RHEX, a novel regulator of human erythroid progenitor cell expansion and erythroblast development

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Ligation of erythropoietin (EPO) receptor (EPOR) JAK2 kinase complexes propagates signals within erythroid progenitor cells (EPCs) that are essential for red blood cell production. To reveal hypothesized novel EPOR/JAK2 targets, a phosphotyrosine (PY) phosphoproteomics approach was applied. Beyond known signal transduction factors, 32 new targets of EPO-modulated tyrosine phosphorylation were defined. Molecular adaptors comprised one major set including growth factor receptor–bound protein 2 (GRB2)–associated binding proteins 1–3 (GAB1–3), insulin receptor substrate 2 (IRS2), docking protein 1 (DOK1), Src homology 2 domain containing transforming protein 1 (SHC1), and sprouty homologue 1 (SPRY1) as validating targets, and SPRY2, SH2 domain containing 2A (SH2D2A), and signal transducing adaptor molecule 2 (STAM2) as novel candidate adaptors together with an ORF factor designated as regulator of human erythroid cell expansion (RHEX). RHEX is well conserved in Homo sapiens and primates but absent from mouse, rat, and lower vertebrate genomes. Among tissues and lineages, RHEX was elevated in EPCs, occurred as a plasma membrane protein, was rapidly PY-phosphorylated >20-fold upon EPO exposure, and coimmunoprecipitated with the EPOR. In UT7epo cells, knockdown of RHEX inhibited EPO-dependent growth. This was associated with extracellular signal–regulated kinase 1,2 (ERK1,2) modulation, and RHEX coupling to GRB2. In primary human EPCs, shRNA knockdown studies confirmed RHEX regulation of erythroid progenitor expansion and further revealed roles in promoting the formation of hemoglobinizing erythroblasts. RHEX therefore comprises a new EPO/EPOR target and regulator of human erythroid cell expansion that additionally acts to support late-stage erythroblast development.

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RESULTS AND DISCUSSION

EPO ligation of dimeric EPOR complexes activates JAK2 kinase and the phosphorylation of nine EPOR cytoplasmic PY sites (Fig. 1A). Certain signal transduction factors (STFs) that dock at EPOR PY sites are well defined (e.g., p85α at PY480, STAT5 at PY344) and some are also direct JAK2 targets (e.g., STAT5, phospholipase Cγ), whereas others couple to downstream signaling modules (Wojchowski et al., 2010; Watowich, 2011; Broxmeyer, 2013). To seek novel EPOR STFs, a PY-phosphoproteomic approach was applied using UT7epo cells as a human EPC model (Komatsu et al., 1993). This involved hematopoietic growth factor withdrawal with or without EPO challenge, tryptic digests of lysates, and PY peptide isolation plus liquid chromatography-tandem mass spectrometry (LC-MS/MS) identification (Fig. 1B). Overall, 54 unique EPO/EPOR-modulated PY proteins were identified with ≥2.5–82.6-fold modulation (Table S1). These included known STFs modulated at known PY sites, known regulator of cell cycle progression (Fang et al., 2007), MASL1 as a RAF-interacting inducer of EPO-dependent erythropoiesis (Kumkahea et al., 2013), and Spi2A as an EPO-induced inhibitor of leached lysosomal executioner cathepsins (Dev et al., 2013).

To provide new insight into EPO/EPOR effects, we presently have applied a global PY-phosphoproteomics approach. One strongly regulated novel EPOR target is designated as regulator of human erythroid cell expansion (RHEX). We first characterize RHEX’s genealogical representation plus stage- and lineage-restricted expression, plasma membrane localization, and EPOR co-association. In functional contexts, loss-of-function investigations then define RHEX effects on EPC growth, ERK1,2 regulation, and growth factor receptor-bound protein 2 (GRB2) association. In primary hEPCs, RHEX is further revealed to modulate the development of maturing erythroblasts. RHEX thus has evolved as an important upstream mediator of EPO/EPOR-dependent human red cell production.

Figure 1. Phosphoproteomic analysis of EPO/EPOR PY-regulated molecular adaptors including RHEX. (A) The hEPOR is depicted, including cytoplasmic PY sites (o), a box1,2 JAK2 binding domain, and JAK2 plus possible protein tyrosine kinase routes to PY targets. (B) PY-phosphoproteomic steps used to define EPO/EPOR-regulated targets in UT7epo EPCs. (C and D) For a major functional subset of EPO/EPOR signal transducers as molecular adaptors, LC-MS/MS data are summarized, including fold regulation at defined PY residues. Validating targets are GAB1-3, SHC1, IRS1, DOK1, and SPRY1 (C). Novel targets are SH2D2A, STAM2, SPRY2, and C1ORF186/RHEX (D). (E) For RHEX (C1ORF186), duplicate LC-MS/MS data are illustrated for EPO modulation at Y132 plus Y141 sites (single tryptic peptide; and are representative of two independent analyses; **, P ≤ 0.01, Student’s t-test).
transcripts (Fig. 2 B). Interestingly, **RHEX** proved to be well conserved in *Homo sapiens* and primates (99% nt conservation) but was not detected in rat, mouse, or lower vertebrate genomes. **RHEX** transcript expression among tissues and blood cells was also investigated and was relatively high level in primary human EPCs and kidney (Fig. 2, C and D). RNA-Seq also indicated elevated **RHEX** levels in CFUe as compared with CD34<sup>+</sup> progenitors (Fig. 2 E). At a protein level, **RHEX**’s predicted domains included an amino-terminal (NT) hydrophobic region and two carboxy-terminal candidate GRB2 binding sites (Neumann et al., 2009; Fig. 2, F and G). **RHEX**, however, is unique and exhibits homology only with limited residues of a recently reported erythrocytic spectrin (NP_003117.2). Basic assessments of **RHEX** levels among STFs modulated at novel sites, known proteins not previously associated with EPO/EPOR signaling, and novel targets encoded by predicted ORFs. As one major functional subset, 11 targets proved to be molecular adaptors. GAB1-3, SHC1, IRS2, DOK1, and SPRY1 comprise known (and validating) targets (Fig. 1 C), whereas SH2D1A, STAM2, and SPRY2, together with a C1ORF186 product, represent novel candidate EPO/EPOR STFs (Fig. 1 D). C1ORF186 (designated as **RHEX**) was up-modulated by EPO ≥20-fold in its phosphorylation at PY132 and PY141 sites (single PY peptide; Fig. 1 E) and is this report’s prime focus.

**RHEX** is encoded at a six-exon C1ORF186 locus (Fig. 2 A) that generates a singular predicted 1.6 kb nt coding transcript. Northern blotting detected major 1.6 kb, and minor <0.5 kb nt transcripts (Fig. 2 B). Interestingly, **RHEX** proved to be well conserved in *Homo sapiens* and primates (99% nt conservation) but was not detected in rat, mouse, or lower vertebrate genomes. **RHEX** transcript expression among tissues and blood cells was also investigated and was relatively high level in primary human EPCs and kidney (Fig. 2, C and D). RNA-Seq also indicated elevated **RHEX** levels in CFUe as compared with CD34<sup>+</sup> progenitors (Fig. 2 E). At a protein level, **RHEX**’s predicted domains included an amino-terminal (NT) hydrophobic region and two carboxy-terminal candidate GRB2 binding sites (Neumann et al., 2009; Fig. 2, F and G). **RHEX**, however, is unique and exhibits homology only with limited residues of a recently reported erythrocytic spectrin (NP_003117.2). Basic assessments of **RHEX** levels among

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**Figure 2.** **RHEX** locus, transcripts, and primary protein structure. (A) C1ORF186/RHEX gene structure. (B) Analyses of putative **RHEX** transcripts (top) and Northern blotting (bottom) defined major 1.6 kb nt (and minor <0.5 kb nt) transcripts in UT7epo cells, and in primary human EPCs. (C and D) RT-PCR assays of **RHEX** transcript expression levels in primary human tissues (C) and among human peripheral blood monocytes, T cells, neutrophils, and platelets (as compared with primary CD71<sup>+</sup> EPCs, D). (For elevated **RHEX** levels in EPCs and kidney, P ≤ 0.01; **, Student’s t test, representative of two independent analyses). (E) RNA-Seq analyses of **RHEX** (and ALAS2) transcript levels in primary human CD34<sup>+</sup> progenitors and CFUe (*, P ≤ 0.05, Student’s t test, single experiment). (F and G) Primary sequence of **RHEX** (F) and candidate functional domains (G). (H) Western blot analysis of **RHEX** protein expression among human hematopoietic cell lines (representative of two independent studies).
To assess function, RHEX knockdown experiments were performed. Three assessed shRNAs each efficiently inhibited RHEX expression (Fig. 4 A and not depicted). As assayed via clonal colony formation, RHEX knockdown limited UT7epo cell growth ≥3-fold (Fig. 4 B, top). Analyses in liquid culture confirmed effects on growth (with no significant effects on survival), and a parallel attenuation of ERK1,2 activity was also observed (Fig. 4, B and C). Possible effects of RHEX on murine EPCs were also studied via lentiviral expression in human hematopoietic cell lines (and 293 cells) using polyclonal antisem to RHEX further revealed expression only in erythroid UT7epo cells (Fig. 2 H).

To analyze RHEX’s subcellular localization and actions, a (PY)RHEX reactive monoclonal antibody was next generated and was used in UT7epo cells to first validate rapid EPO induction of PY-RHEX (Fig. 3 A and not depicted). Human SCF, IL3, GMCSF, TPO, Flt3L, or serum, in contrast, did not detectably stimulate RHEX’s PY phosphorylation (unpublished data). RHEX’s hydrophobic NT region prompted subcellular localization analyses. As indicated by CD71 and wheat germ agglutinin (WGA) markers, (PY)RHEX resided at the plasma membrane (Fig. 3 B). Experiments using JAK2 and SRC kinase inhibitors (TG101348 and Dasatinib, respectively) pointed to RHEX as a JAK2 target (Fig. 3 C). Beyond this, coimmunoprecipitation experiments using an RHEX-flag construct indicated co-association with EPOR and JAK2 complexes (Fig. 3 D). To analyze RHEX’s subcellular localization and actions, a (PY)RHEX reactive monoclonal antibody was next generated and was used in Western blot analyses of EPO-challenged UT7epo cells to validate EPO-induced RHEX PY phosphorylation (representative of three independent analyses). Black lines indicate that intervening lanes have been spliced out. (B) After cytokine withdrawal, UT7epo cells were challenged with EPO (±3 U/ml). At 15 min, cells were cytospun, fixed, and assayed by confocal immunofluorescence microscopy for PY-RHEX localization (right). In top paired panels (− vs. + EPO), CD71 was co-stained. In lower paired panels (+ vs. + EPO), cells were co-stained with AF555-WGA (representative of two independent experiments). RHEX’s predicted N-terminal α-helical transmembrane domain is also diagrammed. (C) UT7epo cells were exposed to TG101348 (0, 5, and 15 nM, indexed as −, +, and ++). After HGF withdrawal, cells were exposed to 2 U/ml EPO. At 10 min, lysates were prepared and analyzed for levels of (PY)RHEX. Effects of SRC inhibition by Dasatinib on EPO-induced PY-RHEX formation also were assessed (+ and + as 50 and 150 nM; bottom). Results are representative of two independent experiments. (D) UT7epo cells were stably transduced with lentivirus encoding a Flag epitope-tagged RHEX (carboxy-terminal tag). In anti-Flag immunoprecipitations, the EPOR (top pair of panels) and JAK2 (center pair of panels) were observed to co-immunoprecipitate with RHEX-(Flag) (representative of two independent experiments). In the bottom panel, the immunoprecipitation of RHEX-(Flag) on its own is shown.
reproducible effects on proliferation or on Gata1-ER–induced late-stage differentiation were observed (unpublished data).

RHEX’s actions were next studied in human CD34pos progenitor–derived EPCs. Endogenous RHEX transcript levels became elevated at days 5–7 of culture (pro- to basophilic erythroblasts; Fig. 5, A and B). In developmentally staged EPCs as isolated via FACS, RHEX was elevated in proerythroblasts (Fig. 5 C and not depicted). In shRNA lentivirus transduction experiments, when RHEX was knocked down, EPC expansion was attenuated (without significant effects on viability; Fig. 5, D and E). Beyond this, glycophorin-A (GPA) levels were skewed and increased when RHEX expression was inhibited (Fig. 5 F). Cytospin morphologies also revealed apparent effects of RHEX knockdown on erythroid development, including a delayed formation of normoblasts (Fig. 5 G). By inspection of cell pellets, Western blotting, and RT-PCR, hemoglobinization was also attenuated, which is further consistent with a role for RHEX in supporting erythroblast development at a normoblast stage (Fig. 5, H and I). Here, EPO exposure moderately increased levels of GRB2 plus RHEX immunoprecipitation.

Recent discoveries of novel EPOR pathways (Broxmeyer, 2013) and new EPOR agonists (Drüeke, 2013) have heightened interest in transducers of EPO’s effects. Our PY-phosphoproteomic analyses first reveal several novel candidate EPOR targets as molecular adaptors, S/T kinases, tyrosine phosphatases, ubiquitin factors, and cell cycle regulators (Supplemental Table S1). A focus on RHEX was prompted by its sharp EPO modulation, ORF novelty, conserved representation in H. sapiens and pri-mates, and stage-specific up-modulation in developing erythroid cells. RHEX proved to be plasma membrane–associated, to coimmunoprecipitate with the EPOR, and to comprise a likely JAK2 PY target. EPO-regulated sites Y132 and/or Y141 were also implicated to bind GRB2. Via SOS plus RAS, GRB2 may mediate RHEX effects on ERK1/2 activity, but GRB2 itself is a versatile adaptor protein that can also couple to SHIP, SHP2, and up to 90 interacting proteins (Bisson et al., 2011). GRB2 can also preassemble with RTKs and repress basal signaling (Lin et al., 2012). Merit therefore exists for future investigations of implicated RHEX PY132, PY141, and GRB2 engaged pathways.

In primary EPCs, loss of function studies confirmed RHEX effects on proliferation but also indicated additional roles during normoblast development in that RHEX knockdown led to persistent erythroblastic features, high-level GPA expression, and decreased HBB transcript plus globin expression. Although this might relate indirectly to observed effects of RHEX on growth, effectors that reinforce EPC growth (e.g., via RAS; Zhang and Lodish, 2007; Blanc et al., 2012) often attenuate erythroid differentiation. Apparent effects of RHEX on normoblast development, therefore, may involve alternative mechanisms (and may point to novel EPO/EPOR effects on erythroid differentiation). RHEX-regulated routes that...
Figure 5. In primary human EPCs, shRNA knockdown reveals positive roles for RHEX during erythroid progenitor expansion and erythroblast maturation. (A) Upon a shift of CD34<sup>+</sup> progenitors to erythroid culture conditions, courses of CD71 and GPA marker expression were defined via flow cytometry. Data are normalized means ± SE, n = 3 (representative of 5 independent experiments). (B) RT-PCR analyses of RHEX expression during EPC development ex vivo (ALAS2, Aminolevulinic Acid Synthetase-2; HBB, Hemoglobin-β; means ± SE, n = 2). (C) HPCs expanded short-term (48 h) proerythroblasts (as CD71<sup>high</sup>CD36<sup>high</sup>GPA<sup>+</sup> EPCs) and erythroblasts (as CD71<sup>high</sup>CD36<sup>low</sup>GPA<sup>+</sup> EBs) were isolated (FACS) and analyzed via Western blotting for RHEX expression levels. (D) Efficient transduction of hCD34 progenitors and persistent GFP marker expression among developing CD71<sup>high</sup>EPCs as demonstrated by co-positive day 7 CD71<sup>high</sup>GFP<sup>+</sup> EPCs (flow cytometry analysis). (E) Decreased EPC growth (at days 6 and 9) due to RHEX knockdown (mean cell numbers ± SE, n = 2; **, P < 0.01, Student’s t test, representative of two independent experiments). (F) Representative expression profiles of CD71 and GPA marker expression are shown for shRNA-NT and shRNA-RHEX transduced EPCs (day 9 after transduction), together with summary data (means ± SE, n = 2; **, P < 0.01, Student’s t test, representative of two independent experiments). (G) Representative EPC morphologies are shown after shRNA-NT and shRNA-RHEX transduction (day 9). (H) Western blot analysis and visualization of hemoglobin levels in developing EPCs after transduction with shRNA-NT or shRNA-RHEX. (I) RT-PCR analyses of HBB, RHEX, and ALAS2 (means ± SE, n = 2; **, P < 0.01, Student’s t test). (J) In UT7epo cells, coimmunoprecipitation assays demonstrated the association of RHEX-Flag with endogenous GRB2. (K) Co-immunoprecipitation of endogenous GRB2 with RHEX was also observed.
support erythroblast development, therefore, should also be of significant interest to define during myeloproliferative disease (Barbui et al., 2013) and the ineffective erythropoiesis of thalassemia (Rivella, 2012).

MATERIALS AND METHODS

Cell lines and primary hematopoietic cells. UT7/epo EPC (Komatsu et al., 1993), 293, JURKAT, RAJ1, K562, HL60, and HEL cell lines were maintained as previously described (Singh et al., 2012b). In cytokine withdrawal experiments, UT7/epo cells were washed three times and cultured for 20 h in 0.2% BSA, 10 µg/ml holo-transferrin, 0.1 mM 2-mercaptoethanol, and IMDM (Singh et al., 2012b). Subsequent EPO (and cytokine) challenges were at the concentrations and time intervals indicated. Primary human hematopoietic cells used in RT-PCR analyses included monocytes, neutrophils, T cells, and platelets (AllCells). Murine G1E-ER4 cells (M. Weiss, Children's Hospital of Philadelphia, Philadelphia, PA) were maintained in EPO plus mSCF as per Welch et al. (2004).

Primary EPCs were generated ex vivo from GCSF mobilized human CD34+ cells. CD34+ progenitors were cultured for 48 h in X-vivo10 medium (Lonza) supplemented with 50 ng/ml each of rhSCF, rhIL3, rhFLT3, and rhTPO (PeproTech). For erythroid cell development, expanded progenitors were plated at (10^5 cells/ml) in StemSpan medium (STEMCELL Technologies) supplemented with 100 ng/ml rhSCF 2 U/ml EPO, 100 µg/ml holo-transferrin, 1.5 µM β-estradiol, 0.5 µM dexamethasone, and 0.1 mM β-mercaptoethanol. On days 2, 6, and 10, cultures received 0.6 volume of medium. On days 4, 8, and 12, cultures were replated in fresh medium (10^5 cells/ml). Erythroid development was assessed via flow cytometry, cytospin, and RT-PCR analyses.

Phospho-proteomics and in silico analyses. In UT7/epo cell PY-phospho-proteomic studies, cytokines were withdrawn for 20 h. Cells were then exposed to EPO (± 4 U/ml for 15 min), and 9 M urea lysates were prepared and used to generate tryptic digests (Sathyanarayana et al., 2012). For duplicate samples, PY-phospho-proteomic LC-MS/MS analyses were then applied using a PhosphoScan approach (Stokes et al., 2012). Gene and transcript analyses used NCBI programs Gene, BLAST, Homologene, Gene Expression Omnibus, and Ensembl Genome browser. RHEX's candidate structural features were assessed using BLAST Conserved Domain plus Cobalt (NCBI), ProSite (ExPASy), TMHMM Server (v 2.0), and TMPnmd (cHEMM Net).

RT-PCR and Northern blotting. For RT-PCR analyses, EPCs (including FACS or MACS isolated subpopulations) were lysed in TRIZOL reagent (Invitrogen) and RNA was prepared (RNeasy; QIAGEN). Reverse transcription (Superscriptase III; Invitrogen) and real-time quantitative polymerase chain reactions (Q SYBR Green, i-Cycler; Bio-Rad Laboratories) were performed as previously described (Dev et al., 2013). The following primer pairs were from Invitrogen: hRHEX/C1ORF186, hALAS2, hHBß, and hBETA-ACTIN. Northern blotting was performed as detailed by Pircher et al. (2001) using a randomly primed 32P-P-RHEX cDNA probe.

Antibodies and Western blot analyses. Rabbit antibodies to RHEX were generated by immunizations with KLH-coupled peptides and immunogen boosts. Responses were evaluated by ELISA (phosphorylated and non-phosphorylated peptides) and by Western blotting (UT7/epo and HL60 cell lysates). Additional antibodies were to P(ERK1,2, GRB2, β-TUBULIN (Cell Signaling Technology), Hemoglobin (Santa Cruz Biotechnology, Inc.), and the EPOP. (Singh et al., 2012b). Western blotting was as described previously (Singh et al., 2012b). In chemiluminescence, HRP-conjugated antibodies (Jackson ImmunoResearch Laboratories, Inc.) and Super-Signal West-Dura reagent were used. In immunoprecipitations, 0.4% Igepal lyses were prepared as per Singh et al. (2012b). Antibodies used (4 µg per immunoprecipitation) were to a RHEX-Flag epitope (Sigma-Aldrich) and the EPOP (Singh et al., 2012b). Immune complexes were retrieved using protein A/G particles (Thermo Fisher Scientific). In analyses of RHEX PY phosphorylation, cell lysates were exposed to Lambda phosphatase before SDS denaturation and Western blotting. In JAK2 and SRC inhibitor studies, UT7/epo cells were cultured for 20 h with TG101348 or Dasatinib at the indicated concentrations (0.5% DMSO, solvent).

Immunofluorescence microscopy. UT7/epo cells were stained with FITC-anti-CD71, collected onto poly-L-lysine slides, exposed to 4% formaldehyde in PBS (20 min, 23°C), and then to 90% methanol (10 min, −20°C). Anti-(PY)RHEX antibody was used at 4 µg/ml (4°C, 60 min) and was monitored using an Alexa Fluor 647 anti-rabbit second antibody. Alternatively, surface staining was with 5 µg/ml AF555-WGA for 10 min at 37°C (Life Technologies). Nuclear staining was with 1 µM ToPro-3 iodide for 20 min (Invitrogen). Preparations were washed three times in PBS and coverslipped (1.5 mm, Corning) in Vectashield (Vector Laboratories). Images were acquired and analyzed using a confocal microscope system (SP1; Leica; Dev et al., 2013).

Lentiviruses, and transductions. shRNAs (as designed with SBI) were cloned to a pGreenPuro vector modified to include an EF1α promoter for GFP plus puromycin resistance cDNA expression. shRNA sequences were as follows: shRNA-RHEX-UTR-#1, 5'-GATCGGGAGAAGACTTCCAGGGTAACCTTCCTGTCAGCTGAGCTTAGTTTTTTTTGAATT-3'; shRNA-RHEX-UTR-#2, 5'-GATCCGAGGGAAAACTGGTGAAATTGATCACCTGACTTCTCGTCACTTTCTATCTGTATTGTTTGAATT-3'; shRNA-RHEX-UTR-#3, 5'-GATCCGGAGAAGACTTCCAGGGTAACCTTCCTGTCAGCTGAGCTTAGTTTTTTTTGAATT-3'; shRNA-RHEX-UTR-#4, 5'-GATCCGACTGACTACATATTGGAAAACTGGTGAAATTGATCACCTGACTTCTCGTCACTTTCTATCTGTATTGTTTGAATT-3'; shRNA-RHEX-UTR-#5, 5'-GATCCCCGACGTTGACACTGGTGAAATTGATCACCTGACTTCTCGTCACTTTCTATCTGTATTGTTTGAATT-3'; shRNA-RHEX-UTR-#6, 5'-GATCCCCTAGGCTGAAATTGATCACCTGACTTCTCGTCACTTTCTATCTGTATTGTTTGAATT-3'. Packaged lentiviruses (prepared using SBI-NI 293 cells) were recovered at 50 h after transduction, 0.4 µm filtered, concentrated (PEG-IT reagent, SBI), stored at −80°C. UT7/epo and G1E-ER4 cell transductions involved plating cells at 3 × 10^3 cells/ml and at 20 h re-plating to 12-well plates at 2 × 10^6 cells/ml (0.4 ml/well). At 4 h of culture, 4 µg/ml polybrene was added, followed by low 40 µl shRNA-encoding lentivirus in IMDM (at defined MOIs). At 20 h after transduction, 0.5 ml of culture medium was added. At 40 h after transduction, 0.5 ml of cells were selected in 2.5 µg/ml puromycin or plated in methylcellulose for colony forming assays (see below).

In the lentiviral knockdown of RHEX in primary human EPCs, pre-expanded cells were transduced with pGreenPuro lentiviruses encoding RHEX-targeting or nontargeting (NT) shRNAs. Specifically, nontreated 12-well tissue culture plates (BD), were coated with 20 µg/ml Retronectin (Takara Bio Inc.), washed, and used to preabsorb lentiviruses at 37°C for 1.5 h. This was followed by centrifugation at 800 g for 25 min at 4°C. Pre-expanded hCD34+ cells were plated at 0.5 × 10^6 cells/ml in Xvivo 10 medium (as described above) and were transduced using 3 µg/ml polybrene (30 min incubation at 37°C) and spinoculation for 30 min at 400 g at 25°C. After culture for 20 h, transduced cells were then plated in StemSpan medium (STEMCELL Technologies) supplemented with 100 ng/ml rhSCF, 100 µg/ml holo-transferrin, 2 U/ml EPO, 1.5 µM β-estradiol, 0.5 µM dexamethasone, and 0.1 mM β-mercaptoethanol. Puromycin was included at 2.5 µg/ml. At the indicated time points, erythroid cell formation was assayed via flow cytometry (c-KIT, TRFR1, GPA), cytoxin morphologies, and RT-PCR. Viable cell counts (ViCell system; Beckman Coulter) and YOPRO3 viability analyses (Molecular Probes) were also performed. Lentiviral expression vectors also were prepared including those expressing RHEX-flag (EF1α) plus puromycin resistance cDNAs (PGK), and RHEX plus GFP cDNAs. Each template, together with empty vector negative control templates, was packaged as above and was titrated using NIH3T3 cells.

Flow cytometry, colony-forming assays, and cytoxin histomorphologies. Flow cytometry analyses of erythroid development used APC–anti-CD117, FITC–anti-CD71, and PE–anti-CD235a antibodies (BD), a FACS caliber cytometer (BD), and CellQuest Pro software. In colony-forming unit assays, 0.5 × 10^4 transduced UT7/epo cells were plated in MethoCult H4434 medium (STEMCELL Technologies) supplemented with 3 U/ml EPO and 4.0 µg/ml puromycin. Colonies were analyzed using a Stemvision system (STEMCELL Technologies), differential interference fluorescence microscopy (DMI6000 B; Brief Definitive Report 1721
Online supplemental material. Table S1 shows EPO/EPOR PV-regulated targets as interrogated in UT7/Epo cells via PhosphoScan LC-MS/MS PV- proteomics. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20130624/DC1.

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