Physicochemical properties that control protein aggregation also determine whether a protein is retained or released from necrotic cells

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Amyloidogenic protein aggregation impairs cell function and is a hallmark of many chronic degenerative disorders. Protein aggregation is also a major event during acute injury; however, unlike amyloidogenesis, the process of injury-induced protein aggregation remains largely undefined. To provide this insight, we profiled the insoluble proteome of several cell types after acute injury. These experiments show that the disulfide-driven process of nucleocytoplasmic coagulation (NCC) is the main form of injury-induced protein aggregation. NCC is mechanistically distinct from amyloidogenesis, but still broadly impairs cell function by promoting the aggregation of hundreds of abundant and essential intracellular proteins. A small proportion of the intracellular proteome resists NCC and is instead released from necrotic cells. Notably, the physicochemical properties of NCC-resistant proteins are contrary to those of NCC-sensitive proteins. These observations challenge the dogma that liberation of constituents during necrosis is anarchic. Rather, inherent physicochemical features including cysteine content, hydrophobicity and intrinsic disorder determine whether a protein is released from necrotic cells. Furthermore, as half of the identified NCC-resistant proteins are known autoantigens, we propose that physicochemical properties that control NCC also affect immune tolerance and other host responses important for the restoration of homeostasis after necrotic injury.

1. Introduction

Proteins must fold into their native conformation to be fully functional. However, as the native conformation is labile, all proteins are also capable of misfolding. Protein misfolding exposes hydrophobic regions that form intra- and inter-molecular associations, which if left unchecked can multimerize into higher-order toxic aggregates [1]. Protein quality control mechanisms adequately deal with most forms and degrees of misfolding. Nonetheless, destabilizing mutations, extreme environments or ageing can overwhelm proteostasis leading to the unwanted accumulation of aggregated proteins. Protein aggregation, and in particular amyloidogenic aggregation, underlies many chronic age-related diseases including Alzheimer’s disease.
and Parkinson’s disease. In these chronic settings, the onset of protein aggregation is widely thought to elicit injury.

Paradoxically, protein aggregation is also a key response to acute tissue injury. Injury-induced protein aggregation was first described in the 1980s [2–5], where aberrant actin aggregation occurred in cells after ATP depletion or oxidant exposure. Subsequent studies showed that actin, myosin, vinculin and HSP70 all aggregated upon ATP depletion [6–9]. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) is another intracellular protein that readily aggregates during stress in vitro [10–14] and in vivo [10,14–16]. Injury-induced protein aggregation has also been observed during hypoxia [17–19].

Recently, we [10,12,20,21] and others [22] have characterized the process of injury-induced protein aggregation. In particular, we found that injury can trigger a specific form of protein aggregation called nucleocytoplasmic coagulation (NCC) [10]. By definition, NCC is a temporally coordinated disulfide-crosslinking reaction that occurs during late-stage cell death. NCC causes many intracellular proteins, including actin, tubulin, GAPDH and HSP90β, to convert from a soluble form into high-molecular-weight insoluble species [10]. In vitro and in vivo data show that NCC-aggregated proteins promote activation of the extracellular protease plasmin on the surface of dead cells [10,20,21]. Surface-bound plasmin then initiates two forms of clearance: direct proteolytic degradation of dead cells [10,20] and signalling nearby dendritic cells to increase their phagocytic capacity [21]. Thus, NCC-aggregated proteins can be considered a novel ‘damage-associated molecular pattern’ that initiates humoral and cellular responses to sterile injury [23].

Despite these recent advancements the mechanisms of injury-induced protein aggregation are poorly understood. Accordingly, to ascertain the basis of injury-induced protein aggregation, we have used quantitative mass spectrometry to profile the insoluble proteome of three cell types across different injury conditions. We find that NCC is a specific consequence of necrosis. Moreover, our data suggest that NCC is the predominant form of injury-induced protein aggregation, which affects a much wider array of proteins than first thought. Surprisingly, a small subset of proteins avoids NCC-mediated aggregation. Comparison of NCC-resistant and NCC-sensitive proteins uncovers two fundamental principles: (i) that injury-induced aggregation is distinguishable from amyloidogenic aggregation, and (ii) that a protein’s inherent physicochemical properties determines whether it is retained or released from necrotic cells.

Altogether, NCC represents a ‘fail-safe switch’ that broadly incapacitates cellular functions during necrosis and selectively retains debris at the site of injury. As a result, we propose that NCC is a beneficial event that pacifies the reactivity of necrotic cells. In support of this notion, and consistent with the fact that defective removal of dead cells is a strong risk factor for autoimmune disease [24–30], we note that many proteins which evade NCC are also recognized autoantigens. Future studies should now assess whether mutations that alter a protein’s propensity to undergo NCC can in turn influence its ability to act as an autoantigen.

2. Results and discussion

2.1. Profiling the insoluble proteome of injured cells

We first characterized injury and protein aggregation in human Jurkat lymphocyte cells treated with etoposide (a topoisomerase-II inhibitor), staurosporine (a pan-kinase inhibitor) or Fas ligand (FasL; a TNF death receptor agonist). All agonists induced apoptosis in Jurkat cells (indicated by an increase in phosphatidyserine exposure and caspase-mediated RIPK1 cleavage; figure 1a; electronic supplementary material, figure S1a,b). More specifically, FasL and staurosporine triggered a similar degree/rate of injury, with early stage apoptosis evident within 3 h of treatment (indicated by phosphatidyserine-positive, TO-PRO-3-negative staining; electronic supplementary material, figure S1a) and transition to a secondary necrotic state within 24 h of treatment (indicated by complete loss of metabolism, plasma membrane disruption, pronounced LDH release and the acquisition of TO-PRO-3-positive staining; figure 1a; electronic supplementary material, figure S1a). Note that neither necroptosis (mediated by full-length RIPK1) nor autophagic death (mediated by LC3 cleavage; electronic supplementary material, figure S1d) was evident after FasL or staurosporine treatment. By contrast, etoposide caused early stage apoptosis after 24 h of treatment with no signs of secondary necrosis (as indicated by LDH retention, plasma membrane impermeability, partial metabolic suppression, and modest elevations in phosphatidyserine exposure and RIPK1 cleavage; figure 1a; electronic supplementary material, figure S1a).

Lastly, we confirmed protein aggregation in Jurkat cells during the later stages of staurosporine-treatment (figure S1d). Disulfide-crosslinked insoluble forms of these same proteins were also observed after FasL treatment, but not after etoposide treatment (not shown). Note that the coordinated loss of protein solubility upon secondary necrosis represents a genuine aggregation event (rather than merely being a consequence of disulfide crosslinking), as these same insoluble proteins were observed in the presence of a reducing agent (electronic supplementary material, figure S1d). Thus, injury-induced protein aggregation in Jurkat cells occurred specifically during necrosis, but not during early stage apoptosis. Moreover, as the vast majority of insoluble material was disulfide-crosslinked (electronic supplementary material, figure S1d), NCC appears to be the predominant form of injury-induced protein aggregation.

Next, we profiled the insoluble proteome of Jurkat cells after 24 h of treatment with vehicle, etoposide, staurosporine or FasL using the quantitative mass spectrometry-based technique of SILAC (stable isotope labelling by amino acids in cell culture [32]; see electronic supplementary material, figure S2 and Material and methods for experimental details). This approach allowed the identification of hundreds to thousands of insoluble proteins per experiment with measurement of their relative abundance between uninjured and injured cells. Figure 1b and electronic supplementary material, table S1 summarize the SILAC results. Proteins with an increased right-shifted SILAC ratio in figure 1b were assumed to have undergone injury-induced aggregation, as this shift was only observed in staurosporine- or FasL-treatedJurkat cells. To verify this assumption, 12 proteins identified as putative aggregators in the SILAC experiments were subjected to immunoblot analysis. As shown in figure 2a, all 12 proteins were confirmed to undergo injury-induced protein aggregation in necrotic Jurkat cells. Consistent with the initial description of NCC in neuronal cells [10], necrosis in Jurkat cells caused all tested proteins to convert into insoluble high-molecular-weight...
NCC-participating subproteome (green colour) and proteins with a log2 ratio proteins underwent NCC in necrotic Jurkat cells (figure 1 population of proteins with log2 SILAC ratios less than portion of the proteome than prior studies suggest. injury-induced aggregation appears to affect a much larger 'NCC-resistant' proteins were decreased in abundance in the

By definition, apoptosis is an energetic process, whereas necrosis coincides with metabolic incompetency. Therefore, because 24 h incubation with staurosporine and necrosis (% LDH and % trypan blue positive) were measured after 24 h of treatment. Data are presented as mean ± s.e.m. from n = 3–7 independent experiments. *p < 0.05, **p < 0.01 and ****p < 0.0001 relative to uninjured cultures as determined by one-way ANOVA with Newman–Keuls correction. Moreover, as 30–46% of the identified proteins within each gate is shown. (c) Partial overlap in the gated subproteomes between staurosporine- and Fas ligand-induced necrotic Jurkat cells. The number of proteins within each gate is shown. (d) Partial overlap in the NCC-participating subproteome between staurosporine-induced necrotic Jurkat cells versus staurosporine-induced necrotic human U937 monocytes.

Figure 1. Diametric changes in the abundance of insoluble intracellular proteins upon secondary necrosis. (a) Jurkat lymphocytes were treated with vehicle (uninjured) or injured with etoposide, staurosporine or Fas ligand. Phosphatidylserine (PS) exposure (relative AnnexinV-signal geomean), metabolism (% MTS) and necrosis (% LDH and % trypan blue positive) were measured after 24 h of treatment. Data are presented as mean ± s.e.m. from n = 3–7 independent experiments. *p < 0.05, **p < 0.01 and ****p < 0.0001 relative to uninjured cultures as determined by one-way ANOVA with Newman–Keuls correction. Therefore, because 24 h incubation with staurosporine or Fas ligand abolished metabolism (in conjunction with high levels of LDH release, trypan blue uptake and TO-PRO-3-staining; electronic supplementary material, figure S1a), we have concluded that these agonists produced a secondary necrotic state by this time point. (b) Jurkat cultures were labelled with either heavy or light lysine isopes and treated with vehicle (uninjured), etoposide, staurosporine or Fas ligand for 24 h. Triton-insoluble fractions were extracted and proteins identified using quantitative mass spectrometry. The abundance of identified proteins relative to that in isotypically different uninjured cultures was then plotted as the log2 ratio of the stable isotope label (SILAC), where an increased ratio indicates more of a specific insoluble protein in injured cells relative to uninjured cells and a decreased ratio indicates less of a specific insoluble protein in injured cells relative to uninjured cells. Proteins with a log2 ratio > 0.6 were gated into the NCC-participating subproteome (green colour) and proteins with a log2, ratio < −0.6 were gated into the NCC-resistant subproteome (pink colour). This gating was also chosen based on the recommendations of Zhang & Neubert [31]. Inserts show the percentage of identified proteins that fall within each gate. (c) Partial overlap in the gated subproteomes between staurosporine- and Fas ligand-induced necrotic Jurkat cells. The number of proteins within each gate is shown. (d) Partial overlap in the NCC-participating subproteome between staurosporine-induced necrotic Jurkat cells versus staurosporine-induced necrotic human U937 monocytes.

dithiobisreitol-sensitive species (as assessed by immunoblot; figure 2b) and to relocate into distinct aberrant subcellular structures (as assessed by super-resolution microscopy; figure 2c) which stained strongly for the presence of oxidized thiols (as assessed by confocal microscopy; figure 2f). Thus, we concluded that proteins that exhibited log2 SILAC ratios greater than +0.6 were indeed ‘NCC-participating’ proteins (green colour in figure 1b). Moreover, as 30–46% of the identified proteins underwent NCC in necrotic Jurkat cells (figure 1b), injury-induced aggregation appears to affect a much larger portion of the proteome than prior studies suggest.

Closer inspection of the SILAC data revealed another population of proteins with log2 SILAC ratios less than −0.6 (11–13% of identified targets; red colour in figure 1b). These ‘NCC-resistant’ proteins were decreased in abundance in the insoluble fraction of necrotic Jurkat cells relative to uninjured cells, suggesting that they were either degraded or avoided aggregation during injury. To assess which scenario was applicable, six putative NCC-resistant proteins were selected for immunoblot analysis. Consistent with the SILAC data, all six tested proteins were decreased in abundance in the insoluble fraction of necrotic Jurkat cells relative to uninjured cells (figure 3e). Notably, the disappearance of these NCC-resistant proteins in necrotic cells did not result in the commensurate appearance of lower molecular weight immunoreactive species, suggesting that they were not being degraded (figure 3e). Next, we reasoned that if a protein avoided both NCC and degradation, then it should instead be released into the extracellular milieu upon necrosis. Consequently, we assayed for the presence of FUS (also called ‘RNA-binding protein FUS’ or
Figure 2. NCC is the main form of injury-induced protein aggregation. (a–d) Jurkat cells were treated with vehicle (uninjured), etoposide, staurosporine or Fas ligand for 24 h. (a) Triton-soluble and -insoluble proteins were isolated and subjected to SDS-PAGE under reducing conditions followed by immunoblot analysis. Results are representative of n = 3–6 independent experiments. (b) Triton-soluble and -insoluble proteins were isolated and subjected to SDS-PAGE in the presence/absence of dithiothreitol (DTT) followed by immunoblot analysis. Results are representative of n = 2–3 independent experiments. (c) Cells were fixed, subjected to immunofluorescence and imaged by super-resolution microscopy with xy-resolution of 70 nm. Shown are representative x,y micrographs, where the bottom two rows are magnified micrographs taken from the boxed areas in the top two rows. Staurosporine-induced necrosis causes NCC-proteins (α-tubulin, EF2 and MCM7) to relocate into aberrant and distinct structures. Arrows indicate EF2 deposits that exhibit little-to-no staining for tubulin or MCM7. Arrowheads indicate MCM7 deposits that exhibit little-to-no staining for oxidized thiols. Note, the procedure used to stain oxidized thiols was not compatible with the sample preparation for super-resolution microscopy. (d) Cells were fixed and subjected to immunofluorescence with maleimide co-staining for oxidized thiols. Shown are representative x,y confocal micrographs. Staurosporine-induced necrosis leads to nuclear breakdown and a marked increase in oxidized thiols that colocalize with aberrantly deposited NCC-proteins (3PDGH, GAPDH, MCM7, EF1, EF2 and α-tubulin).

Translocated in liposarcoma’, TLS; which was the most easily detected NCC-resistant protein) in the conditioned media from injured Jurkat cells. As shown in figure 3b, full-length FUS, but not MCM7 (an NCC-participating protein), was readily detected in the conditioned media of necrotic Jurkat cells. To the best of our knowledge, this is the first report that the ubiquitous protein, FUS, is released from necrotic cells. It remains to be determined whether other NCC-resistant proteins are released from necrotic cells, but this seems highly likely (for reasons provided below with figures 4 and 5). The capacity of FUS and other NCC-resistant proteins to act as paracrine communicators of injury should now be assessed, especially given the influence of NCC on the proteolytic [10,20] and phagocytic [21] removal of dead cells. It will also be interesting to see if FUS released from necrotic cells has a role during FUS-mediated amyotrophic lateral sclerosis [37].

We next examined injury-induced protein aggregation in human U937 monocyte cells, where both secondary necrosis (electronic supplementary material, figure S3a) and NCC (electronic supplementary material, figure S3b) occurred after 24 h of treatment with either etoposide or staurosporine. SILAC experiments showed that similar, albeit more pronounced changes to the insoluble proteome occurred upon secondary necrosis in U937 cells (figure 1d; electronic supplementary material, figure S3c), with 53–61% of proteins participating in NCC and 16–30% of proteins avoiding NCC (electronic supplementary material, figure S3c). The reason for this increased aggregation of the U937 proteome is unknown, but may reflect the heightened sensitivity of U937 cells to injury (cf. figure 1a and electronic supplementary material, figure S3a), or could relate to the extreme differences in the metabolism of monocytes and lymphocytes [38]. It is telling that etoposide elicited neither necrosis nor NCC in Jurkat cells, but strongly induced both secondary necrosis and NCC in U937 cells. This observation emphasizes that necrosis per se is the major cellular correlate of NCC. It is also salient that, despite marked differences in the means by which etoposide, staurosporine and Fasl injure cells, many of the same proteins underwent NCC when these...
treatments caused necrosis (figure 1c; electronic supplementary material, figure S3f). Such significant overlap in the insoluble proteome across different cell types and necrotic conditions denotes that a common underlying mechanism governs the participation/resistance of proteins to NCC.

2.2. Elucidating the physicochemical basis of nucleocytoplasmic coagulation

All proteins are capable of aggregating; however, the propensity, kinetics and conformations of aggregation are largely determined by the inherent physicochemical properties of the protein in question. Therefore, a high-confidence list of NCC-participating, NCC-resistant and intermediary ‘unchanged’ proteins was collated from the Jurkat and U937 SILAC datasets (electronic supplementary material, table S1), and their primary sequences were compared to ascertain the physicochemical basis of NCC. The most prominent feature of NCC-participating proteins was their relatively high cysteine content (figure 4a), with a significant correlation between cysteine content and the degree of injury-induced aggregation (figure 4b). This finding, in conjunction with our earlier publications showing that NCC-aggregates exhibit a prefibrillar oligomeric conformation with no subsequent appearance of fibrillar species [10].

Other characteristics of NCC-participating proteins include a higher representation of aromatic residues (F, Y, W; \( p < 0.001 \) via unpaired two-tailed t-test; figure 4a), aliphatic residues (I, V, L; \( p < 0.0001 \) via unpaired 2-tailed t-test; figure 4a) and increased hydrophobicity (electronic supplementary material, figure S5a)—attributes which explain the predicted high aggregation propensity of NCC-participating proteins relative to NCC-resistant proteins (figure 4f). Curiously, the intrinsic disorder content (figure 4e) and the molecular weight (electronic supplementary material, figure S5b) of NCC-participating proteins were lower than those of NCC-resistant proteins. These collective attributes further distinguish NCC from amyloidogenesis, which instead favours co-aggregation of large hydrophobic proteins with high intrinsic disorder (figure 4f) [34]. That diametrically opposed traits exist between NCC-participating and NCC-resistant proteins shows that the same physicochemical forces control these protein fates during necrosis. Indeed, our data show that whether a protein is retained or released from necrotic cells is in part determined by its propensity to undergo injury-induced aggregation. This conclusion was independently supported by analysing the insoluble...
proteome of a third cell line after necrotic injury (i.e. the mouse AT3 mammary adenocarcinoma cells; electronic supplementary material, figure S6 and table S1).

2.3. Identifying higher-order features of nucleocytoplasmic coagulation

Many factors besides primary sequence affect protein aggregation. Foremost among these factors is the concentration dependency of protein aggregation. In line with this tenet, NCC-participating proteins were found to be more abundant than NCC-resistant proteins (figure 5i). Next, we performed ontology enrichment analysis to gain a broader sense of what molecular functions (figure 5b; electronic supplementary material, table S2) and subcellular compartments (electronic supplementary material, figure S7a and table S3) were affected by NCC. These analyses showed that NCC preferentially affects metabolic (e.g. GTPases), oxidoreductase (e.g. dehydrogenases, perioxidoxins), signalling (e.g. kinases) and proteostatic enzymes (e.g. HSPs, translation factors), as well as nucleotide- and nucleic acid-binding proteins ($p < 0.05$; electronic supplementary material, table S2). NCC was also found to preferentially affect macromolecular complexes in the nucleus (e.g. nuclear pore), cytoplasm (e.g. chaperonin, proteosome, cytoskeleton) and mitochondria (e.g. nuclear pore, cytoplasm (e.g. chaperonin, proteosome, cytoskeleton) and mitochondria (electronic supplementary material, figure S7i and table S3), subcellular compartments with a constitutively redox-reduced state ($p < 0.05$; electronic supplementary material, table S3). By comparison, distinct molecular functions and subcellular compartments were prevalent among NCC-resistant proteins (electronic supplementary material, tables S2 and S3). Thus, numerous energetic and homeostatic functions in the nucleus, cytoplasm and mitochondria are directly affected by NCC.
The node size is proportional to the statistical significance of enrichment. The coloured pie segments that fill each node indicate the percentage of the ontology representation of known autoantigens within the different protein subsets. (c) Graph comparing the abundance across the different protein subsets. Black lines indicate the average abundance of each subset. **p < 0.001 as determined by one-way ANOVA with Tukey’s correction. (b) Network depicting the molecular functions that were significantly over-represented in the NCC-participating subset, relative to the complete human proteome. Nodes (circles) represent different molecular function ontologies. Functionally related nodes are connected by edges (straight lines) and have been grouped using ClueGO [35], depicting the molecular functions that were significantly over-represented in the NCC-participating subset, relative to the complete human proteome. Nodes (circles) are connected by edges (straight lines) and have been grouped using ClueGO [35], depicting the molecular functions that were significantly over-represented in the NCC-participating subset, relative to the complete human proteome.

Figure 5. NCC preferentially affects abundant ancient proteins that perform essential cellular functions and exhibit reduced autoreactivity. (a–e) The same high-confidence protein subsets and the randomly selected control subset from figure 4 were further analysed. (a) Graph comparing the abundance across the different protein subsets. Black lines indicate the average abundance of each subset. **p < 0.001 as determined by one-way ANOVA with Tukey’s correction. (b) Network depicting the molecular functions that were significantly over-represented in the NCC-participating subset, relative to the complete human proteome. Nodes (circles) represent different molecular function ontologies. Functionally related nodes are connected by edges (straight lines) and have been grouped using ClueGO [35], depicting the molecular functions that were significantly over-represented in the NCC-participating subset, relative to the complete human proteome. Nodes (circles) are connected by edges (straight lines) and have been grouped using ClueGO [35], depicting the molecular functions that were significantly over-represented in the NCC-participating subset, relative to the complete human proteome.

Interrogation of the human interactome [41] suggests that NCC causes aggregation of seemingly unrelated proteins (i.e. the NCC-interactome has low connectivity; electronic supplementary material, figure S7b) that participate in a high number of protein–protein interactions (especially, homotypic interactions; electronic supplementary material, figure S7b). That NCC preferentially affects fundamental enzyme complexes agrees well with the observation that NCC-participating proteins are evolutionarily more ancient than NCC-resistant proteins (figure 5c). Altogether, our data suggest that NCC preferentially shuts down a wide array of ancient proteins with essential interactions and functions. NCC is therefore likely to be a rudimentary response that ‘inactivates’ necrotic cells.

2.4. Putative role for nucleocytoplasmic coagulation in attenuating autoimmunity

Maintaining tolerance after injury is a necessary and problematic process, especially given that cell death can allow the unwanted exposure/release of epitopes that would otherwise be sequestered from the immune system. Indeed, cell death is a potent trigger of inflammation, with autoimmune disease often preceded by injury [24–27]. Moreover, defective or inadequate removal of dead cells underlies the onset of many autoimmune disorders including systemic lupus erythematosus [28]. Accordingly, we hypothesized that NCC, by restricting the escape of constituents from necrotic cells and by promoting tolerogenic clearance [21], is a favourable response that mitigates autoimmunity. To support this hypothesis, we assessed the representation of known autoantigens (using a set of 2079 autoantigens from Backes et al. [36]) and found that autoantigens were far more prevalent among the NCC-resistant subpopulation (figure 5d). Furthermore, the primary amino acid composition of autoantigens closely mirrored that of NCC-resistant proteins, with cysteines (p < 0.0001), aliphatic residues (I, V, L; p < 0.0001) and aromatic residues (F, Y, W; p < 0.0001) being under-represented, and charged residues (D, E, K, R; p < 0.0001) being over-represented in autoantigens (figure 5e). Collectively, this profile may allow autoantigens to evade NCC and thereby promote their release into the
extracellular space upon necrosis. Collectively, these findings add support to the hypothesis that NCC is a favourable response that may mitigate autoimmunity. Future studies should now test whether mutations that increase a protein’s tendency to undergo NCC also reduce its autoreactivity following necrosis—an approach that holds great appeal for investigating the basis of lupus and other injury-related autoimmune disorders.

2.5. Concluding remarks

A cascade of events that accounts for both our previous [10,12] and current findings is where NCC relies upon membrane disruption during necrosis (either primary or secondary necrosis). Membrane failure would cause a major loss of redox potential and widespread oxidation, which in turn would preferentially disulfide-crosslink abundant protein complexes within otherwise redox-reduced compartments. Our prior studies also suggest that initial misfolding events (e.g. línchpin residue oxidation) that precede necrosis may be a prerequisite for NCC [10,12]. We postulate that NCC represents a fail-safe switch that ensures irreversible metabolic and proteostatic incompetency during intense injury. The proteomic coverage of NCC is much wider than first thought, with a large number of essential proteins and macromolecular structures entrapped by NCC, thereby preventing their release into the extracellular space. Entrapment by NCC, however, is a selective process because a small proportion of the proteome (including FUS) avoids NCC and is therefore selectively released into the extracellular milieu upon necrosis. As the release of intracellular molecules upon necrosis is deleterious, NCC is likely to be a beneficial response that helps multicellular organisms restore tissue homeostasis after injury. This supposition is somewhat counterintuitive given that amyloidogenic protein aggregation promotes neurodegenerative diseases. However, based on the similar albeit distinct physicochemical signatures of NCC and amyloidogenesis, we hypothesize that NCC may be an ‘offshoot pathway’ that prevents deleterious amyloid formation in necrotic cells (electronic supplementary material, figure S4). Such interrelations between amyloidogenic aggregation and NCC should now be assessed. Beyond this, it is intriguing that features such as cysteine content, hydrophobicity, intrinsic disorder and abundance dictate whether a protein is retained or released from necrotic cells. This understanding broadens the biology of protein aggregation and indicates that reductionist biochemical approaches used in the field of protein misfolding may also be used to investigate the immunogenic and inflammatory nature of necrosis.

3. Material and methods

3.1. Materials

Reagents were from Sigma unless otherwise indicated. Soluble Fas ligand (SuperFasLigand) was from Enzo Life Science. NuPAGE Novex Bis-Tris Mini gel (4–12%), NuPAGE MOPS SDS Running buffer, RPMI media, penicillin and streptomycin, dialysed and non-dialysed fetal calf serum (FCS), TO-PRO-3 and 7-AAD were from Life Technologies. Immobilon-FL polyvinylidene difluoride (PVDF) membrane and Protein G Plus/Protein A agarose (50% suspension) beads were from Merck Millipore. Cell culture SILAC reagents were from Cambridge Isotope Laboratories. InstantBlue protein stain was from CBS Scientific. Sequence Grade Modified Trypsin (34 V511A), CytoTox96 non-radioactive cytotoxicity (LDH) and CellTiter96 aqueous non-radioactive cell proliferation (MTS) assays were from Promega. Odyssey Blocking Buffer and IRdye secondary antibodies (800CW donkey anti-mouse IgG, 680LT donkey anti-mouse IgG, 800CW donkey anti-rabbit IgG, 680LT donkey anti-rabbit IgG, 800CW donkey anti-goat IgG and 680LT donkey anti-goat IgG) were from LI-COR Biosciences. Primary antibodies were from Santa Cruz Biotechnology unless stipulated otherwise: anti-RIPK1 (BD Biosciences, #610458), anti-LC3 (MBT International, #PM036), mouse anti-HSP90β (Abcam, #ab82522), rabbit anti-α-tubulin (Merck Millipore, #04–1117), mouse anti-β-tubulin (Sigma, #T0198), goat anti-β-actin (#sc-1616), goat anti-PRX1 (#sc-7381), rabbit anti-Cyclophilin A (Abcam, #ab41684), goat anti-PKG1 (#sc-17943), mouse anti-3PDGH (#sc-100317), rabbit anti-EF1 (#sc-28578), goat anti-IMPDH2 (Abnova, #PAB19702), mouse anti-TCP1β (#sc-373769), mouse anti-MCM7 (#sc-56324), goat anti-EF2 (#sc-13004), mouse anti-LRP130 (#sc-166178), mouse anti-GAPDH (Merck Millipore, #MAB374), mouse anti-FUS/TLS (#sc-373698), mouse anti-SSRP1 (#sc-56781), mouse anti-Skip (#sc-136546), rabbit anti-ribosomal protein S19 (#sc-134779), mouse anti-Cdc5 L (#sc-81220), mouse anti-C23 (#sc-8031) and rabbit anti-GAPDH (#sc-25778).

3.2. Cell culture

Jurkat and U937 cells were maintained as non-adherent cultures in RPMI supplemented with 10%(v/v) heat-inactivated FCS, 2 mM L-glutamine, 50 U ml⁻¹ penicillin and 50 U ml⁻¹ streptomycin (P/S) under humidified 5% CO₂. AT3 cells were maintained as an adherent culture in DMEM supplemented with 10%(v/v) heat-inactivated FCS, 2 mM L-glutamine, 50 U ml⁻¹ penicillin and 50 U ml⁻¹ streptomycin (P/S) under humidified 5% CO₂. For injury induction, cells were resuspended in RPMI/DMEM supplemented with 1%(v/v) heat-inactivated FCS, P/S and L-glutamine. Cells were then seeded at 0.15 × 10⁶ live cells per well (for Jurkat and AT3 cells) or 0.1 × 10⁶ live cells per well (for U937 cells) in 24-well plates and allowed to equilibrate for 1 h. Injury-causing treatments or an equivalent amount of DMSO (as the vehicle) were then added to their final concentrations: 300 nM staurosporine, 25 µg ml⁻¹ etoposide (for Jurkat cells) or 6 µg ml⁻¹ etoposide (for U937 cells), 40 ng ml⁻¹ soluble Fas ligand and 500 mM doxorubicin. Injury was assessed 3–24 h later (as stipulated in the corresponding figure legend). For stable Lysine isotope incorporation, cells were maintained for more than 10 passages in Lysine- and Arginine-free RPMI/DMEM media supplemented with 10% dialysed FCS, P/S, l-glutamine, 0.1 g l⁻¹ of unlabelled l-Arginine·HCl and either 0.1 g l⁻¹ of unlabelled l-Lysine·2HCl or 0.1 g l⁻¹ of l-Lysine·2HCl (U¹³C/²H, 99%).

3.3. Cell injury assays (LDH, MTS and trypan blue assays)

LDH and MTS assay kits were from Promega and were performed according to the manufacturer’s instructions. Trypan blue positivity was determined using the TC-10 Automated Cell counter (Bio-Rad) according to the manufacturer’s instructions.

3.4. Flow cytometry

Jurkat and U937 cells (seeded and treated as in §3.2) were incubated for the indicated time period. Cells were then
incubated for a further 5 min with 1 mg l⁻¹ AlexaFluor488 conjugated-Annexin V and 250 µM TO-PRO-3 or 7-AAD and subject to flow cytometry (BD Accuri C6 Plus flow cytometer). Data were analysed off-line using FlowJo v. 10.1r5.

3.5. Cell homogenization and fractionation

Cells were collected, washed in PBS, pelleted and resuspended in ice-cold lysis buffer (PBS + 1% Triton X-100 + 1 x complete EDTA-free protease cocktail (Roche) + 1 x PhosSTOP phosphatase inhibitor cocktail (Roche) + 10 mM chloroacetamide). Chloroacetamide was added to the lysis buffer to prevent inadvertent disulfide bond formation during/after cell homogenization and subsequent fractionation. Note, chloroacetamide was chosen over iodoacetamide as it reacts more specifically with cysteine thiols [42]. Homogenates were trituated and then stored at −80°C until further fractionation was necessary; 1 ml of lysis buffer was used per 0.6 × 10⁶ cells (for Jurkat cells) or per 0.4 × 10⁶ cells (for U937 cells). To extract the soluble and insoluble fractions, 1 ml of homogenate was centrifuged (4°C, 16 000 g, 30 min). The supernatant was kept as the ‘Triton-soluble fraction’ and the pellet was washed in 1 ml of PBS + 1% Triton X-100 and incubated for 1 h with gentle mixing at 25°C in 60 µl of 62.5 mM Tris–HCl (pH 6.8), 2% SDS, 5% glycerol with or without 0.1 M DTT. Finally, the sample was centrifuged (25°C, 16 000 g, 30 min) and the supernatant kept as the ‘Triton-insoluble fraction’.

3.6. SDS-PAGE

3.6.1. SDS-PAGE for immunoblotting

Samples (50 µl of Triton-soluble fraction and the entire Triton-insoluble fraction) were boiled in 2% SDS-loading buffer with or without dithiothreitol, subjected to SDS-PAGE, and transferred onto PVDF membranes. Membranes were probed with the appropriate IR-dye-conjugated secondary antibody (LI-COR) and signals revealed with an Odyssey scanner (LI-COR).

3.6.2. SDS-PAGE for trypsic digestion

Samples were boiled in 2% SDS-loading buffer with dithiothreitol and subjected to SDS-PAGE using a NuPAGE Novex Bis-Tris Mini gel (4–12%) with NuPAGE MOPS SDS-Running buffer supplemented with 1 x NuPAGE antioxidant (Life Technologies). After electrophoresis, the gel was stained with InstantBlue protein stain and cut into approximately 2 x 6 mm slices then each slice further cut into 1 mm² pieces. Gel pieces were subjected to in-gel trypsin digestion as in [43].

3.7. Mass spectrometry

Liquid chromatography tandem mass spectrometry was performed as in [44] with minor modifications. Extracted peptides were analysed by Q-Exactive Orbitrap mass spectrometer (Thermo Scientific) coupled online with an RSLC nano-HPLC (Ultimate 3000, Thermo Scientific). Samples were injected onto a Thermo RSLC pepmap100, 75um id, 100 angstrom pore size, 50 cm reversed phase nano column with buffer A (2.5% acetonitrile, 0.1% formic acid) at a flow rate of 300 nl min⁻¹. The peptides were eluted over a 40-min gradient to 40% buffer B (80% acetonitrile 0.1%, formic acid). The eluate was nebulized and ionized using the Thermo nano electrospray source coated silica emitter with a capillary voltage of 1700 V. Peptides were selected for MS/MS analysis using Xcalibur software (Thermo Fisher) in Full MS/dd-MS2 (TopN) mode with the following parameter settings: TopN 10, MSMS AGC target 1e5, 60 ms Max IT, NCE 27 and 2 m/z isolation window. Dynamic exclusion was set to 20 s.

MS data were analysed with MaxQuant [45] software v. 1.3.0.5. Search parameters included: specific digestion with trypsin with up to two missed cleavages, +6 Lysine was set as label, protein N-terminal acetylation and methionine oxidation were set as variable modifications while cysteine alkylation was set as fixed modification. The searches were performed against UniProt human protein sequences (March 2013 version). The remaining search parameters were default settings. The search was then imported into Perseus software [45] for collation (electronic supplementary material, table S1) and for further annotation. Note that the gating of SILAC ratios for analysis is explained in the respective figure legends. These gating strategies were based on the recommendations of Zhang & Neubert [31], with more than 95% of the identified proteins having a ratio that fell within the ‘unchanged’ intermediary gate in the corresponding unijured cell samples.

3.8. Immunoprecipitation

Jurkat cells were treated for 24 h and the conditioned media collected. Fifteen microlitres of Protein-G/-A beads (as a 50% slurry) was added to 1.3 ml of conditioned media and the mixture gently rotated (30 min, 4°C). The beads were centrifuged (2 min, 3000 g) and discarded; 20 µl of Protein-G/-A beads + 1 µg of anti-MCM7 antibody was added to the supernatant and the mixture gently rotated (3 h, 4°C). The beads were centrifuged (2 min, 3000 g) and kept as the ‘MCM7 pulldown’ material; 20 µl of Protein-G/-A beads + 1 µg of anti-FUS antibody was added to the supernatant and the mixture gently rotated (3 h, 4°C). The beads were centrifuged (2 min, 3000 g) and kept as the ‘FUS pulldown’ material. The beads were boiled for 10 min in 35 µl of SDS-loading buffer with dithiothreitol and the supernatants subjected to SDS-PAGE/immunoblot analysis. Quantitation of the immunoprecipitation signal was performed using ImageQuant v. 5.2 software (Molecular Dynamics) and expressed relative to the total amount of FUS present in the Triton-soluble lysate from unijured Jurkat cells.

3.9. Immunofluorescence

3.9.1. Immunofluorescence for maleimide co-staining

Twenty-four hour-treated Jurkat cells were fixed in 2% paraformaldehyde with 40 mM chloroacetamide (4°C, 40 min, in dark). Cells were immobilized onto Superfrost slides (Tharmac Cellspin II, 320 g, 5 min) and incubated in 4% paraformaldehyde with 40 mM chloroacetamide (5 min, in dark). Cells were incubated in TBS-T (100 mM Tris–HCl pH 8.0 + 150 mM NaCl + 0.05% Tween20) with 40 mM chloroacetamide (20 min in dark). Cells were washed then incubated in TBS-T with 20 mM TCEP (20 min in dark). Cells were washed then incubated in TBS-T with 50 nM AlexaFluor488-conjugated maleimide (20 min in dark). Cells were washed then incubated in...
Using the FoldAmyloid [39] and Zyggregator [40] databases, similar analyses (with similar results) were performed in vivo. Penalties are derived from soluble polypeptide [33]. AGGRESCAN’s aggregation propensity is determined using the FoldAmyloid [39] and Zyggregator [40] databases.

3.10. Protein property analysis

3.10.1. Protein sequence analysis

Canonical FASTA sequences for the different protein subsets (electronic supplementary material, table S1) were retrieved from UniProt and entered into the Sequence Manipulation Suite [46] to determine amino acid content, hydrophobicity and molecular weight.

3.10.2. Protein aggregation prediction

Canonical FASTA sequences for the different protein subsets (electronic supplementary material, table S1) were retrieved from UniProt and uploaded to the AGGRESCAN [33] site to ascertain the predicted aggregation propensity. Note, only proteins with less than 1000 residues were analysed. AGGRESCAN provides the relative aggregation propensity of amino acids within both the central hydrophobic cluster of β-amyloid and within the context of a larger full-length soluble polypeptide [33]. AGGRESCAN’s aggregation propensities are derived from in vivo experimental data [33]. Similar analyses (with similar results) were performed using the FoldAmyloid [39] and Zyggregator [40] databases (not shown).

3.10.3. Intrinsic disorder content analysis

UniProt IDs (electronic supplementary material, table S1) were used to query the MobiDB 2.0 database [47]. The percentage of each protein in disorder (either from curated data or from consensus predicted data when no curated data existed) irrespective of segment was determined.

3.10.4. Autoantigen analysis

UniProt IDs (electronic supplementary material, table S1) were cross-referenced against the full set of autoantigens from Backes et al. [36].

3.10.5. Protein abundance analysis

The different protein subsets (electronic supplementary material, table S1) were uploaded to the PaxDB [48] site and their organisinal abundance was retrieved from the Human PaxDB integrated dataset.

3.10.6. Protein age analysis

Protein age was determined using the ProteinHistorian [49] website, whereby protein subsets were individually analysed via the swisspfam-PTH1R7_youngest database and the Dollo parsimony reconstruction algorithm.

3.10.7. Native disulfide bond analysis

Annotations from the UniProt [50] database were used to determine the number and location of native disulfide bonds for each protein of interest.

3.10.8. Ontology enrichment analysis

Cytoscape v. 3.2.0 and the ClueGO v. 2.2.5 and CluePedia v. 1.2.5 plug-ins were used to perform ontology enrichment analysis as in [35]. Molecular function enrichment analysis was performed within GO term levels 3–5. Cellular compartment enrichment analysis was performed within GO term levels 4–10.

3.10.9. Protein–protein interaction analysis

The high-confidence NCC-participating protein subset was entered into the Mentha [51] interactome database. All human interacting proteins were extracted and visualized as a network using Cytoscape. Duplicate interactions were removed and the network rendered directionless. The topology and characteristics of the resultant network was analysed using the Network Analyzer plug-in [52].

3.11. Statistical analyses

All results are based on three independent experiments unless stipulated otherwise. Analyses were performed with GraphPad Prism 6 with p < 0.05 considered statistically significant. The analysis employed for each cohort is stated in the respective figure legend.

Data accessibility. All mass spectrometry files used in the manuscript will be deposited in mzML format on the ProteomeXchange website (http://www.proteomexchange.org/).

Authors’ contributions. A.L.S. conceived, designed, and conducted experiments, analysed data and drafted the manuscript. B.H. conducted experiments, analysed data and helped draft the manuscript. M.J.S. provided research tools and helped draft the manuscript. S.P.B. helped conceive the study. O.K. helped design experiments, conducted experiments, analysed data and helped draft the manuscript. R.L.M. helped conceive the study and helped draft the manuscript. All authors gave final approval for publication.
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