Extracellular domain N-glycosylation controls human thrombopoietin receptor cell surface levels

Roxana I. Albu1,2 and Stefan N. Constantinescu1,2*

1 Ludwig Institute for Cancer Research, Brussels, Belgium
2 de Duve Institute, Université catholique de Louvain, Brussels, Belgium

INTRODUCTION
The thrombopoietin receptor (TpoR) is a type I transmembrane protein that mediates the signaling functions of thrombopoietin (Tpo) in regulating megakaryocyte differentiation, platelet formation, and hematopoietic stem cell renewal. We probed the role of each of the four extracellular domain putative N-glycosylation sites for cell surface localization and function of the receptor. Single N-glycosylation mutants at any of the four sites were able to acquire the mature N-glycosylated pattern, but exhibited a decreased Tpo-dependent JAK2–STAT response in stably transduced Ba/F3 or Ba/F3-JAK2 cell lines. The ability of JAK2 to promote cell surface localization and stability of TpoR required the first N-glycosylation site (Asn117). In contrast, the third N-glycosylation site (Asn298) decreased receptor maturation and stability. TpoR mutants lacking three N-glycosylation sites were defective in maturation, but N-glycosylation on the single remaining site could be detected by sensitivity to PNGaseF. The TpoR mutant defective in all four N-glycosylation sites was severely impaired in plasma membrane localization and was degraded by the proteasome. N-glycosylation receptor mutants are not misfolded as, once localized on the cell surface in overexpression conditions, they can bind and respond to Tpo. Our data indicate that extracellular domain N-glycosylation sites regulate in a combinatorial manner cell surface localization of TpoR. We discuss how mutations around TpoR N-glycosylation sites might contribute to inefficient receptor traffic and disease.

Keywords: cytokine receptor, thrombopoietin, JAK2, N-glycosylation, signal transduction, endoglycosidase H, ER maturation, cell surface traffic

Abbreviations: CAMT, congenital amegakaryocytic thrombocytopenia; FT, familial thrombocytosis; JAK, Janus kinase; STAT, signal transducers and activators of transcription; TpoR, thrombopoietin receptor.
Little is known about regulation of TpoR traffic to the cell surface or about the mechanisms that govern receptor internalization, degradation, and recycling. On the cytosolic side, JAK2 and TYK2 were shown to promote cell surface localization and stability of TpoR (Royer et al., 2005), but the extracellular determinants of endoplasmic reticulum (ER) to cell surface traffic remain unknown.

The role of extracellular domain N-glycosylation for cytokine receptor traffic remains debated. N-glycosylation is apparently not required for induction of gene transcription via PrlR and gp130, but is crucial for receptors’ stability and cell surface localization (Buteau et al., 1998; Bolander, 1999; Waetzig et al., 2010). On the other hand, in the case of leptin and GM-CSF receptors, N-glycans are important for ligand binding (Ding et al., 1995; Niu et al., 2000; Kamikubo et al., 2008). Eliminating individual N-glycans in human β-common subunit of the IL3, IL5, and GM-CSF receptors did not affect ligand binding and receptor activation, but the authors did not exclude the possibility that N-glycans may exert some sort of fine control (Murphy et al., 2008). TpoR presents four putative N-glycosylation sites in its extracellular domain, two in each cytokine receptor module (Figure 1). In this study we aimed to probe the role of each N-glycosylation site in the cell surface expression and function of TpoR.

MATERIALS AND METHODS
PLASMIDS AND GENERATION OF HUMAN TpoR N-GLYCOSYLATION DEFECTIVE VARIANTS

The human TpoR wild-type (WT) was cloned into the pMX-IRES-GFP bicistronic retroviral vector upstream of the IRES as described (Liu et al., 2000). Several mutagenesis reactions with specific primers were carried out on human TpoR WT (Swiss-Prot accession number P40238) with the QuickChange Site-Directed Mutagenesis Kit (Stratagene). Human TpoR variants lacking single N-glycosylation sites were obtained by changing the codon for Asn (AAC or AAT) to the codon for Gln (CAA) in each of the four potential N-glycosylation sequons. The resulting proteins were named (Δ (1), Δ (2), Δ (3), and Δ (4), respectively, according to the particular N-glycosylation site that was removed (Table 1). Other mutants lacking more than one N-glycosylation site were obtained leaving only one N-glycosylated site: (Δ (123), (Δ (134), (Δ (124), (Δ (234) (Table 1). All constructs were verified
Table 1 | Name and description of the engineered human TpoR variants.

| Name of TpoR variant | Mutated N-glycosylation sites* | Remaining N-glycosylation sites* |
|----------------------|---------------------------------|---------------------------------|
| Δ (1234)             | N117, N178, N298, N358          | –                               |
| Δ (1)                | N117                            | N178, N298, N358                 |
| Δ (2)                | N178                            | N117, N298, N358                 |
| Δ (3)                | N298                            | N117, N178, N358                 |
| Δ (4)                | N358                            | N117, N178, N298                 |
| Δ (123)              | N117, N178, N298                 | N358                            |
| Δ (124)              | N117, N178, N358                 | N298                            |
| Δ (134)              | N117, N298, N358                 | N178                            |
| Δ (234)              | N178, N298, N358                 | N117                            |

*Residue number for the Asn in the Asn-X-Ser/Thr sequon, TPOR_HUMAN, Swiss-Prot accession number P40238.

by sequencing. All TpoR constructs contain an HA tag at their N-terminus, after the signal peptide cleavage site. The murine JAK2 WT was subcloned into the pREX-IRES-CD4 vector.

GENERATION OF CELL LINES AND RETROVIRAL TRANSDUCTIONS

HEK293-derived BOSC cells are ecotropic retrovirus packaging cells (Pear et al., 1993). γ2A are JAK2-deficient human fibrosarcoma cells (Pellegrini et al., 1989). Ba/F3 parental and Ba/F3-JAK2 cells are IL3-dependent murine pro-B cells (Palacios and Steinmetz, 1985). Ba/F3-JAK2 is a stably transduced cell line with murine JAK2 WT (Royer et al., 2005) in order to have increased endogenous JAK2 levels of Ba/F3 cells) and the cDNA coding for each individual TpoR mutant, if not mentioned otherwise (in some cases the cDNA of human JAK2 WT was added). Empty vector was added in order to normalize the total transected cDNA. Four hours after transfection the cells were stimulated with recombinant human Tpo (R&D System) or left unstimulated for 20 h. Twenty-four hours after transfection, cells were lysed in 100 μl 1× passive lysis (Promega) and luminescence was recorded on a Glomax 96-well plate luminometer. When performing the assay on Ba/F3 or Ba/F3-JAK2 cells (also stably expressing each human TpoR mutant), cells were starved 3–4 h in RPMI medium with 1 mg/ml BSA, and electroporated (250 V, 75 Ω, 1,500 μF) with the pGRR5 and pRLTK luciferase reporters. The cells were subsequently cultured for 2 h, stimulated with recombinant human Tpo (rhTpo, R&D System) or IL3 or left unstimulated. After 2 h, the cells were lysed in 100 μl 1× passive lysis buffer and their luminescence was recorded. All the luciferase assays were performed using the dual luciferase reporter kit (Promega).

Tpo-DEPENDENT PROLIFERATION ASSAY

Three thousand stably transduced Ba/F3 cells were seeded in 96-well plates and stimulated with different concentrations of rhTpo, or mock. Control for proliferation was performed using 10 ng IL3/ml for each cell line. After 72 h, tritiated thymidine was added to the cells for 4 h. Cells were then collected on microfiltered plates, and thymidine incorporation was measured with Top Count microplate scintillation counter (Canberra-Packard, Meriden, CT, USA).

SURFACE EXPRESSION OF HA TAGGED TpoR MUTANTS

Surface expression of the receptors was measured in 10×10⁶ Ba/F3, Ba/F3-JAK2, or HEK293 cells by flow cytometry using 10 μg/ml monoclonal anti-HA antibody (HA.11, Covance) and 5 μg/ml R-phycocerythrin-conjugated donkey F(ab')₂ anti-mouse IgG secondary antibody (Jackson ImmunoResearch; Huang et al., 2001).

WESTERN BLOTTING, ENDOGLYCOSIDASE-H (ENDOH), AND N-GLYCOSIDASE F (PNGaseF) DIGESTION

Ba/F3, Ba/F3-JAK2, or HEK293 cells expressing stably or transiently each human TpoR mutant were washed in cold PBS, lysed in 200 μl Nonidet P-40 buffer with sodium orthovanadate, sodium fluoride, phenylmethanesulfonyl fluoride, and complete protease inhibitor cocktail (Roche Applied Science). Cell lysates were digested with EndoH or PNGaseF (both from New England Biolabs) or left untreated for 16 h at 37°C. The lyzate supernatants/products of the digestions were then mixed with Laemmli Blue 4×, boiled for 10 min, and centrifuged for 1 min at 14,000 rpm. A volume equivalent to 30 μg protein/well was loaded on 4–12% BisTris precast gels (Invitrogen). After transfer to nitrocellulose membranes and blocking in 20% milk/TBS–Tween, immunoblotting was performed overnight at 4°C with mouse anti-HA tag (HA.11, Covance), mouse anti-β-actin (Sigma), or rabbit anti-JAK2 (Cell Signaling) antibodies in a solution of 3% milk/TBS–Tween with a 1:1,000 antibody dilution. Secondary mouse or rabbit-horseradish peroxidase antibodies (GE Healthcare, UK) were used in a 1:10,000 dilution in 3% milk/TBS–Tweed.

TpoR variant

Table 1 | Name and description of the engineered human TpoR variants.
MEASUREMENTS OF TpOR MUTANT HALF-LIFE AND DEGRADATION

For measuring the contribution of the proteasome- or lysosome-mediated degradation of the TpOR N-glycosylation defective mutant Δ(1234) versus WT, stably transduced Ba/F3 or Ba/F3-JAK2 cells were incubated 3 h at 37°C with 50 μg/ml cycloheximide (Sigma; to block protein synthesis) and proteasome inhibitor MG132 (10 μM; Sigma) or with 50 μg/ml cycloheximide and lysosome inhibitors (200 μM chloroquine or 10 μg/ml leupeptin; Roche). Control cells were kept untreated to have 100% protein expression. Cells were lysed and analyzed by Western blotting with antibodies anti-HA for TpOR levels.

The half-lives of different TpOR variants were determined by treating the Ba/F3 stably transduced cell lines with 50 μg/ml cycloheximide for different time lengths. Cells were then lysed and analyzed by Western blotting with antibodies anti-HA for the levels of immature and mature TpOR. Anti-β-actin was used as a control.

RESULTS

HUMAN TpOR CELL SURFACE LEVELS AND ACTIVITY ARE NOT CRITICALLY DEPENDENT ON INDIVIDUAL N-GLYCOSYLATION

The first question of our study was whether there is a single critical N-glycosylation site of human TpOR that is responsible for the efficiency of cell surface localization. Taking into account that JAK2 is able to promote cell surface localization and to increase TpOR protein stability in the Ba/F3 cell line (Royer et al., 2005), we assessed cell surface levels of single N-glycosylation mutant TpOR variants in Ba/F3 cells with endogenous or elevated JAK2 levels.

Thrombopoietin receptor mutants lacking one out of four putative N-glycosylation sites (Figure 1; Table 1) were obtained by using site-directed mutagenesis on a human TpOR WT template, and then cloned into the bicistronic vector, pMX-IRES-GFP. The constructs were then used to produce retroviral supernatants in HEK293-derived BOSC cells. The retroviral supernatants were subsequently utilized to infect murine Ba/F3 cells expressing or not increased levels of murine JAK2 WT (cloned in bicistronic pREX-IRES-CD4 vector). To avoid clone-specific artifacts in expression or signal transduction, stably transduced Ba/F3 cells pools, with the same level of TpOR construct expression were used. More precisely, the cells were sorted for similar GFP levels 72 h later (correlated with the same mRNA level for each TpOR variant). Sorted cell lines were lysed and the protein for each TpOR variant was analyzed for the correct molecular size using anti-HA in Western blotting (data not shown).

Flow cytometry using HA.11 antibody performed in Ba/F3 cell lines expressing equivalent GFP expression consistently showed that, compared with the WT receptor, Δ(1) cell surface localization was decreased, while Δ(3) cell surface localization was increased (Figure 2A). In the Ba/F3-JAK2 cell line, the four variants lacking single N-glycosylation site are expressed at the same level [Δ(2), Δ(3), Δ(4)] or lower, as is the case for Δ(1) (Figure 2A). We also transiently expressed these variants in HEK293 cells, achieving overexpression. Also in these conditions Δ(1) cell surface localization was decreased, while Δ(3) cell surface localization was increased (data not shown).

The flow cytometry results were verified by Western blots experiments, where we enzymatically analyzed the N-glycosylation pattern. The fact that all the TpOR variants defective in single N-glycosylation site are able to localize at the cell surface is confirmed in Figure 2B. We showed in the HEK293 cell line that these variants presented a strong EndoH resistant band, which correlated with the mature form of each variant. When the first N-glycosylation site is removed, the level of the mature receptor band seems to be weaker compared with the mature band of the WT, correlating very well with the decreased cell surface localization for this mutant.

To test whether the cell surface receptors defective in single N-glycosylation site are as stable as the WT TpOR, we used Ba/F3 cells stably transduced with each of the mutants. The cells were treated with CHX (cycloheximide) over different time periods, in order to block protein synthesis and determine the half-life of each
TpoR variant by using anti-HA and anti-β-actin Western blotting. Compared with the WT, Δ (3) TpoR showed higher and Δ (1) TpoR lower stability (Figure 2C), which correlated well with the cell surface level differences. We were interested to determine the relevance of each of the N-glycans for the TpoR function. For each of the human TpoR variants lacking one N-glycosylation site, we assessed the STAT3 and STAT5 transcriptional activities using luciferase assays in stably transduced Ba/F3 or Ba/F3-JAK2 cell models. These TpoR variants presented a decreased transcriptional activity in the Tpo-stimulated condition, compared with the WT TpoR (Figure 3A). Between the TpoR variants the levels of Tpo-dependent transcriptional activity correlated well with the cell surface levels of the TpoR variants, with Δ (1) having the lowest and Δ (3) the highest Tpo-dependent transcriptional activity.

Our preliminary observation was that Tpo binding to TpoR is not critically dependent on individual N-glycosylation sites. To test this observation in a second set of experiments, we measured Tpo-dependent cell growth of Ba/F3 stably transduced with each of the TpoR variant. Cells were washed to remove IL3, incubated 72 h in Tpo or IL3 or left untreated. As a negative control, we used Ba/F3 cells without any TpoR variant. As a positive control for cell growth, we used IL3 treatment of each cell line. After 72 h, tritiated thymidine was added to the cells for 4 h. The proliferation of the cells was quantified according to the level of the incorporated thymidine (Figure 3B). The order of the cell proliferation level was Δ4 > Δ3 = WT > (Δ2 > Δ1) and the result is consistent with the result obtained in the luciferase test of Figure 3A.

**REMOVAL OF THREE N-GLYCOSYLATION SITES DECREASED CELL SURFACE LEVELS AND ACTIVITY OF HUMAN TpoR**
The Ba/F3 and Ba/F3-JAK2 cells were stably transduced with the variants missing different combinations of N-glycosylation sites (Table 1). The cells were sorted for the same protein expression level for each TpoR variant and tested for cell surface expression of the receptors. All of the variants, with only one N-glycosylation site left [(Δ (123), Δ (124), (Δ (134), Δ (234))] were located at a lower level at the cell surface, compared with the WT TpoR (Figure 4A). JAK2 overexpression did not increase the cell surface level of these variants. We also tested the cell surface levels of these variants in transiently transfected HEK293 cells and found the levels are higher than in Ba/F3 but still lower than the WT or the single N-glycosylation site defective variants.

We next sought to determine the carbohydrate occupancy of the four putative N-glycosylation sites. We performed EndoH (to detect shifts in the case of high mannose glycosylated immature receptors that are localized in ER or early Golgi) or PNGaseF (to remove the N-glycans from the immature and mature proteins) treatments on the transiently transfected HEK293 cell lysates. We analyzed the enzymatic products in electrophoresis under reducing conditions, and Western blotting for HA immunodetection. Because all these TpoR variants have only one putative N-glycosylation site left and all of them showed a faster migration in the presence of EndoH or PNGaseF (Figure 4B) compared with the untreated condition, we concluded that all of the possible sites could be N-glycosylated. Thus sites 1, 2, 3, and 4 at Asn residues 117, 178, 298, and 358, respectively, are fully occupied.

The results from the enzymatic assessment of these TpoR variants on transiently transfected HEK293 cell lysates showed that there is very low or absent EndoH resistant band (mature receptor, cell surface localized receptor) when compared with the WT TpoR (Figure 4B), supporting the observation of the flow cytometry results of Figure 4A.
To test the stability of TpoRs defective in multiple N-glycosylation sites, we used stably transduced Ba/F3 cells with each of the mutants and treated the cells with CHX over different time periods. A very unstable pattern was observed for the three N-glycosylation mutated variants (Figure 4C), indicating that N-glycosylation exerts a protective action on TpoR stability.
We next tested in a more sensitive assay, the ability of these TpoR variants to localize at the cell surface and to bind Tpo. Compared with the WT, a dramatic decrease of JAK2–STAT transcriptional activity for these mutants was observed in stably transduced Ba/F3 or Ba/F3-JAK2 cell models (Figure 4D). In the case of Δ (123) and Δ (234) the Tpo-dependent JAK2–STAT transcriptional activity was decreased compared with the WT but higher than in the case of Δ (124) and Δ (134) variants. We further tested the same mutants in transiently transduced HEK293 cells (see below Figure 6), but this difference between the mutants defective in three glycosylation sites was not statistically significant, suggesting that it might either reflect cell type specific differences, or be relevant at the lower stable expression levels achieved in the Ba/F3 system after sorting. These results pointed to the first (Asn117) or the last N-glycosylation site (Asn358) being important, for cell surface localization (Figure 4A) or stability (Figure 4C) in Ba/F3 cells.

**MAJOR IMPAIRMENT OF TpoR CELL SURFACE LOCALIZATION AND ACTIVITY BY REMOVAL OF ALL N-GLYCOSYLATION SITES**

Although the removal of three N-glycosylation sites dramatically affected the cell surface localization and Tpo-dependent cell signaling in Ba/F3 cells, we also wanted to test the TpoR variant lacking all of the four putative N-glycosylation sites, Δ (1234). This variant was completely absent from the cell surface by flow cytometry both in Ba/F3 or Ba/F3 overexpressing JAK2 (Figure 5A). Our preliminary observation is that, for a cell surface level detectable by flow cytometry, at least one N-glycosylation site needs to be glycosylated (Figures 4A,D).

Due to the fact the Δ (1234) TpoR was not detected at the cell surface, we tested its degradation features compared with the WT. We performed Western blot on the lysates of Ba/F3 and Ba/F3-JAK2 cell lines stably transfected with this mutant or the WT TpoR. Before lysis, these cells were incubated 3 h at 37°C with 50 μg/ml cycloheximide (to block protein synthesis) and proteasome inhibitor MG132 (10 μM) or with 50 μg/ml cycloheximide and lysosome inhibitors (200 μM chloroquine or 10 μg/ml leupeptin). We observed a stabilization effect due to the blocking of the proteosomal degradation (Figure 5B). Thus, the TpoR variant lacking all four putative N-glycosylation sites [Δ (1234)] is a very unstable protein that can be stabilized by blocking proteasome-mediated degradation. It is very likely that this mutant is an ER-associated degradation (ERAD) substrate, but future analysis will be needed to demonstrate this hypothesis.

The removal of all the N-glycosylation sites [Δ (1234) TpoR] dramatically affected the STAT transcriptional activity in stably transfected Ba/F3 and Ba/F3-JAK2 cell lines, compared with WT TpoR (Figure 5C). This result was expected, because the cell surface level of this mutant was undetectable by flow cytometry in Ba/F3 cells. We also tested the Tpo-dependent cell growth of these stable cell lines in a more sensitive assay (thymidine incorporation), confirming the data obtained in the luciferase assay. We could not detect any Tpo-dependent growth even in the presence of high Tpo levels (10 ng Tpo/ml; Figure A1 in Appendix).

**THE DEFECTIVE N-GLYCOSYLATION VARIANTS ARE ABLE TO ACTIVATE Tpo-DEPENDENT STAT ACTIVITY IN OVEREXPRESSION CONDITIONS IN HEK293 AND γ2A CELLS**

In order to assess whether the TpoR variants that exhibit defective localization and stability [Δ (123), Δ (124), Δ (134), Δ (234)] or [Δ (1234)] are able to bind Tpo and activate Tpo-dependent signaling pathways, we tested all the TpoR variants in an overexpression cell system.

In transiently transfected γ2A or HEK293 cells, in the case of single N-glycosylation site [Δ (1), Δ (2), (Δ 3), Δ (4)] defective variants, there was no difference in the Tpo-dependent JAK2–STAT transcriptional activity when compared with the WT receptor (Figure 6). In the case of mutation of multiple N-glycosylation sites [Δ (124), Δ (134), Δ (234)], their Tpo-dependent JAK2–STAT transcriptional activity in transiently transfected was lower than that of WT TpoR (Figure 6). Mutant Δ (124) was similar to Δ (134) with respect to transcriptional activity in γ2A cells (data not shown). For the mutant Δ (1234) TpoR, removal of all N-glycosylation sites led to a significantly decreased, but still detectable, level of transcriptional activity in response to Tpo (Figure 6). These data suggest that overexpression conditions allow for some cell surface localization, which is sufficient to support a response to Tpo ligand in the very sensitive luciferase assays. Importantly, these data indicate that the receptor mutants are not misfolded and, once localized on the surface, they can bind Tpo.

**DISCUSSION**

Our major finding is that N-glycosylation sites on the extracellular TpoR domain exert essential regulatory roles on TpoR cell surface localization and function. Using a combination of mutagenesis and biochemistry approaches in stably transduced hematopoietic cells and in several other cell lines, we show that the four sites exert positive and negative roles on receptor trafficking, that at least one glycosylation site is essential for cell surface transport, and that exit from ER and stability are regulated by N-glycosylation. When three of the four N-glycosylation sites were mutated, we could demonstrate that the remaining N-glycosylation site was occupied.

Previous data presented a differential role of the N-glycans for different cytokine receptor traffic, ligand binding, and function (Ding et al., 1995; Buteau et al., 1998; Bolander, 1999; Niu et al., 2000; Kamikubo et al., 2008; Waetzig et al., 2010). There are no available studies on the importance of the carbohydrate groups on the cell surface expression and function of human TpoR. In order to perform our study, we constructed nine human TpoR variants, defective in one, three, or four out of four putative N-glycosylation sites located of the extracellular domain. Taking into account that TpoR is very important for the cells of hematopoietic lineage, most of the results of this study were obtained in Ba/F3 and Ba/F3-JAK2 cell lines.

The first question we asked was whether the physiologic human TpoR cell surface localization requires the occupancy of a specific N-glycosylation site from the four putative ones (Figure 1A). None of the four single sites was critical for cell surface localization, for Tpo binding or for Tpo-dependent activation of the JAK2–STAT signal pathway. Nevertheless, the TpoR mutant lacking the first N-glycosylation site (Asn117) is the most interesting and it constantly
FIGURE 5 | Removal of all four N-glycosylation sites dramatically affects the cell surface localization of human TpoR in Ba/F3 cells. (A) Ba/F3 or Ba/F3-JAK2 cells stably expressing the WT TpoR or the variant of the receptor defective in all four N-glycosylation sites \(\Delta (1234)\) were tested for cell surface levels using anti-HA antibody based flow cytometry. Similar results were obtained in five independent experiments, on two different cell lines obtained for the \(\Delta (1234)\) variant \(p < 0.0001\). Right panel of the (A) shows a Western blot of the cells used in the experiment, probing the receptor with anti-HA antibodies, the murine JAK2 WT with anti-JAK2 antibodies and using anti-\(\beta\)-actin as a control. (B) Ba/F3 or Ba/F3-JAK2 cells used in (A) were tested for the contribution of the proteasome- or lysosome-mediated degradation of the TpoR N-glycosylation defective mutant \(\Delta (1234)\) versus WT. The cells were incubated 3 h at 37˚C with 50 \(\mu\)g/ml cycloheximide and proteasome inhibitor MG132 (10 \(\mu\)M) or with 50 \(\mu\)g/ml cycloheximide and lysosome inhibitors (200 \(\mu\)M chloroquine or 10 \(\mu\)g/ml leupeptin). Control cells were kept untreated to have 100% protein expression. Cells were lysed and analyzed by Western blotting with antibodies against HA for the levels of TpoR and anti-\(\beta\)-actin as a loading control. (C) Pools from the same cell lines used in (A,B) were starved 3 h in RPMI medium +1 mg/ml BSA and electroporated with pGRR5-luc and pRLTK-luc reporters. The cells were also stimulated with Tpo or left unstimulated during 2 h at 37˚C. Upon treatment the cells were lysed and their luminescence recorded. Results are the mean ± variation of triplicate samples. One of three independent experiments is depicted. The \(t\)-test showed statistically significant differences between the Tpo-dependent transcriptional activities level of the \(\Delta (1234)\) TpoR and that of WT TpoR \(p < 0.0001\) for both Ba/F3 and Ba/F3-JAK2 cell lines.

presents striking differences compared with the WT TpoR. The cell surface level of \(\Delta (1)\) TpoR was decreased in Ba/F3 and Ba/F3-JAK2 cells (Figure 2A). Compared with the WT, the JAK–STAT transcriptional activity of this variant was also decreased in Ba/F3 and Ba/F3-JAK2 (Figure 3A), and we consequently observed a decrease in the Tpo-dependent cell proliferation of Ba/F3 stably transduced with this receptor, compared with Ba/F3 stably transduced with WT TpoR (Figure 3B). Alignments of TpoR from different species (Figure 1A) pointed out that this first N-glycosylation site of human TpoR is highly conserved between
species. We hypothesize that occupancy of this site is important but not crucial for TpoR cell surface expression. Rather, this carbohydrate group might exert a fine control on the cell surface level of TpoR. Consistent with our results, previous observations found that cell surface levels of TpoR are important for setting the correct blood platelet production. Insufficient expression of TpoR at the surface of late megakaryocytes and of platelets can lead to the unexpected phenotype of thrombocytosis, since Tpo ligand was not properly removed from circulation and it could still stimulate early megakaryocyte proliferation and differentiation (Lannutti et al., 2009; Tiedt et al., 2009). Moreover, TpoR defective N-glycosylation and impaired cell surface localization was detected in the megakaryocytes and platelets of Polycythemia Vera patients (Moliterno et al., 1998; Moliterno and Spivak, 1999).

In contrast to the results with Δ (1) TpoR, the Δ (3) TpoR mutant was localized at the cell surface to higher extents than the WT TpoR and showed enhanced maturation and activity. This glycosylation site is not conserved in rodents. Interestingly, its
presence limits receptor function in human TpoR. It is interesting to note that normal platelet levels are higher in mice than in humans. One possibility is that absence of this site allows higher levels of platelet production in mice. Further studies using a knock-in approach of a murine TpoR containing this N-glycosylation site could answer this question.

Several mutations in the first five exons coding for the first cytokine receptor homology domain have been described to affect the cell surface display of TpoR. Mutations located around residue Asn117, possibly affecting the N-glycosylation pattern, are at the basis of two types of hematological diseases, with opposite pathologies: congenital amegakaryocytic thrombocytopenia (CAMT; Ballmaier et al., 2001; Germeshausen et al., 2006; Tijssen et al., 2008; Fox et al., 2009) and familial thrombocytosis (FT; Moliterno et al., 2004; El-Harith El et al., 2009; Teofili and Larocca, 2011; Figure 1B). Extracellular domain mutations in CAMT abolish cell surface receptor localization, while FT extracellular domain mutations severely impair, though do not abolish, cell surface localization. For the latter, since TpoR on platelets performs clearance of Tpo from the circulation, low platelet TpoR levels are associated with increased levels of Tpo in the blood and in the bone marrow, a situation similar to the one reported when TpoR expression levels are decreased at the late megakaryocyte and platelet levels (Lannutti et al., 2009; Tiedt et al., 2009). Our results show that Asn117 occupancy is able to control the cell surface expression of human TpoR and consequently its availability for Tpo. We hypothesize that this first N-glycan is important for efficient cell surface localization and that mutations around Asn117 might either impact the efficiency of glycosylation of this site, or its interaction with ER chaperone proteins. More studies are required in order to determine which scenario is true for each mutation detected in humans.

A major finding of our study is that removal of three out of four N-glycosylation sites did not abolish, but significantly decreased, the cell surface localization, and activity of TpoR (Figure 4). In Ba/F3 cell lines, at least one N-glycosylation site of human TpoR needs to be occupied in order to have detectable cell surface levels and Tpo-dependent gene transcription. Neither JAK2 nor TYK2 were able to restore to normal the cell surface level of these variants (Figure 4). JAK2 levels differ between the cell types of the hematopoietic lineage (Dalal et al., 1998) and JAK2 was shown to promote WT TpoR traffic to the cell surface (Royer et al., 2005). Interestingly, in this study we show that the ability of JAK2 to promote cell surface localization of TpoR requires a normal N-glycosylation, with more than one site occupied. It is possible that JAK2 is not able to rescue the cell surface localization of TpoRs that do not acquire a specific N-glycosylation pattern. In any case, our data suggest that JAK2 binding to the cytosolic juxtamembrane region induces an inside-out conformational change that must be sensed by the extracellular domain Asn117. More studies are required in order to determine whether N-glycosylation places TpoR in a particular region of the ER, or facilitates interaction with export proteins. Such studies are currently underway.

The analysis of the cell surface localization of TpoR mutants with different N-glycosylation patterns is very important, as one of the essential functions of the N-glycans is to secure efficient protein production. The N-glycosylation sites are crucial for the interaction of the receptor with the lectin-like chaperones from ER, in the process of proper folding. The calnexin–calreticulin quality control cycle was previously described, in which these two chaperones are acting in the initial steps of glycoprotein folding. Prolonged interaction with calnexin, due to inefficient folding, results in the substrate being directed to the proteasome for degradation. We hypothesize that the TpoR mutants reported in different hematological diseases and that are defective in cell surface localization, could be retained in early folding compartments due to this prolonged interactions with the lectin-like chaperones. In this model, N-glycans are a limiting event for proper cell surface localization. The nature and expression levels of chaperones differ between cell types or stages of differentiation, thus possibly explaining why a particular mutant can be localized at the surface of some progenitors and not others. Future experiments will have to address the importance of N-glycans for human TpoR interactions with the ER chaperones. Besides its key role in Tpo-dependent signaling, JAK2 acts as a cytosolic chaperone that increases the receptor traffic to cell surface (Royer et al., 2005), but it is possible that the main determinants that govern the ER to plasma membrane traffic of human TpoR to be located in ER and to be influenced by receptor N-glycosylation pattern.

We also observed in our study that the removal of all four putative N-glycosylation sites dramatically affects cell surface localization of TpoR and Tpo-dependent cell proliferation in the Ba/F3 cell model (Figure 5 and data not shown). This observation supports the notion that N-glycosylation is crucial for the ER to cell surface traffic of TpoR. In this model, the presence of N-glycans promotes traffic of the receptor to the cell surface. JAK2 was not able to promote cell surface localization of the mutant defective in all N-glycosylation sites. Along with the results obtained with single and multiple defective glycosylation sites, this result again points to an inside-out conformational change induced by JAK2 binding to the cytosolic domain that is sensed by the extracellular domain via N-glycosylation sites.

Finally, our results showed that all the receptors defective in different combinations of N-glycosylation sites were able to support Tpo-dependent transcriptional activity in an overexpressed system (Figure 6), thus indicating that defective N-glycosylation does not simply induce misfolding and that N-glycosylation defective TpoR mutants are able to bind Tpo, once they reach the cell surface. This is quite important and it can serve as a starting point for targeting ER-associated folding machinery in diseases with TpoR mutants blocked in this compartment, as receptors could be functional once they escape from the ER and localize at the cell surface.

ACKNOWLEDGMENTS

We thank to Dr Alexandra Dusa for critical reading of this manuscript. We thank to André Tonon for support with flow cytometry. Roxana I. Albu currently holds a F.N.R.S Télévée PhD fellowship and was also funded by the Marie Curie ReceptEUR Network PhD fellowship. Stefan N. Constantinescu is a Senior Research Associate of the F.N.R.S., Belgium. We acknowledge support from the Fondation contre le Cancer, the Salus Sanguinis Foundation, the Action de Recherche Concertée MEXP31C1 of the Université catholique de Louvain, the PAI Program BCHM61B5, Belgium and the Atlantic Philanthropies, New York.
thrombocytosis in transgenic mice expressing reduced levels of Mpl in platelets and terminally differentiated megakaryocytes. Blood 113, 1768–1777.

Tijssen, M. R., Di Summa, F., Van Den Oudenrijn, S., Zwaginga, J. J., Van Der Schoot, C. E., Voermans, C., and De Haas, M. (2008). Functional analysis of single amino-acid mutations in the thrombopoietin-receptor Mpl underlying congenital amegakaryocytic thrombocytopenia. Br. J. Haematol. 141, 808–813.

Vigon, I., Florindo, C., Fichelson, S., Guenet, J. L., Mattei, M. G., Souyri, M., Cosman, D., and Gisselbrecht, S. (1993). Characterization of the murine Mpl proto-oncogene, a member of the hematopoietic cytokine receptor family: molecular cloning, chromosomal location and evidence for a function in cell growth. Oncogene 8, 2607–2615.

Vigon, I., Moron, J. P., Cacault, L., Mitjavila, M. T., Tambourin, P., Gisselbrecht, S., and Souyri, M. (1992). Molecular cloning and characterization of MPL, the human homolog of the v-mpl oncogene: identification of a member of the hematopoietic growth factor receptor superfamily. Proc. Natl. Acad. Sci. U.S.A. 89, 5640–5644.

Waetzig, G. H., Chalaris, A., Rosenstiel, P., Suthaus, J., Holland, C., Karl, N., Valles Uriarte, L., Till, A., Scheller, J., Grotzinger, J., Schreiber, S., Rose-John, S., and Seegert, D. (2010). N-linked glycosylation is essential for the stability but not the signaling function of the interleukin-6 signal transducer glycoprotein 130. J. Biol. Chem. 285, 1781–1789.

Yoshimura, A., Zimmers, T., Neumann, D., Longmore, G., Yoshimura, Y., and Lodish, H. F. (1992). Mutations in the Trp-Ser-X-Trp-Ser motif of the erythropoietin receptor abolish processing, ligand binding, and activation of the receptor. J. Biol. Chem. 267, 11619–11625.

Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.
APPENDIX

FIGURE A1 | Removal of all four N-glycosylation sites dramatically affects the Tpo-dependent cell proliferation of human TpoR in Ba/F3 and Ba/F3-JAK2 cells. Ba/F3 and Ba/F3-JAK2 cells stably expressing WT or Δ (1234) human TpoR were washed 3 times and seeded in 96-well plates at a density of 3,000 cells/well. The cells were stimulated with different Tpo concentrations, or with 5 ng IL3/ml or left unstimulated during 72 h. After 72 h, cell proliferation was determined by the DNA synthesis assay reflected by the measurements of the thymidine incorporation as described in Experimental procedures. Results are the mean ± variation of triplicate samples. One of two independent experiments is depicted.