β-thalassemia major (β-TM) is an inherited hemoglobinopathy caused by a quantitative defect in the synthesis of β-globin chains of hemoglobin, leading to the accumulation of free α-globin chains that aggregate and cause ineffective erythropoiesis. We have previously demonstrated that terminal erythroid maturation requires a transient activation of caspase-3 and that the chaperone Heat Shock Protein 70 (HSP70) accumulates in the nucleus to protect GATA-1 transcription factor from caspase-3 cleavage. This nuclear accumulation of HSP70 is inhibited in human β-TM erythroblasts due to HSP70 sequestration in the cytoplasm by free α-globin chains, resulting in maturation arrest and apoptosis. Likewise, terminal maturation can be restored by transduction of a nuclear-targeted HSP70 mutant. Here we demonstrate that in normal erythroid progenitors, HSP70 localization is regulated by the exportin-1 (XPO1), and that treatment of β-thalassemic erythroblasts with an XPO1 inhibitor increased the amount of nuclear HSP70, rescued GATA-1 expression and improved terminal differentiation, thus representing a new therapeutic option to ameliorate ineffective erythropoiesis of β-TM.

**ABSTRACT**

β-thalassemia major (β-TM) is an inherited hemoglobinopathy caused by a quantitative defect in the synthesis of β-globin chains of hemoglobin, leading to the accumulation of free α-globin chains that aggregate and cause ineffective erythropoiesis. We have previously demonstrated that terminal erythroid maturation requires a transient activation of caspase-3 and that the chaperone Heat Shock Protein 70 (HSP70) accumulates in the nucleus to protect GATA-1 transcription factor from caspase-3 cleavage. This nuclear accumulation of HSP70 is inhibited in human β-TM erythroblasts due to HSP70 sequestration in the cytoplasm by free α-globin chains, resulting in maturation arrest and apoptosis. Likewise, terminal maturation can be restored by transduction of a nuclear-targeted HSP70 mutant. Here we demonstrate that in normal erythroid progenitors, HSP70 localization is regulated by the exportin-1 (XPO1), and that treatment of β-thalassemic erythroblasts with an XPO1 inhibitor increased the amount of nuclear HSP70, rescued GATA-1 expression and improved terminal differentiation, thus representing a new therapeutic option to ameliorate ineffective erythropoiesis of β-TM.
**Introduction**

Erythropoiesis is a multistep process leading to red cell production from hematopoietic stem cells. We have demonstrated that entry into the terminal maturation stage of erythroblasts, marked by important morphological modifications, requires a transient activation of caspase-3. However, in this process, some caspase-3 targets including GATA1, the transcription factor involved in the expression of erythroid genes like erythropoietin (EPO) receptor, Glycophorin A or globin chains, remains uncleaved. The chaperone Heat Shock Protein 70 (HSP70), which is constitutively expressed during erythroid differentiation, accumulates in the nucleus and protects GATA-1 transcription factor from caspase-3 cleavage, allowing continued terminal maturation. We have demonstrated that this nuclear accumulation of HSP70 did not occur in human β-TM erythroblasts, resulting in GATA-1 cleavage and thus erythroblast maturation arrest and apoptosis. Indeed, in β-TM, HSP70 is sequestered in the cytoplasm by the excess of free α-globin chains preventing its nuclear localization. Likewise, terminal maturation in human β-TM could be restored by transduction of a nuclear-targeted HSP70 mutant. Thus, any regulation enabling an increase in HSP70 nuclear concentration would be an interesting therapeutic strategy to correct ineffective erythropoiesis of β-TM. However, mechanisms involved in HSP70 trafficking during human erythroid differentiation are still unknown. Thus, we focused our work on elucidating these mechanisms. We have previously shown by both confocal microscopy and immunoblot analyses that, in erythroblasts derived from CD36+ cells, EPO starvation induced a dramatic decrease in both nuclear localization of HSP70 and as a consequence GATA-1 expression. Addition of the nuclear export inhibitor Leptomycin B (LMB) prevented this phenomenon. These previous observations strongly suggested that HSP70 localization could be regulated mainly by its export. Exportins are proteins of the β-karyopherin group that mediate export from the nucleus to the cytoplasm of cellular proteins (cargos) or mRNAs, using the RanGTP/GDP gradient. There are seven known exportins analyzed the expression profile of the seven different human exportins from transcriptomic and proteomic data from erythroid cells at different differentiation stages of maturation, and showed functionality that XPO1 is involved in HSP70 export and could be considered as a good target to ameliorate ineffective erythropoiesis of β-TM.

**Methods**

**Erythroid liquid culture and KPT-251 treatment**

Erythroid cells were generated in vitro from peripheral blood circulating CD34+ cells from adult patients with β0-thalassemia major (β-TM), which were collected before routine transfusion or from cord blood from healthy donors. This study was performed according to the Declaration of Helsinki with the approval from the ethics committee of our institution (Comité de Protection des Personnes (CPP) “Ile de France II”). All patients gave written informed consent. In the first step of culture (“cell expansion”), isolated CD34+ progenitors (Miltenyi CD34 Progenitor Cell Isolation Kit) were grown in the presence of 100 ng/mL IL-6, 10 ng/mL IL-3, and 100 ng/mL SCF for 7 days. At day 7, erythroid progenitors were switched to the second phase of culture, which allows the differentiation and maturation of erythroblasts: cells were cultured in the presence of 10 ng/mL IL-3, 50 ng/mL SCF, and 2 U/mL EPO in IMDM (Gibco cell culture) supplemented with 15% BIT 9500 (Stern Cell Technologies), as previously described. Solid KPT-251 was reconstituted in DMSO. Cells were treated with KPT-251 at 100 or 1000 nM, or with DMSO alone as control condition.

**Cell differentiation**

Erythroid differentiation was assessed by various methods. First, morphological analysis after May-Grunwald-Giemsa (MGG) staining was used. Cells were examined under Zeiss microscope axioplan 2, Camera Qimaging QICAM FAST 1394. The number of immature (proerythroblasts + basophilic), polychromatophilic, and mature (orthochromatic erythroblasts + reticulocytes) erythroblasts was assessed in each experiment by counting approximately 200 cells in consecutive fields and expressed as a percentage of total cells. Additionally, differentiation was assessed by calculating a terminal maturation index (TMI) on MGG, defined by the number of orthochromatic erythroblasts + reticulocytes/100/ number of polychromatophilic cells per slide. This allowed us to better characterize the maturation arrest at the polychromatophilic stage, which is known to be a hallmark of β-TM ineffective erythropoiesis, and its modulation.

Flow cytometry analysis was also performed at several different times during erythroid culture after double labeling with Band3 and 4-integrin, two optimal surface markers to differentiate highly mature erythroblasts. PE-conjugated anti-Band3 (PE-BRIC6 conjugate, Bristol Institute for Transfusion Sciences, UK) and APC-conjugated anti- 4-integrin (Milteny Biotech) antibodies were used for flow cytometry (FACS Gallois). This double labeling allowed us to assess terminal erythroblastic differentiation and purify cell populations by cell sorting (BD FACS Aria II SORP). High Band3 (high band3, low 4-integrin), and low Band3 (low band3, high 4-integrin) gates were defined, and cells from each gate were sorted and analyzed morphologically after MGG staining.

**Statistical analysis**

Statistical analyses were performed with GraphPadPrism (Version 4.0 GraphPad software). Data are expressed as the mean±standard deviation or standard error of the mean. Paired t-test, or Anova Dunn’s, Dunnett’s, and Tukey’s multiple comparison tests were used as appropriate. P<0.05 was considered statistically significant; lower P-values are indicated in the figure legends. Further information is available in the Online Supplementary Appendix.

**Results**

**XP01 can interact with HSP70 nuclear export sequence**

In our previous work, we demonstrated that HSP70 export mediated by EPO deprivation could be repressed by treatment with the nuclear export inhibitor LMB.
Figure 1 (previous page). In human erythroid progenitors, HSP70 is exported from the nucleus by an XPO1-dependent mechanism. Expression profiles for the seven different exportins (XPO1-XPO7) during terminal erythroid differentiation in human: mRNA expression from proerythroblast stage (ProE) to orthochromatic stage (Ortho). Values are extracted from public data and presented as log of reads per kilobase of transcript per million reads (logRPKM); EB: early basophile; LB: late basophile; Poly: polychromatophilic, (A) and protein expression from progenitor stage (ProG) to orthochromatic stage (Ortho). Values are presented as mean of protein copies per cell. ProG1: BFU-E; ProG2: CFU-E; Baso1: early basophile; Baso2: late basophile; Poly: polychromatophilic (B). Data are representative of three independent experiments. (C) Putative XPO1 specific leucin-rich NES in the protein sequence of human HSP70 (NP_005336) at position L394-L403. The interactions between purified XPO1 and WT HSP70 as well as XPO1 with the nuclear-targeted HSP70 mutant (S400A) were analyzed using BLI. WT HSP70 exhibits a much higher signal (i.e. affinity) for the ligand XPO1 compared to the nuclear HSP70 protein bearing a mutation in the NES residue S400A. Data are representative of two independent experiments. (D) Proximity of HSP70 and XPO1 proteins was analyzed in CD36+ erythroid progenitors derived from cord blood, by Duolink assay, using anti-XPO1 and anti-HSP70 antibodies (or anti-GATA1 for negative control). Red spots indicate <40 nm proximity between cellular-bound antibodies. Nuclei are stained with DAPI (blue). Images have been observed by confocal microscopy (x63 oil objective, scale bar= 5 μm). Data are representative of three independent experiments. (E) HSP70 and XPO1 direct interaction was demonstrated by CoIP experiments. HSP70 and XPO1 immunoblot detection is shown in total lysate (TL), in eluate from HSP70 IP and from IgG Control (IgG CTL) IP. The data are representative of three independent experiments in human erythroid cells. (F) Erythroid progenitors from β-thalassemia major (β-TM) patient at day 2 of CD36+ culture were transduced with a shRNA specific for XPO1 or a sh scramble (shCTL). Both constructions express GFP. GFP cells were sorted and stained with anti-HSP70 or anti-XPO1 antibodies, and DAPI, XPO1 and HSP70 nuclear expression (mean pixel) were analyzed at day 2 following transduction, by ImageStream. In addition, HSP70 nuclear translocation was evaluated by measuring the similarity score between HSP70 and DAPI nuclear stainings. Data are presented as mean ± standard error of mean (SEM). On average, 30,000 events were collected in all experiments. P-values are determined by paired t-test. **P<0.0001. Three illustrative images (ImageStream) of shCTL and shXPO1 conditions are presented. Cells were probed for HSP70 expression and run on the ImageStream. Bright field (white), DAPI (purple), HSP70 (green), and HSP70/DAPI composite (scale bar=7 μm). Data are representative of six independent experiments, n=2 different β-TM patients with n=2 different shRNA XPO1.
Here, we validate these previous findings using confocal microscopy and ImageStream analysis for a more quantitative approach. In normal culture conditions of cord blood-derived CD36+ erythroid progenitors at day 3, a 2-hour treatment with LMB at 20nM induced an increase in HSP70 nuclear accumulation (Online Supplementary Figure S1A and B) resulting in an increase in HSP70 nuclear/cyttoplasmic ratio.

In order to determine which specific exportin was involved in HSP70 export, we first analyzed expression of the seven exportins during erythropoiesis. All of them were expressed at the mRNA and protein levels during human erythroid differentiation as shown by transcriptomic and proteomic analysis10,11 Figure 1A and B).

Based on the proteomic database (Figure 1B), we found that exportin-2 (XPO2) is the most expressed exportin at the protein level in human erythroid progenitors. However, since primary function of XPO2 is to mediate re-export of importin-α from the nucleus to the cytoplasm, it is unlikely to be the candidate exportin involved in HSP70 nuclear export. In contrast, XPO1, in addition to exporting RNAs, mediates export of a broad range of cargo proteins bearing a leucine-rich Nuclear Export Sequence (NES),12,13 that include a large variety of tumor suppressor proteins (e.g. p53, p21, FOXO) and thereby fulfills all the criteria to be a good candidate in exporting HSP70. We performed in silico analysis of published data for GATA-1 chromatin immuno-precipitation (ChIP) in human erythroid cells,14 and found twelve annotated peaks for GATA-1 in the promoter of the XPO1 gene suggesting that it might be an erythroid regulated gene (Table 1). In addition, XPO1 was the second most expressed exportin in erythroblasts, and concomitantly with HSP70 nuclear accumulation, its expression was down-regulated both at the mRNA and protein levels along terminal differentiation (black line in Figure 1A and B). This hypothesis was further supported by in silico analysis showing that human HSP70 protein sequence contains a putative leucine-rich NES at position L394-L400 that could interact with XPO1 (Figure 1C, NES human HSP70 protein WT).

In order to validate this hypothesis, we first showed that XPO1 interacted with HSP70 in vitro by protein-protein interaction experiments using BioLayer interferometry (BLI) (Figure 1C, red line). Furthermore, we observed a decreased binding affinity between XPO1 and HSP70 with the mutant Serine 400 Alanine (HSP70S400A) in the putative leucine-rich NES (Figure 1C, blue line). This mutant HSP70 shows restricted nuclear localization when transduced in erythroid cells. Thus, these results validate the importance of putative NES in the interaction of HSP70 with XPO1, and in the cellular localization of HSP70. Next, using proximity ligation assay (PLA/Duolink assay) (Figure 1D) and performing CoIP experiments in normal conditions (Figure 1E), we obtained additional evidence that this interaction is direct and occurs in vivo in CD36+ derived erythroid progenitors. Interestingly, the fraction of XPO1 protein immunoprecipitating with HSP70 upon EPO deprivation is increased as shown by CoIP experiments (data not shown).

Table 1. GATA-1 chromatin immunoprecipitation (ChIP) peak scores in XPO1 gene and three known erythroid genes.

| Gene name | Genomic peak location | Peak score |
|-----------|-----------------------|------------|
| XPO1      | Intragenic intron     | 53         |
|           | Intragenic intron     | 695.35     |
|           | Intragenic intron     | 76.79      |
|           | Intragenic intron     | 94.13      |
|           | Intragenic intron     | 231.83     |
|           | Immediate down. Intron| 62.88      |
|           | Promoter              | 292.11     |
|           | Enhancer              | 110.24     |
|           | Intergenic intron     | 66.67      |
|           | Intergenic intron     | 288.01     |
|           | Intergenic intron     | 73.56      |
|           | Intergenic intron     | 89.17      |
| EPOR      | Intragenic exon       | 101.5      |
|           | Immediate down. Intron| 3100       |
|           | Enhancer              | 72.94      |
|           | Enhancer              | 186.49     |
| HBA1      | Enhancer              | 55.94      |
|           | Promoter              | 898.4      |
|           | Promoter              | 106.72     |
| HBA2      | Enhancer              | 1259.62    |
|           | Enhancer              | 313.4      |
|           | Promoter              | 55.51      |

Annotated peaks extracted from GATA-1 ChIP public data on human erythroid progenitors14 analyzed with Galaxy software. Peak scores and locations for XPO1 and the known erythroid genes EPOR (erythropoietin receptor) HBA1 (Hemoglobin α-1) and HBA2 (Hemoglobin α-2). Peak score in MACS output corresponds to $-10^9 \log_{10} \text{pvalue}$ of peak region. Pvalue corresponds to the probability that the candidate peak (reads enrichment) is random.

**XPO1 is the exporter of HSP70 in human erythroid progenitors**

To document the functional role of XPO1 in human erythropoiesis, and its putative target to treat β-TM, we first repressed XPO1 expression in human β-TM erythroblasts. For this purpose, CD36+ cells from a β-TM patient were transduced with lentiviruses expressing a XPO1 specific shRNA (shXPO1) or a sh scramble used as control (shCTL). Two days after transduction, ImageStream quantification showed that the cells transduced with shXPO1 efficiently repressed the expression of XPO1 [mean pixel nuclear XPO1 556.4±3.517 (shCTL) vs. 520.0±2.767 (shXPO1) (P<0.0001)] (Figure 1F and Online Supplementary Figure 2A). Similar results were obtained in erythroblasts from cord and peripheral blood stem cells (data not shown). In agreement with our hypothesis, XPO1 repression was associated with a significant increase in HSP70 nuclear translocation [similarity score 1.090±0.012 for shCTL vs. 1.378±0.011 for shXPO1 (P<0.0001)] and in HSP70 nuclear accumulation [mean pixel nuclear HSP70 139.1±1.176 for shCTL vs. 159.2±1.173 for shXPO1 (P<0.0001), corresponding to a 14.5% increase] (Figure 1F). Similar results were obtained using two different shRNA specifically targeting XPO1. As a control of efficacy, P53 that is a well-known target of XPO1, was also significantly increased in the nucleus (Online Supplementary Figure 2B and C). Taken together, these data suggest that in human erythroid cells, HSP70 interacts with XPO1 through a leucine-rich NES region, which is required for its nuclear export and may thus be a good target to improve ineffective erythropoiesis of β-TM.
Chemical inhibition of XPO1 using KPT-251 treatment increases the amount of nuclear HSP70 and GATA-1 in β-TM erythroid progenitors

We repressed XPO1 activity using KPT-251 (Merck Millipore), a Selective Inhibitor of Nuclear Export (SINE) that specifically inhibits the formation of XPO1-cargo complex by interacting with the NES binding-groove of XPO1. KPT-251 treatment induces minimal toxicity in non-cancerous hematopoietic cells both in vitro and in vivo,15-19 with almost no effect on cell survival (Figure 2A and C) and a decrease in cell proliferation but not significant (Figure 2B and D). This is in contrast to LMB treatment, an inhibitor of nuclear export which is highly toxic for erythroid progenitors (data not shown). We exposed erythroid progenitors from β-TM patients, at day 4 of CD36+ day of culture, to KPT-251 at 100nM, 1000nM, or to DMSO (control) for 72 hours (h). As evidenced by immunoblot analyses at day 7 of CD36+ culture, the treatment resulted in a significant dose-dependent decrease in the amount of XPO1 protein compared to control (DMSO) in cytoplasmic extracts (CE) (Figure 3A). XPO1 protein decrease following KPT treatment was also observed in erythroid progenitors derived from cord blood, and significant XPO1 protein decrease can be observed in erythroid progenitors from 24 h of treatment (data not shown). This finding is consistent with previous studies using different primary cancer cells and cell lines.20,21 XPO1 repression by the treatment probably acts through a proteasome-dependent mechanism as reported earlier for KPT-185 and KPT-330, two other members of the KPT family.22,23 In β-TM erythroblasts, western blot analysis showed the decrease in XPO1 protein amount is associated with a significant dose-dependent increase in HSP70 protein amount in nuclear extracts (NE) (Figure 3A). Nuclear GATA1 amounts and cytoplasmic HSP70 amounts remain unchanged. These results were further confirmed by confocal analyses (Figure 3B); after a 72-h exposure to KPT-251, increases were observed in HSP70 nuclear concentration [mean fluorescence intensity (MFI) nuclear HSP70 48.27±4.6 (control), 58.85±5.2 (KPT 100 nM) (NS) and 79.06±9.0 (KPT 1000nM) (P<0.01)] and in HSP70 nuclear/cytoplasmic (N/C) ratio [MFI ratio 0.24±0.02 (control), 0.35±0.02 (KPT 100nM) (P<0.01) and 0.57±0.03 (KPT 1000nM) (P<0.01)]. Consistent with an increase in nuclear location of HSP70, GATA-1 nuclear concentration was also increased as assessed by confocal analyses [MFI nuclear GATA-1 48.53±5.9 (control), 44.32±2.9 (KPT 100 nM) (NS) and 74.17±2.8 (KPT 1000nM) (P<0.01)] (Figure 3B). Quantification by ImageStream (Figure 3C and D) further confirmed the increase in HSP70 nuclear accumulation by KPT-251 treatment [mean pixel nuclear HSP70 95.22±0.40 (control), 98.27±0.36 (KPT 100nM) (P<0.01), or 108.2±0.38 (KPT 1000nM) (P<0.01)], and of the HSP70 N/C ratio [mean pixel ratio 1.55±0.02 (control), 1.65±0.02 (KPT 100nM) (P<0.01), and 1.64±0.02 (KPT 1000nM) (P<0.01)] and GATA-1 nuclear expression [mean pixel nuclear GATA-1 730.2±2.1 (control), 756.2±2.2 (KPT100nM) (P<0.01) and 816.5±2.1 (KPT 1000nM) (P<0.01)] induced by KPT treatment, in a dose-dependent manner. In addition, as determined by similarity score, the fraction of erythroblasts with a nuclear translocation of HSP70 was increased fol-

Figure 2. KPT-251 treatment has low effect on cell proliferation and cell death. Cell death and proliferation curves analysis of β-thalassemia major (β-TM) (A and B) and cord blood (C and D) erythroid progenitors, assessed by blue trypan staining at 24, 48 and 72 hours (h) of treatment with KPT100nM, KPT1000nM, or DMSO (control). Daily mean percentage±standard deviation (SD) of dead cells (n=5 independent experiments for β-TM and n=3 independent experiments for cord blood). Daily mean±standard deviation of cell proliferation (n=6 independent experiments for β-TM and n=3 independent experiments for cord blood). P-values are determined by ANOVA Dunnett’s multiple comparison test **P<0.01, NS: not significant.
XPO1 is a target to treat β-thalassemia

Figure 3. KPT-251 treatment increases the amount of nuclear HSP70 and GATA-1 in β-thalassemia major (β-TM) erythroid progenitors. Erythroblasts derived from β-TM peripheral blood cells were treated at day 4 of CD36+ cell culture with 100nM, 1000nM of KPT-251, or with DMSO (control) for 72 hours (H). All data were analyzed at day 7 of CD36+ cell culture (72 hours of treatment). (A) Immunoblot from 10 μg of nuclear extracts (NE) and 30 μg of cytoplasmic extracts (CE), (representative of three independent experiments performed on two different β-TM patient cell cultures). Graph shows optical relative quantity values of XPO1, HSP70 and GATA1 proteins normalized to that of HSP90 for CE and to that of HDAC2 for NE. Conditions KPT 100nM and 1000nM are normalized to that of DMSO condition. Absence of cytoplasmic proteins contamination in nuclear extracts is evidenced by the absence of HSP90 in NE. (B) Graph shows nuclear mean fluorescence intensity (MFI) of HSP70 and GATA-1, and HSP70 nuclear/cytoplasmic (N/C) ratio of MFI in treated (KPT 100 and 1000) and control (DMSO) cells determined by confocal microscopy images analyses. Data are presented as mean±standard error of mean (SEM) (for a minimum of 30 cells per condition), and are normalized on area. P-values are determined by ANOVA Dunnett’s multiple comparison test. Data are representative of three independent experiments. (C) HSP70 and GATA-1 nuclear expression (mean pixel), HSP70 N/C ratio (mean pixel) and HSP70 nuclear translocation (similarity score) were analyzed by ImageStream. Data are presented in histograms as mean±SEM. P-values are determined by ANOVA Dunnett’s multiple comparison test (representative of three independent experiments). On average, 30,000 events were collected in all experiments. (D) Three illustrating images of ImageStream experiments. Cells were probed for HSP70 and GATA-1 expression and run on the ImageStream. Bright field (white), HSP70 (green), GATA-1 (red), DAPI (purple) and HSP70/DAPI composite (scale bar=7μm). Respective similarity scores±SEM are indicated under each group of images. *P<0.05; **P<0.01.
Figure 4. KPT-251 treatment improves terminal erythroid maturation in β-thalassemia major (β-TM) erythroid progenitors in vitro. β-TM erythroid progenitors were treated at day 4 of CD36+ cell culture with 100nM, 1000nM of KPT-251, or with DMSO (control) for 72 hours (day 7 CD36+ cell culture). (A) The mean fluorescence intensity (MFI) of Band 3 was analyzed by flow cytometry after 72 hours of treatment. MFI were normalized on DMSO condition. P-values are determined by ANOVA Dunn’s multiple comparison test **P<0.01, NS: not significant (n=8 independent experiments, n=3 different β-TM samples). (B) (Left) Percentage of high Band3 cell population under the different treatment conditions and (Right) absolute number of high Band3 cells, normalized to DMSO treatment. P-values are determined by ANOVA Dunn’s multiple comparison test, **P<0.01, NS: not significant, n=8 independent experiments, n=3 β-TM patients. (C) Representative flow cytometry plots (α4-integrin and Band3 staining) of β-TM erythroid progenitors treated with KPT1000nM or DMSO. Strategy for cell sorting purification of high Band3 (red box) and low Band3 (blue box) erythroblasts populations after 72 hours (H) of KPT1000nM or DMSO treatment (day 7 CD36+ cell culture). A representative morphological analysis x(25) oil objective, scale bar= 10 μm of purified cells from each gate by May-Grünwald-Giemsa staining. Corresponding graph showing the percentage of mature cells (orthochromatic erythroblasts + reticulocytes) contained in low Band3 and in high Band3 gates (n=3 independent experiments, n=2 different β-TM patients). P-values are determined by paired t-test **P<0.001, NS: not significant. (D) Corresponding TM index for the different conditions of treatment. P values are determined by ANOVA Dunn’s multiple comparison test *P<0.05, NS: not significant (n=5 independent experiments).
Following KPT-251 treatment [similarity score 0.7483±0.006 (control), 0.8587±0.005 (KPT100nM) (P<0.01) and 0.9872±0.005 (KPT1000nM) (P<0.01)].

To further demonstrate that XPO1 is indeed the main protein involved in HSP70 nuclear export, we tested its role in HeLa cells, which upon heat shock at 43°C exhibited nuclear HSP70 localization as a consequence of both an increase in HSP70 nuclear inflow rate due to an increase of Hikeshi expression and a reduction in nuclear outflow by an unknown mechanism.24 After a 6-h recovery phase at 37°C, the outflow rate increases and HSP70 progressively re-localizes in the cytoplasm.24 To demonstrate the role of XPO1 in the nuclear export of HSP70, HSP70 outflow following heat shock was analyzed by ImageStream with or without XPO1 repression (24 h KPT-251 pre-treatment at 1000nM). As expected, heat shock induced increased nuclear HSP70 localization and after 6 h of recovery at 37°C, HSP70 exited the nucleus, which was delayed when cells have been pre-treated with KPT-251 as compared to control treated cells (Online Supplementary Figure S3). This observation suggests that XPO1 is the main exportin of HSP70.

**Chemical inhibition of XPO1 using KPT-251 treatment ameliorates erythroid terminal differentiation of β-TM erythroid progenitors with low cytotoxicity**

As expected, this nuclear accumulation of both HSP70 and GATA-1 was associated with an increase in terminal erythroid differentiation of β-TM erythroblasts as assessed by flow cytometry analysis showing a significant increase in total Band3 MFI [MFI normalized on DMSO treated cells used as a control: DMSO=1, KPT100nM=1.06±0.04 (NS), and KPT1000nM=1.48±0.09 (P<0.01)] (Figure 4A). Consistent with an increase in terminal erythroid maturation, the fraction of cells expressing high Band3 was also increased by KPT treatment in a dose-dependent manner [2.62%±0.39 (control), 3.13%±0.53 (KPT100nM) (NS) and 9.27%±1.44 (KPT 1000nM) (P<0.01)] (Figure 4B, left). The absolute number of high Band3 cells was also increased (Figure 4B, right). To ensure that this high Band3 pool rep-
resented orthochromatric erythroblasts and reticulocytes, as previously described, we sorted them by FACS based on their Band3 and 4-integrin expression (Figure 4C). As expected, mature erythroblast content (orthochromatric erythroblasts+reticulocytes) was enriched in high Band3 gate (62.23%±5.31) compared to low Band3 gate (22.83%±4.49) (P=0.0004).

We further confirmed that in β-TM progenitors, KPT treatment alleviated the maturation arrest at the polychromatophilic stage as it decreased the proportion of polychromatophilic cells and increased the proportion of mature cells (Figure 4D). More importantly, KPT treatment significantly increased the terminal maturation index (TIM), determined as the ratio of orthochromatric erythroblasts+reticulocytes *100/ polychromatophilic cells, at the dose of 1000nM KPT (TIM normalized to control DMSO: DMSO=1, KPT100nM=1.5±0.5 (NS) KPT1000nM=3.7±1.05, (P<0.05)] (Figure 4E). Finally, we show that the significant increase in TIM and in absolute number of mature cells at 1000nM is associated with a high increase in HbF amount in the pool of mature cells, which was clearly observed at lower doses of treatment. This effect was observed with a greater magnitude at higher dose (1500nM). However at this dose, inhibition of cell proliferation and apoptosis were higher (Online Supplementary Figure S4).

In Figure 5, we schematically illustrate the molecular mechanisms modulated by KPT treatment in β-TM erythroid progenitors compared to β-TM and normal erythroid progenitors in normal conditions.

Discussion

Taken together, our data demonstrate that XPO1 is the main nuclear exporter of HSP70 in various cell types, and it may participate in the regulation of HSP70 in human erythroblasts during normal and pathological erythropoiesis. Decreased expression of XPO1 along erythroid differentiation provides an explanation for the augmentation of nuclear localization of HSP70 during normal erythroid maturation. Interestingly, proteins of the Ran Machinery (RAN, RANBP1, RCC1, NUTF1 AND RAN-GAP1) essential for XPO1 activity are also expressed during erythroid differentiation and also down-regulated at the time of caspase activation (data not shown).

Furthermore, we demonstrate that inhibition of XPO1 by KPT-251 treatment improves terminal maturation of β-TM erythroblasts by preventing GATA-1 degradation by caspase-3. Thus, XPO1 inhibitors could be added to the armamentarium of therapeutic options of β-TM to correct ineffective erythropoiesis and its pathological consequences (e.g. iron overload and extramedullary hematopoiesis), to increase hemoglobin level, to reduce transfusion burden and ultimately to increase overall survival. Interestingly, KPT compounds are currently under clinical development and exhibit high efficacy and safety profile in cancer therapy. More recently, a new generation of KPT compounds with reduced toxicity is being developed (clinicaltrials.gov identifier: NCT02649790) and as such, if shown to be not toxic in the long-term, they could be used in β-TM as well. In addition, maintaining HSP70 in the nucleus by blocking XPO1 could be useful in other pathologies of erythropoiesis where HSP70 is not localized in the nucleus, such as myelodysplastic syndrome and congenital erythroblastopenia, but also in other diseases such as some neurodegenerative diseases, in which protein aggregates may also prevent HSP70 nuclear location.

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