within 439 1-hour of injury exhibited significantly enhanced basal NET generation (Figure 1A), a 440 hyperactivity that was accompanied by significantly elevated serum concentrations of 441 HMGB-1 (Figure 1B) and IL-33 (Figure 1C). By the 4-12 and 48-72 hour post-injury time 442 points, a significant reduction in basal NET production was observed (Figure 1A). In 443 response to stimulation with PMA, patient neutrophils released significantly less DNA at all 444 three sampling time-points when compared to HCs (Figure 1D). Fluorescence microscopy

- 445 confirmed the impairment in NET generation (Figure 1E).
- 446

ROS production in response to PMA stimulation is reduced in the acute post-injury phase

448 **I** 449

ROS generation is a non-redundant event in NET formation (Fuchs et al. 2007). Having 450 451 observed trauma-induced alterations in both basal and stimulated NET formation, we examined the effect of injury on ROS production. In the absence of stimulation, patient 452 neutrophils isolated 48-72 hours post-injury exhibited significantly enhanced ROS production 453 when compared to the response of neutrophils from HCs (Figure 2A). No difference in basal 454 ROS generation was seen between HCs and patient neutrophils acquired ≤ 1 -hour or 4-12 455 hours post-injury (Figure 2A). In response to PMA stimulation, there was a significant 456 reduction in ROS production, relative to HCs, for neutrophils isolated from patients only at 457 the 48-72 hour post-injury time point (Figure 2B). 458

459

460 Traumatic injury results in an immediate and sustained elevation in the frequency and 461 absolute number of circulating IGs

462
463 Compared to their mature counterparts, immature neutrophils exhibit impaired *ex vivo* NET
464 production and reduced ROS production upon stimulation with inflammatory agonists
465 (*Hampson et al. 2017; Martinelli et al. 2004*). Relative to the values recorded for HCs,
466 trauma patients presented, at all sampling time points, with a significantly elevated frequency
467 (Figure 2C) and absolute number (Figure 2D) of circulating IGs.

468

469 Traumatic injury is associated with impaired MAPK signalling

470

MAPK signalling is a prerequisite for PMA-induced NET production (*Keshari et al. 2013*).
Due to the significant lymphocytosis that occurs within minutes of traumatic injury

472 *(Hazeldine et al. 2017)*, and the small blood volume collected from patients at the scene of

- 474 injury, we were unable to isolate a sufficient number of neutrophils from pre-hospital blood
- 475 samples to examine MAPK signalling. However, we found neutrophils isolated from patients

476 4-12 and 48-72 hours post-injury exhibited significantly increased phosphorylation of P38
477 MAPK (Figure 3A) but not ERK1/2 (Figure 4A) in the absence of exogenous stimulation.

478

479 In response to treatment with PMA, neutrophils obtained from HCs exhibited a significant increase in P38 phosphorylation (Figure 3B-D). In contrast, no significant PMA-induced 480 increase in P38 phosphorylation was observed for neutrophils isolated from trauma patients 481 482 4-12 or 48-72 hours post-injury (Figures 3B-D). Compared to untreated cells, neutrophils isolated from HCs and trauma patients at the 4-12 and 48-72 hour post-injury time points 483 displayed a significant increase in ERK1/2 phosphorylation following 5, 10 and 15 minutes 484 485 of PMA stimulation (Figure 4B-D). However, across these three stimulation time points, the degree of ERK1/2 phosphorylation was significantly greater in neutrophils isolated from HCs 486 (Figure 4B-D) 487

488

489 Altered neutrophil glucose uptake and metabolism post-trauma

490 491 Confirming the results of a recent study that demonstrated a necessity for exogenous glucose 492 in PMA-induced NET production (Rodriguez-Espinosa et al. 2015), we found neutrophils cultured in glucose free media released significantly less DNA upon PMA stimulation than 493 neutrophils stimulated in glucose containing media (Supplementary Figure 2). Based on our 494 observation of a trauma-induced impairment in ex vivo NET generation following PMA 495 496 treatment, we investigated the effect of injury on neutrophil glucose uptake. Using the fluorescent glucose analogue 2-NBDG, enhanced basal glucose uptake was recorded for 497 498 neutrophils isolated from trauma patients within 1-hour of injury (Figure 5A), but in response 499 to PMA stimulation, a significant trauma-induced impairment in neutrophil glucose uptake was seen at all sampling time points (Figure 5B). 500

501

502 We next examined whether injury impacted upon glucose metabolism, a non-redundant step in NET formation triggered by PMA stimulation (Rodriguez-Espinosa et al. 2015). Using 503 lactate production as a marker of neutrophil glycolytic activity, we measured lactate 504 concentrations in supernatants collected from resting and PMA-stimulated neutrophils 505 following a 3-hour in vitro culture. Compared to HCs, neutrophils isolated from trauma 506 patients at the 48-72 hour post-injury time-point exhibited enhanced basal (Figure 5C) but 507 impaired PMA-induced lactate production (Figure 5D). The increase in basal lactate 508 generation was accompanied by a significant up-regulation in the expression of the glycolytic 509 enzymes pyruvate kinase and lactate dehydrogenase A (Figure 5E-F). 510

511

512 Neutrophils pre-treated with mtDAMPs exhibit impaired NET production but 513 enhanced ROS generation upon secondary stimulation

514

515 Compared to the levels measured in samples from HCs, serum concentrations of the mitochondrial-derived protein ND6 were significantly increased in patients at all post-injury 516 time points, confirming the release of mtDAMPs after trauma (Figure 6A). Demonstrating the 517 immune stimulatory properties of mtDAMPs, we measured significantly reduced CD62L, 518 CXCR1 and CXCR2 expression as well as increased CD11b density on the surface of 519 mtDAMP treated neutrophils (Supplementary Table 1). These changes in neutrophil surface 520 phenotype were accompanied by activation of ERK 1/2 MAPK signalling (Supplementary 521 Figure 3A). The emerging concept of mtDAMP-induced tolerance of neutrophil function is 522 523 based in part on experimental data that has shown prior activation of neutrophils with bacterial-derived N-formylated peptides results in impaired migration upon secondary 524 stimulation (Li et al. 2015). Confirming these findings, we found that neutrophils pre-treated 525

with 40 or 100 μ g/ml preparations of whole mtDAMPs exhibited significantly reduced transmigration towards the chemokine LTB₄ (Supplementary Figure 3B). In contrast, no impairment in migration was witnessed for neutrophils pre-treated with 100 μ g/ml of purified mtDNA (Supplementary Figure 3C).

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To determine whether prior mtDAMP treatment influenced PMA-induced NET production, 531 532 fluorometric analysis was performed on cell-free supernatants collected from cultures of PMA stimulated neutrophils that had been pre-treated with mtDAMPs or vehicle control. 533 Analysis revealed neutrophils pre-exposed to 40 or 100 µg/ml mtDAMPs released 534 535 significantly less DNA following a 3-hour stimulation with PMA than vehicle-treated controls (Figure 6B). Fluorescence microscopy confirmed this mtDAMP-induced inhibition 536 of PMA-induced NET generation (Figure 6C). Interestingly, focussing upon neutrophils pre-537 treated with 100 µg/ml mtDAMPs, images revealed that despite a significant reduction in 538 NET formation, the cells had lost their multi-lobed nuclear morphology following stimulation 539 with PMA, presenting with decondensed nuclear material (Figure 6C). In contrast to whole 540 mtDAMP preparations, neutrophils pre-treated with 40 µg/ml of purified mtDNA prior to 541 PMA stimulation showed no impairment in NET production (data not shown). Interestingly, 542 ROS production, which is a prerequisite for NET formation, was significantly higher upon 543 secondary PMA stimulation for neutrophils pre-treated with 100 µg/ml mtDAMPs 544 545 (Figure 6D).

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547 AMPK is activated by mtDAMP treatment and inhibition of AMPK partially 548 ameliorates the mtDAMP-induced reduction in NET formation 549

In stimulated T cells, elevated intracellular calcium levels activate AMPK, a recently 550 551 described negative regulator of PMA-induced NET formation (Tamas et al. 2006; Wang et al. 2015). As raised intracellular calcium levels are a feature of mtDAMP treated neutrophils 552 (Zhang et al. 2010), we determined the activation status of AMPK in neutrophils following 553 mtDAMP stimulation. To do this, cell lysates, prepared from neutrophils stimulated for 2, 5, 554 10 and 15 minutes with 100 µg/ml mtDAMPs, were probed with a phospho-specific antibody 555 directed against Thr172, a residue within the activation loop of AMPK. As shown in Figure 556 7A, mtDAMP treatment resulted in an immediate and persistent phosphorylation of residue 557 Thr172. Treating neutrophils with the FPR-1 antagonist CsH prior to mtDAMP stimulation 558 resulted in a significant reduction in AMPK phosphorylation, suggesting that N-formyl 559 peptides drive mtDAMP-induced activation of AMPK (Figure 7B). In antigen challenged T 560 cells, phosphorylation of AMPK requires the activation of calcium-calmodulin-dependent 561 protein kinase kinases (CaMKKs), a class of serine/threonine protein kinases activated by 562 increases in intracellular calcium (Tamas et al. 2006). To investigate whether CaMKKs were 563 involved in mtDAMP-induced phosphorylation of AMPK in neutrophils, we treated 564 neutrophils with the CaMKK selective inhibitor STO-609 prior to mtDAMP stimulation. 565 Compared to vehicle control, a significant impairment in mtDAMP-induced activation of 566 AMPK was detected in neutrophils pre-treated with STO-609 (Figure 7C). 567

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AMPK has recently been shown to be a negative regulator of PMA-induced NET formation (*Jiang et al. 2014; Wang et al. 2015*). To investigate whether AMPK signalling was involved in mtDAMP-mediated suppression of NET formation, we treated neutrophils with compound C, an inhibitor of AMPK, prior to mtDAMP exposure. Compared to vehicle control, significantly greater NET production in response to PMA stimulation was recorded for neutrophils pre-treated with compound C (Figure 7D).

575 MtDAMP pre-treatment results in impaired lactate generation by neutrophils upon 576 secondary stimulation with PMA

Confirming results of previous studies that had shown AMPK to be a negative regulator of aerobic glycolysis (Faubert et al. 2013), we measured significantly lower concentrations of lactate in supernatants collected from PMA stimulated neutrophils that had been pre-treated with the AMP mimetic AICAR when compared to vehicle control (Figure 8A). Given that aerobic glycolysis is a key metabolic event in PMA-induced NET formation (Rodriguez-Espinosa et al. 2015) and our observiation of reduced NET generation following PMA stimulation for neutrophils pre-treated with mtDAMPs (Figure 6B-C), we investigated whether AMPK activation triggered by mtDAMP exposure was associated with an impairment in neutrophil glycolysis. Following 1, 2 or 3 hour stimulation with PMA, significantly lower lactate concentrations were measured in supernatants collected from neutrophils pre-treated with 100 µg/ml mtDAMPs (Figure 8B).

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- 625 Discussion
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Here, via the analysis of blood samples acquired from trauma patients within 1-hour of 627 injury, we have shown for the first time that major trauma results in an immediate impairment 628 in neutrophil anti-microbial defences, specifically the formation of NETs. This defect 629 persisted into the acute post-injury phase (4-72 hours) and was accompanied by impaired 630 631 ROS production, MAPK activation and a reduction in both glucose uptake and breakdown. Furthermore, we showed that the trauma-induced reduction in NET formation could be 632 replicated in vitro by treating neutrophils isolated from healthy donors with mtDAMPs prior 633 634 to secondary stimulation. Thus, our data provides support for the emerging concept of mtDAMP-induced tolerance, where the post-injury release of mtDAMPs into the circulation 635 has been proposed to contribute to the neutrophil dysfunction that develops in the aftermath 636 of traumatic injury (Kaczmarek et al. 2018; Li et al. 2015). 637

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In the absence of secondary stimulation, neutrophils obtained from patients within minutes, 639 but not hours, of injury released significantly more DNA into culture supernatants than 640 neutrophils isolated from healthy controls. This immediate trauma-induced increase in basal 641 NET generation is in agreement with the findings of our previous trauma-based study, where 642 in pre-hospital plasma samples, we reported the presence of citrullinated histone H3, a 643 protein that decorates the nuclear DNA backbone of NETs (Hazeldine et al. 2017; Liu et al. 644 645 2012). Thus, taken together, our results suggest that neutrophils are immediately exposed to NET-inducing stimuli post-injury. Supporting this proposal, data presented here and in our 646 647 previous manuscript have shown serum concentrations of known NET inducers, which include TNF-α, IL-8, IL-33 and HMGB-1 are all significantly elevated within minutes of 648 injury (Hazeldine et al. 2017; Keshari et al. 2012; Liu et al. 2012; Tadie et al. 2013; Yazdani 649 650 et al. 2017). Of these agonists, the immediate release of HMGB-1 may be particularly pertinent given that within 60 minutes of *in vitro* co-culture, neutrophils stimulated with this 651 nuclear-derived DAMP have been shown to generate NETs (Huang et al. 2015). Moreover, 652 as HMGB-1 induced NET formation is independent of ROS generation by NADPH oxidase 653 (Tadie et al. 2013), immediate exposure to this DAMP could explain our observation of 654 enhanced *ex vivo* NET formation by neutrophils isolated from pre-hospital blood samples that 655 exhibited no difference in basal ROS production when compared to neutrophils from HCs. In 656 terms of stimulus-induced NET formation, this is the first study to show quantitatively that 657 trauma results in reduced NET generation to PMA stimulation. We confirmed this 658 impairment, which was evident at all three sampling time points, by fluorescent microscopy, 659 660 with our images akin to those presented in a previous study that reported a qualitative postinjury reduction in PMA-induced NET production in a much smaller cohort of trauma 661 patients (Li et al 2015). 662

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The processes that mediate PMA-induced NET generation are well defined, with prominent 664 roles assigned to ROS generation, MAPK activation, glucose uptake and glycolysis (Fuchs et 665 al. 2007; Hakkim et al. 2011; Keshari et al. 2013; Rodriguez-Espinosa et al. 2015). Across 666 our sampling time points, we observed defects in each of these processes, suggesting that 667 multiple impairments rather than a single aberration underlie the post-injury reduction in 668 NET formation. Whilst our data demonstrating a trauma-induced impairment in neutrophil 669 ROS production has been described previously (Hampson et al. 2017; Hazeldine et al. 2017), 670 we are the first to report a post-trauma reduction in MAPK activation, glucose uptake and 671 672 metabolism by neutrophils in response to ex vivo stimulation. Aside from NET production, other anti-microbial mechanisms of neutrophils utilise glucose. For instance, chemotaxis 673 requires the uptake of exogenous glucose (Weisdorf et al. 1982), whilst breakdown of 674

675 endogenous glucose is important for stimulus-induced ROS production and phagocytosis (Rodriguez-Espinosa et al. 2015; Weisdorf et al. 1982). Interestingly, as reported here for 676 NET generation, these three defence strategies have all been shown in ex vivo assays to be 677 significantly impaired following trauma (Hampson et al. 2017; Li et al. 2015; Liao et al. 678 2013; Tarlowe et al. 2003). Thus, the injury-induced reduction we have demonstrated in 679 glucose uptake and metabolism may be a mechanism underlying many facets of post-trauma 680 681 neutrophil dysfunction. Aside from the changes we found in neutrophil signalling and metabolism, trauma patients presented at all three time points with elevated circulating levels 682 of IGs. Shown in vitro to exhibit impaired NET production upon secondary stimulation 683 684 (Martinelli et al. 2004), the immediate and persistent presence of immature cells offers another potential mechanistic explanation for the trauma-induced reduction in NET 685 formation. 686

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Suggesting that traumatic injury modulates cell metabolism, we demonstrated a post-trauma 688 elevation in lactate production by resting neutrophils, an observation that is in agreement with 689 the findings of a previous study. In a cohort of polytrauma patients, Oehler et al reported a 690 higher glycolytic activity, relative to HCs, for neutrophils isolated from subjects between 48 691 and 120 hours post-injury, an enhancement they attributed to increased expression and 692 activity of pyruvate kinase, a glycolytic enzyme that catalyses the conversion of 693 phosphoenolpyruvate to pyruvate (Oehler et al. 2000). Here, we confirmed that traumatic 694 695 injury induces increased expression of pyruvate kinase and showed that this is accompanied by increased expression of lactate dehydrogenase A. Whilst up-regulation of lactate 696 dehydrogenase A, which converts pyruvate to lactate, has been described in transcriptomic 697 analysis of whole blood leukocytes isolated from critically-ill patients (Nalos et al. 2016), 698 this is the first study to demonstrate increased protein expression of lactate dehydrogenase in 699 700 neutrophils post-injury.

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Accompanying the impairment we observed in ex vivo NET formation was a significant 702 trauma-induced elevation in the circulating levels of the mitochondrial-derived N-formylated 703 peptide ND6. In a recent study, suppressed chemotactic responses towards CXCL1 and 704 LTB₄ were reported for neutrophils pre-treated with synthetic ND6 (*Kaczmarek et al. 2018*), 705 706 a finding that mirrored results of previous studies where prior exposure to bacterial-derived N-formylated peptides or ND6 respectively was shown to reduce neutrophil migration and 707 ROS production upon secondary stimulation (Gabl et al. 2018; Li et al. 2015). Adding to this 708 growing body of literature that suggests a tolerising effect for mitochondrial-derived peptides 709 710 on neutrophil function, we demonstrated that neutrophils pre-treated with whole mtDAMP preparations, but not purified mtDNA, exhibited significantly reduced NET production 711 following PMA stimulation. 712

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714 A striking observation of our NET based assays was that despite an absence of NET production, mtDAMP pre-treated neutrophils lost their distinctive multi-lobed nuclear 715 morphology upon PMA challenge. Interestingly, chromatin decondensation in the absence of 716 DNA release was recently reported for PMA stimulated neutrophils pre-treated with the 717 718 glycolysis inhibitor 2-deoxy-glucose (Rodriguez-Espinosa et al. 2015). Using lactate as a readout of glycolytic activity, we measured significantly reduced lactate concentrations in the 719 supernatants of mtDAMP pre-treated neutrophils stimulated with PMA, demonstrating that 720 exposure to mtDAMPs influences the metabolism of immune cells. 721

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723 We found that exposing neutrophils to mtDAMPs resulted in activation of the 724 serine/threonine protein kinase AMPK. In T cells, increases in intracellular calcium levels 725 promote AMPK phosphorylation through activation of CaMKKs (Tamas et al. 2006). Our data demonstrating a significant reduction in mtDAMP-induced phosphorylation of AMPK in 726 neutrophils pre-treated with the selective CAMMK inhibitor STO-506 indicates this 727 signalling pathway is also activated in stimulated neutrophils. As the only component of 728 mtDAMPs that promotes calcium mobilisation in neutrophils (Zhang et al. 2010), signals 729 derived from N-formyl peptides are likely to have driven the mtDAMP-induced 730 phosphorylation of AMPK. Supporting this idea, we observed significantly reduced 731 mtDAMP-induced AMPK phosphorylation in neutrophils pre-treated with CsH, an FPR-1 732 antagonist that prevents calcium mobilisation upon mtDAMP stimulation (Zhang et al. 2010). 733 734

Demonstrating that AMPK is a negative regulator of NET formation, significantly enhanced

- and impaired NET formation has been reported for PMA stimulated neutrophils pre-treated 736 737 with AMPK inhibitors and activators respectively (Jiang et al. 2014; Wang et al. 2015). Supporting these data, we showed significantly greater PMA-induced NET production by 738 neutrophils treated with the AMPK inhibitor compound C prior to mtDAMP exposure. How 739 activation of AMPK inhibits NET formation is currently unknown. Based on published 740 741 literature, we propose two mechanisms, both of which revolve around the ability of AMPK to inhibit the serine/threonine protein kinase mammalian target of rapamycin (mTOR) (Faubert 742 et al. 2013; MacIver et al. 2011). Firstly, inhibition of mTOR has been shown to significantly 743 reduce Glut1 transporter activity (Wieman et al. 2007). Given the importance of extracellular-744 745 derived glucose in NET formation (Rodriguez-Espinosa et al. 2015), reduced glucose uptake, secondary to impaired mTOR and Glut1 activity, could contribute to the reduction in NET 746 747 production and lactate generation that we observed for neutrophils pre-treated with mtDAMPs. Furthermore, this mechanism could contribute to the mtDAMP-induced 748 749 impairment we reported in neutrophil transmigration since chemotaxis also utilises 750 extracellular sources of glucose (Weisdorf et al. 1982). Alongside Glut1, mTOR signalling has been implicated in regulating the expression/activity of hypoxia-inducible factor-1a 751 (HIF-1a) (Faubert et al. 2013). A transcription factor involved in promoting aerobic 752 glycolysis, activation of HIF-1a precedes both NET formation (McInturff et al. 2012) and 753 myeloid cell migration (Cramer et al. 2003). Thus, reduced activity/expression of HIF-1a, 754 secondary to AMPK-mediated inhibition of mTOR, could be an additional/alternative 755 explanation for the mtDAMP-induced reduction in lactate generation, NET generation and 756 neutrophil chemotaxis. Importantly, both these proposed mechanisms could occur in 757 neutrophils without affecting their capacity for ROS production, which we found was 758 significantly increased following mtDAMP treatment. Indeed, as the energy required for ROS 759 760 generation is derived from endogenous sources of glucose (Rodriguez-Espinosa et al. 2015), this anti-microbial function could occur in the background of an AMPK driven reduction in 761 extracellular glucose uptake. Our finding of increased ROS production to PMA stimulation 762 763 for neutrophils pre-treated with mtDAMPs contradicts the post-injury impairment we reported in ROS generation for trauma patients that presented with significantly elevated 764 serum concentrations of mtDAMPs. We believe this discrepancy may be attributable to 765 trauma-induced changes in the composition of the circulating neutrophil pool. For example, 766 traumatic injury results in the emergence into circulation of IGs and CD16^{BRIGHT} CD62L^{DIM} 767 neutrophils, both of which exhibit impaired stimulus-induced ROS generation (Drifte et al. 768 769 2013; Sauce et al. 2017). In contrast, healthy subjects, who served as the cohort for our mtDAMP pre-treatment experiments, possess a homogenous pool of fully-functional mature 770 neutrophils that would exhibit a greater capacity to respond to stimulation.
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Although we have shown that neutrophils pre-treated with compound C exhibit increasedNET generation upon PMA stimulation, inhibition of AMPK only partially ameliorated the

775 reduction in NET formation that occurred with mtDAMP treatment. Other factors aside from AMPK activation must therefore be involved in mediating the mtDAMP-induced suppression 776 of NET production. Recently, through a proposed mechanism of action that involved the 777 prevention of membrane rupture, lactoferrin, an iron-binding glycoprotein stored within the 778 secondary granules of neutrophils, was found to suppress NET release triggered by PMA 779 stimulation (Okubo et al. 2016). Indicative of a defect in the latter stages of NET production, 780 781 we showed chromatin decondensation in the absence of DNA release was a feature of mtDAMP-treated neutrophils. Thus, based on this observation and the fact that exposure to 782 mtDAMPs promotes neutrophil degranulation (Hauser et al. 2010; Hazeldine et al. 2015; 783 784 Zhang et al. 2010), we suggest that a mtDAMP-induced release of lactoferrin could represent an additional mechanistic explanation for the impairment in NET formation that occurs 785 following mtDAMP exposure. 786

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This study has some limitations. Conducted at a single major trauma centre, the results of our 788 prospective observational study are based on the analysis of a small number of patient 789 samples, meaning our findings require validation in larger independent cohorts. This point is 790 particularly pertinent to our analysis of pre-hospital blood samples, where inter-individual 791 variability in immune cell number and volume of blood collected at the scene of injury meant 792 we were unable to perform all assays on each patient at this time point. As a heterogeneous 793 collection of proteins, lipids and DNA, no study to date has quantified the exact concentration 794 795 of mtDAMPs released into the circulation post-injury. Thus, our in vitro treatment of neutrophils with 40 or 100 µg/ml mtDAMPs may not be physiologically relevant. However, 796 797 these doses match those used in previous in vitro based studies that have examined the effect 798 of mtDAMP exposure on neutrophil anti-microbial function (Hauser et al. 2010; Hazeldine et al. 2015; Zhang et al. 2010). Similarly, our decision to use the phorbol ester PMA as an 799 800 agonist may be considered a study limitation. However, as a potent stimulus, it allowed us to study maximal neutrophil responses. Moreover, as the agonist of choice for all previous 801 trauma-based studies that had examined stimulus-induced NET production post-injury 802 (Hampson et al. 2017; Itagaki et al. 2015; Li et al. 2015), our use of PMA enabled us to 803 compare our observations to those in the published literature. 804

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In summary, this is the first study to describe a quantitative post-trauma reduction in NET 806 formation as an immediate on-scene phenomenon, which is accompanied by aberrant 807 intracellular signalling and cell metabolism. In addition, we have shown that the post-injury 808 reduction in NET generation can be recreated *in vitro* by treating neutrophils isolated from 809 810 healthy subjects with mtDAMPs prior to PMA stimulation. Thus, our data support the suggestion that the release of mtDAMPs from damaged tissue is a contributory factor in the 811 reduction in neutrophil function that occurs post-injury (Hampson et al. 2017; Hazeldine et 812 813 al. 2017; Li et al. 2015; Liao et al. 2013; Tarlowe et al. 2003).

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- 825 Acknowledgements

The research was funded by the NIHR Surgical Reconstruction and Microbiology Research Centre (SRMRC). The authors would also like to acknowledge the Queen Elizabeth Hospital Birmingham Charity for funding the purchase of the Sysmex XN-1000 haematology analyser. The views expressed are those of the author(s) and not necessarily those of the NHS, the NIHR or the Department of Health. The authors thank all the research, nursing and administrative staff at the NIHR-SRMRC for their assistance in data collection and management. We also thank the emergency healthcare professionals in the West Midlands Ambulance Service NHS Foundation Trust, the Midlands Air Ambulance Charity and the University Hospitals Birmingham NHS Foundation Trust for their participation in the identification, enrolment and follow up of patients in this study.

838 Author contributions statement

JH designed the study, performed experimental work, analysed data and wrote the
manuscript. RD performed experimental work and analysed data. PH critically appraised the
manuscript and JL conceptualised the study and contributed to writing the manuscript.

844 Conflict of interest statement

The authors declare the submitted manuscript was not carried out in the presence of any
personal, professional or financial relationships that could potentially be construed as a
conflict of interest.

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Figure 1. Effect of traumatic injury on PMA-induced NET formation. (A) Basal NET 1038 1039 generation by resting neutrophils isolated from healthy controls (HC) and trauma patients as assessed by DNA concentration in cell free supernatants following a 3-hour in vitro culture. 1040 (B) HMGB-1 and (C) IL-33 concentrations in serum samples from HC and trauma patients. 1041 1042 IL-33 levels were undetectable (N.D) in serum samples from HC. (D-E) Following a 3-hour in vitro stimulation with PMA, NET production by neutrophils from HC and trauma patients 1043 was compared by measuring DNA concentration in cell free supernatants (D) and 1044 1045 fluorescence microscopy (E). For supernatant analysis, number of samples are shown below 1046 each time-point. For microscope images, HC (n=12), \leq 1H (n=6), 4-12H (n=6) and 48-72H (n=8). *p<0.01, **p<0.001, ***p<0.0001 vs. HC. 1047

- Figure 2. Neutrophil ROS production and immature granulocyte count post-injury. Comparison of basal (A) and PMA-induced (B) ROS production by neutrophils isolated from healthy controls (HC) and trauma patients. Data are presented as area under the curve (AUC) (A) or fold increase above vehicle treated controls (B). (C-D) Frequency (C) and absolute number (D) of immature granulocytes (IG) in peripheral blood samples from healthy controls (HC) and trauma patients. Number of samples analysed are shown below each time-point. p<0.01, ***p<0.001, ****p<0.0001 vs. HC.
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- 1057 Figure 3. Traumatic injury results in impaired PMA-induced activation of p38 MAPK. 1058 Phosphorylation status of P38 in resting and PMA-stimulated neutrophils isolated from 1059 healthy controls (HC) and trauma patients 4-12 and 48-72 hours post-injury. Data are presented as representative Western blots (B) and densitometry analysis of p38 1060 1061 phosphorylation in resting (A) or PMA-stimulated neutrophils at the 4-12H (C) and 48-72H (**D**) post-injury time points. HC (n=7), 4-12h (n=9) and 48-72h (n=4). For (A) *p<0.01, 1062 *p<0.001 vs. HC. For (C) and (D) **p<0.001 vs. Time 0, *p<0.01, **p<0.0001 vs. HC 1063 1064 sample at matched time point.
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Figure 4. Traumatic injury results in impaired PMA-induced activation of ERK 1066 MAPK. Phosphorylation status of ERK 1/2 in resting and PMA-stimulated neutrophils 1067 isolated from healthy controls (HC) and trauma patients 4-12 and 48-72 hours post-injury. 1068 Data are presented as representative Western blots (B) and densitometry analysis of ERK 1069 phosphorylation in resting (A) or PMA-stimulated neutrophils at the 4-12h (C) and 48-72h 1070 1071 (D) post-injury time points. HC (n=4), 4-12h (n=8) and 48-72h (n=6). For (C) and (D) $*^{*}p<0.0001$ vs. Time 0, $*^{**}p<0.001$, $*^{***}p<0.0001$ vs. HC sample at 1072 *p<0.01, **p<0.001, matched time point. 1073

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1075 Figure 5. Effect of traumatic injury on neutrophil glucose uptake and metabolism. Glucose uptake by basal (A) or PMA-stimulated (B) neutrophils isolated from healthy 1076 controls (HC) and trauma patients. MFI, Mean fluorescence intensity. (C-D) Comparison of 1077 lactate concentration in cell-free supernatants collected from resting (C) or PMA stimulated 1078 1079 (D) neutrophils isolated from HC and trauma patients following a 3-hour in vitro culture. Number of samples analysed are indicated below each time-point. * p<0.01, **p<0.001, 1080 ^{*}p<0.0001 vs. HC. (**E-F**) Expression of the glycolytic enzymes pyruvate kinase (PKM2) 1081 and lactate dehydrogenase A (LDHA) in resting neutrophils isolated from HC (n=5) and 1082 1083 trauma patients 48-72 hours post-injury (n=14). Data are presented as representative Western blots (E) and densitometry analysis of collated data for LDHA (F, top panel) and PKM2 1084 (**F**, bottom panel). p<0.01 vs. HC. 1085

Figure 6. Effect of mtDAMP pre-treatment on neutrophil NET generation. (A) Serum concentrations of the mitochondrial-derived N-formylated peptide ND6 in peripheral blood samples acquired from healthy controls (HC) and trauma patients. Number of samples analysed are indicated below each time-point. ****p<0.0001 vs. HC. (B) NET production by PMA stimulated neutrophils pre-treated with 40 or 100 µg/ml mtDAMPs was assessed by measuring DNA content of cell-free supernatants (n=5). p<0.01 vs. Vehicle. (C) MtDAMP-induced inhibition of NET generation was confirmed by fluorescence microscopy (n=5; top panel x20 magnification; bottom panel x40 magnification). (D) ROS generation by neutrophils pre-treated with 100 µg/ml mtDAMPs or vehicle control was measured in response to stimulation with 25 nM PMA using luminol-based chemiluminescence (n=10). Data are presented as area under the curve (AUC) and are mean±SEM. *** p<0.0001 vs. PMA.

- Figure 7. Treatment of neutrophils with mtDAMPs results in phosphorylation of AMPK. (A) Whole cell lysates prepared from purified neutrophils stimulated for 2-15 minutes with 100 µg/ml mtDAMPs were screened for phosphorylated AMPK. Western blot in top panel is representative of 4 independent experiments. For densitometry analysis ^{**}p<0.0001 vs. 0 minutes. (**B**) AMPK phosphorylation in neutrophils treated for 1-hour with the FPR-1 antagonist Cyclosporin H (CsH) or (C) or the CaMKK inhibitor STO-609 prior to a 5-minute stimulation with 100 μ g/ml mtDAMPs. Blots are representative of 5 (**B**) and 10 (C) independent experiments, with densitometric data depicted in the accompanying histogram. **p<0.001, ***p<0.0001 vs. vehicle. (D) Comparison of PMA-induced NET formation by mtDAMP stimulated neutrophils pre-treated with the AMPK inhibitor compound C or vehicle control (n=10). **p<0.01 vs. PMA treated.

Figure 8. MtDAMP pre-treatment results in impaired neutrophil glycolysis. (A) Neutrophils pre-treated for 1-hour with 1 mM AICAR or vehicle control were stimulated for 3 hours with 25 nM PMA, after which lactate concentrations were measured in cell-free supernatants. Data are mean±SEM of 12 independent experiments. **** p<0.0001. (B) Comparison of lactate concentrations in supernatants collected from neutrophils pre-treated with 100 µg/ml mtDAMPs or vehicle-control and subsequently stimulated with 25 nM PMA for 1, 2 and 3 hours. Data are mean±SEM of 10 independent experiments. *p<0.001, ****p<0.0001 vs. vehicle.

1136 Table 1. Cohort demographics.

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	Patients (n=62)
Age, years Male, n (%) Time to pre-hospital sample, minutes post-injury	44 (19-95) 56 (90) 39 (13-59)
ISS NISS	26 (9-57) 38 (9-75)
AIS	
Head, n (%) Face, n (%) Chest, n (%) Abdomen, n (%) Spine, n (%) Pelvis, n (%) Limbs, n (%) Other, n (%)	24 (48) 17 (34) 29 (58) 14 (28) 19 (38) 8 (16) 28 (56) 6 (12)
Mechanism of injury	
Fall, n (%) A/P, n (%) Blunt, n (%) RTC, n (%)	10 (16) 13 (21) 2 (3) 37 (60)
Outcomes	
ICU-free days Hospital-free days Mortality, n (%)	22 (0-30) 13 (0-29) 8 (13)

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1140 Data are expressed as mean (range) unless indicated otherwise.

1141 The number of data points for each clinical variable are: ISS and NISS, n=48; AIS scores, 1142 n=50; ICU and hospital free days, n=53.

1143 ICU-free days and hospital-free days were calculated by 30 minus the number of days the 1144 patient stayed in hospital).

1145 A/P, Assault/penetrating; AIS, Abbreviated injury scale; ISS, Injury severity score; ICU,

1146 Intensive care unit; NISS, New injury severity score; RTC, Road traffic collision.

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PMA-induced NET production



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