The BAF A12T mutation disrupts lamin A/C interaction, impairing robust repair of nuclear envelope ruptures in Nestor–Guillermo progeria syndrome cells

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ABSTRACT

Nestor–Guillermo progeria syndrome (NGPS) is caused by a homozygous alanine-to-threonine mutation at position 12 (A12T) in barrier-to-autointegration factor (BAF). It is characterized by accelerated aging with severe skeletal abnormalities. BAF is an essential protein binding to DNA and nuclear envelope (NE) proteins, involved in NE rupture repair. Here, we assessed the impact of BAF A12T on NE integrity using NGPS-derived patient fibroblasts. We observed a strong defect in lamin A/C accumulation to NE ruptures in NGPS cells, restored upon homozygous reversion of the pathogenic BAF A12T mutation with CRISPR/Cas9. By combining in vitro and cellular assays, we demonstrated that while the A12T mutation does not affect BAF 3D structure and phosphorylation by VRK1, it specifically decreases the interaction between BAF and lamin A/C. Finally, we revealed that the disrupted interaction does not prevent repair of NE ruptures but instead generates weak points in the NE that lead to a higher frequency of NE re-rupturing in NGPS cells. We propose that this NE fragility could directly contribute to the premature aging phenotype in patients.

INTRODUCTION

The nuclear envelope (NE) is a critical double membrane structure that surrounds and encloses the nucleus, maintaining the organization of the chromatin (1,2), controlling nucleocytoplasmic transport and allowing the transduction of mechanical signals from the cytoplasm into the nucleus (3,4). The NE is made of the inner and outer nuclear membranes (INM and ONM). The lamina, which is a meshwork of intermediate filaments of A-type (lamins A and C, encoded by LMNA) and B-type (lamin B1 and B2, encoded by LMNB1 and LMNB2, respectively) lamin proteins, lies at the nucleoplasmic side of the INM (5). Lamin filaments interact with LEM (LAP2-emerin-MAN1) domain proteins that are embedded in the INM, with chromatin and with other nuclear proteins. These interactions play critical roles in maintaining the structural integrity of the nucleus (6,7).

The importance of the lamina is evident by the numerous diseases arising from mutations in LMNA or in genes encoding for NE-associated proteins. These mutations compromise the integrity of the lamina and of the NE, causing a range of laminopathies (8,9) including premature aging syndromes, muscular dystrophies and neuropathies (10–12). One of the consequences of NE destabilization is the appearance of NE ruptures that cause loss of nuclear compartmentalization. This has been observed in laminopathy patient cells and animal models (13–15), in cells undergoing a viral infection (16,17) or lacking specific components of the lamina (13,18). Additionally, mechanical stress can cause NE rupture, for example in vivo when cells are migrating through dense tissues (19,20), or in vitro when cells are cultured in 2D on stiff substrates (22).

A NE rupture is typically preceded by the formation of a gap in the nuclear lamina, generating a weak point at the NE, more prone to deformation by mechanical stress (23,24). This allows the formation of a protrusion of the nuclear membrane that under continued mechanical stress will grow and eventually rupture (19,21,25). The exposure and leakage of the nuclear content—including the chromosomal DNA—into the cytoplasmic compartment can cause DNA damage (19,20,26) and activation of innate immune sig-
nalling pathways, such as eGAS/STING that can trigger inflammation (19,20,27,28). Importantly, the cells are able to detect and seal a NE rupture within minutes, through the recruitment of several proteins to the site of rupture. These include barrier-to-autointegration factor (BAF), LEM domain proteins (including emerin) and ESCRT-III components (19,20,29–31). Finally, lamin A/C accumulates at the sites of ruptures, leaving behind what was defined as a lamin ‘scar’ (20,21,24), whose role remains unclear.

BAF is a small (89 amino acid) protein that localizes to the nucleus, at the NE and in the cytoplasm. BAF dimerization allows for its binding to LEM domain proteins and the Ig-fold domain of lamin A/C (32,33), while each individual subunit can bind dsDNA (34,35). BAF has been previously involved in the regulation of transcription, viral defence and postmitotic nuclear envelope reassembly (5,36–38). More recently, BAF was found to play a direct role in repairing ruptures of the NE by recruiting LEM domain proteins and membranes to the sites of rupture (29,31).

The interest around the function of BAF at the NE has grown since an alanine to threonine homozygous missense mutation at position 12 (Ala12Thr–BAF A12T) was identified as the cause of a premature aging disorder termed Nestor–Guillermo Progeria Syndrome (NGPS) (39–41). So far, only three NGPS patients have been identified and they all carry the same homozygous A12T mutation, inheriting one mutated copy from each of their parents, both carriers of a heterozygous BAF A12T mutation and devoid of disease. On the contrary to the classic Hutchinson–Gilford progeria syndrome (HGPS), caused by heterozygous LMNA mutations (11,42), NGPS patients live over 30 years and do not display any vascular or cardiovascular dysfunction. Instead, in addition to ageing phenotypes including alopecia, lipodystrophy and joint stiffness, they present with severe osteolysis and skeletal deformation that is the primary cause of concern in these patients.

The mechanism by which the BAF A12T mutation causes these detrimental effects remains unclear. Here, we set out to characterize the effects of this mutation on NE integrity in NGPS-derived patient cells. To this aim, we engineered the first NGPS isogenic cell lines by correcting the homozygous BAF A12T mutation in NGPS patient cells using CRISPR-Cas9 mediated genome editing. We observed that the BAF A12T mutation in NGPS patient cells prevents the recruitment of A-type lamins to NE rupture sites, a phenotype that was ameliorated upon correction of the mutation. By combining biophysical analysis and cell biology assays, we showed that while not affecting BAF structure, the A12T mutation significantly decreases the binding affinity of BAF to lamin A/C. Finally, we showed that in a disease context, BAF A12T causes reduced recruitment of A-type lamins to the sites of NE rupture. While this did not prevent or delay NE repair, this led to enhanced nuclear fragility, characterized by more frequent NE re-rupturing.

MATERIALS AND METHODS

Cell culture

Cells were cultured in Dulbecco’s modified Eagle’s medium containing 10% foetal calf serum and penicillin/streptomycin. Cells were maintained at 37°C and 5% CO₂. Control human fibroblast cell line was derived from AG10803 immortalized with SV40LT and TERT, NGPS1 and NGPS2 were derived from NGPS5796 and NGPS5787, respectively, and were immortalized with SV40LT and TERT. These immortalized cell lines were a gift from Carlos López-Otín.

Generation of stable cell lines

Stable cell lines expressing FLAG-BAFWT and FLAG-BAFA12T were generated using a PiggyBac system. First, BAF WT and BAF A12T sequence were PCR-amplified from GFP-BAF vectors (kind gift from Cristina Capanni) and inserted into a FLAG vector. Piggybac vectors containing FLAG-BAF WT and FLAG BAF A12T were subsequently assembled by PCR amplification followed by Gibson assembly using HiFi DNA assembly cloning kit (New England Biolabs, #E5520S). FLAG-BAF was cloned into a BamHI and AfeI digested piggybacV1 CMV, a custom piggyback vector containing a CMV promoter and hygromycin resistance gene (kind gift from Jonathon Nixon-Abell). Stable cell lines expressing GFP-BAF WT and GFP-BAF A12T were generated using a PiggyBac system. EGFP-linker-BAF was first assembled in another vector and contains a short (GGGGG)₂ linker. Piggybac vectors containing GFP-BAF WT and GFP-BAF A12T were subsequently assembled by PCR amplification followed by Gibson assembly into a BamHI and AfeI digested piggybacV1 CMV. Piggybac vector containing GFP-NLS was assembled by PCR amplification of 3xGFP-NLS followed by Gibson assembly using HiFi DNA assembly cloning kit (New England Biolabs, #E5520S). The 3xGFP-NLS was described previously (43) and contains cycle3GFP fused to EGFP, an NLS, and a second EGFP in a pcDNA backbone (kind gift from Emily Hatch). 3xGFP NLS was cloned into a BamHI and AfeI digested piggybacV1 CMV.

Stable cell lines were generated by transfecting cells with the piggybacV1 plasmid together with a second plasmid containing the PiggyBac transposase (Jonathon Nixon-Abell) under an EF1alpha promoter using lipofectamine 3000 or Transit2020, according to manufacturer’s protocol. Cells were grown on hygromycin selection (100 μg/ml) the next day and were maintained in hygromycin containing medium.

Reversion of the BAF A12T mutation using CRISPR-Cas9 gene editing

To reverse the A12T mutation in BANFI, we used NGPS2 cells as they grew clones more easily from single cells. We used a previously described strategy that combines an All-in-One Cas9Δ10A nickase vector with enrichment by fluorescence-activated cell sorting to enrich for transfected cells (44). A pair of guide RNAs (sgRNAs) was designed to target the DNA on opposing strands surrounding the BANFI mutation in intron 1. The sgRNAs were then cloned into the All-in-One Cas9Δ10A nickase vector using DNA oligos (Sigma-Aldrich) using the BsaI and BbsI recognition sites. Several sgRNA pairs were originally designed using the CRISPR design tool in Benchling. After optimization, the sgRNA
sequences: GCCCATGGGGGAGAAGCCAG and GAAAGTCCTCGTGCTTTTGGG were used for CRISPR targeting in combination with a PAGE-purified ssODN of 200 bp long (Integrated DNA Technologies). The ssODN contains the BANF1 WT genomic sequence to resequence the A12T mutation and in addition has silent mutations to mutate the PAM sites to prevent Cas9 recutting and a mutation in an NcoI restriction site to facilitate clone screening by digestion of PCR products. Full ssODN sequence: AGAAGTTCCAGGTCTCCAGCCCTATCTGCTTTTGGGAGATCTGATCAAGATGAACAATCCTGGAAAAGCAACCGAGACTTCGTTGGCAGAGCCTATGGGGAGAAGCCAGTCGGAGCCTGTGGATTTTGGGATTCCTAGATTAAGCCTGTGAAGCTTGAGGCTAAGATGACAACATCCCAAAAAGCACCGAGACTGAGAGCCTTTTTTTGGGATTCCTAGATTAAGCCTGTGAAGCTTGAGGCTAAGATGACAACATCCCAAAAAGCACCGAGACTGAGAG.

RNA isolation and cDNA sequencing

RNA isolation was performed starting with a confluent 10 cm dish and using the Monarch total RNA miniprep kit for RNA isolation (NEB, #T2010S). RNA was stored at -80°C. RT-PCR was performed using a One-Step RT-PCR kit (Qiagen, #210210) using the following primers: Forward: AAGGATCCCGAGACCTTTGTTTGGGAGATCTGATCAAGATGAACAATCCTGGAAAAGCAACCGAGACTTCGTTGGCAGAGCCTATGGGGAGAAGCCAGTCGGAGCCTGTGGATTTTGGGATTCCTAGATTAAGCCTGTGAAGCTTGAGGCTAAGATGACAACATCCCAAAAAGCACCGAGACTGAGAGCCTTTTTTTGGGATTCCTAGATTAAGCCTGTGAAGCTTGAGGCTAAGATGACAACATCCCAAAAAGCACCGAGACTGAGAG.

Immunoblotting

Cells were washed with ice-cold PBS, lysed in Laemmli buffer (4% SDS, 20% glycerol, and 120 mM Tris-HCl [pH 6.8]) and then incubated for 5 min at 95°C. The DNA was sheared by sonicating the lysates 10 times through a 25-gauge needle. Absorbance at 280 nm was measured (NanoDrop; Thermo Fisher Scientific) to determine protein concentration. Samples were prepared using Protein Sample Loading Buffer (LI-COR, #928-40004) and DTT (final concentration 50 mM) and heated at 95°C for 5 min. Proteins were separated using NuPAGE 4-12% Bis-Tris gels (ThermoFisher) and NuPAGE MES SDS running buffer (Thermo Fisher, #NP0002) and transferred to nitrocellulose membranes for immunoblotting. Membranes were blocked in 5% milk PBS and incubated overnight at 4°C with primary antibodies. Next day, membranes were incubated for 1 h at room temperature with IRDye-conjugated secondary antibodies (LI-COR) and scanned on an Odyssey imaging system. The following primary antibodies were used: mouse anti-Tubulin (Sigma Aldrich, #T9026, 1/2000), mouse anti-lamin A/C (Santa Cruz, #sc-7292, 1/500), mouse anti-lamin B1 (Santa Cruz, #sc-365214, 1/1000), rabbit anti-emerin (ProteinTech, #10351-1-AP, 1/1000), rabbit anti-H3K9me3 (Abcam, #ab8898, 1/1000), rabbit anti-BAF (ProSci, #4019, 1/500), mouse anti-γH2AX (Millipore, #05-636-I, 1/500) and rabbit anti-H2AX (Bethyl, A300-083A-T, 1/1000).

Immunofluorescence

Cells were fixed at room temperature for 10 min with 4% PFA. Cells were washed in PBS, permeabilized using 0.2% Triton-X100, and blocked using 3% bovine serum albumin (BSA) in PBS for 30 min. Cells were incubated overnight at 4°C or for 1 h at RT in 3% BSA PBS containing primary antibody. Cells were washed using PBS and incubated for 1 h at room temperature with secondary antibody in 3% BSA PBS. Cells were washed in PBS and mounted using Prolong Gold (Thermo Fischer). The following primary antibodies were used: rabbit anti-BAF (Abcam, ab129184, 1/200), mouse anti-lamin A/C (Santa Cruz, sc-376248, 1/500), mouse anti-lamin A/C (Santa Cruz, sc-7292, 1/500), rabbit anti-emerin (ProteinTech, 10351-1-AP, 1/500), mouse anti-lamin B1 (Santa Cruz, sc-365214, 1/500), mouse anti-γH2AX (Millipore, 05-636-I, 1/200), rabbit anti-sun1 (Abcam, ab124770, 1/200), mouse anti-FLAG (Sigma Aldrich, F1804, 1/1000). Images were taken on Zeiss Axio Imager Z2 using a 63× oil immersion objective (Plan APO, NA 1.4, Zeiss).

Quantitative analysis of nuclear/cytoplasmic ratio’s, nuclear size and recruitment of proteins to blebs was performed using CellProfiler V4.1.3. Custom pipelines will be made available on request.

Image processing and analysis

Quantitative analysis of nuclear/cytoplasmic ratio’s, nuclear size and recruitment of proteins to blebs was performed using CellProfiler V4.1.3. Custom pipelines will be made available on request.

Nuclear/cytoplasmic ratio: Nuclear object was identified based on DAPI channel. The cytoplasmic object was subsequently generated by expansion of the nuclear object by 50 pixels and subtraction of the nucleus to create a cytoplasmic ring. The intensity of additional channels in the nuclear and cytoplasmic object were measured. In addition, the area of the nuclear object was measured. Data were transferred to Microsoft Excel for calculation of the nuclear/cytoplasmic ratio and subsequently plotted and analysed using GraphPad Prism 9.

Blebanalysis (45): Chromatin object was identified based on DAPI staining. A lamin B1 object was identified and subtraction of both areas identifies the bleb object as it is devoid of lamin B1. Subsequently, bleb areas are filtered to exclude small identified regions or pixels at the edge of the nuclei. Size, shape and number of bleb areas are measured as well as intensity of any additional channels at the bleb regions.

Quantitation of γH2AX foci by automated microscopy

Cells (Control, NGPS1, NGPS2, NGPS2 WT clone1 or clone2) were plated in 96-well Viewplates (Perkin Elmer, Beaconsfield, UK) and allowed to grow to 80% confluency over 48 h in a 37°C incubator. All subsequent steps were carried out at room temperature. Cells were washed once with
PBS before fixing with 4% paraformaldehyde (AR1068, Boster) for 15 min, followed by permeabilization with 0.2% Triton X-100 (AppliChem) in PBS for 12 min. Unspecific antibody binding was blocked by incubation in 5% bovine serum albumin (Sigma), 0.2% Tween-20 (Fisher) in PBS for 30 min. Primary and secondary antibodies were diluted in this blocking buffer. Antibody incubations were for 1.5 h. Mouse anti-phospho(S139)-histone H2AX (Millipore, #05-636-1, 1/300) primary antibody was used. DAPI (EMP Biotech, 0.2 µg/ml) was added together with the secondary antibody Alexa Fluor 568 goat anti-mouse IgG (LifeTechnologies, #A21124, 1/500). Cells were washed with PBS once after the primary antibody incubation and twice after the secondary antibody incubation, and stored in PBS at 4°C until imaged. Images were acquired on a CellInsight CX7 high-content microscope (Thermo Fisher) using a 20× objective (NA 0.45, Olympus). DAPI was used for automated focussing and object detection, and 1000 objects (nuclei) were imaged in three wells for each cell line. DAPI intensity thresholds were used to exclude mitotic cells from the analysis. Phospho(S139)-histone H2AX foci were detected using the spot detection algorithm in the HCS Studio software. Spots were detected after background correction using the box method and a fixed intensity threshold. Data were exported to Excel and average number of spots per nucleus were calculated and plotted in Graphpad Prism. Results from the technical repeats were averaged, and three biological repeats were carried out.

Proximity ligation assays (PLA)

Proximity ligation assays (PLA) were used to detect interaction between endogenous lamin A/C and BAF in control, NGPS2 and NGPS2 clone1 cells. Cells were seeded on 12 mm coverslips and fixed for 10 min with 4% PFA at room temperature and permeabilized using 0.2% Triton-X100. Cells were blocked for 1.5 h at room temperature using manufacturers blocking solution and primary antibody incubation, DuoLink Probe incubation, ligation and amplification steps were all carried out according to manufacturer’s protocol (DuoLink PLA assay kit, #DUO92008, Sigma Aldrich). Primary antibodies used were rabbit anti-BAF (Abcam, ab129184, 1/200) and mouse anti-lamin A/C (Santa Cruz, sc-376248, 1/1000). Duolink In Situ PLA probe anti-rabbit PLUS, Duolink In Situ PLA probe anti-mouse MINUS and Duolink Amplification Red were used. Cells were mounted in Duolink Mounting Media with DAPI. Confocal microscopy image acquisition was performed using LSM880 Laser scanning microscope (Zeiss) using a 63× oil immersion objective (Plan Apo, NA 1.4, Zeiss). Single plane images were taken except for analysis of PLA foci location, Z-stacks were recorded, and the middle slice was eventually used for analysis.

Quantitative analysis of PLA foci was performed using CellProfiler V4.1.3. In brief, first the nuclear object was detected based on DAPI signal after which PLA foci were detected in the nuclear object using thresholding. The number of PLA foci per nucleus was plotted for three independent experiments and the mean was indicated. Statistical analysis was performed using GraphPad PRISM 9. For analysis of the location of PLA foci, the middle slice from the recorded Z-stack was used. In brief, the nucleus was detected based on DAPI signal after which two outer rings were identified by shrinking the nucleus by 10 pixels. The defined outer nuclear circles and inner objects were used to identify the number of PLA foci in these areas. These numbers were then normalized for the surface area, and the fraction of foci in the outer rings and the centre was calculated.

Live cell imaging and analysis

Live-cell imaging of spontaneous NE ruptures was performed with an AxioObserver Z1 Inverted microscope (Zeiss) equipped with a HXP 120V high-pressure metalhalide lightsource. For imaging of spontaneous ruptures, cells were imaged in 4-well μ-slides (Ibidi, #80426) in FluoroBrite DMEM + FBS + PSQ and were maintained at 37°C and 5% CO₂. Cells stably expressing GFP-NLS were imaged every 3 min for 8 h using a 20× air objective (APOCROMAT, NA 0.8, Zeiss) and an ORCA-Flash4 v2 sCMOS camera (Hamamatsu). 1/2000 SPY650 DNA (COMPANY) was added 1 h before start imaging to allow for detection of nuclei. Image analysis was performed in FIJI. For analysis of nuclear GFP-NLS intensity image sequences were cropped to individual nuclei based on the presence of rupture events. Cells that were not visible for the entire time-lapse, underwent mitosis or showed severe nuclear shape abnormalities were excluded from the analysis. Cells were then automatically tracked using a dedicated script (trackRuptures_V5, (30)). Nuclei were automatically tracked based on SPY650-DNA signal. The nuclear NLS-GFP mean intensities were transferred to Microsoft Excel for further analysis. First, background subtraction was performed based on a background Region Of Interest (ROI), followed by normalization based on 5 frames before rupture occurs if available. First frame of rupture was aligned at t = 0. Ruptures were identified based on decreased NLS-GFP intensity and manually verified in time lapes. Ruptures were classified as major rupture events when the mean intensity loss in the nucleus was >30%. Time between ruptures after a major rupture was identified based on decrease in NLS-GFP intensity (including minor ruptures) and manually verified in time lapes. Ruptures were only included if they could be followed for at least 30 frames after the major rupture event so that they could be pooled in the overflow bin. For assessing differences in the probability of re-rupturing between Control and NGPS cell line, a Fisher’s exact test was performed. The time interval between rupture events from all replicates were pooled and we compared the number of ruptures in the >90 min category versus all other ruptures. Pooling was required due to low numbers of values across categories.

For imaging of GFP-BAF recruitment to sites of ruptures. GFP-BAF expressing cells were seeded in 35 mm Fluorodishes (#FD35, World Precision Instruments). One hour before the start of experiments, medium was replaced with CO₂ independent medium supplemented with 1/2000 SPY650-DNA (#SC501, Spirochrome) and cells were maintained at 37°C. Compression was performed using a Dynamic Cell Confiner (4D Cell). The confinement slides (0.2 µm pillar height) and suction cup were incubated in medium 1 h before start of the experiment. Live cell imaging of GFP-BAF expressing cells was performed.
on the same AxioObserver Z1 Inverted microscope using a 40× objective (APOCHROMAT, NA 0.95, Zeiss). Stoc-
tion cup with confinement slide was placed into the Fluo-
rodish, and pressure was lowered to −30 mbar, which en-
ures the device sticks to the dish without compression. At
this point, 30 s timelapse imaging of GFP and SPY650-
DNA was started and at least three frames were recorded
before compression. Pressure was decreased slowly to −100
mbar allowing compression of cells to 0.2 μm and was held
for 1 min after which compression was slowly released by
bringing the pressure back up to −30 mbar. After compres-
sion release, cells were imaged for 1 h.

**Protein constructs and expression vectors**

For protein purification, we used our BAF WT construct
that codes for an N-terminal tag containing 8 histidines,
a TEV cleavage site, and the human BAF sequence from
which all cysteines were mutated into alanines to allow
for protein resistance to oxidation and thus, aggregation
(32). After cleavage of the tag, the purified protein corre-
sponds to human BAF containing the following muta-
tions: M1G, C67A, C77A, C80A and C85A. The gene coding for
BAF WT was synthesized by Genscript after codon opti-
mization for expression in *Escherichia coli* and cloned in a
pETM13 vector providing kanamycin resistance to bacte-
ria. The vector used for BAF A12T expression was obtained
by mutagenesis of the BAF WT expression vector using the
Quikchange Site-Directed Mutagenesis kit (Agilent). The
lamin A/C Ig-fold domain, after sonication at
4°C, the supernatant was incubated 20 min at room tem-
perture with benzonase for 20 min at 37°C (room
ature, then the protein was recovered in the flow-through
sample was centrifuged again and the soluble fraction was
Tris, pH 8.0, 150 mM NaCl, 5 mM DTT). The
GST tag was cleaved with thrombin (commercial thrombin,
Sigma Aldrich), at 200 units per ml for 2 h at room tem-
perture, then the protein was recovered in the flow-through
and separated from thrombin and last contaminants using
gel filtration (Superdex 200 pg HiLoad 16/600 column, GE
healthcare). The final yield was typically about 0.6 mg (LB) or 0.1
mg (M9) of purified protein per liter of bacterial culture for
BAF WT and twice more for BAF A12T.

For the lamin A/C Ig-fold domain, after sonication at
10°C, the supernatant was incubated 20 min at room tem-
perature with benzonase and centrifuged at 50 000 g for 15
min at 4°C. The soluble extract was then supplemented with
5 mM DTT and loaded on to gluthathione beads. After 1 h
of incubation at 4°C, glutathione beads were washed first
with 1 M NaCl buffer and then with the purification buffer
(50 mM Tris, pH 7.5, 150 mM NaCl, 5 mM DTT). The
GST tag was cleaved with thrombin (commercial thrombin,
Sigma Aldrich), at 200 units per ml for 2 h at room tem-
perture, then the protein was recovered in the flow-through
and separated from thrombin and last contaminants using
gel filtration (Superdex 200 pg HiLoad 16/600 column, GE
healthcare). The final yield was typically 20 mg (LB) or 6
mg (M9) of purified protein per liter of bacterial culture.

In the case of VRK1, after sonication, the soluble extract
was incubated with benzonase for 20 min at 30°C (room
temperature). The lysate was then centrifuged at 50 000 g
for 15 min at 4°C and loaded on to a 5 ml Ni-NTA col-
umn (FF crude, GE-Healthcare). The column was washed
with washing buffer (50 mM Tris, pH 8.0, 1 M NaCl), re-
equilibrated with purification buffer (50 mM Tris, pH 8.0,
150 mM NaCl) and eluted with an imidazole gradient (0–
500 mM). After concentration to 5 ml, the histidine tag was
claved by the TEV protease (from a batch of TEV purified in the lab) overnight
at 4°C. The protein was separated from the TEV protease
(containing a histidine tag) and its Tag by Ni-NTA affinity
chromatography. Finally, a gel filtration was performed using
a Superdex 200 pg HiLoad 16/600 column (GE healthcare).
The final yield was typically 28 mg (LB) or 6
mg (M9) of purified protein per liter of bacterial culture.

**Protein expression and purification**

All vectors were transformed in *E. coli* BL21* (DE3). The
transformations were carried out by adding 100 ng of plasmid,
on to about one billion of bacteria (30 μl). Vectors entry
in the cells was triggered by a heat shock at 42°C during
45 s. Finally, the cells were spread on LB agar medium supple-
minted with the appropriated antibiotic.

Bacteria grew either in LB (Lysogeny Broth) or M9 (Min-
um 9) medium depending on the needs. The M9 media
were supplemented with either 15NH4Cl or natural abun-
dance glucose. Precultures of bacteria containing the vec-
tor coding for the protein of interest were prepared in LB
with antibiotic and incubated overnight at 37°C under ag-
itation (180 rpm). Then, 20 ml of preculture was used to
inoculate 800 ml of culture (LB or M9). When the OD600nm
reached 0.8 ± 0.1, protein expression was triggered using
IPTG (isopropyl-β-D-thiogalactoside). All proteins were
expressed overnight at 20°C. Cells were finally harvested by
centrifugation, flashed frozen in 30 ml of lysis buffer (50
mM Tris HCl pH 8, 300 mM NaCl, 5% glycerol, 0.1% Triton
X-100, 1 mM PMSF) and stored at −20°C during max-
imum 1–2 months.

Both BAF constructs are insoluble after overexpression in
*E. coli*, so purification was performed in urea and fol-
lowed by a refolding step. After sonication in lysis buffer
(50 mM Tris, pH 8, 300 mM NaCl, 5% glycerol, 0.1% Triton
100X), and centrifugation at 50 000 g for 15 min at 4°C, the
pellet was resuspended in urea purification buffer (50 mM
Tris, pH 8.0, 150 mM NaCl, 8 M urea) for 20 min. Then, the
sample was centrifuged again and the soluble fraction was
incubated on Ni-NTA beads pre-equilibrated with urea pur-
ification buffer, for 30 min at room temperature. Ni-NTA
beads were washed with the purification buffer, and the pro-
etin was eluted in 50 ml of the same buffer supplemented
with 1 M imidazole. Proteins were then refolded by dialy-
sis in BAF buffer (50 mM Tris, pH 8, 150 mM NaCl). Af-
ter concentration, the histidine-tag was cleaved by the TEV
protease (from a Batch of TEV purified in the lab) overnight
at 4°C. The protein was separated from the TEV protease
(containing a histidine tag) and its Tag by Ni-NTA affinity
chromatography. Finally, a gel filtration was performed using
a Superdex 200 pg HiLoad 16/600 column (GE healthcare).
The final yield was typically about 0.6 mg (LB) or 0.1
mg (M9) of purified protein per liter of bacterial culture for
BAF WT and twice more for BAF A12T.

**X-ray crystallography**

The BAF A12T-lamin A/C Ig-fold complex was recov-
ered after ITC experiments (in 50 mM Hepes, pH 7.4, 150
mM NaCl) and concentrated to about 20 mg/ml. Crys-
tallization experiments were carried out at the HTX Lab (EMBL Grenoble) (47). Crystals were obtained by sitting drop vapor diffusion at room temperature against reservoir containing 0.1 M bicine (pH 9) and 3 M ammonium sulfate. They were flash-frozen in liquid nitrogen and prepared for X-ray diffraction experiments using the CrystalDirect technology (48). Diffraction data were collected on the MASSIF-1 beamline (ESRF synchrotron, Grenoble, France). The 3D structure was solved by molecular replacement with Molrep software in CCP4 using the 6GHD.pdb coordinates file as starting model (49,50). The resulting model was iteratively improved by alternating manual reconstruction with the COOT software (51) and refinement with the BUSTER (52) and PHENIX REFINEM softwares (53) (Supplementary Table S1). Structure coordinates were deposited to the PDB, with entry 7Z21. All structure representations and Co RMSD calculations were performed with PyMOL (Schrodinger, LLC).

**Liquid-state nuclear magnetic resonance spectroscopy**

NMR experiments were performed on 600 and 700 MHz spectrometers equipped with triple resonance cryogenic probes. The data were processed using Topspin v. 4.0.2 to v. 4.0.8 (Bruker) and analysed using Topspin 4.1.3 (Bruker) and CCPNMR 2.4 (54). Sodium trimethylsilylpropanesulfonate (DSS) was used as a chemical shift reference.

For monitoring phosphorylation of BAF WT and BAF A12T by NMR, 2D 1H-15N HSQC spectra were recorded at 303 K on a 700 MHz spectrometer. The 3 mm-diameter NMR sample contoured 150 μM of BAF (either WT or A12T) in kinetics 40 mM HEPES, pH 7.2, 150 mM NaCl, 5 mM ATP, 5 mM MgSO4, 1 mM TCEP, 1× antiproteases (Roche), 95:5 H2O:D2O, and 150 nM of VRK1 kinase (molar ratio relatively to BAF: 0.1%). 2D 1H-15N NMR spectra were acquired every 15–25 min, and 1D 1H spectra were recorded in between to report for potential pH drifts.

To detect an interaction between the lamin A/C Ig-fold domain and BAF (either WT or A12T), 2D 1H-15N HSQC spectra were recorded on a sample containing 80 μM of 15N labelled lamin A/C Ig-fold and non-labelled BAF at different ratios. For the interaction with BAF WT, two spectra were acquired by adding 80 or 160 μM of BAF WT corresponding to molar ratios of 1:1 and 1:2, respectively. For the interaction with BAF A12T, two spectra were acquired: the first with 80 μM lamin A/C Ig-fold domain and 160 μM of BAF A12T, corresponding to a ratio of 1:2; the second with the lamin A/C Ig-fold domain at 160 μM and BAF WT at 160 μM, corresponding to a ratio of 1:1. The number of scans during the NMR acquisition was adjusted to obtain similar signal-to-noise ratios. All these experiments were performed in 50 mM HEPES, pH 7.4, 150 mM NaCl, 95:5 H2O:D2O, DSS, in 3 mm diameter tubes, at 293 K. Control experiments were carried out with the labelled protein at 80 μM, in the same experimental conditions.

To calculate the NMR intensity ratios I/Io, we started by assigning the 1H-15N HSQC signals to their corresponding lamin residues. Using a previous assignment of the Ig-fold domain (BMRB: 5224), we managed to assign the 1H-15N HSQC signals of approximately 80% of the lamin residues. The volume of each peak was measured using CCPNMR 2.4 and the reference lamin A/C Ig-fold 1H-15N HSQC spectrum at 80 μM used to obtain Io values. When plotting intensity ratios as a function of the residue number, values corresponding to residues that could not be assigned were calculated as the average between the ratios of the previous and next residues.

**Isothermal titration calorimetry (ITC) binding assays**

The interaction between lamin A/C Ig-fold domain and BAF (either WT or A12T) was assessed by ITC using a VP-ITC calorimetry system (MicroCal-Malvern Panalytical, Malvern, UK). Calorimetric titrations were performed with either 100 or 200 μM lamin A/C Ig-fold domain in the injecting syringe and either 20 μM BAF WT or 40 μM BAF A12T in the calorimetric cell, all in 50 mM Hepes, pH 7.4 and 150 mM NaCl. All measurements were performed at 288 K in order to increase the signal-to-noise ratio. For each titration, a sequence of 29 times 10 μl injections was programmed, with reference power of 10 mcal/s and spacing between injections of 180 s. Each experiment was performed twice. Data were analysed using Origin (OriginLab, Northampton, MA, USA) by setting the stoichiometry to 0.5 (assuming that a dimer of BAF binds to a monomer of lamin A/C Ig-fold domain, as observed by X-ray crystallography).

**Statistics**

Statistical analysis was done using GraphPad Prism v9. Individual data points are plotted from three independent experiments. Mean or median is indicated by the black bar as indicated in the figure legends. Post-testing was performed as indicated to correct for multiple comparison. Details of statistical tests are included in the figure legends. Sample size was not predetermined and experiments were not randomized.

**RESULTS**

The phenotypes of NGPS cells are distinct from the ones of HGPS cells

To characterize NGPS patient cell phenotypes, we used hTERT immortalized skin fibroblasts of a healthy age-matched donor (control) and two NGPS patients (NGPS1 and NGPS2) (gift from C. Lopez-Otin). Microscope visualization of the nuclei in both NGPS patient cells showed clear differences between the cell lines. Cells from NGPS patients were previously reported as showing nuclear abnormalities including nuclear blebs and delocalization of emerin from the NE to the cytoplasm (40,55). Here, we observed that NGPS1 cells displayed smaller nuclei with nuclear membrane folding (white arrowheads), while NGPS2 cells had bigger nuclei, and discontinuity of the lamin A/C network (magenta arrowheads) without the presence of membrane folds (Figure 1A, B). The control cells looked more homogeneous even though they also showed some nuclear shape abnormalities with areas of lamin A/C accumulation at the NE (green arrowheads).

We further assessed several NE constituents known to be altered in HGPS, in senescent cells or in cells from normal aged individuals. We first assessed the lamin B1 expression
Figure 1. Characterization and comparison of fibroblast cell lines derived from two Nestor–Guillermo progeria patients. (A) Representative lamin A/C immunofluorescence images of immortalized fibroblasts obtained from an unrelated healthy donor (Control) or from two NGPS patients (NGPS1 and NGPS2). The arrows indicate areas of increased lamin intensity (green), nuclear envelope folding (white) and lamina gaps (magenta); scale bar: 20 μm. (B) Quantification of nuclear area based on DAPI images as presented in (A). Mean from \( n = 469, 563 \) and 413 nuclei for Control, NGPS1 and NGPS2, respectively. The data were collected in three independent experiments and analysed using a one-way ANOVA analysis with Šídák’s multiple comparisons test (**\( P < 0.0001 \)). (C) Representative immunofluorescence images of immortalized NGPS patient and control cell fibroblasts stained for laminB1, and Sun1; scale bar: 20 μm. (D) Immunoblot analysis of the indicated proteins from whole-cell lysate in Control and NGPS cells. Tubulin was used as a loading control.

level, as its downregulation has previously been associated with senescence (56) and observed in HGPS cells (57,58). However, NGPS cells did not show any reduction in lamin B1 (Figure 1C, D). Similarly, we did not observe any change in the expression levels of lamin A/C or emerin (Figure 1D) in these cells. In HGPS cells, the LINC (Linker of Nucleoskeleton and Cytoskeleton) complex protein SUN1 has been shown to accumulate at the NE and in the Golgi, contributing to HGPS pathogenicity (59). On the contrary, the expression and localization of SUN1 was not affected in NGPS cells (Figure 1C). Apart from NE changes, DNA damage accumulation is another marker frequently observed in various age-related diseases including HGPS (60). Perhaps surprisingly, we did not find increased DNA damage in NGPS cells—as assessed by immunofluorescence staining and immunoblotting of γH2AX, a marker of DNA double-strand breaks (Supplementary Figure S1). Finally, we investigated other potential ageing-associated changes in chromatin organization by probing for H3K9me3, a marker of transcriptionally silent heterochromatin, loss of which is another hallmark of HGPS and aging (57,61). However, H3K9me3 level was similar in control and NGPS patient cells (Figure 1D).

Together, these observations reinforce the fact that, despite the existence of premature aging features in NGPS patients, the effects of the BAF A12T mutation at the cellular level are very different from the ones caused by LMNA mutations in HGPS. One potential caveat here is the use of immortalized cells, which could impact on the expression level of some of these markers, however, primary NGPS cells could not be obtained due to their inability to grow in culture.
The BAF A12T mutation modifies BAF and emerin localization in NGPS cells

Through these initial studies, we thus did not find highly consistent phenotypes associated with aging in the available patient cell lines derived from the two originally identified NGPS patients. Therefore, we decided to engineer an isogenic cell line using CRISPR-Cas9 gene editing to reverse the BAF A12T homozygous mutation in NGPS cells using an all-in-one plasmid strategy containing a Cas9 nickase, two sgRNA targeting BAF around the mutation site, and GFP for cell sorting (Figure 2A) (44). We identified two separate clones with a homozygous reversion of the A12T mutation (NGPS2 WT clone1-2) (Figure 2B and Supplementary Figure S2a). Using these isogenic cell lines, we observed that reversing the BAF A12T mutation significantly improves the emerin nuclear/cyttoplasmic ratio being down to 50% of the control cells' ratio in the WT clone (Figure 2C, D; Supplementary Figure S2h, c), confirming that this phenotype is a direct consequence of the BAF mutation. One possible explanation for the partial rescue observed between the WT clone and the control cells could be due to differences in the basal ratio observed in cells from different individuals, as the control cells are derived from a healthy, unrelated donor.

Interestingly, BANF1 sequencing revealed the presence of an additional and similar deletion in intron 2 in both NGPS patient cell lines (Supplementary Figure S3a). To check whether this deletion might affect RNA splicing and therefore result in an altered protein, we isolated RNA from both patient cell lines. Sequencing of the generated cDNA showed that this deletion did not affect the mRNA sequence (Supplementary Figure S3b).

In addition to the previously described delocalization of emerin from the nucleus to the cytoplasm (Figure 2C, D), we found that both NGPS cell lines showed a decrease in nuclear BAF enrichment compared to control cells, as observed by immunofluorescence, being down to 69% in the NGPS2 cells and rescued to 84% by the mutation reversion (NGPS2 WT clone 1) (Figure 3A, B). This suggested that BAF levels might be decreased in NGPS patient cells. However, immunoblot analysis instead showed that BAF levels are increased in both NGPS patient cell lines, going down upon mutation reversion (Figure 3C). Phosphorylation of BAF is known to affect its subcellular localization and to regulate its interaction with different protein partners. The decrease of nuclear BAF we observed in NGPS patient cells could thus suggest altered BAF phosphorylation upon A12T mutation. Since BAF is phosphorylated by the vaccinia-related kinase 1 (VRK1) on Serine 4 (Ser4) and then Threonine 3 (Thr3) (62,63), both being close to the mutated A12T site, we speculated that phosphorylation of BAF could be affected by the NGPS mutation.

The A12T mutation does not affect BAF phosphorylation in vitro

We therefore monitored the phosphorylation of BAF WT and BAF A12T in vitro over time using nuclear magnetic resonance (NMR) spectroscopy (Figure 3D-E and Supplementary Figure S4). On the recorded $^{1}$H-$^{15}$N HSQC NMR spectra, each peak reports on the chemical environment of one residue. A change of this residue's chemical environment, due to a phosphorylation for example, will change the position of the corresponding NMR peak on the spectrum. Peaks of phosphorylated serine and threonine appear on the very left side of the spectrum: in general, at >9 ppm in the hydrogen dimension. Therefore, we focused our analysis on this area, where very little difference was observed between BAF WT and BAF A12T before VRK1 kinase addition (Figure 3E, top panel). After 15 min of in vitro phosphorylation reaction of BAF by VRK1, the signal corresponding to phosphorylated Ser4 was visible on both spectra (Figure 3E, middle panel). Similarly, after 8 h of phosphorylation, the signal corresponding to the phosphorylated Thr3 was visible on both spectra (Figure 3E, bottom panel). This phosphorylation caused a global change in both spectra (Supplementary Figure S4). Finally, no signal corresponding to another phosphorylated threonine appeared on the BAF A12T spectrum at the end of the reaction, showing that the introduced Thr12 is not phosphorylated by VRK1.

Altogether, this analysis showed that BAF WT and BAF A12T are phosphorylated in a similar way by VRK1 in vitro: they are both phosphorylated on Ser4 and Thr3 only, with similar kinetics, and phosphorylation of Thr3 triggers a similar global change in their $^{1}$H-$^{15}$N HSQC NMR spectra, reflecting the compaction of BAF 3D structure upon phosphorylation (62).

BAF A12T binds to lamin A/C with a ten-fold weaker affinity

In order to further characterize the effect of the A12T mutation on BAF 3D structure and binding properties, we initiated a structural analysis of BAF A12T by X-ray crystallography. The only interaction known to directly involve alanine 12 is the interaction of BAF with the Ig-fold domain of lamin A/C (32). Therefore, we determined the 3D structure of BAF A12T bound to the lamin A/C Ig-fold domain at 1.6 Å resolution (Figure 4A). We compared the structure of this complex to that of the wild-type complex (PDB code: 6GHD) and found that they nicely overlapped (CαRMSD = 0.4 Å) with Ala12 or Thr12 being largely exposed to the solvent (Figure 4A). Only slight changes in the relative positioning of the BAF monomers, as well as in the structure of some lamin A/C loops (especially at Ser433-Thr434) were observed. We thus concluded that the A12T mutation does not affect BAF 3D structure. It has been proposed that the A12T mutation perturbs the interaction with the Ig-fold domain of lamin A/C (32). Additionally, this interaction is important for the retention of BAF in the nucleus (64,65).

We investigated the effect of the BAF A12T mutation on the binding affinity to the lamin A/C Ig-fold domain in vitro. Despite an apparent similarity between the 3D structures of the WT and mutated complexes, prediction of the binding energies from these 3D structures using the PDBBind server (66) suggested that the A12T mutation changes the thermodynamical equilibrium of the system, with the mutant binding with a free energy $\Delta G$ that is 1.8 kcal/mol less favourable compared to the WT (Supplementary Figure S5a). To confirm this prediction experimentally, we first showed by NMR that the A12T mutation causes a significant decrease in BAF affinity for lamin A/C Ig-fold.
Figure 2. Reversion of the homozygous BAFA12T mutation in NGPS patient cells using CRISPR/Cas9. (A) Graphical representation of the CRISPR-Cas9 strategy showing: (top) *BANF1* sequence around the A12T mutated codon (orange), the PAM sites for Cas9 recognition (purple boxes) and the annealing sites of the single guide RNAs (sgRNAs: grey bars). Left: the all-in-one vector carrying the Cas9 nickase (orange), the sgRNA pairs targeting the *BANF1* mutation site (grey) and the Green Fluorescent Protein (GFP: used for cell sorting) was used in combination with an ssODN (bottom) carrying the correct wild-type *BANF1* sequence as a template for mutation correction through homologous recombination. (B) DNA sequencing traces of the *BANF1* gene around the mutation site showing the wild-type GCA codon (Alanine) in control cells, the mutated ACA (Threonine) codon in NGPS2 cells and the corrected mutation (GCA, Alanine) in one of the successfully NGPS2 derived wild-type clones (NGPS2 WT clone1). Boxes indicate the A12T mutation site (black) and additional silent mutations introduced in the PAM motifs (purple) and in the restriction enzyme NcoI site for screening (grey). (C) Representative immunofluorescence images of emerin in control, NGPS patient cells and NGPS2 WT clone1; scale bar: 20 μm. (D) Quantification of the mean emerin nuclear to cytoplasmic ratio measured in 345, 465, 334 and 301 cells for control, NGPS1, NGPS2 and NGPS2 WT clone1 respectively. Data are presented from three independent experiments, and the *P* value (** ** ** ** *P*<0.0001) was calculated using a one-way ANOVA analysis with Šidák’s multiple comparisons test.
Figure 3. The BAF A12T mutation causes BAF mislocalization in NGPS patient cells without affecting its phosphorylation by VRK1 in vitro. (A) Representative immunofluorescence images showing endogenous BAF staining in Control, NGPS patient cells and NGPS2 WT clone1; scale bar: 20 μm. (B) Quantification of the mean BAF nuclear to cytoplasmic ratio in 425, 495, 401 and 363 cells for Control, NGPS1, NGPS2 and NGPS2 WT clone1, respectively. Data were obtained from three independent experiments and quantified using a one-way ANOVA analysis with Sidák’s multiple comparisons test (****P < 0.0001). (C) Representative immunoblot showing the expression level of BAF in whole cell lysates from the indicated cell lines. Tubulin was used as a loading control. (D–E) 2D NMR $^1$H-$^{15}$N Heteronuclear Single Quantum Coherence (HSQC) spectra were recorded on purified BAF WT (in shades of blue) and A12T (in shades of red) upon phosphorylation by the VRK1 kinase in vitro. Six spectra are superimposed, corresponding to: non-phosphorylated BAF WT and A12T ($t = 0$ min), in dark and light grey respectively, mono-phosphorylated BAF WT and A12T ($t = 15$ min), in light blue and pink respectively, and di-phosphorylated BAF WT and A12T ($t = 8$ h), in dark blue and red, respectively. The zoom in (E) shows the spectral regions of the $^1$H-$^{15}$N HSQC of BAF WT and A12T spectra where signals of phosphorylated serines and threonines are commonly observed. The signals of the two BAF residues phosphorylated by VRK1 (Ser4 and Thr3) are annotated, and the arrow indicates the shift of the peak corresponding to phosphorylated Ser4 upon phosphorylation of Thr3.
Figure 4. The BAF A12T mutation disrupts BAF binding to lamin A/C in vitro and in NGPS cells. (A) Superimposition of the 3D structure of BAF A12T (in light pink) bound to the lamin A/C Ig-fold domain (in light blue), on to the previously reported structure of BAF WT bound to this same lamin A/C domain (in grey; PDB code: 6GHD). Binding free energies calculated from these 3D structures are indicated in Supplementary Figure S5a. (B) ITC curves reporting binding of BAF (either WT or A12T) to the lamin A/C Ig-fold domain. The experiments were performed twice and the mean dissociation constant ($K_d$) values are shown under one representative curve. The duplicated experiments are shown in Supplementary Figure S5b. The thermodynamic parameters deduced from all experiments are summarized in Supplementary Figure S5c. (C) Representative confocal images of the signal obtained by PLA using either BAF or lamin A/C antibodies alone as negative controls or as a combination in the indicated cell lines. The top row shows merged pictures of the PLA signal (green) and DAPI (magenta); scale bar: 10 μm. (D) Quantification of the number of PLA foci per nucleus. The data represent the mean PLA signal measured in individual nuclei in $n = 214$ (Control BAF), 187 (Control lamin A/C), 229 (Control lamin A/C+BAF), 213 (NGPS2 lamin A/C+BAF) and 219 cells (NGPS2 WT clone1, lamin A/C + BAF). One-way ANOVA analysis with Sidak’s multiple comparisons test ($** P < 0.001$). (E) Analysis of the nuclear distribution of PLA foci in the middle slice of a z-stack obtained from confocal images in control cells using CellProfiler. The nucleus was subdivided in concentric circles going from the nuclear periphery to the nuclear interior and the fraction of PLA foci in each of these circles was counted per nucleus ($n = 10$ nuclei). (F) Example confocal images (from images as shown in C) showing the BAF-lamin A/C PLA signal accumulation at nuclear bleb sites in Control cells. Scale bar: 10 and 3 μm (zoom).
domain (Supplementary Figure S5d–k). Therefore, we measured the intensities of the $^1$H–$^{15}$N HSQC peaks of the Ig-fold domain in the presence of different ratios of BAF WT and BAF A12T. After binding to BAF WT dimer at 1:1 and 1:2 ratio, only 27% and 4% of the initial NMR signals were observed, respectively (in light and dark blue in Supplementary Figure S5f, h, k). In the case of BAF A12T dimer, using the same ratios (1:1 and 1:2), we measured lower intensity decreases: 43 and 66% respectively (in pink and red in Supplementary Figure S5g, i, k). This indicated that significantly less complexes are formed when BAF alanine 12 is mutated into threonine. We then determined the dissociation constant ($K_D$) of these interactions using Isothermal Titration Calorimetry (ITC). As the affinities were on the (high) micromolar range, the solubility limits of our proteins prevented reaching saturation. To improve the quality of our data analysis, we decided to fit our ITC curves assuming a stoichiometry of two BAF molecules for one lamin A/C Ig-fold domain. Indeed, we already reported that BAF A12T eluted from the size-exclusion chromatography column at the same volume as BAF WT, which demonstrates that BAF A12T is a dimer in solution (32). Also, both BAF WT and A12T dimers bind to a single Ig-fold domain in the crystal structures. We obtained $K_D$ values of 2.7 ± 1.2 μM and 33 ± 4 μM for the interactions involving BAF WT and BAF A12T, respectively (Figure 4B and Supplementary Figure S5b, c), showing that the A12T mutation decreases the affinity of BAF for lamin A/C Ig-fold by about tenfold.

To assess whether the reduced interaction between BAF A12T and the lamin A/C Ig-fold in vitro was also translated in a cellular context, we used a PLA. While each of the BAF or lamin A/C antibodies on their own gave very little PLA signal as expected, specific PLA foci were detected upon combining the two antibodies, confirming the proximity of the two proteins in control cells. Furthermore, we observed a strong reduction in the number of PLA foci in NGPS2 cells, reflecting a decreased interaction between BAF-lamin A/C, which was rescued by reversion of the mutation (Figure 4C, D). Even though BAF and lamin A/C are both found throughout the nucleus, we observed a clear enrichment of PLA foci at the nuclear periphery. Indeed, by measuring the number of PLA foci in concentric rings within the nucleus, starting from the nuclear periphery, we confirmed that most of the PLA foci were present in the outer nuclear ring (Figure 4E), indicating that these proteins mostly interact at the NE. In addition, we noticed enrichment of PLA foci in nuclear blebs of a subset of control cells, indicating that blebs could be a site of increased interaction between BAF and lamin A/C (Figure 4F). Together these data show that the BAF A12T mutation reduces the affinity of BAF for lamin A/C both in vitro and in NGPS cells.

Reduced binding of BAF A12T to lamin A/C in NGPS cells prevents lamin A/C recruitment to NE ruptures

Recruitment of lamin A/C by BAF at sites of nuclear ruptures has recently been shown to be an early event in the repair of nuclear envelope ruptures and ultimately leaves behind a lamin ‘scar’ (65,67). We observed these lamin scars at the NE of control cells (Figure 1A, green arrowheads) but in contrast, often observed gaps in the lamina of NGPS patient cells (Figure 1A, magenta arrowheads). Gaps in the lamina can generate weak spots in the NE leading to bleb formation and ultimately NE rupture. We therefore investigated the effect of the BAF A12T mutation on the recruitment of lamin A/C to nuclear blebs, which can be recognized on immunofluorescence images by a lack of B-type lamins (20). In control cells, we observed a clear accumulation of lamin A/C to a subpopulation of blebs (Figure 5A, green arrowheads). NGPS cells on the other hand did not show lamin A/C recruitment to blebs (Figure 5A, magenta arrowheads). Automated quantification using a CellProfiler based pipeline (45) of either lamin A/C intensity at blebs or enrichment of lamin A/C at blebs, confirmed a lack of lamin A/C recruitment in NGPS cells (Figure 5B). Reversion of the BAF A12T mutation in NGPS2 cells was able to fully rescue the lamin A/C recruitment defect (Figure 5A [green arrowheads] and 5b, Supplementary Figure S6a, b), demonstrating that this phenotype is caused by the mutation. Additionally, we examined the recruitment of emerin to blebs, as emerin is also known to accumulate at rupture sites in the NE repair process and because emerin localization is altered in NGPS patient cells. We found a decrease in emerin intensity at NE blebs in NGPS cells, that was fully rescued by the mutation reversion, which could be partially explained by an overall decrease in emerin levels at the NE (Figure 5C, D, Supplementary Figure S6c, d).

Wild-type BAF overexpression in NGPS cells is sufficient to restore lamin A/C recruitment to NE ruptures

To assess whether overexpression of wild-type BAF in NGPS cells could rescue the observed defects in the recruitment of lamin A/C or emerin to blebs, we generated control and NGPS patient cells stably expressing FLAG-BAF wild-type (WT) or FLAG-BAF A12T. This mimics a situation where BAF A12T is present as a heterozygous mutation, as it occurs in NGPS patients’ parents, who are devoid of any disease phenotype. First, we assessed FLAG-BAF (WT and A12T) localization in these cells and observed that both proteins localized to the nucleus as expected (Figure 6A). We additionally checked whether BAF A12T recruitment to rupture sites was affected, as contradictory results on the affinity of BAF A12T for DNA were previously published (62,68). We observed that both FLAG-BAF WT and A12T localized to blebs identified by lack of lamin B in both Control and NGPS2 background (Figure 6A, B). Around 30% of all blebs showed FLAG-BAF enrichment for both the WT and the A12T protein (Figure 6C). Together, these results suggested that the recruitment of BAF to DNA at nuclear rupture sites is not affected by the A12T mutation, consistent with in vitro data (62) and with recent reports by others in cells (67). In addition, we observed that emerin nuclear localization was rescued by overexpression of FLAG-BAF WT in NGPS patient cells (Supplementary Figure S7a, b). Finally, we showed that overexpression of FLAG-BAF WT in NGPS patient cells was sufficient to rescue lamin A/C and emerin recruitment to NE blebs (Figure 6D–F). Similarly, overexpression of BAF A12T in a wild-type background did not cause defects in the recruitment of these proteins. Thus the presence of wild-type BAF prevents or reverses the observed phenotypes which is consis-
Figure 5. The BAF A12T mutation in NGPS patient cells prevents the recruitment of lamin A/C and emerin to nuclear blebs. (A) Representative immunofluorescence images of endogenous lamin B1, and lamin A/C at nuclear blebs in Control, NGPS patient cells and NGPS2 WT clone1 cells. Arrows indicate the enrichment (green) or lack of (magenta) lamin A/C at blebs; scale bar: 10 μm. (B) Quantification of the normalized lamin A/C intensity at blebs and of the enrichment of lamin A/C at nuclear bleb regions compared to the levels measured in the rest of the nucleus. Blebs were identified by the lack of lamin B1 staining. Data points represent 117 (Control), 81 (NGPS1), 235 (NGPS2), and 122 (WT clone1) individual blebs from three independent experiments, the median is indicated. One-way ANOVA analysis with Šidák’s multiple comparisons test (**P < 0.0001). (C) Representative immunofluorescence images of lamin B1 and emerin in nuclear blebs of Control, NGPS patient cells and NGPS2 WT clone1 cells. Arrows indicate the enrichment (green) or lack of (magenta) emerin at blebs; scale bar: 10 μm. (D) Quantification of the normalized emerin intensity at blebs and the enrichment of emerin at bleb regions compared to levels in the rest of the nucleus. Data points represent 113 (Control), 87 (NGPS1), 163 (NGPS2) and 134 (WT clone1) individual blebs from three independent experiments, and the median is indicated. The data were analysed with a one-way ANOVA analysis with Šidák’s multiple comparisons test (**P < 0.0001, ***P < 0.005, *P < 0.05).

tent with the lack of disease phenotypes in NGPS parents who carry a BAF A12T heterozygous mutation (39, 40).

Since we identified that BAF A12T accumulates at NE ruptures in a similar way as BAF WT, we next sought to investigate the dynamics of BAF recruitment to these ruptures. Therefore, we performed live cell imaging on control cells stably expressing GFP-BAF WT and on NGPS2 cells stably expressing GFP-BAF A12T. To study the dynamics with a high temporal resolution, nuclear rupture was induced by cellular compression. Both GFP-BAF WT and GFP-BAF A12T were rapidly recruited to rupture sites (Figure 7A) after compression. Recruitment to chromatin herniations (white arrowheads) occurred within 30 s. After release of cellular compression, GFP-BAF A12T in NGPS2 and GFP-BAF WT in control cells dissociated from the site of rupture at a similar rate. Therefore, these data suggest that the recruitment to and persistence of BAF at sites of ruptures is not affected by the A12T mutation.
Figure 6. The defective recruitment of lamin A/C to nuclear blebs is restored in NGPS cells by BAF wild-type overexpression. (A) Representative immunofluorescence images of FLAG and lamin B1 staining in Control and NGPS2 cells stably expressing FLAG-BAF WT or A12T. Green arrows indicate FLAG-BAF accumulation at nuclear blebs. Scale bar: 20 µm. (B) Quantification of FLAG-BAF WT and A12T enrichment at bleb regions compared to the levels measured in the rest of the nucleus. Blebs were identified by the lack of lamin B1 staining. Data points represent \( n = 56 \) (Control FLAG-BAF), 82 (Control FLAG-BAFA12T), 112 (NGPS2 FLAG-BAF) and 156 (NGPS2 FLAG-BAFA12T) individual blebs from three independent experiments, and median is indicated. One-way ANOVA analysis with Šidák’s multiple comparisons test showed no significant (ns) differences. (C) For each experiment in (B) the percentage of cells showing >1.5 FLAG-BAF enrichment at blebs was calculated. Mixed-effects analysis with Šidák’s multiple comparisons test showed no significant (ns) differences, and mean is indicated. (D) Representative immunofluorescence images of lamin B1, lamin A/C and emerin staining in Control and NGPS2 cells stably expressing FLAG-BAF WT or A12T. Arrows point to nuclear blebs with accumulation (green arrowheads) or lack of (magenta arrowheads) lamin A/C. Scale bar: 10 µm. (E) Quantification of the normalized lamin A/C intensity at blebs and of the enrichment of lamin A/C at bleb regions compared to the levels measured in the rest of the nucleus. Blebs were identified by the lack of lamin B1 staining. Data points represent \( n = 61 \) (Control FLAG-BAF), 79 (Control FLAG-BAFA12T), 108 (NGPS2 FLAG-BAF) and 153 (NGPS2 FLAG-BAFA12T) individual blebs from three independent experiments, median is indicated. One-way ANOVA analysis with Šidák’s multiple comparisons test (****\( P < 0.0001 \), **\( P < 0.01 \)). (F) Quantification of the normalized emerin intensity at blebs and of the enrichment of emerin at bleb regions compared to the levels measured in the rest of the nucleus. Data points represent \( n = 61 \) (Control FLAG-BAF), 79 (Control FLAG-BAFA12T), 108 (NGPS2 FLAG-BAF) and 153 (NGPS2 FLAG-BAFA12T) individual blebs from three independent experiments. One-way ANOVA analysis with Šidák’s multiple comparisons test (****\( P < 0.0001 \)).
Figure 7. The NE rupture repair kinetic is unaffected in NGPS cells but nuclei are more prone to re-rupturing. (A) Still images of Control GFP-BAF WT and NGPS2 GFP-BAF A12T cells showing nuclear rupture upon compression (white arrowheads). Time is indicated in min:sec relative to the first time point of compression; scale bar: 20 μm. (B) Example of still images from a time lapse obtained in NGPS2 cells stably expressing NLS-GFP and showing several rupture and repair events originating from the same nuclear bleb. Time is indicated in hr:min; scale bar: 20 μm. (C) Graph showing the mean GFP-NLS fluorescence intensity over time in the nucleus shown in (B). The repetitive ruptures depicted in (A) are indicated on the graph by green arrows. (D) Analysis of the recovery kinetics of nuclear GFP-NLS after major NE rupture events (>30% loss of nuclear GFP-NLS fluorescence intensity). Mean and standard deviation is indicated for control (black, n = 62 ruptures), NGPS1 (dark grey, n = 7 ruptures), NGPS2 (light grey, n = 110 ruptures). (E) Histogram showing the proportion of ruptures with the indicated duration between rupture events after an initial major rupture occurred. Control (black, n = 41 ruptures), NGPS1 (dark grey, n = 6 ruptures) and NGPS2 (light grey, n = 103 ruptures) cells were imaged every 3 min and were binned to every 15 min. Fisher’s exact test (**P<0.01).
The reduced accumulation of lamin A/C to NE ruptures in NGPS cells does not affect NE rupture repair but causes NE re-rupturing

The specific role of lamin A/C in nuclear envelope rupture repair is yet to be identified. No associated defect in NE repair and integrity has been reported yet as a consequence of a lack of lamin A/C recruitment to sites of nuclear rupture. We, therefore, monitored NE rupture and repair kinetics following spontaneous rupture events by imaging of stably expressed fluorescently tagged nuclear localization signal (GFP-NLS) in control and NGPS cells. Upon NE rupture, the GFP-NLS reporter leaks into the cytoplasm. Once the rupture has been repaired, GFP-NLS reaccumulates into the nucleus. Using live cell imaging of GFP-NLS, we observed spontaneous ruptures in all cell lines. Some cells showed repetitive NE rupture events during the time of observation, which was surprisingly not associated with cell death and appeared to be repaired quickly (Figure 7B, C). As BAF-dependent NE repair was proposed to be more important for large membrane gaps (31), we decided to focus our analysis on major NE rupture events, that we defined as being the ones leading to loss of >30% of the mean nuclear GFP-NLS intensity. We found that the lack of lamin A/C recruitment did not interfere with the repair process as we did not find a delay in the repair kinetics when comparing both NGPS cell lines to the control (Figure 7D).

We then analysed the re-occurrence of ruptures after an initial major rupture event and observed that both NGPS cell lines were more likely to re-rupture within 90 min compared to control cells (Figure 7E). These data showed the same trend for both NGPS cell lines, statistical analysis did however not find a significant difference for NGPS1 compared to the control, probably due to the low number of recorded ruptures in this cell line overall. Altogether, our data show that the lack of lamin A/C recruitment to sites of NE rupture causes NE instability through recurrent rupturing of the NE. This is associated with the persistence of a lamina gap after initial NE repair has occurred.

DISCUSSION

In the last ten years, the interest in the small DNA binding protein BAF has grown due to two main discoveries. First, a recessive mutation in BAF, namely A12T, causes Nestor–Guillermo Progeria Syndrome. Second, BAF plays a role in the repair of nuclear envelope ruptures. BAF directly binds to two well-characterized nuclear envelope proteins: A-type lamins and emerin. A recent report showed that, in fibroblasts overexpressing GFP-BAF A12T, recruitment of A-type lamins to laser-induced nuclear ruptures is impaired (67). However, until now, the mechanism behind this phenotype was unknown. Here, we revealed a mechanism for the defective recruitment of A-type lamins at the nuclear envelope ruptures and we further characterized the consequences of this disease-associated mechanism on the nuclear envelope integrity in the context of NGPS cells.

Using a combination of in vitro and cell-based experiments, we showed that, while the A12T mutation did not affect BAF 3D structure nor its phosphorylation by VRK1, it significantly reduces its binding affinity for the Igfold domain of lamin A/C. As a consequence, we found that the accumulation of A-type lamins at nuclear rupture sites—known to be dependent on the presence of BAF (31)—was strongly reduced in NGPS patient cells. We confirmed that this was indeed a specific effect of the BAF mutation by correcting the BAF A12T homozygous mutation using CRISPR-Cas9. We observed that the A12T mutation also causes delocalization of emerin from the NE to the cytoplasm in NGPS cells, even though this mutation is not expected to impair BAF-emerin binding directly. Both BAF and lamin are important for the nuclear enrichment and dynamics of emerin (69,70). Therefore, we speculate that inefficient immobilization of emerin in BAF-emerin-lamin A/C tertiary complexes at the NE caused by the reduced binding of BAF to lamin A/C could affect emerin nuclear localization. This in turn could have additional effects on downstream emerin functions such as mechanotransduction processes at the NE. Interestingly, NGPS cells show an increase in BAF levels which could be a mechanism to compensate for a decrease in BAF–lamin A/C interaction and a defect in associated cellular processes.

While BAF is important for recruiting A-type lamins to NE ruptures, the presence of lamin A has been shown to be reciprocally important for the accumulation of BAF in the nucleus (65,69) and A-type lamins have been suggested to be involved in BAF accumulation at rupture sites (65). However, we and others (67) did not find an obvious defect in BAF A12T recruitment to sites of nuclear ruptures. In addition, we did not find any change in the NE repair kinetics between NGPS and control cells. As BAF A12T can still localize to the exposed DNA at the rupture site, it is likely to still be able to recruit downstream proteins required for the repair process, especially as the mutation is not predicted to interfere with LEM domain protein interactions. We did observe a reduction in emerin recruitment to NE ruptures even though this did not seem to interfere with the repair process. We propose that either multiple LEM domain proteins contribute to NE repair or the partial recruitment of emerin in NGPS cells is sufficient for its repair function at the NE. We also showed that overexpression of FLAG-BAF WT in NGPS patient cells was sufficient to rescue lamin A/C and emerin recruitment, and overexpression of BAF A12T in a wild-type background did not cause defects in the recruitment of these proteins, which explains the lack of disease phenotypes in NGPS parents who carry a BAF A12T heterozygous mutation (39,40).

Finally, our results revealed that the NE of NGPS cells was more likely to re-rupture after an initial rupture event, supporting a role for A-type lamins in strengthening the ruptured NE. So far, it was hypothesized that A-type lamin recruitment to sites of NE ruptures stabilize the nuclear envelope after it had ruptured, thereby protecting its integrity (20). However, experimental evidence for the role of A-type lamins recruitment by BAF at the NE was lacking. Here, we demonstrated that, in the presence of BAF A12T, lack of A-type lamins accumulation at sites of ruptures does not affect the dynamics of NE repair but leaves areas of the NE devoid of lamin A/C that are more prone to re-rupture. Our data suggest that the role of the BAF–lamin A/C interaction is to increase the robustness of the NE at the sites of rupture. Several mutations in the Ig-fold domain of lamin A/C found in progeroid laminopathies have
recently been shown to impair binding to BAF (32), leading to reduced recruitment of A-type lamins to rupture sites (65,67), with unknown consequences. This suggests a common mechanism for these diseases, in which lack of A-type lamins at the rupture sites creates fragile sites in the NE that are more prone to re-rupture. This mechanism could contribute to the tissue-specific severity observed in progeria and in other laminopathies. Indeed, the most severely affected tissues are the ones subjected to higher mechanical stress (heart and vascular tissues in HGPS) and/or very dense tissues (bone tissues in NGPS patients). These tissues are therefore likely to be more prone to NE ruptures.

Altogether, our work, as well as these recent studies, suggest that, once exposed to continuous mechanical forces and therefore more frequent re-ruptures, NGPS cells might over time accumulate DNA damage and/or activate inflammatory pathways that could contribute to the premature aging disease pathology.

DATA AVAILABILITY

Structure coordinates were deposited to the PDB, with entry 7Z21.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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