Neonatal expression of RNA-binding protein IGF2BP3 regulates the human fetal-adult megakaryocyte transition

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Hematopoietic transitions that accompany fetal development, such as erythroid globin chain switching, play important roles in normal physiology and disease development. In the megakaryocyte lineage, human fetal progenitors do not execute the adult morphogenesis program of enlargement, polyploidization, and proplatelet formation. Although these defects decline with gestational stage, they remain sufficiently severe at birth to predispose newborns to thrombocytopenia. These defects may also contribute to inferior platelet recovery after cord blood stem cell transplantation and may underlie inefficient platelet production by megakaryocytes derived from pluripotent stem cells. In this study, comparison of neonatal versus adult human progenitors has identified a blockade in the specialized positive transcription elongation factor b (P-TEFb) activation mechanism that is known to drive adult megakaryocyte morphogenesis. This blockade resulted from neonatal-specific expression of an oncofetal RNA-binding protein, IGF2BP3, which prevented the destabilization of the nuclear RNA 7SK, a process normally associated with adult megakaryocytic P-TEFb activation. Knockdown of IGF2BP3 sufficed to confer both phenotypic and molecular features of adult-type cells on neonatal megakaryocytes. Pharmacologic inhibition of IGF2BP3 expression via bromodomain and extraterminal domain (BET) inhibition also elicited adult features in neonatal megakaryocytes. These results identify IGF2BP3 as a human ontogenic master switch that restricts megakaryocyte development by modulating a lineage-specific P-TEFb activation mechanism, revealing potential strategies toward enhancing platelet production.

Introduction

During fetal development, i.e., ontogeny, mammalian hematopoiesis goes through multiple phases of reconfiguration. This reconfiguration encompasses both lineage output and phenotypic features within lineages. The lineage most dramatically affected by ontogenic stage consists of human megakaryocytes, long known to display distinct morphologies in the fetus and adult. In situ analysis of primary tissue has shown fetal megakaryocytes to be smaller and less polyploid than adult counterparts (1, 2). Size differences are even discernible between children less than 2 years of age and those more than 4 years of age, indicating postnatal persistence of ontogenic influence (3). Ontogenic stage also affects platelet function, with moderate to marked hyporesponsiveness observed in full-term and premature neonates, respectively (4, 5). These infantile properties have clinical significance, as they underlie the common clinical problems of neonatal thrombocytopenia and hemorrhage (6).

The molecular basis for ontogenic regulation of megakaryopoiesis remains unknown but consists of a cell-intrinsic mechanism initiating within multipotent progenitors. Thus, transplant recipients of neonatal, cord blood–derived hematopoietic stem and progenitor cells (CB HSPCs) have smaller megakaryocytes and slower platelet recovery compared with age-matched recipients of adult HSPCs, despite having equal megakaryocyte numbers (7). In ex vivo HSPC cultures, CB megakaryocytes show a greater than 10-fold enhancement in proliferation and markedly diminished morphogenesis compared with adult counterparts (8). The morphogenetic impairment correlates directly with diminished capacity for platelet release (9). Fetal liver (FL) HSPCs and embryonic stem cell progenitors have even more limited morphogenetic potential than CB HSPCs (10), indicating a graded influence of ontogenic stage. Importantly, the ontogenic influence on megakaryopoiesis, while affecting morphogenesis and proliferation, does not affect all aspects of the megakaryocyte differentiation program (8).
Recently, megakaryocyte morphogenesis was found to depend on sustained, high-amplitude activation of the positive transcription elongation factor b (P-TEFb) kinase complex, consisting of CDK9 and cyclin T (11, 12). P-TEFb promotes RNA polymerase II (RNAPII) elongation on stimulus-responsive genes through phosphorylation of pausing factors (5,6-dichloro-1-β-D-ribofuranosylbenzimidazole [DRB] sensitivity–inducing factor [DSIF] and negative elongation factor [NELF]) and the carboxy terminal domain of RNAPII large subunit (RNAPII CTD) (13). In most cells, a feedback loop maintains the majority of P-TEFb sequestered in an inactive state within the 7SK small nuclear ribonucleoprotein (snRNP) complex, which contains the kinase repressor hexamethylene bisacetamide inducible 1 (HEXIM1) scaffolded by the small nuclear RNA (snRNA) 7SK. This complex also contains the 7SK-stabilizing proteins methylphosphate capping enzyme (MePCE) and La ribonucleoprotein domain family member 7 (LARP7). Conventional activation occurs through stimulus-induced release of P-TEFb from the 7SK snRNP, promoting HEXIM1 transcription, which ultimately drives resequestration of P-TEFb (13). The megakaryocytic differentiation program, in contrast, employs a specialized activation pathway initiated by MePCE and LARP7 downregulation, leading to 7SK destruction and global, irreversible P-TEFb release (11). This mode of activation upregulates a cohort of cytoskeletal remodeling factors, such as megakaryoblastic leukemia (translocation) 1 (MLKL), filamin A (FLNA), and α-actinin 1 (ACTN1), which function as morphogenesis effectors during differentiation (11, 14, 15).

The current studies address whether the diminished morphogenesis of fetal megakaryocytes reflected impairment in this specialized program of P-TEFb activation. Supporting this concept, CB megakaryocytes failed to upregulate P-TEFb–dependent cytoskeletal remodeling factors and demonstrated deficiency in P-TEFb activation. Despite lineage-appropriate downregulation of the 7SK stabilizing factors MePCE and LARP7, 7SK snRNA levels remained paradoxically elevated in CB megakaryocytes. These findings suggested the existence of a fetal-specific 7SK-stabilizing protein. Functional screening of candidate factors identified IGF2BP3 as a fetal-specific, 7SK-associated factor whose ectopic expression in adult HSPCs elicited fetal features. Knockdown of IGF2BP3 in CB HSPCs enabled adult-type megakaryopoiesis both phenotypically and in terms of P-TEFb activation, supporting a role as an ontogenic master switch within this lineage. Known regulation of IGF2BP3 by pluripotency circuits prompted a strategy to target its expression through bromodomain and extraterminal domain (BET) inhibition. This approach successfully downmodulated IGF2BP3 in CB megakaryocytes and elicited adult patterns of differentiation and P-TEFb activation, indicating accessibility of this switch to exogenous manipulation.

Results

Neonatal megakaryocytes display decreased morphogenesis, enhanced proliferation, and incomplete erythroid silencing. To analyze molecular mechanisms involved in ontogenic regulation of megakaryopoiesis, we used unilineage culture of purified CD34+ HSPCs derived from normal neonatal umbilical CB or adult peripheral blood (PB). Gating strategies for the flow cytometric studies are illustrated in Supplemental Figure 1A (supplemental material available online with this article; https://doi.org/10.1172/JCI88936DS1). This system reliably modeled decreased enlargement (forward scatter [FSC] in Figure 1A), enhanced proliferation (dilution of PKH dye in Figure 1B), and decreased polyploidization (propidium iodide [PI] peaks in Figure 1C) of neonatal/fetal CD41+ megakaryocytes. Control studies showed no differences in initial PKH loading between sample types (Supplemental Figure 1B).

Woo et al. previously generated transcriptional profiles comparing fetal and adult murine megakaryocyte progenitors; in the top 122 fetal-upregulated transcripts, 7 erythroid genes were represented (16). Their results suggest an ontogenic influence on
HSPCs consistently produced subpopulations of CD41+GPA+ and CD41−GPA+ cells (Figure 1D), consistent with a diminished capacity for lineage consolidation.

Neonatal progenitors fail to execute the megakaryocytic P-TEFb activation pathway. In adult megakaryopoiesis, constitutive P-TEFb activation drives upregulation of a cohort of cytoskeletal remodeling factors that function as critical effectors in morphogenesis (11). In neonatal megakaryopoiesis, in contrast, minimal lineage consolidation of megakaryocytes emerging from bipotent megakaryocyte-erythroid progenitors, a notion further supported by the hybrid CD41+ glycophorin A+ (GPA) cells observed in pluripotent stem cell (PSC) hematopoiesis (17). We therefore determined whether human neonatal CB HSPCs manifested impairment in megakaryocytic lineage consolidation. In megakaryocytic unilineage medium, which lacks erythropoietin, adult PB HSPCs yielded essentially no erythroid GPA+ cells, while CB HSPCs consistently produced subpopulations of CD41+GPA+ and CD41 GPA+ cells (Figure 1D), consistent with a diminished capacity for lineage consolidation.

Neonatal progenitors fail to execute the megakaryocytic P-TEFb activation pathway. (A) Neonatal failure to upregulate megakaryocytic P-TEFb targets. Left panel: primary adult and neonatal progenitors, either undifferentiated (Un) or cultured 6 days in erythroid (Ery) or megakaryocytic (Mk) medium were immunoblotted for indicated factors. Top arrow, full-length FLNA; bottom arrow, approximately 190-kDa cleavage fragment. Right panel: densitometry comparing adult and neonatal megakaryocyte levels from 3 independent experiments conducted as in the right panel. Graphs show mean ± SEM for signals normalized to tubulin, with PB values set at 1. *P < 0.05; **P < 0.01; ***P < 0.005, t test. Rel, relative. (B–D) Evidence for diminished P-TEFb activation in neonatal versus adult megakaryocytes. Adult and neonatal progenitors cultured 6 days in megakaryocyte medium underwent IB for RNAPII subunit RPB1 (B), SPT5 phospho-threonine 806 (pT806) and total (SPT5) (C), and histone H2Bub1 and total H2B (D). I10 and IIA designate hyper- and hypophosphorylated forms for RNAPII, respectively. Graphs show mean ± SEM for scanning densitometry values derived from 3 independent experiments. *P < 0.05; **P < 0.01, t test. (E) Neonatal megakaryocytes downregulate 7SK-stabilizing factors to a degree similar to that of adult megakaryocytes. (F) Neonatal block in megakaryocytic downregulation of 7SK snRNA. 7SK levels relative to CB megakaryocytes in progenitors cultured as in A. Graphs show mean ± SEM of 7SK normalized to GAPDH in 3 independent experiments. *P < 0.05, t test. Note that panels A and E as well as Figure 3A and Supplemental Figure 4A all derive from the same IB membrane and therefore share the same tubulin control. Note that panel D and Supplemental Figure 4B derive from the same IB membrane and share the same tubulin control. See also Supplemental Figures 2 and 3.
functions as a master regulator of megakaryocyte morphogenesis. MKL1 markedly diminished expression in neonatal versus adult megakaryocytes cultured ex vivo. For this reason, and because MKL1 is a direct and specific P-TEFb substrate, SPT5 (18), showed 2.5-fold diminished phosphorylation in the neonatal versus adult megakaryocytes (Figure 2C). An epigenetic mark downstream of P-TEFb activity consists of monoubiquitination of histone H2B on K120, H2Bub1 (24, 25). K120 is the amino acid position on the CTD phosphorylation, is best reflected by the relative abundance of the more slowly migrating II0 isoform (18, 19). RNAPII phosphorylation on S2 does not provide a reliable readout, as CDK9 phosphorylates multiple positions on the CTD and multiple kinases phosphorylate the S2 position (20–23). Our studies consistently revealed a marked difference between neonatal and adult megakaryocytes in RNAPII isoform distribution (Figure 2B). In addition, a direct and specific P-TEFb substrate, SPT5 (18), showed 2.5-fold diminished phosphorylation in the neonatal versus adult megakaryocytes (Figure 2C). An epigenetic mark downstream of P-TEFb activity consists of monoubiquitination of histone H2B on K120, H2Bub1 (24, 25). K120 is the amino acid position on the protein H2B which is modified by attachment of mono-ubiquitin to K120, H2Bub1 (24, 25). K120 is the amino acid position on the protein H2B which is modified by attachment of mono-ubiquitin to yield the modified protein H2Bub1. Notably, neonatal megakaryocytes displayed global deficiency in H2Bub1 as compared with adult cells (Figure 2D). Thus, assessment of multiple independent targets confirmed an ontogenically determined block in megakaryocytic P-TEFb activation.

As illustrated in Figure 2A, the P-TEFb target MKL1 showed markedly diminished expression in neonatal versus adult megakaryocytes cultured ex vivo. For this reason, and because MKL1 functions as a master regulator of megakaryocyte morphogenesis (14), we further assessed its expression in human marrow biopsies by immunohistochemistry. Within the limited samples available, megakaryocytes from 2 adult stem cell transplant recipients showed strong staining for MKL1, while megakaryocytes from 3 CB stem cell transplant recipients showed negative to weak staining (Supplemental Table 1 and Supplemental Figure 2, A–D). In addition, megakaryocytes in marrow samples from 3 neonates (2 to 9 weeks of age) lacked detectable MKL1 (Supplemental Table 1). Neonatal megakaryocytes downregulate 7SK-stabilizing factors, but fail to downregulate 7SK. The specialized pathway of P-TEFb activation in adult megakaryocytes involves destabilization of 7SK snRNA through downregulation of its binding partners, the stabilizing factors LARP7 and MePCE (11). In this process, MePCE undergoes proteolysis due to megakaryocytic upregulation of calpain 2, which also cleaves the cytoskeletal remodeling factor FLNA. The appearance of FLNA cleavage fragments in both neonatal and adult megakaryocytes suggested intact calpain 2 regulation and function in the neonatal cells (Figure 2A). Indeed, both of the 7SK-stabilizing factors underwent full downregulation during neonatal megakaryopoiesis (Figure 2E and Supplemental Figure 3A). Numerous studies have documented tight coupling of 7SK snRNA levels with its stabilizing factors (26–30). However, the stabilizing factors’ decline in neonatal megakaryocytes occurred in the absence of a proportionate drop in 7SK snRNA levels, which remained 2.5-fold higher than adult levels (Figure 2F).
An ontogenically regulated RNA-binding factor confers the neonatal phenotype. The relative stability of 7SK in neonatal megakaryocytes raised the possibility that fetal-specific RNA-binding factors might provide stabilizing functions in the absence of LARP7 and MePCE. To address this hypothesis, candidate factors underwent assessment first for ontogenic regulation and then for phenotypic influence on adult megakaryocytes. Candidates consisted of 7SK-associated or general RNA-binding factors that displayed upregulation in CB versus adult marrow HSPCs, based on available gene-expression profiles (31). One of the most compelling candidates consisted of IGF2BP3, a known oncofetal RNA-binding factor that participates in an interactome network containing 7SK-binding proteins (32, 33). IGF2BP3 protein levels markedly differed according to ontogenic stage, with highly abundant levels in neonatal hematopoietic cells, including megakaryocytes, and completely absent levels in adult counterparts (Figure 3A). An antibody recognizing both IGF2BP1 and IGF2BP2 showed their expression in both neonatal and adult megakaryocytes, although with differences in isoform patterns (Supplemental Figure 4A). Fetal-selective protein expression in megakaryocytes was also observed for the candidates LIN28B and HMGAI (Supplemental Figure 4, B and C).

Ectopic expression in adult megakaryocytes of IGF2BP3, but not of LIN28B or HMGAI, blocked cellular enlargement (Figure 3B and Supplemental Figure 4D). Furthermore, IGF2BP3 expression caused adult megakaryocytes to undergo additional rounds of cell division, as reflected by enhanced PKH dye dilution, and promoted expression of the erythroid marker GPA (Figure 3, C and D; see Supplemental Figure 4E for initial PKH dye loading). These effects were specific for adult progenitors, as similar transduction of neonatal progenitors minimally altered their megakaryocytic phenotype (Supplemental Figure 5). Thus, IGF2BP3 fulfills multiple criteria for a megakaryocytic ontogenic regulator.

To confirm these findings using an in vivo model system, IGF2BP3 expression was enforced in adult murine marrow by retroviral transduction followed by transplantation. Prior studies in mice have shown that IGF2BP3 transcript levels are much higher (4- to 16-fold) in fetal versus adult hematopoietic stem cells and megakaryocytic progenitors (MkPs) (16, 34). Animals receiving IGF2BP3-transduced marrow retained transgene expression at 6 weeks after transplant. The chimeras were studied under steady-state conditions and after stimulation of megakaryopoiesis with thrombopoietin (TPO). Under both conditions, enforcement of IGF2BP3 interfered with megakaryocytic enlargement, reflected by diminished FSC in gated GFP±CD41+ cells from marrow and spleen (Figure 4, A and B, and Supplemental Figure 6A). In addition, ectopic IGF2BP3 increased the marrow frequency of CD41+ cells and of CD41+Ter119+ double-positive cells (Figures 4C and Supplemental Figure 6B). These latter findings were consistent with the enhanced megakaryocytic proliferation and erythroid antigen retention seen with ectopic IGF2BP3 in adult human progenitors (see Figure 3, C and D).

IGF2BP3 participates in 7SK snRNP and contributes to 7SK stabilization. Prior studies have shown a capacity for IGF2BP3, but not IGF2BP1 or IGF2BP2, to undergo nuclear localization (35).
studies, neonatal megakaryocytes displayed both nuclear and cytoplasmic IGF2BP3 localization, as assessed by confocal immunofluorescence microscopy and subcellular fractionation (Supplemental Figure 7, A and B). Association with the 7SK snRNP was supported by endogenous co-IP of IGF2BP3 and HEXIM1 (Figure 5A and Supplemental Figure 7C). In addition, 7SK snRNA coprecipitated with epitope-tagged IGF2BP3 in 293 transfectants (Figure 5B). This experiment employed ectopic expression of epitope-tagged IGF2BP3 in 293 cells because of the lack of available antibodies suitable for native IP and because of the minimal endogenous IGF2BP3 in 293 cells. Mining of iCLIP-seq data examining endogenous IGF2BP3 interactions (36) revealed 7SK to be one of the top targets (7th out of 2185), far ahead of the canonical targets MYC and CD44 (Figure 5C). Notably, the iCLIP-seq data mapped IGF2BP3 binding to 7SK hairpin 4 (HP4), adjacent to a stabilizing binding site for LARP7 (37) (Figure 5D).

To further assess effects on the 7SK snRNP, 293 transfectants underwent biophysical characterization of LARP7, a dedicated component of this complex (29). Enforced IGF2BP3 expression redistributed LARP7 into high–molecular weight complexes by glycerol gradient analysis (Figure 5E) and out of the high-salt–extractable nuclear fraction (Supplemental Figure 7D). To determine whether IGF2BP3 influences 7SK snRNA stability, neonatal progenitors underwent lentiviral shRNA knockdown with puromycin selection and megakaryocytic differentiation of transduced cells. This approach yielded isoform-selective repression of IGF2BP3 (Supplemental Figure 7E) and consistently diminished its levels by approximately 80% (see summary graph in Figure 6E). Notably, the knockdown of IGF2BP3 caused significant 7SK downregulation during neonatal megakaryopoiesis (Figure 5F). These results support the notion that IGF2BP3 participates in fetal/neonatal 7SK snRNP complexes and may thereby enhance 7SK stability in neonatal megakaryocytes.

IGF2BP3 contributes to the phenotypic and molecular features of neonatal megakaryocytes. As isoform-selective knockdown of IGF2BP3 promoted 7SK downregulation in neonatal megakaryocytes, additional studies examined the phenotypic consequences of this manipulation. As anticipated, decreasing the levels of IGF2BP3 conferred several adult-type features, such as enhanced cellular enlargement, growth arrest, polyploidization, and eryth-
roid suppression (Figure 6, A–C, and Supplemental Figure 8, A and B). At the molecular level, this knockdown enhanced P-TEFb activation, as reflected by SPT5 phosphorylation as well as levels of HEXIM1, RNAPII0, and H2Bub1 (Figure 6, D and E, and Supplemental Figure 8C). This activation was further associated with upregulation of the P-TEFb–dependent cytoskeletal remodeling factors, particularly the morphogenetic regulator MKL1 (Figure 6E). As a specificity control, adult megakaryocytes transduced with the same lentiviral shRNA constructs underwent comprehensive phenotypic and molecular characterization. In these cells, the IGF2BP3-targeting shRNA caused no significant alterations in any parameters (Supplemental Figure 9), confirming a neonatal-specific effect. These loss-of-function studies thus reveal an influence of IGF2BP3 levels on P-TEFb activation and megakaryocytic morphogenesis, consistent with a role as a critical ontogenic determinant.

BET factor regulation of IGF2BP3 enables pharmacologic modulation of ontogenic influence

Recently published data demonstrate induction of IGF2BP3 by a cancer stem cell pluripotency program (40). In addition, a factor or factors in the BET family have been shown to participate in pluripotency reprogramming of somatic cells (41), suggesting potential BET regulation of IGF2BP3. For these reasons, effects of BET inhibition on neonatal megakaryopoiesis were tested. As shown in Figure 7, 2 different inhibitors, JQ1 and I-BET151, both downregulated the erythroid marker GPA while enhancing expression of megakaryocytic markers CD41 and CD42 (Figure 7, A and B, and Supplemental Figure 11A). These treatments also promoted platelet release, cellular enlargement, and growth arrest (Figure 7, C and D, and Supplemental Figure 11B). At the molecular level, BET inhibition suppressed IGF2GP3 expression while inducing RNAPII hyperphosphorylation, upregulation of cytoskeletal remodeling factors, and HEXIM1 upregulation (Figure 8, A and B). MKL1 and FLNA levels showed particularly robust responsiveness to BET inhibition. In control experiments, treatment of adult progenitors with BET inhibitors failed to upregulate the cytoskeletal remodeling factors or HEXIM1, failed to induce RNAPII hyperphosphorylation, and did not induce cellular enlargement (Supplemental Figure 11, C–E). During the preparation of this manuscript, Palanichamy et al. also showed repression of IGF2BP3 expression by BET inhibition (36).

To further ascertain that the effects of BET inhibition on neonatal megakaryopoiesis resulted from repressing IGF2BP3 expression, retroviral transduction was used to enforce IGF2BP3 expression in neonatal progenitors. Enforced IGF2BP3 significantly blunted the capacity of BET inhibitors to induce enlargement of neonatal megakaryocytes (Figure 9A). Exogenous IGF2BP3 expression basally
upregulated GPA and rendered both GPA and CD41 less responsive to BET inhibition (Figure 9, B and C). These findings confirm that some of the phenotypic effects of BET inhibitors on neonatal megakaryocytes result from downregulation of IGF2BP3.

**Discussion**

It has long been appreciated that the ontogenic stage exerts a profound influence on human megakaryocyte morphogenesis. In addition to the clinical implications for neonatal infants and CB HSC transplant recipients, this influence affects the feasibility of ex vivo, donor-independent platelet production. Due to their capacity for unlimited expansion and genetic manipulation, PSCs have become a leading candidate for ex vivo megakaryocyte generation. However, PSC-derived megakaryocytes display poor platelet yields (42), and extensive profiling has shown these cells to be even more ontogenically primitive than human FL megakaryocytes, likely reflecting a yolk-sac stage (10). Thus, ontogenic manipulation has the potential to improve both yield and quality of ex vivo–derived platelets.

While a megakaryocyte ontogenic regulatory mechanism has not previously been identified, contributory factors have been described. Klussmann et al. identified fetal-specific hyperactivation of the IGF/AKT/mTOR pathway, which primed MkPs for leukemic transformation (43). These findings were supported by a subsequent study in which the proliferative phenotype of neonatal megakaryocytes was linked to mTOR hyperactivation (8). Impor...
Importantly, the ex vivo differentiation of murine FL MkPs is associated with reproducible downregulation of \textit{Igf2bp3} (see GEO GDS1316 from ref. 44) in contrast with cultured human neonatal progenitors, which show no change in \textit{IGF2BP3} during maturation of committed MkPs (see BloodSpot, DMAP data set; ref. 45). Thus there may be differences in the regulation of the ontogenic program in murine versus human hematopoiesis, with murine progenitors possessing more plasticity in responding to milieu influences.

The current results identify IGF2BP3 as an ontogenic master switch that regulates megakaryocyte morphogenesis, at least in part, through modulation of a specialized P-TEFb activation program (see Figure 8C). Among the notable targets in this pathway is MKL1, a factor previously implicated as a key regulator of

tantly, this latter study demonstrated confinement of the neonatal phenotype to morphogenesis and proliferation, with no alterations in other aspects of megakaryocyte differentiation, including expression of platelet-surface glycoproteins (e.g., CD42), granule maturation, transcription factor expression, and TPO receptor function. The known participation of IGF2BP3 in inducing IGF/ AKT/mTOR signaling during oncogenesis (40) suggests that its abundant levels in neonatal megakaryocytes (see Figure 3A) may contribute to mTOR hyperactivation.

Comparisons of murine FL and adult marrow MkPs have also identified a microenvironmental influence, consisting in part of increased type I interferon in the adult milieu (16). Interestingly, murine FL MkPs, when cultured ex vivo, acquire the capacity for adult-type morphogenesis (44), unlike findings with human progenitors (10). Importantly, the ex vivo differentiation of murine FL MkPs is associated with reproducible downregulation of \textit{Igf2bp3} (see GEO GDS1316 from ref. 44) in contrast with cultured human neonatal progenitors, which show no change in \textit{IGF2BP3} during maturation of committed MkPs (see BloodSpot, DMAP data set; ref. 45). Thus there may be differences in the regulation of the ontogenic program in murine versus human hematopoiesis, with murine progenitors possessing more plasticity in responding to milieu influences.

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release factors that locally liberate CDK9/cyclin T, and the super elongation complex (SEC), which delivers the kinase to a recipient complex including RNAPII, polymerase-associated factor (PAF), and DSIF (51). Compared with other hematopoietic lineages, megakaryocytes appear to be deficient in some components of the SEC, PAF, and DSIF complexes (ref. 45 and unpublished data, A.N. Goldfarb laboratory). They may therefore require higher thresholds and alternative pathways of P-TEFb release to insure sufficient delivery to those genes most susceptible to RNAPII pausing. Thus lineage- and gene-specific mechanisms most likely converge to translate specific thresholds of free P-TEFb into a coordinated transcriptional output.

The upregulation of P-TEFb–dependent factors by BET inhibition appears paradoxical, given the role of bromodomain-containing 4 (BRD4) in the recruitment of P-TEFb to target genes. However, 2 mechanisms may contribute to this phenomenon. First, the threshold for IGF2BP3 repression by BET inhibitors may fall below the threshold for repression of megakaryocytic genes, thus providing a window for indirect activation. Second, BRD4 may not be required or may play a negative role in the activation of some megakaryocytic P-TEFb target genes, as has been described for HIV transcription (52). The results shown in Figure 9 indicate that several of the promegakaryocytic effects of BET inhibition occur through IGF2BP3.

Figure 9. Phenotypic effects of BET factor inhibition occur via IGF2BP3 repression. (A–C) Enforced expression of IGF2BP3 blunts the effects of BET factor inhibition in neonatal megakaryocytes. CB progenitors retrovirally transduced with control or IGF2BP3 expression vectors underwent 6 days of culture in megakaryocyte medium ± BET inhibitors as in Figure 7A. Cells were analyzed by flow cytometry for size (FSC) within the viable GFP+CD41+ gated population (A). In addition, viable GFP+ cells were analyzed for expression of the erythroid marker GPA and the megakaryocytic marker CD41 (B and C). Graphs in A–C represent mean ± SEM for 4 independent experiments. Analysis between the 2 groups employed 2-way ANOVA. Asterisks above the long line indicate row factor x column factor interactions. *P < 0.05; **P < 0.01; ***P < 0.005.

megakaryocyte morphogenesis. MKL1 undergoes upregulation, nuclear translocation, and transcriptional activation during mega-karyopoiesis (46). MKL1-deficient murine megakaryocytes display defects in polyploidization, enlargement, proliferation control, and proplatelet production (14, 47, 48), thus phenocopying normal human neonatal megakaryocytes. Gene-expression profiles indicate that MKL1-regulated genes constitute a specific, small subset of megakaryocytic genes, encoding cytoskeletal remodeling factors and the ploidy regulator GEF-H1 (14, 15, 48). Two MKL1-regulated factors, FLNA and HIC-5, bind to MKL1, promote its activation, and thus induce feed-forward loops (49, 50). Notably, our results show that human neonatal megakaryocytes have significant deficiencies of MKL1, FLNA, and HIC-5 and augment these factors in response to IGF2BP3 knockdown (see Figure 5E). MKL1 and FLNA also show striking upregulation in neonatal megakaryocytes in response to BET inhibition (Figure 6D). Therefore, MKL1 likely represents a proximal mediator in the ontogenic regulation of morphogenesis by IGF2BP3–P-TEFb.

How changes in the levels of P-TEFb liberation can exert specific effects on the megakaryocytic morphogenesis program remains unknown. In many cell types, developmentally regulated genes undergo activation through a guided recruitment of active P-TEFb. This process is mediated by a series of steps involving chromatin adaptor factors that recruit the 7SK snRNP, P-TEFb...
repression. However, blockade of BET factor recruitment to acetylated chromatin also disrupts mediator recruitment to super enhancers, repressing hundreds of associated genes (53). Thus, complicated network perturbations and off-target effects are expected with this particular intervention.

The current results elucidate a transcriptional mechanism underlying the ontogenic regulation of megakaryopoiesis. Because IGF2BP3 is broadly expressed during embryogenesis (54), its stabilizing influence on the 7SK snRNP could potentially alter P-TEFB regulation in a variety of developing fetal tissues and provide a general principle for ontogenic regulation. However, its effects are expected to be most prominently manifested in cells that normally downregulate the 7SK-stabilizing factors LARP7 and MePCE, e.g., megakaryocytes. The role of this regulatory pathway does not exclude additional mechanisms for IGF2BP3 in ontogenic regulation, nor does it exclude contributions from unrelated pathways. Our results do show though that manipulation of this pathway offers a promising strategy for enhancing megakaryocyte platelet production capacity.

Methods

**Cell culture.** Purified human adult CD34+ PB cells were purchased from the Fred Hutchinson Cancer Research Center (Seattle, Washington, USA), and purified human umbilical CD34+ CB cells were purchased from AllCells. The cells were cultured in serum-free unlineage media as described previously (11). Briefly, undifferentiated cells were expanded for 72 hours in prestimulation medium containing Iscove modified Dulbecco medium with 20% BBS 9500 (StemCell Technologies), 2 mM l-glutamine, 100 ng/ml stem cell factor (SCF), 100 ng/ml FLT3-ligand (FLT3L), 100 ng/ml TPO, and 20 ng/ml IL-3. Erythroid and megakaryocytic cells were cultured in respective media as described previously (11). For human progenitor transduction, MIG vector and MIG-IGF2BP3 were packaged in HEK293T cells as described (11). For subcloning into the MIG retroviral expression construct pDESTmycIGF2BP3, which expresses an standardized culture (11, 12).

**Transduction experiments enforcing IGF2BP3 expression, analyses** were conducted on viable GFP+ gated cells. Assessment of platelet release was conducted as previously described (38, 39). Studies on transplanted mice were conducted 6 weeks after transplant. Marrow and spleens were disaggregated by passing through 70-μm mesh and subjected to lysis of red cells using hypotonic ammonium chloride (Gibco ACK lysis buffer, Thermo Fisher Scientific). Samples were obtained with APC-conjugated rat anti-CD41 and PE-conjugated rat anti-TER119 or isotype controls (BD Biosciences) and analyzed on a BD LSRFortessa with initial gating on viable GFP+ cells.

**Flow cytometry.** Expression of CD41 and GPA, ploidy analysis, and PKH26 dye dilution assays in human progenitors were performed as described previously (11, 12). For the human progenitor retroviral transduction experiments enforcing IGF2BP3 expression, analyses were conducted on viable GFP+ gated cells. Assessment of platelet release was conducted as previously described (38, 39). Studies on transplanted mice were conducted 6 weeks after transplant. Marrow and spleens were disaggregated by passing through 70-μm mesh and subjected to lysis of red cells using hypotonic ammonium chloride (Gibco ACK lysis buffer, Thermo Fisher Scientific). Samples were obtained with APC-conjugated rat anti-CD41 and PE-conjugated rat anti-TER119 or isotype controls (BD Biosciences) and analyzed on a BD LSRFortessa with initial gating on viable GFP+ cells.

**IP and immunoblot.** IP of HEXIM1 from K562 cellular extracts followed by immunoblot (IB) analysis was performed as described (11, 12). The RNA-binding protein IP assay (RIP) was performed according to the protocol described by Jain et al. (58) with some modifications. Briefly, HEK293T cells transiently transfected with pDESTmycIGF2BP3 were incubated 5 minutes in ice-cold polysome lysis buffer (10 mM Hepes, pH 7.0, 100 mM KCl, 5 mM MgCl2, 0.5% Nonidet P-40 [NP-40], 1 mM DTT) supplemented with EDTA-free protease inhibitor cocktail (Roche), 20 μM calpain inhibitor III, and 40 U/ml RNase inhibitor (Promega) and then stored at −80°C. Rapidly thawed extracts were cleared by centrifugation at 16,000 g at 4°C for 10 minutes followed by dilution with NET-2 buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 mM MgCl2, 0.05% NP-40, 20 mM EDTA, pH 8.0, 1 mM DTT) supplemented with protease and RNase inhibitors as above. For IP, diluted extracts were incubated overnight at 4°C with anti-Myc-tag or isotope-matched control antibody, followed by capture with protein A/G magnetic beads and multiple washes with buffer NT-2 (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 1 mM MgCl2, 0.05% NP-40). One portion of the beads was boiled in 1× SDS-PAGE loading buffer for IB assessment of IP efficiency. The other portion was processed for RNA analysis by treatment first with DNase, then addition of 10% Chel-ex-100 in water, which was boiled for 10 minutes, followed by addi-
tion of proteinase K with digestion at 55°C for 30 minutes. Direct IB analysis of sheared whole cell lysates was conducted as described (12). Densitometry data were acquired on a GS800 calibrated densitometer (Bio-Rad) and analyzed with Quantity One software (Bio-Rad).

**RNA quantitation.** For quantitation of 7SK snRNA, total cellular RNA isolated using the RNeasy Plus Mini Kit (QiAGEN) or RNA from RFP was converted to cDNA using the iScript kit (Bio-Rad) and analyzed by quantitative PCR (qPCR) on the iCycler platform using iQ SYBR Green Supermix (Bio-Rad). Primers and cycling conditions were described previously (11). Relative transcript levels were calculated using GAPDH normalization with the ΔΔCt formula.

**Glyceral gradient analysis.** Transfected HEK293T cells were harvested and resuspended in ice-cold extraction buffer (10 mM HEPES, 150 mM NaCl, 2 mM MgCl2, 0.5% NP-40, 1 mM DTT, 1 mM PMSE, 20 μM calpain inhibitor III, EDTA-free protease inhibitor cocktail, 40 U/ml RNase inhibitor). Extracts were loaded on 10%-45% glycerol gradients prepared in extraction buffer. The gradients were spun on an SW-41 rotor at 40,000 g for 16 hours at 4°C, and resultant fractions underwent SDS-PAGE IB.

**Statistical analysis.** For quantification of 7SK snRNA, total cellular RNA isolated using the RNeasy Plus Mini Kit (QiAGEN) or RNA from RFP was converted to cDNA using the iScript kit (Bio-Rad) and analyzed by quantitative PCR (qPCR) on the iCycler platform using iQ SYBR Green Supermix (Bio-Rad). Primers and cycling conditions were described previously (11). Relative transcript levels were calculated using GAPDH normalization with the ΔΔCt formula.

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