Erythropoietin signaling regulates heme biosynthesis

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Erythropoietin signaling regulates heme biosynthesis

Jacky Chung1, Johannes G Wittig1†, Alireza Ghamari2, Manami Maeda1‡, Tamara A Dailey3,4, Hector Bergonia5, Martin D Kafina1, Emma E Coughlin6, Catherine E Minogue7, Alexander S Hebert8, Liangtao Li9, Jerry Kaplan8, Harvey F Lodish5, Daniel E Bauer2,10, Stuart H Orkin2,10, Alan B Cantor2,10, Takahiro Maeda1‡, John D Phillips5, Joshua J Coon6,7,11, David J Pagliarini12, Harry A Dailey3,4, Barry H Paw1,2,10

1Division of Hematology, Brigham and Women’s Hospital, Harvard Medical School, Boston, United States; 2Division of Hematology-Oncology, Boston Children’s Hospital, Harvard Medical School, Boston, United States; 3Department of Microbiology, University of Georgia, Athens, United States; 4Department of Biochemistry and Molecular Biology, University of Georgia, Athens, United States; 5Division of Hematology and Hematologic Malignancies, University of Utah School of Medicine, Salt Lake City, United States; 6Genome Center of Wisconsin, Madison, United States; 7Department of Chemistry, University of Wisconsin-Madison, Madison, United States; 8Department of Pathology, University of Utah School of Medicine, Salt Lake City, United States; 9Whitehead Institute for Biomedical Research, Massachusetts Institute of Technology, Cambridge, United States; 10Department of Pediatric Oncology, Dana-Farber Cancer Institute, Harvard Medical School, Boston, United States; 11Department of Biomolecular Chemistry, University of Wisconsin-Madison, Madison, United States; 12Department of Biochemistry, University of Wisconsin-Madison, Madison, United States

Abstract Heme is required for survival of all cells, and in most eukaryotes, is produced through a series of eight enzymatic reactions. Although heme production is critical for many cellular processes, how it is coupled to cellular differentiation is unknown. Here, using zebrafish, murine, and human models, we show that erythropoietin (EPO) signaling, together with the GATA1 transcriptional target, AKAP10, regulates heme biosynthesis during erythropoiesis at the outer mitochondrial membrane. This integrated pathway culminates with the direct phosphorylation of the crucial heme biosynthetic enzyme, ferrochelatase (FECH) by protein kinase A (PKA). Biochemical, pharmacological, and genetic inhibition of this signaling pathway result in a block in hemoglobin production and concomitant intracellular accumulation of protoporphyrin intermediates. Broadly, our results implicate aberrant PKA signaling in the pathogenesis of hematologic diseases. We propose a unifying model in which the erythroid transcriptional program works in concert with post-translational mechanisms to regulate heme metabolism during normal development.

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Introduction

Heme biosynthesis is a fundamental biological process that is highly conserved and involves eight enzymatic reactions that occur both in the cytosol and mitochondria (Severance and Hamza, 2009). In vertebrates, the most recognized role of heme is to serve as the oxygen-binding moiety in...
Heme is an iron-containing compound that is important for all living things, from bacteria to humans. Our red blood cells use heme to carry oxygen and deliver it throughout the body. The amount of heme that is produced must be tightly regulated. Too little or too much heme in a person’s red blood cells can lead to blood-related diseases such as anemia and porphyria. Yet, while scientists knew the enzymes needed to make heme, they did not know how these enzymes were controlled.

Now, Chung et al. show that an important signaling molecule called erythropoietin controls how much heme is produced when red blood cells are made. The experiments used a combination of red blood cells from humans and mice as well as zebrafish, which are useful model organisms because their blood develops in a similar way to humans. When Chung et al. inhibited components of erythropoietin signaling, heme production was blocked too and the red blood cells could not work properly.

These new findings pave the way to look at human patients with blood-related disorders to determine if they have defects in the erythropoietin signaling cascade. In the future, this avenue of research might lead to better treatments for a variety of blood diseases in humans.

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Figure 1. PKA activity regulates heme biosynthesis. (A) A schema detailing the preparation of samples enriched for mitochondria from undifferentiated and differentiated MEL cells is shown. Following preparation of samples from both cohorts, the samples were trypsin digested and labeled with different tandem mass tags (TMTs), followed by mass spectrometry analysis. (B) Enrichment of undifferentiated (undiff) and differentiated (diff) mitochondrial samples was confirmed by western analysis for mitochondrial (PDHA1 and HSPD1) and cytosolic (GAPDH and TUBA1A) markers prior to mass spectrometry analysis. (C) The upregulation of PKA regulatory (PKAr) and catalytic (PKAc) subunits as well as previously identified heme metabolism proteins in mitochondria-enriched fractions of differentiated MEL cells but not several housekeeping proteins is presented in logarithmic scale. Please see Figure 1—source data 1 for precise changes. (D and E) Immunoblot analyses of the expression of PKA subunits in mitochondrial fractions (D) and whole cell lysates (E). All immunoblots were performed twice. Undiff—undifferentiated; Diff—differentiated; IB—immunoblot.

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The following source data is available for figure 1:

Figure 1 continued on next page
As expected, erythroid differentiation was associated with the elevated mitochondrial expression of a number of mitochondrial proteins known to have a role in erythropoiesis such as FECH, ATPIF1, and ABCB10 while the expression of housekeeping proteins such as components of the mitochondrial transport and protein translation machinery were relatively unchanged (Figure 1C). Strikingly, we also found increased expression of PKA regulatory (PKAr; PRKAR1A, PRKAR2A, PRKAR2B) and catalytic (PKAc; PRKACA and PRKACB) subunits in mitochondria-enriched fractions (Figure 1C). We independently confirmed these results using immunoblotting with isoform-specific antibodies where we also found increased total expression of these PKA subunits in maturing erythroid cells (Figure 1D and E). We failed to detect PRKACG because it is only present in humans and not found in our murine model (Kirschner et al., 2009). In addition, we also could not detect PRKAR1B since its expression is restricted to neurons (Kirschner et al., 2009). Together, our results suggest that select PKA subunits become highly expressed in mitochondria of developing erythrocytes and that MEL cells are a good model that accurately recapitulates the expected PKA expression pattern.

**Mitochondrial PKA is localized to the outer mitochondrial membrane via AKAP10**

We wondered whether increased mitochondrial PKA was specific for a particular suborganellar compartment, and next, performed a series of experiments to determine their submitochondrial expression. First, intact mitochondria isolated from maturing erythroid cells were treated with proteinase K that would digest all proteins exposed on the outer mitochondrial membrane. Immunoblot analysis of untreated and treated mitochondria revealed that the majority of PKA subunits were sensitive to proteinase K activity similar to TOM20 while VDAC1, a mitochondrial outer membrane (OM) marker known to be resistant to proteinase K digestion, remained largely unaffected (Figure 2A) (Rapaport, 2003; Shirihai et al., 2000). Biochemical fractionation of the mitochondria OM, intermembrane space (IMS), and mitoplast (MP) followed by immunoblotting confirmed the predominant presence of PKA subunits in the OM fraction of maturing erythroid cells (Figure 2B).

A great deal of work has demonstrated that PKA is localized to subcellular compartments through interactions with a family of anchoring proteins called AKAPs (a kinase anchoring proteins) (Wong and Scott, 2004). The majority of AKAPs recruit PKA-RII subunits but not RI (Sarma et al., 2010). However, a subclass of AKAPs can bind to both RI and RII with high affinity to regulate their subcellular distribution and have been referred to as ‘dual-specificity AKAPs’ (Huang et al., 1997a, 1997b; Li et al., 2001; Sarma et al., 2010; Wang et al., 2001). In particular, PAP7, AKAP1, and AKAP10 are three such AKAPs capable of localizing to mitochondria (Huang et al., 1997a, 1997b; Li et al., 2001; Wang et al., 2001; Wong and Scott, 2004). Interestingly, although we failed to detect PAP7 and AKAP1 in our proteomics analysis, we found a pronounced increase in the mitochondrial expression of AKAP10 (Figure 2C and D). Similar to our earlier data with PKA subunits, mitochondrial AKAP10 in maturing erythrocytes is sensitive to proteinase K digestion and primarily found in the OM fraction of maturing erythroid cells (Figure 2E and F).

High-throughput expression analysis has previously shown that AKAP10 expression increases in maturing erythroid cells and is a downstream target of the GATA1 erythroid lineage master transcription factor (Fujiwara et al., 2009; Zhang et al., 2003). To date, it has no known role in erythropoiesis or heme metabolism. However, our results thus far led us to wonder if it was responsible for regulating PKA localization in maturing RBCs, and we tested this by using CRISPR/Cas9-mediated genome editing to introduce null mutations into the endogenous AKAP10 loci. Genotyping and sequencing showed that for one AKAP10 allele [AKAP10\(\text{Cas9(ex1-2)}\)], parts of exons 1 (Ex1) and 3 (Ex3) along with all of exon 2 including the ATG start codon were deleted by our targeting strategy (Figure 2G and H). The second allele [AKAP10\(\text{Cas9(184delTG)}\)] had a 2 base-pair deletion that resulted in a premature stop codon (Stop) (Figure 2G and H). Neither allele gave rise to full-length AKAP10 protein (KO) as shown by immunoblotting (Figure 2H and I).
Figure 2. Mitochondrial PKA is localized to the outer membrane during erythropoiesis by AKAP10. (A) Intact mitochondria isolated from maturing MEL cells (day 3) were untreated or treated with proteinase K and subsequently analyzed by immunoblotting with antibodies specific for the indicated proteins. (B) Mitochondria from day 3 maturing MEL cells were fractionated into the indicated compartments and 5 μg of protein were analyzed with immunoblotting. (C and D) A heat map demonstrating the increased mitochondrial expression of AKAP10 similar to PRKAR2B in maturing erythroid Figure 2 continued on next page
Figure 2 continued

Total expression of PKA subunits in maturing KO cells was similar to wild-type (WT) cells (Figure 2J). However, KO cells had reduced the levels of mitochondrial PKA subunits both in intact preparations as well as in OM-specific fractions (Figure 2J and K). These results strongly suggest that AKAP10 recruits PKA to the outer mitochondrial membrane during red cell development and connects the GATA1 transcriptional program to the PKA signaling pathway.

Mitochondrial outer membrane PKA signaling regulates hemoglobinization and erythropoiesis

Mitochondria are the site of heme production required for hemoglobin synthesis and its physiology is a crucial part of RBC maturation (Nilsson et al., 2009; Shah et al., 2012). Given the increase in mitochondrial expression of PKA subunits in maturing erythroid cells, we wondered whether PKA activity had an influence on heme production. We addressed this by first using pharmacologic agents to toggle PKA function. Compounds that activate PKA such as 8-Br-cAMP and forskolin both caused an increase in the proportion of hemoglobinized cells as shown by o-dianisidine staining, which can be blocked by PKA antagonists H-89 or PKI (14-22) (Figure 3A). An increase in the proportion of hemoglobinized cells was also observed when we treated MEL cells with dimethyl-prostaglandin E2 (dmPGE2) (Figure 3B), which is a more stable analog of prostaglandin E2 (PGE2) that has a physiologic role during multiple aspects of hematopoiesis (Goessling et al., 2009; North et al., 2007). The effects of dmPGE2 can be similarly inhibited by PKI (14-22) (Figure 3C), underscoring the specificity of dmPGE2 signaling via PKA. In contrast to PKA inhibition, the PKC inhibitor, bis-indoyl-maleimide II, could not block the effects of PKA activation suggesting that the observed changes in heme synthesis are specific for PKA (Figure 3—figure supplement 1A).

Although our results from pharmacologic experiments suggest that widespread modulation of PKA has an impact on heme production on maturing erythroid cells, they do not explicitly examine the role of mitochondrial OM PKA. The precise contributions of distinct mitochondrial pools of PKA have been a topic of controversy (DiPilato et al., 2004; Lefkimmiatis et al., 2013). Emerging evidence suggest that PKA agonists such as forskolin and cAMP cannot diffuse into the mitochondrial matrix (Acin-Perez et al., 2009; Lefkimmiatis et al., 2013). Thus far, our pharmacologic data involved the use of a high-dose of forskolin (Figure 3A–C), and when the dose was titrated down to one that was previously shown to not activate matrix PKA, we also failed to detect an effect on hemoglobinization (Figure 3—figure supplement 1B) (Acin-Perez et al., 2009).

It is difficult to rely solely on pharmacologic data to unambiguously dissect the contributions of intracellular PKA pools since dose responses are known to vary form one cell type to another (Humphries et al., 2007; Lefkimmiatis et al., 2013). However, the reduction in the levels of PKA subunits in the mitochondrial OM of AKAP10-KO maturing erythroid cells allowed us to genetically and biochemically examine the functional role of this PKA signaling compartment. Compared to wild-type controls, AKAP10-KO maturing erythroid cells exhibited a deficit in hemoglobinization.
Figure 3. Mitochondrial outer membrane PKA signaling is required for erythropoiesis. (A–C) o-dianisidine staining for hemoglobinized MEL cells treated with several pharmacologic modulators of PKA activity. (A) PKA activation with 8-Br-cAMP or high-dose forskolin (50 μM) triggers an increase in heme production that is blocked by H89 or PKI (14-22) treatment at day 3 of DMSO induction. (B and C) A similar increase in hemoglobinization was observed with dMPGE₂ that was also inhibited by PKI (14-22). (D) Wild-type (WT) or AKAP10-knockout (KO) erythroid cells at day 4 of differentiation were stained with o-dianisidine. (E and F) AKAP10 expression was knocked-down using two different shRNAs (E) that lead to reduced hemoglobinization (F). (G–I) akap10-specific morpholinos (MOs) were used to inhibit akap10 expression in zebrafish embryos (G). These morphants were anemic with reduced hemoglobinization (H) and red blood cell counts (I). *p-value<0.05, Mean ± SEM, n = 3. All immunoblots were performed twice. 8-Br-cAMP-8-bromoadenosine 3',5'-cyclic monophosphate; dMPGE₂-dimethyl-prostaglandin E₂; WT-wild-type; KO-knockout; shRNA-short hairpin RNA; MO-morpholino; IB-immunoblot. DOI: 10.7554/eLife.24767.007

Figure 3 continued on next page
We next examined the in vivo significance of AKAP10 and mitochondrial OM PKA signaling by using morpholinos to block akap10 expression in zebrafish (Danio rerio) embryos (termed morphants) (Figure 3G). For over two decades, the zebrafish has been an invaluable model for the study of hematopoiesis and drug discovery (Jing and Zon, 2011; Zon and Peterson, 2005). Remarkably, akap10 morphants were anemic with decreased hemoglobinization (Figure 3H, red arrowheads) compared to control embryos. We quantified these changes in red cell parameters by performing similar experiments on a transgenic zebrafish line in which all erythroid cells are marked by eGFP expression [Tg(globin-LCR:eGFP)] (Ganis et al., 2012). Flow-cytometry analysis revealed that akap10 morphants had reduced RBC counts (Figure 3I). Together, our data suggest that mitochondrial OM PKA signaling is required for proper heme production and RBC development in vivo.

The terminal heme enzyme, ferrochelatase, is directly phosphorylated by PKA

Next, we asked whether mitochondrial OM PKA signaling directly regulated heme biosynthesis by phosphorylating mitochondrial heme enzymes. Of the mitochondrial enzymes ALAS2, PPOX, CPOX, and FECH, the only enzyme with a predicted high-confidence PKA site (R/K-R/K-X-S/T-Z, where X is any amino acid and Z is an uncharged residue) is FECH at Thr116 (human FECH and Thr115 for murine FECH) (Figure 4A). This residue is evolutionarily conserved and is present on one of the lips of the active site pocket positioned in the middle of a long α-helix (Figure 4B–D). In its unphosphorylated form, the side chain hydroxyl group of Thr116 (colored fuchsia) is sandwiched between His86 (colored blue) and Leu87 on an adjacent α-helix, forming a hydrogen bond with His86 (Figure 4C and D) (Wu et al., 2001). Structural modeling suggests that the bulk of the added phosphate on the side chain of Thr116 would cause movement of the Thr116 α-helix away from the His86 α-helix, and thereby, shift the Thr116-containing α-helix closer inwards towards the active site pocket opening and the porphyrin ring (shown in red) (Figure 4C and D). Such a modification may also destabilize the structure sufficiently to allow for more efficient movement of the active site lip during catalysis.

To test if FECH is directly targeted by PKA for phosphorylation, we performed an in vitro kinase assay by mixing together purified His-tagged human FECH and PKAc, followed by western analysis. This experiment showed that FECH is directly phosphorylated by PKA in an ATP-dependent fashion (Figure 4E). Using [γ-32P]-ATP labeling, we calculated that, in vitro, 9.8 ± 3.2% of purified FECH was phosphorylated after 30 min. Substitution of Thr116 with Ala (T116A) abolished this phosphorylation (Figure 4F). In addition, consistent with the preference of PKAc for positively charged residues at the +3 and +2 positions (Smith et al., 2011), mutation of either Lys113 (K113L) or Arg114 to Leu (R114L) similarly reduced FECH phosphorylation (Figure 4F), strongly indicating that Thr116 of human FECH constitutes a bona fide PKA target. We also examined FECH phosphorylation in erythroid cells by performing similar immunoblot analysis. Immunoprecipitated FECH from differentiated MEL cells was detected by two different phospho-threonine antibodies—one targeting the Lys-X-X-pThr motif and another recognizing the pThr-Pro sequence (Figure 4G). High-dose forskolin treatment also increased phosphorylation of FECH in differentiating MEL cells (Figure 4—figure supplement 1). Conversely, inhibition of OM PKA with loss of AKAP10 resulted in diminished FECH Thr115 phosphorylation (Figure 4H). In toto, our results support a model where PKA becomes localized at the mitochondria OM of maturing erythroid cells and directly phosphorylates FECH.
Figure 4. FECH is directly phosphorylated by PKA. (A) The motif surrounding Thr116 in human FECH constitutes a PKA phosphorylation site with a canonical Lys (K) and Arg (R) at positions −3 and −2, respectively, and an uncharged residue at the +1 position. (B) This PKA phosphorylation motif is highly conserved in FECH proteins across vertebrate species. (C) The FECH homodimer is shown with the transparent surface in green and the subunits in solid green ribbon. PPIX is shown as a red space filling model, the [2Fe-2S] clusters as solid rust and yellow balls, and the highlighted Thr116 (site of PKA-mediated phosphorylation) shown as solid violet spheres. The α-helix in which Thr116 resides, highlighted with lemon green, forms one lip of the opening to the active site where porphyrin is bound in this structure (PDB 2QD1). (D) The structure surrounding Thr116 (shown in fuchsia), which is situated in the middle of a long α-helix, is enlarged. Thr116 is in close proximity and hydrogen bonded to His86 (dark blue) and adjacent to Val85 (dark green). The Thr116-containing α-helix is highlighted with lemon green and the protoporphyrin behind in the active site is purple. Structural modeling suggests that phosphorylation of Thr116 would result in a shift of the Thr116-containing α-helix away from the His86-containing α-helix. (E) Purified recombinant His-tagged human FECH was phosphorylated by purified PKAc only in the presence of ATP at a 1:1 ratio. The phosphorylated form of FECH was detected by immunoblotting with an anti-phosphothreonine antibody. (F) An in vitro kinase assay was also performed with wild-type and variant forms of purified His-tagged human FECH. Disruption of the lysine or arginine at the −3 and −2 positions, respectively, similarly abolishes phosphorylation in vitro as a T116A mutation as shown by immunoblotting. (G) Differentiated MEL cells were immunoprecipitated with the indicated antibodies and immunoblotting analysis was performed. Immunoprecipitated FECH can be recognized by two different anti-phosphothreonine antibodies directed against the upstream positive residues and the proline immediately following the threonine. (H) Wild-type (WT) or AKAP10-KO (KO) were lysed and immunoprecipitated with the indicated antibodies. Bound proteins were analyzed by immunoblotting. All immunoblots were performed twice. ATP-adenosine triphosphate; PPIX-protoporphyrin IX; IB-immunoblot; IP-immunoprecipitate.

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Figure 4 continued on next page
**FECH phosphorylation is required for full activity during erythroid development**

Past work has demonstrated that FECH is phosphorylated by protein kinase C (PKC) (Sakaino et al., 2009). PKC-mediated FECH phosphorylation occurs in a domain buried within an inaccessible hydrophobic fold that did not directly impact enzyme catalysis (Sakaino et al., 2009). In contrast, the position of Thr116 (Thr115 in mice) that is modified by PKA suggests that it would have a direct effect on FECH activity (Wu et al., 2001). We first examined this by using an in vitro $^{55}$Fe-based assay to measure and compare the amount of radiolabeled deuteroporphyrin-IX (DP) that can be produced by unmodified and modified FECH. DP, as a more soluble analog of the naturally occurring heme precursor protoporphyrin-IX (PPIX), is frequently employed in such measurements and is similarly metalated by FECH to generate deuteroheme (Najahi-Missaoui and Dailey, 2005). Significantly more radioactivity can be detected when purified His-tagged human FECH was added alone to the metalation reaction (Figure 5A) and the addition of purified PKAc to the reaction to catalyze the phosphorylation of FECH resulted in an approximately two-fold increase in $^{55}$Fe measurements (Figure 5A). This increase in activity is not attributable to PKAc per se, which has no ferrochelatase activity, and the reaction is completely DP-substrate dependent (Figure 5A). More detailed analysis on enzyme kinetics revealed that phosphorylation had a pronounced effect on maximum velocity ($v_{max}$) but did not significantly change the Michaelis-Menten ($K_m$) constant (Figure 5B and Figure 5—figure supplement 1A), suggesting that it has no major influence on substrate binding. An important caveat to these kinetic measurements is that they were performed at 25°C while all other in vitro assays were performed at 37°C and, thus, may not fully reflect enzyme kinetics both in vivo as well as other single time-point experiments. We also examined FECH activity in intact mitochondria isolated from maturing erythroid cells treated with a high dose of PKA-activating forskolin. Mitochondria from differentiating MEL cells exposed to high-dose forskolin catalyzed higher DP metalation compared to mitochondria derived from untreated cells (Figure 5C). In contrast, performing the assay with N-methyl mesoporphyrin-IX (NMMP)—a PPIX analog that is an inhibitor of FECH and not subject to metalation (Dailey and Fleming, 1983)—instead of DP, resulted is very low $^{55}$Fe extraction that was refractory to forskolin treatment (Figure 5C). Conversely, FECH activity was reduced in AKAP10-KO cells that had compromised FECH phosphorylation (Figures 4H and 5D). These data indicate that phosphorylation of FECH at Thr116 by OM PKA increases FECH catalytic activity.

EPP patients harboring FECH mutations retain residual FECH activity (Balwani and Desnick, 2012), suggesting that subtle changes in FECH function have important biological implications. Thus, to examine the in vivo implications of FECH phosphorylation under physiological conditions, we used CRISPR/Cas9-directed homology repair to knock-in a T115A substitution into the endogenous Fech gene in murine erythroid cells (Figure 5E). Genotyping and subsequent sequencing confirmed that mutant cells possessed only the Fech<sup>T115A</sup> allele (Figure 5F). Compared to wild-type protein, FECH<sup>T115A</sup> mutant protein was similarly induced upon erythroid differentiation and was phosphorylation defective (Figure 5G and H). Direct measurements of enzyme activity from intact mitochondria isolated from wild-type and mutant maturing erythroid cells demonstrated that FECH<sup>T115A</sup> had diminished ferrochelatase activity (Figure 5I). Furthermore, o-dianisidine staining and high-performance liquid chromatography (HPLC) analysis revealed that erythroid cells expressing only FECH<sup>T115A</sup> protein had reduced hemoglobinization as well as lower intracellular hemin levels (Figure 5J and Figure 5—figure supplement 1B). Conversely, FECH<sup>T115A</sup>-expressing cells had concomitantly elevated accumulation of the upstream, free protoporphyrin IX (PPIX) precursor (Figure 5K). Clinically, excess erythroid PPIX accumulation is only found in EPP cases where it serves as a diagnostic marker (Balwani and Desnick, 2012; Whatley et al., 2004). The build-up of PPIX in FECH<sup>T115A</sup>-expressing cells strongly argues that this mutation specifically affects FECH function while the upstream heme biosynthetic pathway remains unaffected.
Figure 5. Phosphorylation of FECH is required for its full activity. (A) Single-point FECH activity was determined in vitro by measuring the amount of $^{55}$Fe incorporation into the protoporphyrin IX analog, deuteroporphyrin (DP). FECH has basal activity that is significantly increased by PKA-mediated phosphorylation. This increase in activity was dependent upon both DP and ATP, highlighting the substrate specificity of the assay for DP and the dependence on phosphorylation. (B) Kinetic analyses were subsequently performed with 0.1 μM FECH, 3 μM DP, and 0.2–100 μM $^{55}$FeCl$_3$ at 25°C. Figure 5 continued on next page
Phosphorylation of FECH leads to a statistically significant increase in maximum velocity ($v_{\text{max}}$). There was no significant difference in the $K_m$ values. (C and D) FECH activity was measured in isolated intact mitochondria. Samples from high-dose forskolin (FSK)-treated differentiating MEL cells have higher FECH activity (C) (*p-value<0.05, Mean ± SEM, n = 5). In contrast, AKAP10-KO mitochondria had less FECH activity (D). Very little activity was detected in samples in which DP was substituted with NMMP. (E) A schematic showing the intron 3 and exon 4 sequences of wild-type murine Feg as well as the CRISPR oligo and the single-stranded DNA (ssDNA) that were introduced as a template for DNA repair. Intronic and exonic sequences are shown in lower and upper cases, respectively. Highlighted in yellow are the three PAM (protospacer adjacent motif) sequences closest to the T115A mutation site that facilitates the potential use of multiple CRISPR oligos. The missense mutations necessary to generate the T115A substitution are in orange. Shown in blue are synonymous substitutions designed to either disrupt the PAM sequences to prevent cleavage of the newly introduced mutant allele or to facilitate genotyping using allele-specific primers near the T115A mutation site. (F) Genomic DNA was isolated from the individual clones of MEL cells and used for PCR analysis with allele-specific primers. The parental MEL cells only had the wild-type allele. The intron 3 and exon 4 sequences of these cells were sequenced to confirm these agarose gel electrophoresis results. (G) Undifferentiated and differentiated parental and mutant cells expressing only the FECH$^{T115A}$ allele were lysed and subjected to western analysis to examine the induction of FECH protein during erythroid maturation. FECH$^{T115A}$ protein had very similar up regulation with differentiation. (H) Differentiated MEL cells were lysed, immunoprecipitated with the indicated antibodies, and bound proteins were subjected to western analysis. Cells expressing only mutant FECH were phosphorylation defective at Thr115. (I–K) Mitochondria isolated from differentiated MEL cells expressing only endogenous FECH$^{T115A}$ (Mut), generated by genome editing, has lower FECH activity than wild-type (WT) control (I). These cells expressing non-phosphorylated FECH also have reduced hemoglobinization by o-dianisidine staining (J) and increased accumulation of PPX substrate (K) as demonstrated by HPLC analysis (*p-value<0.05, Mean ± SEM, n = 11). (L) Wild-type or mutant differentiated MEL cells treated with vehicle (MOCK) or FSK were stained with o-dianisidine. Cells expressing mutant FECH were refractory to the effects of FSK. *p-value<0.05, Mean ± SEM, n = 3, unless otherwise specified. All immunoblots were performed twice. IB-immunoblot; $v_i$-initial velocity; $K_m$-Michaelis-Menten constant; FSK-forskolin; NMMP-N-methyl-mesoporphyrin-IX.

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The following source data and figure supplement are available for figure 5:

**Source data 1.** Maximum velocity ($v_{\text{max}}$) and Michaelis-Menten ($K_m$) constants.

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**Figure supplement 1.** Phosphorylation is required for full FECH activity.

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Erythropoietin signaling activates PKA to phosphorylate FECH

During normal and stress erythropoiesis, erythropoietin (EPO) signaling through its cognate receptor tyrosine kinase (EPOR) regulates survival and proliferation of erythroid progenitors (Kuhrt and Wojchowski, 2015; Testa, 2004). However, there is evidence to suggest that EPO/EPOR signaling regulates other key aspects of erythropoiesis. Unfortunately, the study of such mechanisms has been hampered by the requirement of EPOR signaling in the early stages of the erythropoietic hierarchy (Beale and Chen, 1983; Chida et al., 1999; Socolovsky et al., 1999; Testa, 2004). Nevertheless, uncovering EPOR effectors in later differentiation stages has important clinical relevance. For example, limiting erythroid iron uptake dramatically ameliorates Polycythemia Vera (PV) symptoms in a murine model expressing constitutively active V617F JAK2 (Ishikawa et al., 2015). This is likely due to STAT5-mediated transcriptional up-regulation of the transferrin receptor (Zhu et al., 2008). Despite this, neither PV murine models nor EPO overexpressing transgenic mice show any signs of iron overload (Li et al., 2011; Vogel et al., 2003), indicating that EPO signaling not only promotes iron uptake in RBCs but also coordinates its physiologic assimilation into heme without excessive iron accumulation.

EPOR activates several pathways including the janus kinase 2 (JAK2)/signal transducer and activator of transcription 5 (STAT5), mitogen activated protein kinase (MAPK), and phosphatidylinositol kinase (PI3K) pathways (Kuhrt and Wojchowski, 2015), leading us to question whether PKA is also activated by EPO. We tested this by treating primary murine fetal liver erythroblasts with EPO to determine if EPOR activation had any effect on PKA signaling by monitoring phosphorylation of the prototypical PKA target, CREB, on Ser133 (Altarejos and Montminy, 2011; Taylor et al., 2012). As expected, EPO activation triggered Tyr694 STAT5 phosphorylation, which was similarly observed with EPO treatment of human UT7 erythroid cells (Figure 6A–C). CREB$^{\text{Ser133}}$ phosphorylation but not STAT5$^{\text{Tyr694}}$ phosphorylation was blocked by PKI (14-22) (Figure 6D), indicating that EPO signaling specifically activates PKA. In contrast, co-treatment of UT7 cells with the JAK2 inhibitor, Ruxolitinib, robustly inhibited both STAT5$^{\text{Tyr694}}$ and CREB$^{\text{Ser133}}$ phosphorylation (Figure 6E), strongly suggesting that PKA lies downstream of the EPO/JAK2 pathway. Co-immunoprecipitation
Figure 6. PKA links EPO signaling with heme production during erythropoiesis. (A) Primary murine erythroblasts were cultured from the E13.5 fetal liver, starved for 2 hr and treated with EPO (50 U/mL) for the indicated times. Western analysis showed that EPO triggered increased phosphorylation of STAT5²⁹⁷ and CREB³³³, indicating increased JAK2 and PKA activity, respectively. (B and C) Human UT7 erythroid cells were serum-starved overnight, treated with EPO (2 U/mL) for the indicated times, and subjected to immunoblot analysis. EPO treatment increased both STAT5²⁹⁷ and CREB³³³ phosphorylations similar to (A). A representative blot is shown in (B) and densitometry quantification from three independent experiments (n = 3) where the phospho-CREB signal is normalized to total CREB signal is shown in (C). Time points within the first 60 min were significantly different.
than time 0. (D and E) UT7 cells were treated with the indicated compounds, and immunoblot analysis was performed. CREB<sup>Ser133</sup> phosphorylation can be blocked by both the PKA inhibitor, 14–22, (D) and the JAK2 inhibitor, Ruxolitinib (E). In contrast, 14–22 had no effect on STAT5 phosphorylation (D). (F) UT7 cells untreated or treated with EPO were lysed and immunoprecipitated with the indicated antibodies. Western analysis showed that EPO stimulation resulted in dissociation of PKAc from PKAr. (G) Following PHZ treatment, primary erythroblasts were harvested from the adult murine spleen, starved for 2 hr, and stimulated with EPO (50 U/mL) for 30 min. Lysates were immunoprecipitated and subjected to immunoblot analysis that showed increased FECH phosphorylation with EPO-mediated activation. (H and I) HEL cells induced to hemoglobinize by L-ALA supplementation were treated with Ruxolitinib for 2 hr. Western blot analysis demonstrated reduced CREB<sup>Ser133</sup> phosphorylation (H) and FECH phosphorylation (I) with inhibition of the constitutively active JAK2 mutant. All immunoblots were performed twice unless otherwise specified. *p-value<0.05, Mean ± SEM, n = 3. IB-immunoblot; IP-immunoprecipitate; EPO-erythropoietin; PHZ-phenylhydrazine; HEL-human erythroleukemia; L-ALA-α-aminolevulinic acid.

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Phosphorylated STAT5 forms a complex with PKAc

The role of PKA in erythropoiesis has remained enigmatic. This is because early studies failed to detect changes in cAMP levels despite its stimulatory effect on iron incorporation during RBC maturation that is likely mediated by the synergistic effects of CREB on STAT5 transcription (Boer et al., 2002, 2003; Gidari et al., 1971; Schooley and Mahlmann, 1975). However, recent evidence indicates that PKAc activation is much more complex involving direct protein-protein interactions with other cell signaling regulators and feedback mechanisms independent of cAMP (Taylor et al., 2012; Wong and Scott, 2004; Yang et al., 2013, 1995; Zakhary et al., 2000). Thus, we next asked whether PKAc interacts with proteins in the EPOR/JAK2/STAT5 signaling pathway, leading to its activation.

Co-immunoprecipitation experiments performed using lysates from HEL cells that have constitutive JAK2<sup>V617F</sup> signaling revealed that STAT5 formed a complex with PKAc (Figure 7A). In contrast, STAT5 could not be detected when the immunoprecipitation was performed with a control antibody (Figure 7A). The STAT5-PKAc complex formation was sensitive to pharmacologic inhibition of JAK2 function with Ruxolitinib (Figure 7B). We also tested whether EPO can trigger the formation of the STAT5-PKAc complex. STAT5 only co-precipitated with PKAc in lysates harvested from EPO-stimulated human UT7 erythroid cells (Figure 7C). This fraction of STAT5 was phosphorylated on Tyr694 (Figure 7C). In contrast, STAT5 did not co-precipitate with PKAr subunits (Figure 7D), suggesting that phospho-STAT5/PKAc form a distinct complex apart from PKAr. Lastly, the phospho-STAT5/PKAc complex can only be found in cell lysates derived from the cytosol but not the mitochondria (Figure 7E) and indicates that at least a fraction of active PKAc can diffuse freely to phosphorylate nuclear CREB (Hagiwara et al., 1993; Mayr and Montminy, 2001). Based on our results, we propose a model in which PKA signaling components are localized to the outer mitochondrial membrane during erythropoiesis by AKAP10 where it becomes activated by EPO signaling to regulate heme biosynthesis (Figure 7F).

Discussion

Heme metabolism genes are downstream of GATA1 during RBC development and regulation of heme production has always been thought to occur at the level of gene transcription (Fujiwara et al., 2009; Handschin et al., 2005; Phillips and Kushnir, 2005). However, GATA1 is required for very early stages of erythroid development when the demand for heme and hemoglobin is still low (Fujiwara et al., 1996). This raises the possibility that heme metabolism during cell
Figure 7. Phosphorylated STAT5 forms a molecular complex with mitochondrial PKAc. (A) HEL cells were lysed and immunoprecipitated with the indicated antibodies and subjected to immunoblot analysis. (B) HEL cells treated with MOCK or Ruxolitinib for 2 hr were subjected to similar analysis as (A). (C and D) UT7 cells starved overnight and treated with MOCK or EPO for 10 min were lysed and immunoprecipitated with the indicated antibodies. Immunoblot analysis was performed with the indicated antibodies. (E) Cytosolic and mitochondrial extracts were isolated from HEL cells and immunoprecipitated with the indicated antibodies. Bound proteins were resolved on SDS-PAGE and analyzed by immunoblotting with the indicated antibodies. (F) Model of how, during erythropoiesis, EPO signaling activates PKA at the mitochondrial OM that is localized by the GATA1-target, AKAP10. PKA phosphorylates FECH, which is required to achieve full FECH activity necessary to accommodate the vast heme demand for hemoglobin assembly. All immunoblots were performed twice. IB-immunoblot; IP-immunoprecipitate.

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differentiation requires coordinated metabolic alterations dictated by extracellular signaling cues. EPO signaling is a critical regulator of erythropoiesis and elucidating downstream signaling pathways has been an active area of research. While it is dispensable for early erythroid specification, EPO is critical for the proliferation and survival of early erythroid progenitors (Kuhrt and Wojchowski, 2015) and promotes their differentiation by increasing iron uptake and reducing 'stemness' potential (Decker, 2016; Ishikawa et al., 2015; Park et al., 2016; Zhu et al., 2008). However, nothing is known regarding how EPO signaling can influence heme metabolism. Our work supports a unifying model linking the erythroid transcriptional program with a novel PKA-dependent mechanism downstream of EPO that sheds light into how heme metabolism is coupled to development.

There is a growing body of evidence that, in the absence of intrinsic apoptotic signals, mitochondrial PKA signaling within the matrix is compartmentalized due to the impermeability of the inner mitochondrial membrane (Acin-Perez et al., 2009; Lefkimmiatis et al., 2013). Under this scenario, phosphorylation of matrix proteins can only be achieved by activating signals within the matrix (Acin-Perez et al., 2009; DiPilato et al., 2004; Lefkimmiatis et al., 2013). However, our results support an alternative explanation in which proteins are modified prior to transport. The import of nuclear-encoded mitochondrial proteins requires the maintenance of these proteins in an unfolded state (Lodish et al., 2012), which would allow greater accessibility of target motifs. In support, mitochondrial membrane embedded BAX is, first, phosphorylated by OM PKA (Danial et al., 2003; Harada et al., 1999) and the requirement for protein unfolding during mitochondrial import is consistent with the low level of FECH phosphorylation in vitro (approximately 10%) that we observed. It is also very possible that binding of the FECH substrate to the PKA kinase induces conformational changes that would render the target motif more favorable to modification. This mechanism has previously been proposed as a means to prevent 'promiscuous' phosphorylation (Dar et al., 2005; Dey et al., 2011). Regardless the mechanism, given its exposure to the cytosol and access to proteins destined for mitochondrial localization, the OM is a prime location for such modifications to occur that would coordinate mitochondrial physiology with overall cellular behavior. Accordingly, studies have shown that OM PKA has unique signaling properties in that it is similarly responsive to cytosolic activation mechanisms but remains active much longer and is largely subject to cAMP-independent regulation (Lefkimmiatis et al., 2013). Our data where loss of AKAP10 and OM PKA signaling results in defective FECH modification and activity is consistent with this idea (Figures 2 and 3).

The regulation of PKA activity involves an intricate signaling network more complex than the canonical cAMP pathway (Lefkimmiatis et al., 2013; Manni et al., 2008; Yang et al., 2013). Our work implicates phosphorylated STAT5 as a novel PKAc binding protein that can displace it from autoinhibitory PKAr subunits (Figures 6 and 7) and is corroborated by recent work showing that phospho-STAT proteins, particularly STAT3 and STAT5, localize to mitochondria (Carbognin et al., 2016; Gough et al., 2009; Meier and Larner, 2014; Wegrzyn et al., 2009). Interestingly, in our analysis, mitochondrial expression of PKAc subunits was not as robustly increased as PRKAR2B (Figure 1C and D). Studies have demonstrated that, upon activation, mitochondrial OM PKAc begins to gradually diffuse throughout the cell (Webb et al., 2008). Thus, the non-stoichiometric increase in PKAc compared to PKAr expression is consistent with a dynamic signaling event.

To date, there have been no reported EPP-associated FECH mutations at the PKA target motif. However, the low prevalence of this disease has made discerning genotype-phenotype correlations difficult. Our findings that PKA is an effector of EPO/JAK2 signaling implicate PKA activity not only in the pathogenesis of EPP but also a spectrum of hematologic diseases (Ishikawa et al., 2015). PRKAR1A inactivating mutations are associated with metabolic syndromes in which anemia is prevalent (Stratakis and Cho-Chung, 2002). This is in agreement with murine models where PRKAR1A deletion has the most widespread effect and is the only knockout of the PKA family with embryonic lethality (Stratakis and Cho-Chung, 2002). Paradoxically, we found that it is the mitochondrial expression of PRKAR2B, and not PRKAR1A, that is most dramatically increased in maturing erythroid cells (Figure 1C–E). The complex nuances of PKA signaling make it very difficult to reconcile these findings. Regulatory subunits restrict both the localization as well as the activation of PKA (Wong and Scott, 2004), making it very challenging to distinguish the relative contributions of these two mechanisms. The dual role of PKAr is highlighted by the incomplete rescue of PRKAR1A−/−-associated embryonic defects with PRKACA ablation (Amieux and McKnight, 2002). Further complicating matters is the well-documented instances of compensatory responses (Kirschner et al., 2009), raising the possibility that prominent roles for other PKA isoforms may simply be masked. Indeed,
there is both genetic and biochemical evidence supporting a pivotal and specific role for PRKAR2B in blood development and disease. Global transcriptome analysis has shown that PRKAR2B mRNA is selectively high in CD71+ early erythroid cells (Su et al., 2004) and PRKAR2B binds with higher affinity than PRKAR1A to AKAP10 (Burns et al., 2003). The latter point is particularly important given the recent correlation of AKAP10 polymorphisms with human blood traits in genome-wide association studies (Gieger et al., 2011). It is also notable that AKAP10 encodes many isoforms that may localize to different subcellular compartments (Eggers et al., 2009; Huang et al., 1997a). Although our CRISPR targeting strategy was designed to specifically disrupt the N-terminal mitochondrial-targeting motif (Figure 2G), non-mitochondrial AKAP10 isoforms may function in a variety of contexts both in hematopoiesis and in other aspects of development including the cardiovascular system (Kammerer et al., 2003; Tingley et al., 2007). Our work, here, provides further evidence that perturbations in PKA signaling have significant impact on human health (Kammerer et al., 2003) including the pathogenesis of hematologic diseases that, to date, has been unappreciated and warrants further investigation.

Materials and methods

Cell culture

The DS19 murine erythroleukemia (MEL) subclone (RRID:CVCL_2111) was kindly provided by Arthur Skoultchi (Albert Einstein Medical College, Bronx, NY, USA). Parental human UT7 erythroid cells (RRID:CVCL_5202) were kindly provided by Meredith S. Irwin (Hospital for Sick Children, Toronto, ON, Canada). Human erythroleukemia (HEL) cells (RRID:CVCL_2481) were kindly provided by Ann Mullally (Brigham and Women’s Hospital, Boston, MA, USA). All cells are mycoplasma negative and the International Cell Line Authentication Committee lists none of them as a commonly misidentified cell line. The identities of all cells were confirmed by their labs of origin since none of them are commercially available and have no standard authentication reference sample.

DS19 MEL and primary murine erythroid progenitors from E13.5 fetal liver were cultured and differentiated as previously described (Chung et al., 2015). Differentiating DS19 MEL cells at day 3 of 2% DMSO differentiation were treated with 10 μM (low-dose) or 50 μM (high-dose) forskolin (ThermoFisher, Waltham, MA), or 50 μM 8-Br-cAMP (Sigma-Aldrich, St. Louis, MO) for 30 min or 10 μM dmPGE2 (Cayman Chemicals, Ann Arbor, MI) for 60 min and stained with o-dianisidine as described below. Inhibition with 20 μM H-98 (Tocris Bioscience, Minneapolis, MN), 100 nM PKI (14-22) (ThermoFisher Scientific), or 30 nM bis-indolylmaleimide II (Tocris Bioscience) were performed by pre-treating the cells for 30 min prior to PKA pharmacologic activation.

Bulk murine erythroid progenitors from adult spleen (splenocytes) were prepared and EPO-stimulated as previously described (Maeda et al., 2009; Socolovsky et al., 2001). After two hours of rhEPO (50 U/mL) stimulation, cells were harvested and subjected to immunoprecipitation and western analysis.

Human UT7 erythroid cells were cultured in αMEM (Gibco, Gaithersburg, MD) supplemented with 20% heat-inactivated fetal bovine serum (Serum Source International, Charlotte, NC) and 10 ng/mL GM-CSF (Peprotech, Rocky Hill, NJ). For stimulation experiments, UT7 cells were starved overnight without GM-CSF. The next morning, rhEPO (2 U/mL) was added for the indicated times prior to harvesting.

Human erythroleukemia cells (HEL) were cultured in RPMI supplemented with 10% heat-inactivated fetal bovine serum and treated with 1 μM Ruxolitinib, which was a kind gift of Dr. Ann Mullally (Brigham and Women’s Hospital) and added to the cells for the indicated times.

shRNA experiments

Akap10-targeting and NT9 control shRNAs in the pLKO.1-puro vector were purchased from Sigma-Aldrich. MEL cells were electroporated and monoclonal populations expressing these shRNAs were isolated as previously described (Chung et al., 2015). The sequences of the Akap10 shRNAs were: shRNA-1, 5'-CCGGCCAAAGCTATGTGGCAGATTCGAGATTGATCGAAATGACCTTTGTTTGTGTTTGG-3' and shRNA-2, 5'-CCGGGCAAGAGCCTTTTATGAGTTTCTCGAGAACACTCAAAGTGCATCTTGTTTTTGG-3'.
Isolation of mitochondria-enriched fractions

1 × 10⁶ DS19 MEL cells of each condition or 5 × 10⁶ HEL cells were collected, washed once in cold PBS, and resuspended in 1 mL of MSHE buffer (220 mM mannitol, 70 mM sucrose, 5 mM potassium HEPES pH 7.4, 1 mM EGTA pH 7.4, supplemented with Complete EDTA-free protease inhibitor tablets [Roche, Indianapolis, IN]). Samples were then dounce homogenized and pelleted by spinning at 1000 g for 10 min at 4°C. The supernatant was separated and the leftover pellet was resuspended in 200 μL of MSHE buffer. All samples were centrifuged again at 1000 g for 10 min at 4°C. All supernatants from each undifferentiated and differentiating samples were collected and combined and centrifuged again at 1000 g for 10 min at 4°C. This supernatant was then transferred to a new tube and centrifuged at 8000 g for 20 min at 4°C. The resulting supernatant containing cytosolic proteins was transferred to another tube and flash frozen. The pellet was resuspended and washed twice more in 200 μL of MSHE buffer. The final pellet, containing mitochondrially enriched membrane fractions, was flash frozen until subjected to mass spectrometry analysis. For western analyses, the pellet was lysed in NP-40 lysis buffer. For ferrochelatase activity assays, the Mitochondrial Isolation Kit for Mammalian Cells (ThermoScientific) was used according to manufacturer’s instructions. The mitochondrial pellet was resuspended in 200 μL of reaction buffer and immediately used.

Submitochondrial fractionation and proteinase K digestion experiments

Intact mitochondria were isolated as described above without protease inhibitors and resuspended in MSHE buffer containing 5 mg/mL digitonin that was prepared fresh at 4% in water immediately before each experiment. Samples were incubated on ice for 15 min with vortexing at maximum setting every few minutes for 10 s intervals. After 15 min, samples were centrifuged at 10,000 g for 10 min at 4°C. The supernatant was transferred to a second tube and the leftover pellet was re-extracted twice more with 80 μL of MSHE containing 5 mg/mL digitonin, leaving a final pellet enriched for MPs. The supernatant from the second and third extractions were discarded. The supernatant containing the IMS was removed and kept in a separate tube. The pellet contained the OM fraction. Proteinase K protection assays were performed as previously described (Shirihai et al., 2000).

Proteomics analysis

Protein was extracted from purified mitochondria by dissolution in 8 M urea, 50 mM Tris pH 8.0, followed by probe sonication. Extracted protein was reduced and alkylated with dithiothreitol and iodoacetamide, respectively. Alkylated protein samples were first digested with endoproteinase LysC for 4 hr at ambient temperature with an enzyme to protein ratio of 200:1. Each sample was diluted with 50 mM Tris to 1.5 M urea and further digested with trypsin at an enzyme to protein ratio of 50:1, overnight and at ambient temperature. Peptides from each sample were desalted over a C18 solid phase extraction cartridge and dried down. Each sample was resuspended in 0.2 M triethylammonium bicarbonate pH 8.5, and labeled with tandem mass tag reagents (TMT) as previously described (Hebert et al., 2013). Labeled samples were pooled, dried, and fractionated across a strong cation exchange column (Polysulfoethyl A). Each fraction was dried, desalted, and resuspended in 0.2% formic acid.

All nano UPLC separations were performed on a nanoAcquity system. From each fraction, approximately 2 μg of peptides was injected onto a 75 μm inner diameter, 30 cm long, nano column packed with 1.7 BEH C18 particles. The mobile phases were as follows: A) 0.2% formic acid and B) 100% acetonitrile with 0.2% formic acid. Peptide were eluted with a gradient of increasing B from 0%30% over the course of 100 min, followed by a wash with 100% B and re-equilibration at 0% B. Eluting peptides were electro spray ionized and analyzed with an Orbitrap Elite mass spectrometer. The mass spectrometry analysis cycle was as follows. First a survey scan was performed with Orbitrap analysis at 60,000 resolving power at 400 m/z. Peptide precursors in the survey scan were sampled for ms/ms analysis by data dependent top 15 selection with dynamic exclusion turned on. Each peptide precursor selected for sampling was isolated in the ion trap, fragmented by higher energy collisional dissociation (HCD) at 35 NCE, followed by mass analysis of the fragments in the Orbitrap at resolving power 15,000 at 400 m/z.

All data analysis was performed in the COMPASS software suite (Wenger et al., 2011). Spectra were searched against a tryptic target-decoy mouse Uniprot database including protein isoforms.
Methionine oxidation and TMT on tyrosine were searched as variable modifications. Cysteine carboxymethylation, TMT on lysine and TMT on the peptide N-terminus were searched as fixed modifications. The tolerance was set to 0.01 Da for matching fragments to the database. Matching spectra were filtered to 1% FDR at the unique peptide level based on spectral matching score (E-value) and peptide precursor ppm mass error, followed by reporter ion quantitation, protein grouping according to parsimony, and filtering to 1% FDR. Reporter ion quantitation normalization was performed essentially as previously described with the following changes (Grimsrud et al., 2012). First, the dataset was annotated using the MitoCarta1.0 database (Pagliarini et al., 2008). All proteins annotated as mitochondrial for undifferentiated and differentiated samples were averaged and linear regression was performed using Microsoft Excel. The conversion factor corresponding to the slope of the linear regression was applied to all proteins in the database regardless of MitoCarta1.0 status and the log2 fold change was calculated.

CRISPR/Cas9-directed homology repair and Akap10 targeting
CRISPR oligos were cloned into the px335 vector as previously described (Chung et al., 2015). The sequences of the CRISPR oligos were 5’-CACCGAATTTGGGTGTCGGGT-3’ and 5’-AAA-CAACGCAGAACCCTTCTAC-3’. The single-stranded DNA oligo used to direct homology repair was 5’-TTGAGTTTCGAAAGGTGAATTTACCCACCTCTTTATGTGTCGTGATGTTAG TAAGCTTGCACCATCTTGGAGATCCGACGTGCAGCCAATTTCAAGAGCATATCGCAGAA TCGAGGTTGATCCCCATCAAGATGTTGACT-3’. Targeting of Akap10 (NM_019921.3) was performed as previously described with modifications (Chung et al., 2015). The CRISPR oligos were: exon-1, 5’-CTGCATAGTCCGAAAAACAG-3’ and exon-3, 5’-GCAAGGCTGAGTTTGTGAGT-3’.

DS19 MEL cells were electroporated and cultured as previously described (Chung et al., 2015) along with 5 μL of 10 mM ssDNA oligo. Single clones were grown in 96-well plates and screened for the presence of the FechT115A allele using genomic DNA PCR followed by antisense oligonucleotide hybridization with [γ32P]-ATP-labeled (10 mCi/mL, specific activity = 6000 Ci/mmol, Perkin Elmer) wild-type (5’-CATCGCCAAACGCCGAACC-3’) and mutant (5’-CATTGCGAAGCGACGTGC-3’) oligos as described elsewhere (Hildick-Smith et al., 2013). Clones harboring the mutant allele were expanded and characterized by allele specific PCR (see below) and sequenced to confirm the presence of only the mutant allele.

High performance liquid chromatography
HPLC analysis was performed as previously described and statistical significance was determined using two-way ANOVA (Yien et al., 2014).

Polymerase chain reaction and site-directed mutagenesis
Genomic DNA was isolated from DS19 MEL cells according to manufacturer’s instructions (Qiagen DNAeasy kit, Germantown, MD). For screening FechT115A knock-in clones using dot blots, the following primers were used to generate the amplicon of interest: 5’-CTGTTTGGCTCTCCTTAG-3’ and 5’-GAGTCCTACTGTAACGAG-3’. For allele-specific PCRs, the latter primer from above was used with either the wild-type, 5’-CATCGCCAAACGCCGAACC-3’, or mutant, 5’-CATTGCGAAGCGACGTGC-3’ oligos as described elsewhere (Hildick-Smith et al., 2013). Clones harboring the mutant allele were expanded and characterized by allele specific PCR (see below) and sequenced to confirm the presence of only the mutant allele.

Site-directed mutagenesis was performed using the Agilent QuikChange Lightning Multi Site-Directed Mutagenesis Kit (La Jolla, CA) according to the manufacturer’s instructions. The following mutagenesis primers were used: T116A, 5’-CATCGCCAAACGCCGAAGGCCCAAGATTCAAAG-3’ and 5’-CTTGAATCTTGGGGCTCGGCGATTGCCGATG-3’; K113L, 5’-GCAACCATCCTAGTCCGAGTTCTGACAA-3’ and 5’-CTGTTTGGCTCTCCTTAG-3’; reverse-1 (R1) primer, 5’-CCCTGACAAAACCCTTGCACTG-3’; and reverse-2 (R2) primer, 5’-CATTGCGAAGCGACGTGC-3’.

Protein purification
Recombinant His-tagged human FECH proteins were expressed and purified as previously described (Burden et al., 1999).
Structural modeling
The 2.0 Å structure of human FECH (PDB 2QD4) was visualized by PyMOL (The PyMOL Molecular Graphics System, Version 1.8 Schrödinger, LLC).

In vitro kinase assay
One microgram of His-tagged wild-type or variant human FECH was mixed with purified PKAc according to manufacturer’s instructions (Promega, Madison, WI). 20% of the kinase reaction was subjected to western blot analysis. The stoichiometry of the kinase reaction was kept at 1:1. For kinase assays using [γ-32P]-ATP (specific activity 6000 Ci/mmol, Perkin Elmer, Boston, MA), the reaction was performed with an incubation time of 30 min instead of 10 min and scaled down to where 0.2 pmol of purified FECH and PKAc were used. Four-fold excess [γ-32P]-ATP was added. After 30 min, each reaction was immunoprecipitated with anti-FECH antibodies (see below). The incubation time for [γ-32P]-ATP labelling was three-times longer than for other assays to ensure maximal phosphorylation in vitro. The amount of radioactivity was quantified in a scintillation counter (Chung et al., 2015) and normalized to immunoprecipitation efficiency, which was calculated by determining the percent of FECH protein that was recovered in the immunoprecipitation relative to input using western blotting followed by densitometry with ImageJ (Schneider et al., 2012). The average and standard error was calculated from three independent experiments.

Western blotting and immunoprecipitation
All immunoblots were performed according to manufacturer’s instructions and as previously described except that all phosphothreonine immunoblots were performed with an HRP-conjugated protein A secondary antibody (Chung et al., 2015). Anti-PDHA1 (ab67592) and anti-AKAP10 (ab97354) rabbit polyclonal antibodies were purchased from Abcam (Cambridge, MA). Mouse monoclonal anti-TUBA1A (DM1A), rabbit polyclonal anti-STAT5 (C-17), anti-PRKACB (C-20), anti-PRKAR2A (M-20), and goat polyclonal anti-FECH (C-20) and anti-HSPD1 (K-19) antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Mouse monoclonal anti-pSer133-CREB (1B6) and anti-pThr-Pro (9391S), rabbit monoclonal anti-pTyr694-STAT5 (D46E7), anti-PRKACA (D38C6), anti-STAT5 (C-17), anti-PRKAR2A (M-20), and goat polyclonal anti-Arg-X-X-pThr (9621S) and anti-PRKARIA (D54D9) antibodies were purchased from Cell Signaling Technology (Danvers, MA). Anti-GAPDH (MAB374) mouse monoclonal and anti-PRKARIIB (ABS14) rabbit polyclonal antibodies were purchased from Millipore (Billerica, MA). Anti-TIM23 mouse monoclonal antibody was purchased from BD Biosciences (Woburn, MA). Immunoprecipitations were performed as previously described (Chung et al., 2010). All immunoblots were performed two independent times except when otherwise specified. Densitometry was performed as previously described and analyzed by one-way ANOVA (Chung et al., 2015).

Ferrochelatase activity assay
Ferrochelatase activity assays using purified His-tagged wild-type human FECH was performed by using 250 ng of purified protein in assay reaction buffer (0.6 M sorbitol, 40 mM HEPES pH 7.4 pH with KOH, 50 mM KCl, 1 mM MgSO4). Ascorbic acid and NADH were then added to final concentrations of 0.4 mg/mL and 2 nM, respectively. The solution was incubated at 37°C for 5 min. After the incubation period, 0.2 mL of 55FeCl3 (38 mCi/mL, specific activity = 54.5 mCi/mol, Perkin Elmer) was added to each sample along with either 5.5 μL of 200 μM DP or NMMP. The samples were then incubated for another 10 min at 37°C and 55Fe-radiolabeled heme was extracted as previously described and counted in a liquid scintillation counter (Chung et al., 2015). For activity assays using intact mitochondria, the protocol was modified to use 50 μg of freshly isolated mitochondria. All experiments were performed in duplicate and three independent experiments were performed followed by statistical analysis using student’s t-test.

For kinetic assays, the conditions of the experiments are described elsewhere using 0.1 μM FECH, 3 μM DP, and 0.2–100 μM 55FeCl3 except the reactions were carried out at 25°C (Hunter et al., 2008). At 30, 60, 120, 180, 300, 600, and 900 s, 10% of each reaction were removed and immediately mixed with cold FeCl3 at a final concentration of 1.25 mM to stop the reaction. All samples were extracted and counted in a scintillation counter as described above. The amount of
product over time was plotted and analyzed using regression analysis with GraphPad Prism v5.0 software. Michaelis-Menten analyses with the initial velocities \( (v_i) \) were subsequently performed using GraphPad Prism v5.0 that calculated the maximum velocity \( (v_{max}) \) and Michaelis-Menten \( (K_m) \) constant. Experiments were performed three times and statistical analysis was performed on GraphPad Prism v5.0.

**Zebrafish experiments**

Injections, o-dianisidine staining, and flow cytometry analysis were performed as described elsewhere (Chung et al., 2015). MOs were purchased from Gene Tools, LLC (Philomath, OR). Zebrafish embryos at the one-cell stage were injected with MOs targeting the exon-1/intron-1 (MO1) and exon-2/intron-2 (MO2) junctions of *D. rerio akap10* (XM_690206). The MO sequences were as follows: MO1, 5’-TGGAGCGGCCCCACTTACCTTTTC-3’; MO3, 5’-TTTAGCAGCAGACACTTACC TTTC-3’.

**Vertebrate animal study approvals and ethics statement**

All zebrafish (RRID:ZIRC_ZL1) and mouse experiments were performed in full compliance with the approved Institutional Animal Care and Use Committee (IACUC) protocols at Boston Children’s Hospital (Protocol #15-07-2974R) and Brigham and Women’s Hospital (Protocol #2016N000117). These studies were approved by local regulatory committees in accordance with the highest ethical standards for biomedical research involving vertebrate animals.

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JC, Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Writing—original draft, Writing—review and editing; JGW, CEM, ASH, LL, Data curation, Formal analysis, Investigation, Methodology; AG, Data curation, Validation, Investigation, Methodology; MM, TAD, Data curation, Investigation, Methodology; HB, Data curation, Formal analysis, Investigation; MDK, Formal analysis, Validation, Investigation, Methodology; JDP, Data curation, Validation, Investigation, Methodology; JK, Conceptualization, Data curation, Formal analysis, Investigation, Methodology; HFL, DEB, ABC, TM, HAD, Formal analysis, Supervision, Investigation, Methodology; SHO, Supervision, Investigation, Methodology; JJC, Formal analysis, Supervision, Validation, Investigation, Methodology; DJP, Formal analysis, Supervision, Investigation, Methodology, Project administration; BHP, Conceptualization, Resources, Supervision, Investigation, Methodology—review and editing

Author ORCIDs
Johannes G Wittig, http://orcid.org/0000-0002-0598-2897
David J Pagliarini, http://orcid.org/0000-0002-0001-0087
Barry H Paw, http://orcid.org/0000-0002-0492-1419

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References
Acín-Pérez R, Salazar E, Kamenetsky M, Buck J, Levin LR, Manfredi G. 2009. Cyclic AMP produced inside mitochondria regulates oxidative phosphorylation. Cell Metabolism 9:265–276. doi: 10.1016/j.cmet.2009.01.012, PMID: 19254571
Altarejos JY, Montminy M. 2011. CREB and the CRTC co-activators: sensors for hormonal and metabolic signals. Nature Reviews Molecular Cell Biology 12:141–151. doi: 10.1038/nrm3072, PMID: 21346730

Chung et al. eLife 2017;6:e24767. DOI: 10.7554/eLife.24767
Amieux PS, McKnight GS. 2002. The essential role of RI alpha in the maintenance of regulated PKA activity. *Annals of the New York Academy of Sciences* **968**:75–95. doi: 10.1111/j.1749-6632.2002.tb04328.x, PMID: 12119269

Amigo JD, Yu M, Troade MB, Gywenn B, Cooney JD, Lambert AJ, Chi NC, Weiss MJ, Peters LL, Kaplan J, Cantor AB, Paw BH. 2011. Identification of distal cis-regulatory elements at mouse mitoferrin loci using zebrafish transgenesis. *Molecular and Cellular Biology* **31**:1344–1356. doi: 10.1128/MCB.00110-10, PMID: 21248200

Bardwell JV, Desnick RJ. 2012. The porphyrias: advances in diagnosis and treatment. *Blood* **120**:4496–4504. doi: 10.1182/blood-2012-05-423186, PMID: 22791288

Bauer DE, Kamran SC, Lessard S, Xu J, Fujiwara Y, Lin C, Shao Z, Canver MC, Smith EC, Pinello L, Sabo PJ, Vierstra J, Voit RA, Yuan GC, Porteus MH, Stamatoyannopoulos JA, Lettre G, Orkin SH. 2013. An erythroid enhancer of BCL11A subject to genetic variation determines fetal hemoglobin level. *Science* **342**:253–257. doi: 10.1126/science.1242088, PMID: 24115442

Beale SJ, Chen NC. 1983. N-Methyl mesoporphyrin IX inhibits phycocyanin, but not Chlorophyll synthesis in Cyanidium caldarium. *Plant Physiology* **71**:263–268. doi: 10.1104/pp.71.1.263, PMID: 16662815

Boer AK, Drayer AL, Rui H, Vellenga E. 2002. Prostaglandin-E2 enhances EPO-mediated STAT5 transcriptional activity by serine phosphorylation of CREB. *Blood* **100**:467–473. doi: 10.1182/blood.V100.2.467, PMID: 12091337

Boer AK, Drayer AL, Vellenga E. 2003. cAMP/PAK-mediated regulation of erythropoiesis. *Leukemia & Lymphoma* **44**:1893–1901. doi: 10.1080/1042819031000116175, PMID: 14738140

Burden AE, Wu C, Dailey TA, Busch JL, Dhawan IK, Rose JP, Wang B, Dailey HA. 1999. Human ferrochelatase: crystallization, characterization of the [2Fe-2S] cluster and determination that the enzyme is a homodimer. *Biochimica Et Biophysica Acta (BBA) - Protein Structure and Molecular Enzymology* **1435**:191–197. doi: 10.1016/S0167-4838(99)00196-X, PMID: 10561552

Burns LL, Canaves JM, Pennybacker JK, Blumenthal DK, Taylor SS. 2003. Isoform specific differences in binding of a dual-specificity A-kinase anchoring protein to type I and type II regulatory subunits of PKA. *Biochemistry* **42**:5754–5763. doi: 10.1021/bi0265729, PMID: 12741833

Canver MC, Bauer DE, Dass A, Yen YY, Chung J, Masuda T, Maeda T, Paw BH, Orkin SH. 2014. Characterization of genomic deletion efficiency mediated by clustered regularly interspaced palindromic repeats (CRISPR)/Cas9 nuclease system in mammalian cells. *Journal of Biological Chemistry* **289**:21312–21324. doi: 10.1074/jbc.M114.56625X, PMID: 24907273

Carbognin E, Betto RM, Soriano ME, Smith AG, Martello G. 2016. Stat3 promotes mitochondrial transcription and oxidative respiration during maintenance and induction of naïve pluripotency. *The EMBO Journal* **35**:618–634. doi: 10.15252/embj.201592629, PMID: 26930601

Chen W, Paradkar PN, Li L, Pierce EL, Langer NB, Takahashi-Makise N, Hyde BB, Shirihi OS, Ward DM, Kaplan J, Paw BH. 2009. Abcb10 physically interacts with mitoferrin-1 (Sic2a3α7) to enhance its stability and function in the erythroid mitochondria. *PNAS* **106**:16263–16268. doi: 10.1073/pnas.0904519106, PMID: 19805291

Chida D, Miura O, Yoshimura A, Miyajima A. 1999. Role of cytokine signaling molecules in erythroid differentiation of mouse fetal liver hematopoietic cells: functional analysis of signaling molecules by retrovirus-mediated expression. *Blood* **93**:1567–1578. PMID: 10029585

Chung J, Lau J, Cheng LS, Grant RI, Robinson F, Ketela T, Reis PP, Roche O, Kamel-Reid S, Moffat J, Ohh M, Perez-Ordazone B, Kaplan DR, Irwin MS. 2010. SATB2 augments enhancer of BCL11A subject to genetic variation determines fetal hemoglobin level. *PNAS* **106**:9133–9138. doi: 10.1073/pnas.0904519106, PMID: 19805291

Chung J, Chen NC. 1983. N-Methyl mesoporphyrin IX inhibits phycocyanin, but not Chlorophyll synthesis in Cyanidium caldarium. *Plant Physiology* **71**:263–268. doi: 10.1104/pp.71.1.263, PMID: 16662815

Chung J, Chen CW, Paw BH. 2012. Heme metabolism and erythropoiesis. *Current Opinion in Hematology* **19**:156–162. doi: 10.1097/MOH.0b013e328351c48b, PMID: 22406824

Chung J, Bauer DE, Ghamari A, Nizzi CP, Deck KM, Kingsley PD, Yen YY, Huston NC, Chen C, Schultz IJ, Dalton AJ, Wittig JG, Palis J, Orkin SH, Lodish HF, Eisenstein RS, Cantor AB, Paw BH. 2015. The mTORC1/4E-BP pathway coordinates hemoglobin production with L-leucine availability. *Science Signaling* **8**:ra34. doi: 10.1126/scisignal.aaa5903, PMID: 25872869

Dailey HA, Fleming JE. 1983. Bovine ferrochelatase. kinetic analysis of inhibition by N-methylprotoporphyrin, manganese, and heme. *The Journal of Biological Chemistry* **258**:11453–11459. PMID: 6688622

Danial NN, Gramm CF, Scorrano L, Zhang CY, Krauss S, Ranger AM, Datta SR, Greenberg ME, Licklider LJ, Lowell BB, Gygi SP, Korsmeyer SJ. 2003. BAD and glucokinase reside in a mitochondrial complex that integrates glycolysis and apoptosis. *Nature* **424**:952–956. doi: 10.1038/nature01825, PMID: 12931191

Dar AC, Dever TE, Sicheri F. 2005. Higher-order substrate recognition of eIF2alpha by the RNA-dependent protein kinase PKR. *Cell* **122**:887–900. doi: 10.1016/j.cell.2005.06.044, PMID: 16179258

Decker T. 2016. Emancipation from transcriptional latency: unphosphorylated STAT5 as guardian of hematopoietic differentiation. *The EMBO Journal* **35**:555–557. doi: 10.15252/embj.201693974, PMID: 26893391

Dey M, Velyvis A, Li JJ, Chiu E, Chiavotti D, Kay LE, Sicheri F, Dever TE. 2011. Requirement for kinase-induced conformational change in eukaryotic initiation factor 2alpha (eIF2alpha) restricts phosphorylation of Ser51. *PNAS* **108**:4316–4321. doi: 10.1073/pnas.1014872108, PMID: 21368187

DiPilato LM, Cheng X, Zhang J. 2004. Fluorescent indicators of cAMP and epac activation reveal differential dynamics of cAMP signaling within discrete subcellular compartments. *PNAS* **101**:16513–16518. doi: 10.1073/pnas.0405973101, PMID: 15545605

Eggers CT, Sacher JC, Goldenring JR, Taylor SS. 2009. D-AKAP2 interacts with Rab4 and Rab11 through its RGS domains and regulates transferrin receptor recycling. *Journal of Biological Chemistry* **284**:32869–32880. doi: 10.1074/jbc.M109.022582, PMID: 19797056
Fujisawa Y, Browne CP, Cunniff K, Goff SC, Orkin SH. 1996. Arrested development of embryonic red blood cell precursors in mouse embryos lacking transcription factor GATA-1. PNAS 93:12355–12358. doi: 10.1073/pnas.93.22.12355, PMID: 8901585

Fujitani T, O’Geen H, Keles S, Blahnik K, Linnemann AK, Kang YA, Choi K, Farnham PJ, Bresnick EH. 2009. Discovering hematopoietic mechanisms through genome-wide analysis of GATA factor chromatin occupancy. Molecular Cell 36:667–681. doi: 10.1016/j.molcel.2009.10.024, PMID: 19941826

Ganis JJ, Hsia N, Trompouki E, de Jong JL, DiBiase A, Lambert JS, Jia Z, Sabo PJ, Weaver M, Sandstrom R, Stamatoyannopoulos JA, Zhou Y, Zon LI. 2012. Zebrafish globin switching occurs in two developmental stages and is controlled by the LCR. Developmental Biology 366:185–194. doi: 10.1016/j.ydbio.2012.03.021, PMID: 22537491

Gidari AS, Zanjani ED, Gordon AS. 1971. Stimulation of erythropoiesis by cyclic adenosine monophosphate. Life Sciences 10:895–900. doi: 10.1016/0024-3205(71)90202-5, PMID: 4327815

Gieger C, Radhakrishnan A, Cvejic A, Tang W, Porcu E, Pistis G, Serbanovic-Canic J, Elling U, Goodall AH, Labrune Y, Lopez LM, Magi R, Meacham S, Okada Y, Pirastu N, Sorice R, Teumer A, Voss K, Zhang W, Ramirez-Solis R, et al. 2011. New gene functions in Megakaryopoiesis and platelet formation. Nature 480:201–208. doi: 10.1038/nature10659, PMID: 22139419

Goessling W, North TE, Loewer S, Lord AM, Lee S, Stoick-Cooper CL, Weidinger G, Puder M, Daley GQ, Moon RT, Zon LI. 2009. Genetic interaction of PGE2 and wnt signaling regulates developmental specification of stem cells and regeneration. Cell 136:1136–1147. doi: 10.1016/j.cell.2009.01.015, PMID: 19303855

Gough DJ, Corlett A, Schlessinger K, Wegryn J, Larner AC, Levy DE. 2009. Mitochondrial STAT3 supports Ras-dependent oncogenic transformation. Science 324:1713–1716. doi: 10.1126/science.1171721, PMID: 19556058

Grimsrud PA, Carson JJ, Hebert AS, Huber SL, Niemi NM, Bailey DJ, Jochem A, Stapleton DS, Keller MP. Westphall MS, Yandell BS, Attie AD, Coon JJ, Pagliarini DJ. 2012. A quantitative map of the liver mitochondrial phosphoproteome reveals posttranslational control of ketogenesis. Cell Metabolism 16:672–683. doi: 10.1016/j.cmet.2012.10.004, PMID: 23140645

Hagiwara M, Brindle P, Haroutunian A, Armstrong R, Rivier J, Vale W, Tsiens R, Montmny MR. 1993. Coupling of hormonal stimulation and transcription via the cysc AMP-responsive factor CREB is rate limited by nuclear entry of protein kinase A. Molecular and Cellular Biology 13:4852–4859. doi: 10.1128/MCB.13.8.4852, PMID: 8336722

Handschin C, Lin J, Rhee J, Peyer AK, Chiu S, Wu PH, Meyer UA, Spiegelman BM. 2005. Nutritional regulation of hepatic heme biosynthesis and Porphyria through PGC-1alpha. Cell 122:505–515. doi: 10.1016/j.cell.2005.06.040, PMID: 16122419

Harada H, Becknell B, Wilm M, Mann M, Huang LJ, Taylor SS, Scott JD, Korsmeyer SJ. 1999. Phosphorylation and inactivation of BAD by mitochondria-anchored protein kinase A. Molecular Cell 3:413–422. doi: 10.1016/S1097-2765(00)80469-4, PMID: 10230394

Hebert AS, Dittenhafer-Reed KE, Yu W, Bailey DJ, Selen ES, Boersma MD, Carson JJ, Tonelli M, Balloon AJ, Higbee AJ, Westphall MS, Pagliarini DJ, Prolla TA, Assadi-Porter F, Roy S, Denu JM, Coon JJ. 2013. Calorie restriction and SIRT3 trigger global reprogramming of the mitochondrial protein acetylome. Molecular Cell 49:186–199. doi: 10.1016/j.molcel.2012.10.024, PMID: 23201123

Hildick-Smith GJ, Cooney JD, Garone C, Kremer LS, Haack TB, Thon JN, Miyata N, Lieber DS, Calvo SE, Akman HO, Yien YY, Huston NC, Branco DS, Shah DJ, Freedman ML, Koehler CM, Italiano JE, Merkenschlager A, Beblo S, Strom TM, et al. 2013. Macrocystic Anemia and mitochondriopathy resulting from a defect in sideroflexin 4. The American Journal of Human Genetics 93:906–914. doi: 10.1016/j.ajhg.2013.09.011, PMID: 24119684

Huang LJ, Durick K, Weiner JA, Chun J, Taylor SS. 1997a. D-AKAP2, a novel protein kinase A anchoring protein with a putative RGS domain. PNAS 94:11184–11189. doi: 10.1073/pnas.94.21.11184, PMID: 9326583

Huang LJ, Durick K, Weiner JA, Chun J, Taylor SS. 1997b. Identification of a novel protein kinase A anchoring protein that binds both type I and type II regulatory subunits. Journal of Biological Chemistry 272:8057–8064. doi: 10.1074/jbc.272.12.8057, PMID: 9065479

Humphries KM, Pennypacker JK, Taylor SS. 2007. Redox regulation of CAMP-dependent protein kinase signaling: kinase versus phosphatase inactivation. Journal of Biological Chemistry 282:22072–22079. doi: 10.1074/jbc.M702582200, PMID: 17548350

Hunter GA, Sampson MP, Ferreira GC. 2008. Metal ion substrate inhibition of ferrochelatase. Journal of Biological Chemistry 283:23685–23691. doi: 10.1074/jbc.M803372200, PMID: 18593702

Ishikawa Y, Maeda M, Pasham M, Aguet F, Tacheva-Grigorova SK, Masuda T, Yi H, Lee SU, Xu J, Teruya-Feldstein J, Ericsson M, Mullally A, Heuser J, Kirchhausen T, Maeda T. 2015. Role of the clathrin adaptor PICALM in normal hematopoiesis and polycythemia vera pathophysiology. Haematologica 100:439–451. doi: 10.3324/haematol.2014.119537, PMID: 25552701

Jing L, Zon LI. 2011. Zebrafish as a model for normal and malignant hematopoiesis. Disease Models & Mechanisms 4:433–438. doi: 10.1242/dmm.006791, PMID: 21708900

Kammerer S, Burns-Hamuro LL, Ma Y, Hamon SC, Canaves JM, Shi MM, Nelson MR, Sing CF, Cantor CR, Taylor SS, Braun A. 2003. Amino acid variant in the kinase binding domain of dual-specific A kinase-anchoring protein 2; a disease susceptibility polymorphism. PNAS 100:4066–4071. doi: 10.1073/pnas.2628028100, PMID: 12646697

Kardon JR, Yien YY, Huston NC, Branco DS, Hildick-Smith GJ, Rhee KY, Paw BH, Baker TA. 2015. Mitochondrial ClpX activates a Key Enzyme for Heme Biosynthesis and Erythropoiesis. Cell 161:858–867. doi: 10.1016/j.cell.2015.04.017, PMID: 25957689
Kirschner LS, Yin Z, Jones GN, Mahoney E. 2009. Mouse models of altered protein kinase A signaling. Endocrine Related Cancer 16:773–793. doi: 10.1677/ERC-09-0068, PMID: 19470615

Kuhrt D, Wojchowski DM. 2015. Emerging EPO and EPO receptor regulators and signal transducers. Blood 125:3536–3541. doi: 10.1182/blood-2014-11-753557, PMID: 25887776

Langendonk JG, Balwani M, Anderson KE, Bonkovsky HL, Anstey AV, Bissell DM, Bloomer J, Edwards C, Neumann NJ, Parker C, Phillips JD, Lim HW, Hamzavi I, Debych JC, Kauppinen R, Rhodes LE, Frank J, Murphy GM, Karstens FP, Sijbrands EJ, et al. 2015. Afamelanotide for Erythropoietic Protoporphyria. New England Journal of Medicine 373:48–59. doi: 10.1056/NEJMoa1414811, PMID: 26123941

Lefkkmimatie K, Leronni D, Hofer AM. 2013. The inner and outer compartments of mitochondria are sites of distinct CAMP/PKA signaling dynamics. The Journal of Cell Biology 202:453–462. doi: 10.1083/jcb.201303159, PMID: 23897891

Li H, Degenhardt B, Tobin D, Yao ZX, Tasken K, Papadopoulos V. 2001. Identification, localization, and function in steroidogenesis of PAP7: a peripheral-type benzodiazepine receptor- and PKA (Rlalpha)-associated protein. Molecular Endocrinology 15:2211–2228. doi: 10.1210/mend.15.12.0736, PMID: 11731621

Li J, Kay DG, Chen E, Green AR. 2011. Mouse models of myeloproliferative neoplasms: jak of all grades. Disease Models & Mechanisms 4:311–317. doi: 10.1242/dmm.006817, PMID: 21558064

Lodish HF, Berk A, Kaiser CA, Krieger M, Bretscher A, Ploegh H, Amon A, Scott MP. 2012. Molecular Cell Biology. 7th Edn. New York: W H Freeman and Company.

Maeda T, Ito K, Mergobh T, Polisen L, Hobbs RM, Wang G, Dong L, Maeda M, Dore LC, Zelent A, Luzzatto L, Teruya-Feldstein J, Weiss MJ, Pandolfi PP. 2009. LRF is an essential downstream target of GATA1 in erythroid development and regulates BIM-dependent apoptosis. Developmental Cell 17:527–540. doi: 10.1016/j.devcel.2009.09.005, PMID: 19853566

Manni S, Maubhatjn HW, Bond M. 2008. Phosphorylation of the cAMP-dependent protein kinase (PKA) regulatory subunit modulates PKA-AKAP interaction, substrate phosphorylation, and calcium signaling in cardiac cells. Journal of Biological Chemistry 283:24145–24154. doi: 10.1074/jbc.M802278200, PMID: 18550536

Mayr B, Montminy M. 2001. Transcriptional regulation by the phosphorylation-dependent factor CREB. Nature Reviews Molecular Cell Biology 2:599–609. doi: 10.1038/35085068, PMID: 11483993

Meier JA, Lamer AC. 2014. Toward a new STATe: the role of STATs in mitochondrial function. Seminars in Immunology 26:20–28. doi: 10.1016/j.smim.2013.12.005, PMID: 24434063

Najahi-Missaoui W, Dailey HA. 2005. Production and characterization of erythropoietic protoporphyrin heterodimeric ferrochelatases. Blood 106:1098–1106. doi: 10.1182/blood-2004-12-4466, PMID: 15831704

Nilsson R, Schultz IJ, Pierce EL, Solts KA, Narangaratr A, Ward DM, Bawghman JM, Paradkar PN, Kingsley PD, Culotta VC, Kaplan J, Palis J, Paw BH, Mootha VK. 2009. Discovery of genes essential for heme biosynthesis through large-scale gene expression analysis. Cell Metabolism 10:119–130. doi: 10.1016/j.cmet.2009.06.012, PMID: 19653490

North TE, Goessling W, Walkley CR, Lengerke C, Kopani KR, Lord AM, Weber G, Bowman TW, Jang IH, Grosser T, Fitzgerald GA, Daley GQ, Orkin SH, Zon L. 2007. Prostaglandin E2 regulates vertebrate haematopoietic stem cell homeostasis. Nature 447:1007–1101. doi: 10.1038/nature05883, PMID: 17581586

Pagliarini DJ, Calvo SE, Chang B, Sheth SA, Vafai SB, Onge SE, Walford GA, Sugiana C, Boneh A, Chen WK, Hill DE, Vidal M, Evans JG, Thoburn DR, Carr SA, Mootha VK. 2008. A mitochondrial protein compendium elucidates complex I disease biology. Cell 134:112–123. doi: 10.1016/j.cell.2008.06.016, PMID: 18614015

Paradkar PN, Zumbrennen KB, Paw BH, Ward DM, Kaplan J. 2009. Regulation of mitochondrial iron import through differential turnover of mitoferrin 1 and mitoferrin 2. Molecular and Cellular Biology 29:1007–1016. doi: 10.1128/MCB.01685-08, PMID: 19705006

Park HJ, Li J, Hannah R, Biddie S, Leah-Cervantes AI, Kirschner K, Flores Santa Cruz D, Sexl V, Gottgens B, Green AR. 2016. Cytokine-induced megakaryocytic differentiation is regulated by genome-wide loss of a uSTAT transcriptional program. The EMBO Journal 35:580–594. doi: 10.15252/embj.201592383, PMID: 26702099

Phillips JD, Kushner JP. 2005. Fast track to the porphyrias. Nature Medicine 11:1049–1050. doi: 10.1038/nm1005-1049, PMID: 16211036

Repaport D. 2003. Finding the right organelle. targeting signals in mitochondrial outer-membrane proteins. EMBO Reports 4:948–952. doi: 10.1038/sj.embor.embor937, PMID: 14528265

Sakaino M, Ishigaki M, Ohgari Y, Kitajima S, Masaki R, Yamamoto A, Taketani S. 2009. Dual mitochondrial localization and different roles of the reversible reaction of mammalian ferrochelatase. FEBS Journal 276:5559–5570. doi: 10.1111/j.1742-4658.2009.07248.x, PMID: 19691493

Sarma GN, Kinderman FS, Kim C, von Daake S, Chen L, Wang BC, Taylor SS. 2010. Structure of D-AKAP2:pka RI complex: insights into AKAP specificity and selectivity. Structure 18:155–166. doi: 10.1016/j.str.2009.12.012, PMID: 20159461

Schneider CA, Rasband WS, Eliceiri KW. 2012. NIH image to ImageJ: 25 years of image analysis. Nature Methods 9:671–675. doi: 10.1038/nmeth.2089, PMID: 22930834

Schooley JC, Mahlimann J. 1975. Adenosine, AMP, cyclic AMP, theophylline and the action and production of erythropoietin. Experimental Biology and Medicine 150:215–219. doi: 10.3181/00377927-150-39005, PMID: 1716882

Severance S, Hamza I. 2009. Trafficking of heme and porphyrins in metazoa. Chemical Reviews 109:4596–4616. doi: 10.1021/cr9001116, PMID: 19764719

Shah Di, Takahashi-Makise N, Cooney JD, Li L, Schultz IJ, Pierce EL, Narla A, Seguin A, Hattangadi SM, Medlock AE, Langer NB, Dailey TA, Hurst SN, Faccenda D, Wiwczar JM, Heggars SK, Vogin G, Chen W, Chen C,
Developmental Biology and Stem Cells  |  Human Biology and Medicine

Campagn DR, et al. 2012. Mitochondrial Atp1f1 regulates haem synthesis in developing erythroblasts. Nature 491:608–612. doi: 10.1038/nature11536, PMID: 23135403

Shaw GC, Cope JJ, Li L, Corson K, Hersey C, Ackermann GE, Gwynn B, Lambert AJ, Wingert RA, Traver D, Trede NS, Barut BA, Zhou Y, Minet E, Donovan A, Brownle A, Balzan R, Weiss MJ, Peters LI, Kaplan J, et al. 2006. Mitofermin is essential for erythroid iron assimilation. Nature 440:96–100. doi: 10.1038/nature04512, PMID: 16511496

Shirai O, Gregory T, Yu C, Orkin SH, Weiss MJ. 2000. ABC-me: one mitochondrial transporter induced by GATA-1 during erythroid differentiation. The EMBO Journal 19:2492–2502. doi: 10.1093/emboj/19.11.2492, PMID: 10835348

Smith FD, Samelson BK, Scott JD. 2011. Discovery of cellular substrates for protein kinase A using a peptide array screening protocol. Biochemical Journal 438:103–110. doi: 10.1042/BJ20110720, PMID: 21644927

Socolovsky M, Fallon AE, Wang S, Brugnara C, Lodish HF. 1999. Fetal Anemia and apoptosis of red cell progenitors in Stat5a/−/−/−b/− mice: a direct role for Stat5 in Bcl-X(L) induction. Cell 98:181–191. doi: 10.1016/S0092-8674(00)81013-2, PMID: 10428030

Socolovsky M, Nam H, Fleming MD, Haase VH, Brugnara C, Lodish HF. 2001. Ineffective erythropoiesis in Stat5a (−/−)/b(b) mice due to decreased survival of early erythroblasts. Blood 98:3261–3273. doi: 10.1182/blood.v98.12.3261, PMID: 11719363

Stratakis CA, Cho-Chung YS. 2002. Protein kinase A and human disease. Trends in Endocrinology & Metabolism 13:50–52. doi: 10.1016/S1043-2760(01)00560-4, PMID: 11854015

Su AJ, Witshire T, Batalov S, Lapp H, Ching KA, Block D, Zhang J, Soden R, Hayakawa M, Kreiman G, Cooke MP, Walker JR, Hogben N. 2004. A gene atlas of the mouse and human protein-encoding transcriptomes. PNAS 101:6062–6067. doi: 10.1073/pnas.0400782101, PMID: 15075390

Taylor SS, Ilovitz R, Zhang P, Kornev AP. 2012. Assembly of allosteric macromolecular switches: lessons from PKA. Nature Reviews Molecular Cell Biology 13:646–656. doi: 10.1038/nrm3432, PMID: 22992589

Testa U. 2004. Apoptotic mechanisms in the control of erythropoiesis. Leukemia 18:1176–1199. doi: 10.1038/sj.leu.2403383, PMID: 15208642

Tingley WG, Pawlikowska L, Zaroff JG, Kim T, Nguyen T, Young SG, Vranizan K, Kwok PY, Woolley MA, Conklin BR. 2007. Gene-trapped mouse embryonic stem cell-derived cardiac myocytes and human genetics implicate AKAP10 in heart rhythm regulation. PNAS 104:8461–8466. doi: 10.1073/pnas.0610393104, PMID: 17485678

Vogel J, Kiessling I, Heinicke K, Stallmach T, Osent P, Vogel O, Aulmann M, Fritsch T, Schmid-Schonbein H, Kuschnisky W, Gassmann M. 2003. Transgenic mice overexpressing erythropoietin adapt to excessive erythrocytosis by regulating blood viscosity. Blood 102:2278–2284. doi: 10.1182/blood-2003-01-0283, PMID: 12750170

Wang L, Sunahara RK, Krumins A, Perkins G, Crochiere ML, Mackey M, Bell S, Ellisman MH, Taylor SS. 2001. Cloning and mitochondrial localization of full-length D-AKAP2, a protein kinase A anchoring protein. PNAS 98:3220–3225. doi: 10.1073/pnas.051633398, PMID: 11248059

Webb RJ, Tinworth L, Thomas GM, Zaccolo M, Carroll J. 2008. Developmentally acquired PKA localisation in mouse oocytes and embryos. Developmental Biology 317:36–45. doi: 10.1016/j.ydbio.2008.01.045, PMID: 18367160

Wegryn J, Potra R, Chwae YJ, Sepuri NB, Zhang Q, Koeck T, Derecka M, Szczepanek K, Szlag M, Gornicka A, Moh A, Moghaddas S, Chen Q, Bobbili S, Cichy J, Dulak J, Baker DP, Wolfman A, Stuehr D, Hassan MO, et al. 2009. Function of mitochondrial Stat3 in cellular respiration. Science 323:793–797. doi: 10.1126/science.1164551, PMID: 19131594

Wenger CD, Phanstiel DH, Lee MV, Bailey DJ, Coon JJ. 2011. COMPASS: a suite of pre- and post-search proteomics software tools for OMSsSA. PROTEOMICS 11:1064–1074. doi: 10.1002/pmic.201000616, PMID: 21298793

Whitley SD, Mason NG, Khan M, Zamiri M, Badminton MN, Missaoui WN, Dailey TA, Dailey HA, Douglas WS, Wainwright NJ, Elder GH. 2004. Autosomal recessive erythroprotopoietic poorphophaenia in the United Kingdom: prevalence and relationship to liver disease. Journal of Medical Genetics 41:e105. doi: 10.1136/jmg.2003.016121, PMID: 15286165

Wingert RA, Galloway JL, Barut B, Footh H, Fraenk P, Axel JL, Weber GJ, Dooley K, Davidson AJ, Schmidt B, Schmidt B, Paw BH, Shaw GC, Kingsley P, Banja JS, Schubert H, Chen O, Kaplan J, Zon LI, Tubingen 2000 Screen Consortium. 2005. Deficiency of glutaredoxin S mutants Fe-S clusters are required for vertebrate haem synthesis. Nature 436:1035–1039. doi: 10.1038/nature03887, PMID: 16110529

Wong W, Scott JD. 2004. AKAP signalling complexes: focal points in space and time. Nature Reviews Molecular Cell Biology 5:959–970. doi: 10.1038/nrm1527, PMID: 15573134

Wu CK, Dailey HA, Rose JP, Burden A, Sellers VM, Wang BC. 2001. The 2.0 A structure of human ferrochelatase, the terminal enzyme of heme biosynthesis. Nature Structural Biology 8:156–160. doi: 10.1038/48152, PMID: 11175906

Yang S, Fletcher WH, Johnson DA. 1995. Regulation of cAMP-dependent protein kinase: enzyme activation without dissociation. Biochemistry 34:6267–6271. doi: 10.1021/bi00019a002, PMID: 7756252

Yang H, Li G, Wu JJ, Wang L, Uhler M, Simeone DM. 2013. Protein kinase A modulates transforming growth factor-β signaling through a direct interaction with Smad4 protein. Journal of Biological Chemistry 288:8737–8749. doi: 10.1074/jbc.M113.485675, PMID: 23362881

Yien YY, Robledo RF, Schultz IJ, Takahashi-Makise N, Gwynn B, Bauer DE, Dass A, Yu G, Li L, Hillidick-Smith GJ, Cooney JD, Pierce EL, Mohler K, Dailey TA, Miyata N, Kingsley PD, Garone C, Hattangi SM, Huang H, Chen...
W, et al. 2014. TMEM14C is required for erythroid mitochondrial heme metabolism. *Journal of Clinical Investigation* 124:4294–4304. doi: 10.1172/JCI76979, PMID: 25157825

Zakhary DR, Moravec CS, Bond M. 2000. Regulation of PKA binding to AKAPs in the heart: alterations in human heart failure. *Circulation* 101:1459–1464. doi: 10.1161/01.CIR.101.12.1459, PMID: 10736293

Zhang J, Socolovsky M, Gross AW, Lodish HF. 2003. Role of ras signaling in erythroid differentiation of mouse fetal liver cells: functional analysis by a flow cytometry-based novel culture system. *Blood* 102:3938–3946. doi: 10.1182/blood-2003-05-1479, PMID: 12907435

Zhu BM, McLoughlin SK, Na R, Liu J, Cui Y, Martin C, Kimura A, Robinson GW, Andrews NC, Hennighausen L. 2008. Hematopoietic-specific Stat5-null mice display microcytic hypochromic Anemia associated with reduced transferrin receptor gene expression. *Blood* 112:2071–2080. doi: 10.1182/blood-2007-12-127480, PMID: 18552213

Zon LI, Peterson RT. 2005. In vivo drug discovery in the zebrafish. *Nature Reviews Drug Discovery* 4:35–44. doi: 10.1038/nrd1606, PMID: 15688071