Endoplasmic reticulum polymers impair luminal protein mobility and sensitise to cellular stress in α₁-antitrypsin deficiency

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Abstract

Point mutants of α₁-antitrypsin form ordered polymers that are retained as inclusions within the endoplasmic reticulum (ER) of hepatocytes in association with neonatal hepatitis, cirrhosis and hepatocellular carcinoma. These inclusions cause cell damage and predispose to ER stress in the absence of the classical unfolded protein response (UPR). The pathophysiology underlying this ER stress was explored by generating cell models that conditionally express wildtype α₁-antitrypsin, two mutants that cause polymer-mediated inclusions and liver disease (E342K [the Z allele] and H334D) and a truncated mutant (Null Hong Kong, NHK) that induces classical ER stress and is removed by ER associated degradation. Expression of the polymeric mutants resulted in gross changes in the ER luminal environment that recapitulated the changes seen in liver sections from individuals with PI*ZZ α₁-antitrypsin deficiency. In contrast expression of NHK α₁-antitrypsin caused electron lucent dilatation and expansion of the ER throughout the cell. Photobleaching microscopy in live cells demonstrated a decrease in the mobility of soluble luminal proteins in cells that express E342K and H334D α₁-antitrypsin when compared to those that express wildtype and NHK α₁-antitrypsin (0.34±0.05, 0.22±0.03, 2.83±0.30 and 2.84±0.55 μm²/s respectively). There was no effect on protein mobility within ER membranes indicating that cisternal connectivity was not disrupted. Polymer expression alone was insufficient to induce the UPR but the resulting protein overload rendered cells hypersensitive to ER stress induced by either tunicamycin or glucose depletion.

Conclusion—Changes in protein diffusion provide an explanation for the cellular consequences of ER protein overload in mutants that cause inclusion body formation and α₁-antitrypsin deficiency.

Keywords
cirrhosis; serpins; UPR; diffusion; FRAP-FLIP
Introduction

Alpha₁-antitrypsin deficiency results from point mutations that lead to the formation of large, ordered polymers that accumulate within the ER of hepatocytes. The resulting inclusions cause a toxic-gain-of-function that is associated with neonatal hepatitis, cirrhosis and hepatocellular carcinoma. This is best exemplified by the Z (E342K), King’s (H334D), Siiyama (S53F) and Mmalton (Δ52F) alleles. Plasma efficiency of α₁-antitrypsin deficiency can also result from truncating mutations such as the Null Hong Kong (NHK) allele that target the protein for ER-associated degradation (ERAD). This reduces secretion of α₁-antitrypsin but does not cause the gain-of-function liver disease. In both cases the lack of circulating α₁-antitrypsin predisposes the homozygote to early onset emphysema.

A wide range of signalling pathways are activated to restore the ER homeostasis in response to an overload of misfolded proteins. An important stress response is the Unfolded Protein Response (UPR) that combines transient attenuation of protein translation with enhancement of the ER’s protein-folding capacity, thus preventing uncontrolled aggregation of proteins within its lumen. This conserved stress response is tightly regulated by the activation of three ER stress sensors operating in parallel: IRE1, PERK, and ATF6. Under non-stressed conditions, each sensor is held in an inactive state by binding of the ER chaperone BiP to its luminal domain. When misfolded protein levels increase during ER stress, the UPR is activated, at least in part, through the titration of BiP away from these sensor molecules.

We aimed to determine the impact of polymers and truncated α₁-antitrypsin on the ER environment. We show that the accumulation of ordered polymers of α₁-antitrypsin leads to gross expansion of ER cisternae and impairment of ER luminal mobility. This hinders the organelle’s capacity to deal with otherwise minor perturbations of protein folding and thus leads to enhanced sensitivity to ER stress. Our findings provide one of several potential mechanisms for the toxicity associated with ER overload in α₁-antitrypsin deficiency related liver disease.

Material and Methods

Plasmid constructs

cDNA for human wildtype α₁-antitrypsin and two polymeric mutants (E342K and H334D) were subcloned into the pTRE2hyg vector (Clontech) for the generation of stable CHO-K1 Tet-On α₁-antitrypsin cell lines. The NHK truncated α₁-antitrypsin variant (L318fsX17) was generated with the QuickChange™ Site-Directed Mutagenesis kit (Stratagene) using wildtype α₁-antitrypsin as the template and then subcloned into the pTRE2hyg plasmid.

Generation and characterization of stable cell lines expressing α₁-antitrypsin

The CHO-K1 cell line was purchased from Clontech and stable cell lines were generated by following the manufacturer’s instructions. The cells were maintained in DMEM supplemented with 10% v/v Tet-FBS, 1% w/v non-essential amino acids, 1% w/v penicillin/streptomycin, 200μg/ml Geneticin, and 500μg/ml Hygromycin B (both selective antibiotics from Invitrogen) at 37°C and 5% v/v CO₂. α₁-Antitrypsin was typically induced with 1μg/ml doxycycline for 48h. Whole cell lysates, SDS and non-denaturing polyacrylamide gel electrophoresis (PAGE) followed by immunoblotting, ELISA analysis and metabolic labelling were performed as detailed previously (see Supporting Materials and Methods).
**Immunofluorescence, electron microscopy and immunocytochemistry**

Immunofluorescence, electron microscopy and immunocytochemistry of CHO-K1 cells and paraffin tissue slides from human subjects were undertaken as described in the Supporting Materials and Methods.

**Photobleaching analysis in live cells**

CHO-K1 Tet-On cells were transiently transfected with an ER-RFP plasmid or a CytERM msfGFP membrane reporter. Live cells were imaged on a 37°C environmental controlled stage of a confocal microscope system (Duoscan; Carl Zeiss Inc.) with a 63x/1.4NA oil objective and a 40 mW 561nm diode laser with a 565nm bandpass filter. For FRAP/FLIP experiments, a small region of interest was photobleached at full laser power in the 561 line and fluorescence recovery or loss was monitored over 0.2s time intervals. Diffusion \(D_{eff}\) measurements were calculated as described previously. Fluorescence loss curves were obtained by transforming the fluorescence intensities into a percentage scale in which the first bleach time point represents 100% of fluorescence intensity. Statistical significance was assessed using Student’s t test.

**Detection of the UPR markers**

Immunoblots for three main ER luminal chaperone proteins (GRP94, BiP/GRP78, PDI), the ATF6-reporter luciferase assay and the RT-PCR XBP1 splicing assay were performed as described previously.

**Results**

**Conditional expression of α₁-antitrypsin**

Stable CHO-K1 cell lines were developed using the Tetraycline-On (Tet-On) inducible system that enables tight regulation of α₁-antitrypsin expression. Four clonal cell lines were generated to express four single variants of α₁-antitrypsin: the wildtype human α₁-antitrypsin (M); the polymeric Z [E342K] mutant; the polymeric King’s [H334D] mutant (both form ordered polymers in the ER of hepatocytes in association with liver disease); and a NHK truncated variant that does not fold properly, preventing the formation of ordered polymers, and instead activates the UPR. Expression of each variant was readily detected on treating cells with doxycycline. Wildtype α₁-antitrypsin was almost completely secreted, compared with Z and H334D α₁-antitrypsin, which were largely retained as polymers. In contrast, NHK α₁-antitrypsin failed to reach the culture medium despite being cleared from the cell lysates at 4h, consistent with its degradation by ER associated degradation (ERAD).

**Accumulation of α₁-antitrypsin polymers causes disruption of the ER**

Immunofluorescence confocal microscopy revealed that both Z and H334D α₁-antitrypsin co-localised with the ER marker BiP, although in many cells the ER structure was distorted from its reticular morphology into a punctate pattern. No gross morphological changes were observed in cells expressing either M or NHK α₁-antitrypsin. Of note, only M α₁-antitrypsin co-localised with the Golgi marker p115, indicating efficient exit from the ER. Electron microscopy was used to visualise cells expressing wildtype and mutant α₁-antitrypsin variants and to examine cell morphology and the location of the polymers.
α₁-antitrypsin. The ultrastructure in cells expressing M α₁-antitrypsin was normal (Fig. 2B; a–b) when they were compared with mock-transfected cells (Fig. 2B; i–k). The inclusions in cells expressing either Z or H334D α₁-antitrypsin were associated with the replacement of normal tubular and lamellar ER by electron-dense vesiculated structures (Fig. 2B; c–f). In contrast, expression of the NHK variant induced a markedly different phenotype with electron lucent dilatation and expansion of the ER throughout the cell (Fig. 2B; g–h). In order to elucidate whether the alterations observed could be due to a massive overexpression of α₁-antitrypsin in our model, we also assessed the distribution and signal of total α₁-antitrypsin in paraffin-embedded liver sections from a PI*ZZ α₁-antitrypsin homozygote and control human liver by immunocytochemistry (Fig. 2C). The PI*ZZ homozygote human liver demonstrated levels of polymer accumulation at least as dramatic as that seen in cultured cells, suggesting that the CHO-K1 Tet-On cells represent a valid model of the accumulation of α₁-antitrypsin polymers within the ER. Taken together, these data show that the accumulation of ordered polymers within the ER leads to gross changes in the organelle’s morphology and importantly that these changes differ markedly from those induced by the NHK mutant that causes classical ER stress.

Polymeric mutants of α₁-antitrypsin impair ER protein mobility in live cells

The impact of polymer formation on the ER luminal environment was determined by monitoring changes in the mobility of the ER-RFP fluorescence marker in live cells. ER-localised RFP is a protein (27kDa and 2.3 nm hydrodynamic radius) with no known interacting partners that can rapidly sample the entire ER lumen, reporting on both crowdedness and interconnectivity. The ER-RFP distribution revealed two distinct ER architectures in cells expressing α₁-antitrypsin variants. Each cell line displayed a typical ER reticular network in the absence of doxycycline, as did cells expressing M and NHK α₁-antitrypsin when treated with doxycycline (Fig. S2). In contrast, approximately 30–40% of cells expressing Z and H334D α₁-antitrypsin developed a characteristic vesiculated architecture when treated with doxycycline (Fig. S2), as detected by staining for either α₁-antitrypsin or BiP (Fig. 2A). Within the same clonal cell lines there remained a proportion of cells expressing Z and H334D α₁-antitrypsin that retained the normal reticular pattern of ER, suggesting that formation of inclusion bodies is a stochastic phenomenon (Fig. S3).

We then assessed the ER lumen continuity by fluorescence loss in photobleaching (FLIP), in which a small region of the cell ER-RFP fluorescence was repeatedly photobleached. If the protein is mobile within a continuous compartment, the total fluorescence within this compartment will be depleted. Over time, ER-RFP fluorescence in the whole cell was steadily and uniformly lost in cells displaying a tubular ER network, whereas fluorescence depletion was substantially slower in cells containing Z α₁-antitrypsin polymer (Fig. 3A and Movies S1 and S2). This did not reflect an effect of transfected ER-RFP expression levels, since normalised plots showed that the remaining whole cell fluorescence intensity after 200 s was 23.5% and 61.1% for cells retaining a tubular ER network or a polymer-filled ER, respectively (Fig. 3A). Similar results were obtained for cells expressing H334D α₁-antitrypsin (Fig. 3B). Thus, the presence of vesiculated structures is associated with marked loss of intra-ER luminal communication (slower ER-RFP mobility between inclusions) that is likely to adversely increase the heterogeneity of the ER luminal environment.

FRAP analysis was used to quantify the change in mobility of the ER-RFP reporter (Fig. 4A). This provides information on local rates of protein diffusion, $D_{eff} (\mu m^2/s)$ by bleaching a small defined region of a single cell. $D_{eff}$ is affected by local viscosity, size of the reporter protein and its protein-protein interactions and is inversely proportional to crowdedness of the environment. Moreover, small changes in $D_{eff}$ are likely to be biologically significant. No significant differences in the mobility of ER-RFP were detected in the different cell lines.
in the absence of α₁-antitrypsin expression (Fig. 4B). ER-RFP $D_{\text{eff}}$ values in cells expressing NHK and M α₁-antitrypsin were identical (2.84±0.30 vs. 2.83±0.55 μm²/s). However, cells containing polymers of either Z or H334D α₁-antitrypsin exhibited significantly lower ER-RFP $D_{\text{eff}}$ values compared with M α₁-antitrypsin (0.34±0.05 and 0.22±0.03 μm²/s respectively; Figs. 4A and B, and Movies S2 and S3).

The striking effect of polymers on the ER-RFP diffusion within the lumen of the ER, prompted an assessment of their effect on the diffusion of an ER transmembrane reporter. We used the CytERM msfGFP plasmid, which encodes a signal domain required to anchor the fluorescent protein into the ER membrane with the sfGFP exposed to the cytoplasm. There is no luminal domain and consequently the luminal content has no effect on membrane protein diffusion. There were no significant differences in CytERM msfGFP $D_{\text{eff}}$ values in cells containing polymers of H334D α₁-antitrypsin compared to cells in which expression was not induced (0.35±0.04 vs. 0.40±0.02 μm²/s) (Fig. 4C).

Taken together, these data demonstrate that inclusions of α₁-antitrypsin polymers have a remarkable effect on the ER environment reducing luminal protein diffusion. However protein mobility within ER membranes is unaffected, indicating that cisternae connectivity is not disrupted.

**Increased susceptibility to ER stress in polymer expressing cells**

Previous work has shown that the ER accumulation of ordered polymers of mutant α₁-antitrypsin and another member of the same protein family, neuroserpin, fail to activate the UPR. However, a recent report has suggested that Z α₁-antitrypsin polymer accumulation triggers UPR activation in peripheral blood monocytes. We hypothesised that the alteration of the ER luminal environment might destabilise ER homeostasis sufficiently to sensitise the cell to a second insult and so trigger activation of the UPR. The activation of multiple reporters of UPR signalling was monitored following treatment with tunicamycin, an inhibitor of N-linked glycosylation. Immunoblot analysis for three ER luminal chaperones (BiP/GRP78, GRP94 and PDI) confirmed that ordered polymers of α₁-antitrypsin alone failed to activate the UPR (Fig. 5A; lanes 1–3). In contrast and as expected, synthesis of the truncated NHK α₁-antitrypsin led to robust activation of the UPR (Fig. 5A; lane 4). Remarkably, both polymerogenic α₁-antitrypsin variants showed a hypersensitive response in UPR activation following treatment with low concentrations of tunicamycin (5–25 ng/ml) when compared to the wildtype protein (Fig. 5A).

A more quantitative reporter assay was used to assess the activation of the ATF6 branch of the UPR. Only cells expressing NHK α₁-antitrypsin showed measureable activation of ATF6 under non-stressed conditions (Fig. 5B). However cells expressing polymeric variants were more susceptible to activation of ATF6 on treatment with moderate concentrations of tunicamycin (60–300 ng/ml). A similar response was seen when cells were treated with low concentrations of glucose (10-0.2 mM), which represents a more physiological ER stress (Fig. 5C).

IRE1 activation was assessed by the splicing of XBP1 mRNA. None of the cell lines showed spliced XBP1 mRNA under basal conditions. However, tunicamycin (300 ng/ml) efficiently triggered IRE1 in all cell lines as reported by the appearance of a smaller spliced XBP1 mRNA band (Fig. 5D). The failure to detect splicing of XBP1 mRNA in cells induced to express the NHK variant in the absence of tunicamycin reflects the transitory nature of IRE1 activation. Following the induction of ER chaperones, which occurs rapidly on expressing this truncated mutant, ER stress is alleviated and IRE1 returns to its inactive state. Indeed, the elevated level of ER chaperones in NHK-expressing cells served to protect them from subsequent ER stress, as evidenced by the blunted response of XBP1 splicing seen in these
cells (Fig. 5D). In contrast, on treatment with tunicamycin, M, Z- and H334D-expressing cells showed rapid splicing of XBP1 mRNA. These data indicate that the presence of polymers does not affect the intrinsic activity of IRE1. They do, however, impair the cell’s ability to restore ER homeostasis, since higher levels of spliced XBP1 mRNA persisted after 16h treatment with tunicamycin in cells expressing both Z and H334D α1-antitrypsin compared to the wildtype protein.

It is possible that differences in the sensitivity to tunicamycin might be responsible for the enhanced UPR activation in cells that express polymers of α1-antitrypsin. However this was not the case as treatment with tunicamycin resulted in equal levels of a slower migrating glycosylated form of α1-antitrypsin and a faster migrating non-glycosylated form in each α1-antitrypsin expressing cell line (Fig. S4).

Discussion

It is well recognised that point mutations of α1-antitrypsin form ordered polymers that are retained as inclusions within the ER, resulting in ER dysfunction and liver disease. These inclusions form in the face of effective disposal of misfolded monomer and polymers by ERAD and autophagy. One of the cellular consequence of the accumulation of these ordered polymers is the ER overload response (EOR), which results in the release of ER calcium and the activation of NF-κB, a central mediator of inflammation. This pathway is distinct from the canonical UPR, which is triggered by the accumulation of misfolded protein and results in the inhibition of protein synthesis, induction of ER chaperones and ER stress. The ER overload response and UPR pathways often occur together but mutants of α1-antitrypsin provide ideal tools to probe these pathways as the polymer forming mutants activate only the EOR, whilst truncated mutants activate only the UPR.

We have used two polymeric mutants of α1-antitrypsin (E342K and H334D) to determine the mechanism by which ER overload can impact on the organization of the ER environment and the cell’s susceptibility to ER stress. Our data reveal that polymers of α1-antitrypsin accumulate in spherical, apparently vesiculated ER, concomitantly with loss of the normal branching tubule network. Similar ER vesiculation has been reported to be associated with other cellular stresses, including mechanical injury and elevated cytosolic calcium concentration, although the physiological relevance of this remains unclear.

Live-cell imaging revealed that accumulation of polymers in vesiculated ER leads to a marked and specific impairment of luminal protein mobility due to an increased ER heterogeneity, which might reflect a change in either the luminal viscosity and/or in the tortuosity for ER-RFP diffusion within and between spherical inclusions. Interestingly, the truncated NHK mutant that causes classical ER stress (UPR) and is efficiently degraded by the proteasome showed a different ultrastructural change characterised by gross expansion of ER cisternae. This had no effect on luminal protein mobility. Current technology limits the area of photobleaching to slightly larger than the size of an inclusion. However, since effects of photobleaching were examined by FLIP throughout the cell, we are confident in these findings. It will be valuable, when the technology has improved sufficiently, to revisit the diffusion between individual inclusions.

In agreement with previous studies, we report that cells expressing polymers of either Z or H334D α1-antitrypsin do not show constitutive activation of the UPR. However these cells do exhibit a hypersensitivity to both pharmacological and physiological ER stressors. To date, most studies that have attempted to detect ER stress in cells expressing mutant α1-antitrypsin have made use of methods such as the up-regulation of ER chaperones. Although providing important insights, such approaches have been unable to address how ER overload with polymers of α1-antitrypsin might impact on the molecular organization of the ER.
organelle. Our observation that enhanced sensitivity to ER stress following the expression of polymerogenic α₁-antitrypsin mutants correlates with the presence of marked changes in the biophysical features of the ER suggests a mechanism for this phenomenon (Fig. 6). Chaperones can readily diffuse to sites of protein misfolding in cells possessing a reticular and highly interconnected ER. In contrast, in cells experiencing ER overload, a comparable level of misfolded client protein cannot diffuse freely, decreasing their accessibility to quality control proteins required for folding and transport. Therefore, we propose a model in which decreased mobility or availability of ER chaperones due to changes in the diffusive features of the ER, or obstruction caused by protein overload, sensitises the cell to subsequent activation of the UPR by a second hit that further increases the unfolded secretory protein burden. Our findings are directly relevant to the liver disease associated with polymerogenic mutants of α₁-antitrypsin. However, although polymer accumulation involves a gain-of-function associated with liver dysfunction, epidemiological studies have shown that only 10% of patients with α₁-antitrypsin deficiency develop clinically apparent liver disease. It appears that additional, incompletely understood genetic and environmental factors contribute to this heterogeneity in hepatic phenotype. Our results suggest that polymer accumulation increases the vulnerability of the cell to a “second hit” that ultimately initiates liver disease in individuals with α₁-antitrypsin deficiency. Moreover, the mechanism proposed is likely to be relevant to other serpinopathies, such as the neuronal toxicity associated with polymers of neuroserpin in the dementia FENIB.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations

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<tr>
<td>α₁AT</td>
<td>α₁-antitrypsin</td>
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<tr>
<td>D&lt;sub&gt;eff&lt;/sub&gt;</td>
<td>Effective diffusion coefficient</td>
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<td>Dox</td>
<td>Doxycycline</td>
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<td>EOR</td>
<td>Endoplasmic overload response</td>
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<td>ER</td>
<td>Endoplasmic reticulum</td>
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<td>FLIP</td>
<td>Fluorescence loss in photobleaching</td>
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<td>FRAP</td>
<td>Fluorescence recovery after photobleaching</td>
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<td>Tm</td>
<td>Tunicamycin</td>
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<td>UPR</td>
<td>Unfolded protein response</td>
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References


Fig. 1. Characterisation of CHO-K1 Tet-On cell lines expressing different variants of α₁-antitrypsin (α₁AT): wildtype (M), Z (E342K), HD (H334D) and the truncated Null Hong Kong (NHK) variant

A) Cell lysates and medium were collected after induction of α₁-antitrypsin with 1μg/ml doxycycline (dox) for 48h and analysed by 10% w/v SDS and non-denaturing PAGE followed by immunoblot for α₁-antitrypsin. B) Percentage of secreted total α₁-antitrypsin and quantification of intracellular polymers for M, Z and HD α₁-antitrypsin were quantified by sandwich ELISA using the 9C5 (left panel) or 2C1 (right panel) monoclonal antibodies, that recognise all forms of α₁-antitrypsin or only polymers, respectively. The graphs show mean±SEM (n=4). C) CHO-K1 Tet-On cells expressing M, Z, HD and NHK α₁-antitrypsin were pulse-labeled with 35S-Met/Cys for 15min and chased for the times indicated (0 – 4h). α₁-Antitrypsin from cell lysates and medium was immunoprecipitated with a polyclonal antibody (total) or the 2C1 monoclonal antibody (polymers) and then samples were resolved by 10% w/v SDS PAGE and detected by autoradiography. Gels are representative of three independent experiments.
Fig. 2. Intracellular accumulation of α₁-antitrypsin polymers in inclusion bodies disrupts ER structure
A) Representative fluorescence images of Tet-On CHO-K1 cells induced to express M, Z, HD and NHK α₁-antitrypsin with dox for 48h, fixed and co-immunostained with an α₁-antitrypsin polyclonal antibody and either an ER protein marker (BiP) or a Golgi protein marker (p115). White arrows in the merge panels show a punctate pattern of co-localised polymerogenic forms of α₁-antitrypsin (Z and HD) with the ER marker (left panel) or wildtype α₁-antitrypsin co-localised with the Golgi marker (right panel). B) Electron microscopy analysis for ER structure in Tet-On CHO-K1 cells expressing α₁-antitrypsin variants. Upper panels represent whole cells, with higher power images shown in lower panels. Black arrows indicate the ER cisterns. Inclusion bodies were only identified in micrographs of the polymerogenic mutants (Z and HD α₁-antitrypsin). C) Immunostaining of paraffin-embedded liver sections from a PI*ZZ α₁-antitrypsin individual and a normal control liver with a polyclonal antibody to total α₁-antitrypsin (white arrow).
Fig. 3. Luminal α₁-antitrypsin inclusion bodies impair protein mobility within the ER lumen
A) FLIP analysis of Z α₁-antitrypsin cells expressing ER-RFP within both a tubular ER
network cell (without dox) and an α₁-antitrypsin polymer-filled ER (with dox for 48h). FLIP
normalised plots setting the pre-bleach intensity as 100%. B) FLIP of H334D α₁-antitrypsin
cells expressing ER-RFP within both a tubular ER network cell and an α₁-antitrypsin
polymer-filled ER from the same sample after treatment with dox for 48h. Normalised plots
from FLIP analysis as described in 3A. Fluorescence loss was specific, as adjacent cells
remained fluorescent (white arrow). White boxes represent the region subjected to
photobleaching.
Fig. 4. Polymers of α1-antitrypsin modify the ER luminal environment decreasing ER protein mobility

A) Left: FRAP series of Tet-On CHO-K1 cells induced to express M, Z, HD and NHK α1-antitrypsin with dox for 48h and transiently co-transfected with ER-RFP. Cells expressing either Z or HD α1-antitrypsin show a vesiculated pattern in which the fluorescence recovery was significantly reduced. All images were captured immediately before (pre-bleach), immediately after (post-bleach) and at times after photobleaching in the white box. Scale bar: 10 μm. Right: $D_{eff}$ values from FRAP analysis in individual cells. B) Mean values±SEM for each condition and variant in 4A. C) Upper panel: $D_{eff}$ values from FRAP analysis in individual Tet-On HD CHO-K1 cells treated with or without dox for 48h and transiently co-transfected with both ER-RFP and CytERM msfGFP membrane reporter. Lower panel: Mean±SEM for each condition and construct. *** p<0.0001 in a two-tailed Student $t$ test.
Fig. 5. Differential activation of three UPR sensors in polymeric forms of α1-antitrypsin reveals an increased sensitivity to UPR following a second hit

A) Immunoblot of whole cells lysates from CHO-K1 Tet-On cells induced with dox for 48h and treated with the indicated doses of tunicamycin (Tm) for 16h. Grouping of separated lanes within the same gel was performed and indicated by dividing white lines. Histogram representation of four independent replicates of experiment 5A showing fold increase of GRP94, BiP/GRP78 or PDI normalised to loading control and then to wildtype variant for each concentration of tunicamycin (mean±SEM). B and C) ATF6 activation during induction of ER stress. α1-Antitrypsin expression was induced with dox for 48h. Twenty-four hours prior to lysis, the cells were co-transfected for 6h with a p5xATF6-Luc encoding firefly luciferase under the control of a UPR response element and with pRL-TK Renilla luciferase as a normalization control, and then treated with the indicated doses of tunicamycin (B) or glucose (C) for 16h. The graphs show firefly luciferase normalised to Renilla luciferase, as mean±SEM (n=3). C) XBP1 processing during ER stress induction. Cells were induced with dox for 48h and subjected to a time course of tunicamycin
treatment (300 ng/mL). RT-PCR was used to amplify \textit{XBP1} mRNA and PCR products were resolved by 2% w/v agarose gel electrophoresis. The ratio of spliced/unspliced \textit{XBP1} mRNA for each \textalpha{}\textsubscript{1}-antitrypsin variant after 16h tunicamycin is represented graphically as mean ±SEM (n=3). *\textit{p}<0.05, **\textit{p}<0.01, or ***\textit{p}<0.001 according to ANOVA test followed by a Bonferroni \textit{post hoc} test.
Fig. 6. Model for the enhanced sensitivity to ER stress seen in cells experiencing ER overload

An insult resulting in misfolded client protein is effectively buffered by chaperone diffusion in healthy cells. This prevents ER stress ("resolved"). In contrast, in cells that accumulate polymerised protein, the global ER environment is affected that impairs chaperone access to misfolded proteins thereby increasing the cell’s propensity to experience ER stress ("unresolved"). Yellow represents a resolved ER stress, while orange corresponds to an unresolved ER response to ER overload.