Ikaros interacts with P-TEFb and cooperates with GATA-1 to enhance transcription elongation

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ABSTRACT
Ikaros is associated with both gene transcriptional activation and repression in lymphocytes. Ikaros acts also as repressor of human γ-globin (huγ-) gene transcription in fetal and adult erythroid cells. Whether and eventually, how Ikaros can function as a transcriptional activator in erythroid cells remains poorly understood. Results presented herein demonstrate that Ikaros is a developmental-specific activator of huγ-gene expression in yolk sac erythroid cells. Molecular analysis in primary cells revealed that Ikaros interacts with Gata-1 and favors Brg1 recruitment to the human β-globin Locus Control Region and the huγ-promoters, supporting long-range chromatin interactions between these regions. Additionally, we demonstrate that Ikaros contributes to transcription initiation and elongation of the huγ-genes, since it is not only required for TBP and RNA Polymerase II (Pol II) assembly at the huγ-promoters but also for conversion of Pol II into the elongation-competent phosphorylated form. In agreement with the latter, we show that Ikaros interacts with Cyclin-dependent kinase 9 (Cdk9), which contributes to efficient transcription elongation by phosphorylating the C-terminal domain of the large subunit of Pol II on Serine 2, and favors Cdk9 recruitment to huγ-promoters. Our results show that Ikaros exerts dual functionality during gene activation, by promoting efficient transcription initiation and elongation.

INTRODUCTION
The transcription factor Ikaros is widely expressed in hematopoietic cells where it regulates various aspects of hematopoiesis (1,2). Ikaros has been associated with gene activation (3–5), potentiation (6), priming (7) and transcriptional repression (8,9). In proliferating T cells, a high fraction of Ikaros co-purifies with Mi-2, a core component of the NuRD complex (10,11). In addition, Ikaros and Mi-2 control transcriptional regulation of the Cd4 locus during T cell differentiation (12). Nevertheless, a significant fraction of Ikaros is associated with a Brg1-based SWI/SNF-like complex (10). The relevance of the latter interaction has been indicated by several observations: (i) Ikaros has been associated with gene activation mediated by SWI/SNF-like complexes in T cells (3); (ii) Ikaros co-fractionates and co-immunoprecipitates with Brg1 (10,13); and (iii) Ikaros and Brg1 are also components of a SWI/SNF-like complex in mouse erythroleukemia (MEL) cells (14).

The human β- (huβ-) globin locus has been widely used as a model to explore the effects of transcription factors and co-factors on tissue- and developmental-specific gene expression. The huβ-globin locus contains five developmentally regulated genes (e.g.,γ-γ-γ-β-β). The locus control region (βLCR), which is located upstream of the globin genes, provides high-level globin gene expression in erythroid cells (EryC). The βLCR is composed of five DNase I hypersensitive sites (HSS) which are particularly rich in transcription factor binding sites (15). In EryC the βLCR favors high-level transcription through close interaction with gene promoters, and is a major determinant of locus chromatin organization (16). The transcription factors KLF (17), Ikaros (18), BCL11A (19), GATA-1 and its co-factor FOG-1 (Friend of GATA-1) (20), as well as the nuclear factor NLI/Ldb1 (21) and the chromatin remodeling co-regulator Brg1 (22) have all been shown to be required for efficient long-range chromatin interactions between the βLCR and β-like globin gene promoters.

Beside transcription initiation, modulation of transcription elongation is also likely implicated in globin gene regulation. Indeed, it has been shown that the βLCR
enhances the transition from transcription initiation to elongation for the β major globin (bmaj) gene (23). Additionally, globin gene expression can be reversibly inhibited by the ATP analog 5,6-dichloro-1-β-ribofuranosylbenzimidazole (DRB) (24,25), an inhibitor of cyclin-dependent kinase 9 (Cdk9) (26). Cdk9 is the catalytic subunit of the positive transcription elongation factor (P-TEFb). Cdk9 contributes to efficient transcription elongation by phosphorylating the C-terminal domain (CTD) of the large subunit of RNA Polymerase II (Pol II) on Ser2 (Pol II phospho Ser2). Several gene-specific regulators interact with P-TEFb but only a very limited number of transcription activators have been shown to recruit it to gene promoters (27). Two Cdk9 isoforms are found in mammalian cells and classified according to apparent molecular weight, namely Cdk9Δ2 and Cdk9Δ6. In general Cdk9 is essential for definitive but not primitive erythropoiesis in zebrafish (28), and deregulation of the Cdk9 pathway is implicated in the establishment and/or maintenance of a transformed cell phenotype, as documented for a number of human tumors and lymphomas (26).

Primitive hematopoiesis is the transient production of large, nucleated erythroid and megakaryocyte progenitors, and takes place in the yolk sac (29). Mouse primitive EryC express the embryonic globin genes εγ and βh1, whereas the human β-like globin genes expressed at the same stage of development in transgenic mice (carrying the complete huβ-globin locus) are the embryonic ε- (huc-) and the fetal γ- (hug-) genes (30,31). Hug-genes are transcriptionally silenced in adult definitive EryC by the contribution of several transcription regulators, such as Gata-1 (32,33), Ikaros (18,34), the TR2/TR4 heterodimer (35) that are present throughout primordial development as well as definitive erythropoiesis or the recently characterized Bcl11a protein, which however appears to be expressed only in definitive but not primitive, yolk sac-derived EryC (30,36). We have recently shown that Ikaros and Gata-1 cooperate to silence the hug-genes in EryC at the fetal stage of development when hug- and huf-genes are expressed (18). To investigate if Ikaros can also act as transcriptional activator in EryC and to define the molecular mechanisms of Ikaros-dependent transcriptional control, we studied the expression profile of hug-genes in embryonic day 10.5 (e10.5) yolk sac EryC. We used line 2 (ln2) mice, which carry a 70 Kb DNA fragment containing the entire huβ-globin locus and express the human β-like globin genes in a developmental-specific manner (31), as well as Ikaros null (Iknull) mice that are null for Ikaros proteins due to a deletion in the last exon that leads to protein instability (37). Our results reveal that Ikaros is a transcriptional activator of hug-gene expression in primitive EryC. We provide the evidence that Ikaros contributes to efficient recruitment of Brg1, Gata-1, Pol II and Cdk9 to the hug-promoters, thus supporting active chromatin re-organization, transcription initiation and elongation of the hug-genes.

**MATERIALS AND METHODS**

**Transgenic mouse lines**

All mutant embryos were compared with wild type littermates, to reduce difference in genetic background or developmental stage. Embryos were genotyped by PCR as described in (37). Homozygous ln2 (ln2+/−) (31): heterozygous Ikaros (Iknull+/−) (37) mice were bred with ln2+/−:Iknull+/− females in order to avoid contamination of embryos with adult blood of the mother, and ln2+/−:Iknull+/− (ln2-Iknull) e10.5 yolk sacs were isolated. Animals were sacrificed by cervical dislocation. Yolk sacs were rinsed several times with phosphate-buffered saline (PBS) to remove maternal blood. Blood was collected by piercing and compressing the yolk sac with forceps. Animal experiments were conducted in accordance with the Canadian Council on Animal Care (CCAC) guidelines and approved by the Maisonneuve-Rosemont Hospital animal care committee.

**Wright-Giemsa staining**

This protocol was carried out as previously described (18).

**Chromatin immunoprecipitation and quantitative real-time PCR analyses**

Chromatin immunoprecipitation (ChIP) procedure and quantification were as previously described (18) starting with 5×10⁵ e10.5 yolk sac EryC. Cells were fixed with 1% formaldehyde for 10 min at 37°C. Chromatin was reduced in size by sonication in order to obtain fragments of 400–500 bp. About 1/30th of immunoprecipitated and unbound (input) material was used as template for quantitative real-time PCR (qPCR) with SYBR Green (Invitrogen) on an iCycler iQ™ (Bio-Rad) system; the kidney-specific Tamm-Horsfall gene promoter (Thp) was used as internal control. All data shown are the results of at least four independent ChIP experiments with qPCR reactions from each ChIP performed in triplicate; the data are plotted as the mean ± standard deviation (SD) of the measurements.

**Protein analysis**

Western blot (WB) and co-immunoprecipitation (co-IP) analyses were basically as previously described (18). For co-IP, 5×10⁶ e10.5 yolk sac EryC, 1×10⁶ COS-7 cells or 10×10⁶ Jurkat cells were lysed in 1 ml/10⁷ cells of ice-cold lysis buffer (20 mM Hepes pH 8.0, 25% glycerol, 0.2 mM EDTA, 150 mM NaCl, 1.5 mM MgCl₂, 0.5% NP-40) containing protease inhibitors (Protease Inhibitor Cocktail; Sigma). Where detailed, co-IP washing buffers contained higher salt (200 mM) or NP-40 (1%) concentration. Ikaros proteins were not detected by WB in yolk sac EryC co-IP input samples, which represents 2% volume of the total lysate, i.e. ~10⁵ yolk sac EryC. Therefore, more concentrated yolk sac protein lysates, i.e. 2.5×10⁶ yolk sac EryC per lane, were loaded onto SDS-PAGE in order to identify Ikaros-specific proteins.
Cell transfection

Cells were transfected with murine Ikaros 2 cDNA, which is, together with Ikaros-1, the most abundantly expressed isoform in hematopoietic cells (4) and/or murine Gata-1. Stable Jurkat cell clones were obtained by retroviral transduction of wild type Jurkat cells with the pOZN-FH-Ik expression vector (18), which allows expression of Ikaros in double N-terminally epitope-tagged (Flag and Hemoagglutinin -FH-) fusion protein. Plasmid characteristic, cell transfection and selection were carried out as described in Nakatani et al. (38). COS-7 cells were transiently transfected with Flag/HA Ikaros (pCMV-FH-Ik) or with Gata-1 (pCMV-Gata) CMV-based expression vector using the FuGene system (FuGene 6, Roche), according to the manufacturer’s instructions. Cells were harvested 36 h post-transfection.

Immunofluorescence

For IF studies, transiently transfected COS-7 cells were seeded on glass slide 24 h post-transfection and allowed to adhere for additional 12 h. Cells were then fixed in 4% formaldehyde for 30 min at RT, washed once in PBS and subjected to heat-mediated antigen retrieval treatment. Slides were heated three times to 90°C for 2 min in the retrieval solution (10 mM citric acid pH 6.0, 0.05% Tween-20) and then allowed to cool down for 20 min at RT. Next, slides were treated with PBS/0.5% Triton for 20 min at RT, washed twice in PBS and blocked in PBS/0.5% NFDM. Primary (listed in Supplementary Figure S1) as well as secondary antibodies (all from Jackson ImmunoResearch) were diluted in PBS containing 0.1% Bovine Serum Albumin and 0.01% Triton. Slides were dehydrated by treatment with 70, 90% and 100% ethanol and covered with Dapi-Dabco: Vectashield solution (1:1) in glycerol. Images were acquired with a LSM 510 META Zeiss confocal microscope with z-resolution of 200 μm equipped with Zeiss LSM Image Browser. Red–green–blue (RGB) profiles were generated by the WCIF ImageJ® program (NIH).

Quantitative reverse transcription–PCR

Total RNA was extracted with Trizol (Invitrogen) and treated with DNaseI–RNase free (Invitrogen). Reverse transcription reactions were performed with oligo(dT)15 or random primers and SuperScript III reverse transcriptase (Invitrogen). qPCR was carried out on an iCycler iQ™ (Bio-Rad) system using specific primer sets; quantification was carried out as previously reported (18). Data shown are the results of at least five independent experiments with qPCR reactions from each cDNA performed in triplicate; the data are expressed as the mean ± SD of the measurements.

Chromosome conformation capture

The chromosome conformation capture (3C) protocol was basically as previously described (18) using 1–2 × 10^6 e10.5 yolk sac EryC; data shown are the results of five independent experiments with qPCR reactions from each 3C reaction performed in triplicate; the data are plotted as the mean ± SD of the measurements.

Statistical analysis

Unpaired ‘Student’s t-test’ was used to determine the level of statistical significance (P-value).

Primer sequences and antibodies

Primer sequences and antibodies are published as Supplementary Figure S1.

RESULTS

Ln2-Iknul e10.5 yolk sacs do not have gross developmental defects

Primitive EryC are characterized by their large size and expression of embryonic β-like globin genes, i.e. εy and βh1 in mice and ε in humans. In transgenic mice carrying the huβ-globin locus (Figure 1A), such as ln2 mice, both huγ- and huε-genes are expressed in yolk sac EryC (30,31). Ikaros is expressed throughout development in EryC and can be detected in yolk sac and fetal liver at e11 (2). The DNA-binding isoforms Ikaros-1, -2/3 and -4 are the predominant isoforms expressed in hematopoietic cells (4). To assess the expression pattern of Ikaros isoforms in yolk sac EryC, total RNA was retro-transcribed and cDNA amplified using Ikaros exon 1- and 7-specific primers. As shown in Figure 1B, Ikaros-1, -2/3 and -4 are the predominant isoforms expressed in hematopoietic cells (4). To assess the expression pattern of Ikaros isoforms in yolk sac EryC, total RNA was retro-transcribed and cDNA amplified using Ikaros exon 1- and 7-specific primers. As shown in Figure 1B, Ikaros-1, -2/3 and -4 isoforms are expressed in yolk sac EryC at a relative molecular ratio resembling that observed in adult thymus cells. We then investigated whether erythroid development and differentiation are affected in ln2-Iknul yolk sac EryC. First, we evaluated yolk sac cellularity. Keeping in mind the difficulties associated with quantitative analysis of e10.5 blood cells, no direct correlation between Ikaros genotype and total number of nucleated EryC were not visibly anemic, had apparently normal morphology, and were indistinguishable from wild-type counterparts (data not shown). Second, we performed Wright-Giemsa staining of ln2 and ln2-Iknul yolk sac EryC. In both genetic backgrounds, embryonic blood was predominated by large basophilic erythroblasts (Figure 1C), indicating similar erythroid differentiation behaviour in ln2 versus ln2-Iknul yolk sac EryC. Third, we evaluated transcript levels of the erythrocyte cell-surface molecule Gpa (Glycophorin A) and the transcription factors Eklf, Gata-1, Fog-1 and Sel/Tal-1, which are co-regulators critical for erythroid differentiation and globin gene expression (39). Quantitative reverse transcription–PCR (qRT–PCR) analysis revealed that Gpa, Eklf, Gata-1, Fog-1 and Sel/Tal-1 expression levels do not significantly vary between ln2 and ln2-Iknul yolk sac EryC (n ≥ 4) whereas, as observed in e12.5 fetal liver EryC (18), loss of Ikaros is associated with increased Gata-2 levels (Figure 1D; P = 0.03). Overall, these
junction sequence-specific primers for qPCR and Actin as internal control. Loss of Ikaros leads to reduced huγ-gene expression levels in yolk sac EryC (Figure 1D; \( P = 0.005 \)). Moreover expression of the embryonic huα- and mouse cε-, ey- and βH1-globin genes is unchanged in ln2-Iknull yolk sac EryC (\( n \geq 4 \)), and in these cells we detected only a slight but significant reduction of huβ- and mouse βmaj/βmin globin gene expression (\( n = 6 ; P = 0.05 \)). Similar results were obtained when Gpa was used as internal control. For instance, in ln2 versus ln2-Iknull cells, the relative level of huγ-gene expression was 5.9 (SD 1.2; \( n = 6 ; P = 0.05 \)). Gpa is an erythroid-specific membrane marker gene used as control to ensure that variations in globin gene expression are not merely due to changes in the number of viable EryC. Altogether, the above data suggest that Ikaros specifically contributes to huγ-gene transcriptional activation in yolk sac EryC.

Ikaros promotes efficient recruitment of Brg1 to βLCR HS3 and huγ-promoters in ln2 yolk sac EryC

It has recently been demonstrated that Ikaros is recruited to βLCR HS3 (H3), huγ-promoters and a pyrimidine stretch (Pyr region) located 1 kilobase upstream of the huδ-gene that contributes to huγ-gene silencing in adult-type erythroid cells; however, Ikaros does not bind to HS2 nor to huε- and huβ-promoters even though the Ikaros consensus sequence, TGGGAA, is found in the latter [data not shown and (1,14,18,40)]. To investigate occupancy by Ikaros of these regulatory regions in yolk sac EryC in vivo, we performed chromatin immunoprecipitation (ChIP) analysis using chromatin isolated from ln2 or ln2-Iknull yolk sac EryC and Ikaros antibodies (18). Ikaros is efficiently recruited to HS3, huγ-promoters and Pyr region in yolk sac EryC and to CD4 enhancer 1 region (CD4E1, used as control) in adult thymus cells (12), whereas no significant recruitment was detected at HS2, huε- or huβ-promoter (Figure 1E).

Ikaros co-fractionates and co-immunoprecipitates with Brg1 (10,13,14), and furthermore acts as a potentiator of gene expression during T-cell development possibly by recruiting Brg1-containing chromatin remodelling complexes to appropriate lineage-specific target genes (4,6). In addition, Brg1 is recruited to mouse HS3, is required for embryonic and adult β-globin gene transcription, erythroblast survival (41,42), and long-range chromatin interactions at the β-globin locus (22). Therefore, we studied the influence of Ikaros, when acting as transcription activator, on long-range chromatin interactions. As such we exploited the 3C assay to determine the physical proximity between chromosomal regions that are normally located many kilobases apart in vivo. Using the βLCR HS4-HS2 region as ‘fixed’ fragment (43), we observed that long-range chromatin interactions between the βLCR and the huγ-promoters are decreased in ln2-Iknull yolk sac EryC (Figure 2A; \( n = 5; P = 0.05 \)). Next, we analyzed in vivo recruitment of Brg1 to several regions across the huβ-globin locus by ChIP. Brg1 recruitment to HS3 and huγ-promoters is reduced in ln2-Iknull EryC but is not affected at HS2, huε- or huβ-promoters or mouse HS2 and βmaj.
Figure 2. BrgL1 and Mi-2 recruitment to the hufβ-globin locus in ln2 and ln2-\textit{Iknull} yolk sac EryC. (A) Ln2 and ln2-\textit{Iknull} yolk sac EryC were subjected to 3C assay; nuclei were digested with EcoRI and genomic DNA was ligated and used as template for qPCR; βLCR HS4-HS2 EcoRI fragment was used as ‘fixed’ point and primer sets were designed in order to amplify the genomic regions corresponding to the \(ε, ε\)-gene; \(\text{ie-γ}, \text{inter-ε-γ}\) region; \(γ\)-gene and \(β\)-gene region; relative crosslinking frequencies (\(γ\)-axis) of the ‘fixed’ fragment with globin fragments were defined using naked DNA encompassing the whole hufβ-globin as control and normalized to endogenous mouse \(Actin\) expression by RT–qPCR. We next examined Mi-2 gene expression by RT–qPCR. We observed that Mi-2 expression levels are the same in ln2 and ln2-\textit{Iknull} yolk sac EryC, whereas the lower half was probed with Ikaros antibodies; asterisks indicate non-specific bands; WCE: yolk sac EryC total cell lysate. In (B) ChIP on ln2 and ln2-\textit{Iknull} yolk sac EryC by ChIP. As depicted in Figure 2E, no significant recruitment of Mi-2 to either HS3, HS2 or to hu\textit{-}, hu\textit{-γ}, hufβ-promoters could be detected. Nonetheless under the same experimental conditions, in confirming the technical feasibility of Mi-2 ChIP, we could demonstrate efficient Mi-2 recruitment to the –2.8 kb region upstream of the \(Gata-2\) promoter in yolk sac EryC (Figure 2E) (45), the CD4E1 region (12) in freshly-isolated thymic cells (Figure 2E), and several regions of the hufβ-globin locus in EryC isolated from e12.5 fetal livers (18). Due to the possibility that absence of significant Mi-2 recruitment to critical regulatory regions of the hufβ-globin locus might depend upon lower levels of Mi-2 expression in ln2-\textit{Iknull} yolk sac EryC, we next examined Mi-2 gene expression by RT–qPCR. We observed that Mi-2 expression levels are the same in ln2 and ln2-\textit{Iknull} yolk sac EryC, whereas the lower half was probed with Ikaros antibodies; asterisks indicate non-specific bands; WCE: yolk sac EryC total cell lysate.

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Ikaros is also known to interact with the nucleosome remodelling ATPase Mi-2 (a subunit of the NuRD chromatin remodeling complex) (10,11), and recruits this co-factor to specific chromatin regions (18). Even though Mi-2 is widely perceived as a transcriptional repressor, the Mi-2-NuRD complex has recently been linked to both gene activation and repression in the case of certain \(Gata-1/Fog-1\) target genes (44). For this reason, we investigated \textit{in vivo} recruitment of Mi-2 to the hufβ-globin locus in ln2 as well as ln2-\textit{Iknull} yolk sac EryC by BrgL1. As depicted in Figure 2E, no significant recruitment of Mi-2 to either HS3, HS2 or to hu\textit{-}, hu\textit{-γ}, hufβ-promoters could be detected. Nonetheless under the same experimental conditions, in confirming the technical feasibility of Mi-2 ChIP, we could demonstrate efficient Mi-2 recruitment to the –2.8 kb region upstream of the \(Gata-2\) promoter in yolk sac EryC (Figure 2E) (45), the CD4E1 region (12) in freshly-isolated thymic cells (Figure 2E), and several regions of the hufβ-globin locus in EryC isolated from e12.5 fetal livers (18). Due to the possibility that absence of significant Mi-2 recruitment to critical regulatory regions of the hufβ-globin locus might depend upon lower levels of Mi-2 expression in ln2-\textit{Iknull} yolk sac EryC, we next examined Mi-2 gene expression by RT–qPCR. We observed that Mi-2 expression levels are the same in ln2 and ln2-\textit{Iknull} yolk sac EryC, whereas the lower half was probed with Ikaros antibodies; asterisks indicate non-specific bands; WCE: yolk sac EryC total cell lysate.

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and ln2-\textit{Ik}\textsuperscript{null} yolk sac EryC (Figure 2F). However, under the same stringency conditions that allowed co-IP of Ikaros and Brg1 (150 mM NaCl, 0.5% NP-40), Ikaros and Mi-2 could not be co-immunoprecipitated in yolk sac EryC with specific antibodies (Figure 2G). These results argue against a direct role of Mi-2 for \textit{hu}\textgamma-\textsuperscript{gene} activation in yolk sac EryC, and rather suggest that Ikaros-dependent recruitment of Brg1 to \textit{hu}\textgamma-promoters and HS3 is required for efficient \textit{hu}\textgamma-\textsuperscript{gene} expression in these cells.

**Ikaros facilitates Gata-1 recruitment to \textit{hu}\textgamma-regulatory regions in yolk sac EryC**

We previously reported that Ikaros physically interacts with Gata-1 and facilitates Gata-1 recruitment to HS3 and \textit{hu}\textgamma-promoters in e12.5 fetal liver EryC (18). To better define Ikaros-Gata-1 protein interaction, we first analyzed the subcellular localization of Ikaros and Gata-1 at the single-cell level by immunofluorescence (IF) and confocal microscopy. COS-7 cells, which lack any Ikaros or Gata-1 protein, were transfected with Flag/HA-Ikaros (pCMV-FH-Ik) together with Gata-1 (pCMV-Gata) expression vectors, fixed with formaldehyde, permeabilized and immunolabeled with antibodies specific for HA (i.e. Ikaros detection) or Gata-1. Ikaros and Gata-1 were detected using secondary antibodies conjugated to Texas red (TR) or Fluorescein isothiocyanate (FITC) fluorochromes, respectively. The use of cross-absorbed and affinity-purified secondary antibodies minimized background and non-specific reactivity to undetectable fluorescent levels (data not shown). Confocal microscopy analysis revealed that both Ikaros (red signals) and Gata-1 (green signals) have a dominant nuclear localization (Figure 3A), whereas COS-7 cells transfected with empty vectors (mock control) showed undetectable levels of green or red stain (Supplementary Figure S2A). Interestingly, the merged image obtained in COS-7 cells that express both Ikaros and Gata-1, leads to the emergence of several yellow nuclear signals per focal plane (Figure 3A). Physical overlap between Ikaros and Gata-1 was further defined by the ImageJ\textsuperscript\textregistered program, which revealed that the line profile plots and fluorescent peaks of Ikaros and Gata-1 often overlap whereas the fluorescence profiles of Ikaros and Ape1, a nuclear protein that does not interact with Ikaros used as negative control (46), are discordant.

Next, Ikaros-Gata-1 protein interaction was investigated by protein co-IP. Protein lysates of ln2 yolk sac EryC were immunoprecipitated with Ikaros or Gata-1 antibodies as well as isotype-matched rabbit (for Ikaros co-IP) or rat (for Gata-1 co-IP) immunoglobulins (Ig) and immunoblots were probed with Ikaros antibodies. The results show that Ikaros can be immunoprecipitated by Gata-1 antibodies but not by control Ig (Figure 3B), suggesting that Ikaros and Gata-1 can be part of the same protein complex also in yolk sac EryC.

Ikaros-Gata-1 cooperative binding to the \textit{hu}\beta-globin locus was then investigated by ChIP. In ln2-\textit{Ik}\textsuperscript{null} yolk sac EryC, Gata-1 binding is diminished at \textit{hu}\gamma-promoters and Pyr region whereas is normal at HS3 and HS2 (Figure 3C). In order to investigate whether loss of Ikaros specifically decreases Gata-1 binding or induces a more general alteration of transcription factor recruitment, we analyzed p45/Nf-e2 and Yy1 occupancy since these proteins bind to \textit{hu}\gamma-promoters. No significant difference in p45/Nf-e2 (Figure 4A) or Yy1 (Figure 4B) recruitment was detected at tested chromatin regions in ln2 versus ln2-\textit{Ik}\textsuperscript{null} yolk sac EryC. We then examined whether the Ikaros-Gata-1 complex could modulate recruitment of the Gata-specific cofactor Fog-1 by ChIP. Fog-1 is recruited to HS3, HS2, \textit{hu}\gamma- and \textit{hu}\beta-promoters in ln2 and, unexpectedly, ln2-\textit{Ik}\textsuperscript{null} yolk sac EryC (Figure 4C), even though Gata-1 binding to these promoters is close to background levels (Figure 3C). This outcome suggests that Fog-1 recruitment to \textit{hu}\gamma- and \textit{hu}\beta-promoters in ln2-\textit{Ik}\textsuperscript{null} cells does not depend solely on Gata-1. Since Gata-1 and Gata-2 (i) bind to the same DNA motifs \textit{in vivo} and \textit{in vitro} (47–49); (ii) function redundantly to promote primitive erythroblast development (50); (iii) exert their respective functions in a dose-dependent fashion (39,51); and (iv) bind Fog-1 protein (52), we set out to investigate Gata-2 occupancy at globin promoters in ln2-\textit{Ik}\textsuperscript{null} EryC. Loss of Ikaros correlates with increased Gata-2 recruitment to HS2, \textit{hu}\gamma- and \textit{hu}\beta-promoters, whereas Gata-2 recruitment does not significantly change at the \textit{Gpu} promoter, which was used as a control for Gata-2 binding (Figure 4D) (53). These results suggest that Ikaros contributes to Gata-1 binding at \textit{hu}\gamma-promoters and Pyr region in ln2 yolk sac EryC. They also indicate that increased Gata-2/Gata-1 ratio, as observed in absence of Ikaros (Figure 1D), might enhance the interaction between Fog-1 and Gata-2, thus allowing Gata-2 to efficiently support Fog-1 recruitment to \textit{hu}\beta-promoters in yolk sac EryC.

**Ikaros contributes to efficient PIC assembly and transcription elongation of \textit{hu}\gamma-promoters in ln2 yolk sac EryC**

A critical step during eukaryotic gene transcription is the assembly of the PIC consisting of Pol II and general transcription factors (27). Both chromatin conformation and gene-specific activators influence PIC assembly and hence, transcriptional efficiency. We proceeded to define Ikaros contribution to PIC assembly at the \textit{hu}\gamma-promoters by ChIP. As expected, TBP and Pol II were each efficiently detected at \textit{hu}\gamma- and \textit{hu}\beta-promoters in ln2 yolk sac EryC; in ln2-\textit{Ik}\textsuperscript{null} cells, TBP and Pol II recruitment is affected at \textit{hu}\gamma-promoters and Pol II recruitment is also diminished at the HS2 region in ln2-\textit{Ik}\textsuperscript{null} yolk sac EryC (Figure 5A and B). Since Pol II recruitment to \textit{hu}\gamma-promoters is only moderately affected by loss of Ikaros, we examined whether Ikaros could also control the efficiency of transcription elongation by studying: (i) levels of Pol II phospho Ser2 (PCTD) recruitment to the \textit{hu}\gamma-genes, because it is known that the levels of this phosphorylated residue is required and increase during transcription elongation (27); (ii) the relative levels of \textit{hu}\gamma-globin 5'- and 3'-primary transcripts, because primary transcript levels accurately reflect the rate of transcription. ChIP assays revealed that PCTD recruitment to...
HS2 as well as huγ-genes and -promoters is significantly affected in absence of Ikaros (Figure 5C). Primary transcript expression was determined by RT-qPCR using random primers for cDNA synthesis and huγ-gene and mouse Gapdh intronic sequence-specific primers for qPCR. In line with reduced PCTD occupancy at huγ regions, there was an accumulation of transcripts corresponding to the 5’-end of the gene and significant reduction of full-length transcripts (Figure 5D). These results are reminiscent of a gene that initiates transcription without completing elongation, thus supporting a possible role of Ikaros during productive huγ-gene transcription elongation.

Ikaros interacts with Cdk9

In higher eukaryotes, Pol II Ser2 phosphorylation is mediated primarily by the Cdk9 catalytic subunit of the P-TEFb complex (27). Considering that the Cdk9 inhibitor DRB is known to affect transcription elongation of huγ-genes (24,25), we investigated whether the P-TEFb complex is involved in huγ-gene transcription elongation and whether Ikaros influences the recruitment of P-TEFb to the huγ-genes. Interestingly, ChIP revealed that Cdk9 is recruited to huγ promoters in ln2 yolk sac EryC, and that Cdk9 occupancy is significantly reduced at huγ but not huγ-promoters in ln2-Iknull yolk sac EryC (Figure 6A). To evaluate the significance of the 3-fold enrichment observed in ln2 EryC, we performed ChIP with Cdk9 antibodies in DMSO-induced MEL cells since it has recently been shown that Cdk9 is recruited to the βmaj promoter in these cells (54). The level of Cdk9 enrichment at βmaj promoter in MEL cells is very similar to the occupancy levels of Cdk9 at huγ-promoters in ln2-Iknull yolk sac EryC, thus confirming the
significance of the results obtained in ln2 yolk sac EryC. Importantly, loss of Ikaros does not modulate the level of Cdk9 transcripts (Figure 6B) or proteins (Figure 6C) in yolk sac EryC.

Cdk9 interacts with a number of transcription factors (27). To investigate whether Ikaros and Cdk9 interact in yolk sac EryC, we performed protein co-IP with Cdk9 antibodies or control Ig. Immunoblots probed with Ikaros antibodies revealed that Ikaros can be immunoprecipitated by Cdk9 antibodies but not by control Ig, hence suggesting that these proteins physically interact (Figure 6D). To assess Ikaros-Cdk9 interaction at single-cell level, we analyzed the subcellular localization of these proteins by IF and deconvolution microscopy.

Figure 4. Recruitment of transcriptional regulators to the huβ-globin locus in yolk sac EryC. (A–D) ChIP on ln2 and ln2-Iknull yolk sac EryC carried out with p45/Nf-e2 (p45), Yy1, Fog-1 and Gata-2 antibodies; fold enrichments (y-axis) were calculated as described in Figure 1E and are plotted as the mean ± SD of the measurements; a value of 1 (dashed line) indicates no enrichment; *P < 0.05 by Student’s t-test; hu, huγ-promoter; huγ, huγ-promoters; huβ, huβ-promoter; m2, mouse HS2; Gpa, Glycophorin A promoter; dark gray bars: ln2 yolk sac EryC; light gray bars: ln2-Iknull yolk sac EryC; white bars: isotype-matched Ig (Ig ctl).

Figure 5. PIC assembly at huγ-genes in ln2 and ln2-Iknull yolk sac EryC. (A–C) ChIP on ln2 and ln2-Iknull yolk sac EryC carried out with TBP, Pol II and Pol II phospho Ser2 (PCTD) antibodies; fold enrichments (y-axis) were calculated as described in Figure 1E and are plotted as the mean ± SD of the measurements; a value of 1 (dashed line) indicates no enrichment; *P < 0.05 by Student’s t-test; hu, huγ-promoters; huγ, huγ-promoters; huβ, huβ-promoter; Gpa, Glycophorin A promoter; dark gray bars: ln2 yolk sac EryC; light gray bars: ln2-Iknull yolk sac EryC; white bars: isotype-matched Ig (Ig ctl).
Figure 6. Ikaros-mediated recruitment of Cdk9 to the huy-genes in yolk sac EryC. (A) ChIP on ln2 and ln2-Iknull yolk sac EryC carried out with Cdk9 antibodies; fold enrichments (y-axis) were calculated as described in Figure 1E and are plotted as the mean ± SD of the measurements; a value of 1 (dashed line) indicates no enrichment; *P ≤ 0.05 by Student’s t-test; huc, huc-promoter; huy, huy-promoters; huβ, huβ-promoter; jma, jmajor promoter; dark gray bars: ln2 yolk sac EryC; light gray bars: ln2-Iknull yolk sac EryC; dark gray dashed bars: MEL cells; white bars: isotype-matched Ig (Ig ctl); (B) RT–qPCR of Cdk9 gene transcripts performed on equal amounts of ln2 and ln2-Iknull yolk sac EryC; transcript quantification was calculated according to Pfaffl (73) using mouse Actin cDNA as internal control; ln2/ln2-Iknull ratios are plotted as the mean ± SD of the measurements; n = 6; (C) WB performed on total cell lysates prepared from wild type or Iknull yolk sac EryC with Cdk9 or Actin antibodies; (D) protein co-IP of total cell lysates prepared from ln2 yolk sac EryC; cell lysates were immunoprecipitated with Cdk9 antibodies or isotype-matched Ig and WB were carried out with Ikaros antibodies; WCE: yolk sac EryC total cell lysate; (E) Confocal immunofluorescence of COS-7 cells expressing Flag/HA-Ikaros protein; cells were stained with mouse antibodies to the HA tag and with rabbit antibodies to Cdk9; the secondary staining was carried out with FITC-conjugated anti-mouse as well as TR-conjugated anti-rabbit antibodies; a single COS-7 cell is shown where Ikaros is detected as green stain (Supplementary Figure S2B). Confocal microscopy analysis revealed that both Ikaros (green signals) and Cdk9 (red signals) are distributed throughout the nucleus and that these proteins significantly overlap, even though they do not colocalize at each and every nuclear sites, in accordance with known evidence that Cdk9 is present in several other protein complexes (26).

Since Ikaros is known to act as transcriptional repressor and activator of specific genes in T lymphocytes, it was intriguing to know whether Ikaros could possibly interact with Cdk9 in the T lymphocyte-like Jurkat cell line. Jurkat cells were stably transfected with the pOZN-FH-Iκκ expression vector that contains a bicistronic transcriptional unit, which allows expression of two proteins from a single transcript. This design ensures tight coupling between expression of the gene of interest and the selection marker, the interleukin-2 receptor α chain (IL2Rα) (18,38). Three rounds of magnetic affinity sorting with antibodies against the IL-2α receptor were sufficient to obtain a pure population of Jurkat cells (namely, Ju/Ik clone) that express Flag/HA-Ikaros at lower levels than endogenous Ikaros (Figure 7A). Total cell lysates were immunoprecipitated with Flag, Cdk9 antibodies or control Ig and WB performed with HA or Cdk9 antibodies. As shown in Figure 7B, Flag/HA-Ikaros is immunoprecipitated by both Flag and Cdk9 antibodies. As well, Cdk9 is immunoprecipitated by both Flag and Cdk9 antibodies, indicating that Ikaros and Cdk9 are part of the same protein complex. These results suggest that the Ikaros-Cdk9 interaction can also occur in lymphoid cells and does not require Ikaros overexpression because Flag/HA-Ikaros is expressed at lower levels than endogenous Ikaros.

Since Ikaros interacts with Gata-1 (18) (Figure 3B), and Gata-1 has been shown to interact with the P-TEFb component Cyclin T1 (CycT1) in megakaryocytes (55), we sought to investigate whether Gata-1 could interact with Cdk9 in mouse erythroleukemic MEL cells. Protein co-IP revealed that Cdk9 can be efficiently immunoprecipitated by one out of two antibodies tested directed against Gata-1 (clone M20) and that Gata-1 is immunoprecipitated, even though weakly, by Cdk9 antibodies (Figure 7C), thus indicating that Gata-1 and Cdk9 are part of the same protein complex in MEL cells. Next, we asked whether Gata-1 could either interfere with or enhance the interaction between Ikaros and Cdk9. To this end, COS-7 cells were transfected with pCMV-FH-Iκκ, pCMV-Gata-1 or pCMV-FH-Iκκ together with pCMV-Gata-1 expression plasmid. Cell lysates were immunoprecipitated with Gata-1 (N6 as well as M20 clone), Cdk9, Flag antibodies or control Ig and precipitated proteins were analyzed by WB with Cdk9, HA or Gata-1 antibodies. As expected, Ikaros and Gata-1 interact in COS-7 transfected cells that express both proteins, as revealed by Gata-1 co-IP and WB
detection with anti-HA antibodies (Figure 7D, bottom panel). However, in COS-7 cells that express either Gata-1 or Gata-1 together with Flag/HA-Ikaros, Gata-1 is not detected upon Cdk9 co-IP nor is Cdk9 detected upon Gata-1 co-IP (Figure 7D), thus suggesting that Gata-1 and Cdk9 do not stably interact in non-hematopoietic, COS-7 cells. Using the same transfected cells and under the same stringency conditions, Flag/HA-Ikaros is detected upon Cdk9 co-IP and Cdk9 is detected upon Flag co-IP (Figure 7D and E), indicating that Ikaros and Cdk9 co-immunoprecipitate also in COS-7 cells and that additional hematopoietic-specific proteins are not required for this interaction. It is to notice that the interaction between Ikaros and Cdk9 can be maintained at higher stringency conditions, i.e. 200 mM NaCl or 1% NP-40 (Supplementary Figure S3A). Interestingly, Gata-1 appears to increase the degree of Ikaros-Cdk9 interaction, since the intensity of the immunoprecipitated bands is proportional to the levels of Gata-1 expression (Figure 7E). Altogether, these data indicate that (i) Ikaros and Gata-1 interact also in COS-7 transfected cells; (ii) Gata-1 interacts with Cdk9 in erythroid cells; (iii) Ikaros interacts with Cdk9 in erythroid, lymphoid as well as COS-7 transfected cells; and (iv) Ikaros-Cdk9 interaction does not require Gata-1 even though Gata-1 appears to enhance the strength of this interaction.

DISCUSSION

In the present study, we have shown that Ikaros acts as a stage-specific activator of huf-genes in yolk sac EryC by (i) favoring Brg1 and Gata-1 recruitment to huf-promoters and contributing to chromatin interaction between the βLCR and these promoters and (ii) supporting efficient PIC assembly and Cdk9 recruitment to huf-promoters and hence, Pol II Ser2 phosphorylation which is required for transcription elongation. This suggests that through combined action with transcription factor and co-factor partners, Ikaros can exert control over both transcription initiation and elongation.
Ikaros and mouse embryonic development

Previous analysis of Iknull mice highlighted the importance of Ikaros for the regulation of hematopoietic genes (7). In particular, Scl/Tal-1, Flk-2 receptor and c-kit receptor gene expression is affected in Iknull bone marrow hematopoietic stem cells (HSC) (56). However, in Iknull fetal liver EryC, Scl/Tal-1 expression levels are unchanged, whereas Eklf and Gata-2 expression levels are, respectively, decreased and increased (18, 34). Finally, in Iknull yolk sac EryC, we report that Eklf, Gata-1, Fog-1 and Scl/Tal-1 expression levels are unchanged, whereas Gata-2 is overexpressed. Therefore, Ikaros, likely in association with specific chromatin regulators, can activate or repress a variety of hematopoietic-specific genes in a cell type- and developmental stage-specific manner.

Ikaros is essential for specification in lymphoid lineages, and inactivation of this protein exerts a profound effect on the production and differentiation of fetal lymphocyte progenitors (37). However, recent studies have identified Ikaros as an important regulator of lympho-myeloid lineage decisions in early hematopoietic progenitor cells (7). Ikaros can also influence erythroid cell differentiation. For example, adult Iknull mice display extramedullary hematopoiesis, bone marrow hypoplasia, and slight reduction in the number of BFU-E (Burst-Forming Unit Erythroid) and CFU-E (Colony-Forming Unit Erythroid) progenitors; however, reduced CFU-E and BFU-E activity is reported to be mainly due to a deficit in the HSC population rather than aberrant erythroid differentiation (56). Accordingly, erythropoiesis appears to be normal at earlier stages of development in fetal liver (18, 37) and yolk sac (this work) isolated from Iknull mice where erythroid differentiation progresses to the point of producing Gpa transcripts, normal numbers of maturing proerythroblasts and typical levels of murine globin gene transcripts. Based on these findings, we conclude that reduced levels of huyγ-gene expression in ln2-Iknull cells is not the consequence of impaired erythroid differentiation or maturation, and hence that Ikaros is recruited to, and directly contributes to regulation of, the huyγ-genes in yolk sac EryC.

Ikaros, Brg1 and Gata-1 cooperate to control huyγ-gene expression in yolk sac EryC

We have recently shown that Ikaros participates in huyγ-gene silencing at the time of γ-to-β globin switching which takes place at ~e12 in transgenic ln2 fetal liver by targeting a Gata-1/Fog-1/Hdac1/Mi-2-containing multiprotein complex to huyγ-promoters (18). Here, we have shown that Ikaros, along with Gata-1 and Brg1, rather contributes to huyγ-gene expression at an earlier developmental stage, i.e. in e10.5 yolk sac EryC. The observation that Ikaros can act as transcription activator and repressor for the same gene in a stage- and cell-specific manner is reminiscent of previous studies showing that Ikaros represses Il4 expression in mast cells (57), but activates that gene in Th2 cells, and also appears to exert distinct effects on Il4 and Ifny gene expression in the same cell type (Th1 and Th2) (58). The ability of Ikaros to preferentially interact with Mi-2, Brg1 or other known Ikaros co-regulators, and hence, to direct them to distinct gene regulatory regions, might depend on Ikaros post-translational modifications. Indeed, sumoylated Ikaros preferentially interacts with Brg1 versus Mi-2 (13). Thus, increased sumoylation levels in yolk sac relative to fetal liver EryC could account for the tendency of Ikaros to interact with Brg1, and possibly act as transcription activator. This could not be directly addressed since sumoylation is a highly dynamic process and a suitable cellular model for human primitive EryC is currently unavailable. A second, non-mutually exclusive explanation for different levels of Ikaros interaction with Mi-2 versus Brg1 is that expression of these co-regulators might fluctuate during development. In accordance with such hypothesis, we observed that Brg1 is expressed at higher levels in yolk sac than fetal liver EryC, whereas Mi-2 expression is higher in fetal liver EryC (Supplementary Figure S3B). Brg1 is the principal catalytic subunit of a SWI/SNF-like complex in Ter119+ EryC, and is essential for viability in mice (59). The use of hypomorphic (41) and conditional Brg1null mutations (42) have demonstrated that Brg1 is required for chromatin remodelling during both embryonic and adult β-globin gene transcription, and for maximal Pol II occupancy at the mouse βmaj promoter (60). A number of transcription factors are known to bind and recruit Brg1 to the globin locus (40, 61–63); nonetheless using a Gata-1-inducible cell line, Kim et al. (22) reported that Gata-1 recruits Brg1 to the adult βmaj promoter more rapidly than other transcription factors tested.

The current results, together with previous published data (18), indicate that Gata-1 is a relevant Ikaros-interacting transcription factor and their combined effect is required for globin gene regulation in EryC. Indeed, Gata-1 and Ikaros are recruited in vivo to the huβ-globin locus, influence long-range chromatin interactions, contribute to huyγ-gene regulation and co-immunoprecipitate in vivo in primary yolk sac (herein) and fetal liver (18) EryC. Gata-1 has been associated to both huyγ-gene activation and repression, depending on the experimental model. For instance, Gata-1 contributes to huyγ-gene silencing at the time of γ-to-β globin switching in mouse fetal liver (18) or in adult (32, 33) EryC. By cooperating with BCL11a and SOX6, GATA-1 has recently been shown to contribute to huyγ-gene silencing in adult human erythroid progenitors (19) and interestingly, BCL11a is not expressed either in wild-type (30) or Iknull primitive EryC (Supplementary Figure S3C). However, GATA-1 has also been associated with huyγ-gene activation (64). Finally, GATA-1 overexpression in K562 cells does not change huyγ-gene expression (65).

Despite the fact that Ikaros does not affect Gata-1 occupancy at βLCR HS2, recruitment of Gata-2 and Pol II to this region is significantly increased and decreased, respectively, in ln2-Iknull yolk sac EryC. It has recently been reported that Gata-1 occupancy at the well-defined ~1.8 kb site of the Gata-2 locus is associated with Pol II expulsion and stable gene repression (66). Therefore, it is possible that, similarly to Gata-1 at the ~1.8 kb site of the Gata-2 locus, increased Gata-2 occupancy at βLCR HS2
of the hβ-globin locus could partially display Pol II at this distal regulatory element.

In conclusion, even though we cannot exclude that Ikaros may also facilitate hγ-gene expression indirectly, e.g. by contributing to Gata-2 gene repression in yolk sac EryC, our data indicate that Ikaros directly contributes to hγ-gene expression mainly by conferring Gata-1 occupancy at hγ-promoters (see below) and, together with Brg1, by stabilizing hγ-promoter chromatin activation.

**Ikaros promotes efficient hγ-gene transcription elongation**

Here, we show that lack of Ikaros affects TBP and Pol II recruitment to hγ-promoters, and reduces Pol II Ser2 phosphorylation levels. Ikaros can interact with components of the basal transcription machinery such as TBP and TFIIIB (67), and with Brg1, which is critical for long-range chromatin interaction and, as such reported (60), Pol II transfer. Even though transcription elongation has been regarded as an important regulatory step for β-like globin gene regulation (23,68,69), the severe reduction of Pol II Ser2 phosphorylation levels observed at hγ-promoters in ln2-Iκnull yolk sac EryC was unanticipated in light of known properties of Ikaros. Our results indicate that Ikaros not only participates in PIC assembly but in addition contributes to Pol II Ser2 phosphorylation by recruiting the P-TEFb component Cdk9 to hγ-gene promoters.

Cdk9 expression levels increase in terminally differentiated cells (26). However, in the human erythroleukemic K562 cell line, Cdk9 knockdown induces erythroid differentiation (55), whereas erythroid differentiation of MEL cells does not lead to Cdk955 downregulation and in fact, engenders upregulation of Cdk955 (28). The reason for these discrepancies is not known, but it might depend on the intrinsic limitations of studying erythroid differentiation using transformed cell lines. Here we show that levels of Cdk9 expression are very similar in ln2 and ln2-Iκnull yolk sac EryC and also in yolk sac EryC when compared to fetal liver EryC (Supplementary Figure S3D). We studied Ikaros-Cdk9 protein interaction in cells expressing predominantly Cdk942 (yolk sac EryC and COS-7 cells) or both isoforms (Jurkat cells) and based on our results we conclude that Ikaros preferentially interacts with Cdk942.

Lack of Ikaros leads to increased and reduced Cdk9 recruitment to HS3 and hγ-promoters respectively, and correlates with reduced long-range chromatin interactions between the BLCR and the hγ-promoters. It is therefore possible that, as proposed for Pol II (70), Cdk9 first accumulates at the BLCR and then, is transferred to the hγ-promoters when the two regulatory regions are in closer proximity. It is widely accepted that developmental control of gene expression can be associated with poised Pol II and the distribution of Pol II across *Drosophila* and human genomes have provided evidence that elongation control might direct the pattern of gene expression during development (71). In mammalian cells, the P-TEFb complex accounts for most of Pol II Ser2 phosphorylation activity (26) and therefore is the best-known activity capable of reactivating poised Pol II (27). Thus, Ikaros-dependent Cdk9 transfer to hγ-promoters might be important for the transition from poised Pol II to productive elongation at the hγ-genes.

Recruitment of P-TEFb to highly acetylated gene promoters can be mediated by the bromodomain protein Brd4, which interacts directly with CycT1 (27). However, certain transcription factors, e.g. c-Myc, CIITA, NF-kB, MyoD, STAT3 and the androgen receptor can recruit P-TEFb to defined promoter regions by interacting with either CycT1 or Cdk9 ([26,72] and references herein) and recently Ldb1 has been shown to recruit P-TEFb to multiple regions of the mouse β-globin locus (54). Interestingly, GATA-1 is reported to interact with CYCT1 in a megakaryocyte-like cell line whereas GATA-1 interaction with CDK9 in the same cells was not addressed (55). Gata-1 and Cdk9 co-immunoprecipitate in MEL cells but not in COS-7 cells expressing Gata-1 or Gata-1 together with Ikaros, under the same stringency conditions that allow co-IP of Ikaros and Cdk9. Thus, it appears that Ikaros can interact with Cdk9 in several cell lines, whereas Gata-1 interaction with Cdk9 would rather necessitate an erythroid environment. Interestingly, even in absence of a detectable interaction between Gata-1 and Cdk9 in COS-7 cells, the presence of Gata-1 can enhance the strength of the interaction between Ikaros and Cdk9. Whether the erythroid-specific association of Ikaros, Gata-1 and Cdk9 depends upon their relative level of expression or post-translational modifications or the presence of additional erythroid-specific proteins is thus far unknown. Nevertheless, it is tempting to assume that in vivo cooperation between Ikaros and Gata-1 is designed to facilitate Pol II Ser2 phosphorylation, possibly by targeting two distinct components of the P-TEFb complex to hγ-gene promoters.

Based on our results, we propose that the recruitment of Ikaros to hγ-promoters in yolk sac EryC, promotes stable PIC formation and contributes to Cdk9 recruitment, which leads to efficient transcription elongation of the hγ-genes. Therefore, unlike Ldb1-dependent recruitment of P-TEFb that occurs at multiple regions of the mouse β-globin locus (54), our results suggest that Ikaros contributes to Cdk9/P-TEFb recruitment specifically to promoters of the transcriptionally active hγ-genes. According to the observation that Ikaros and Cdk9 co-immunoprecipitate also in lymphoid-like cells (Figure 7B), it is likely that Ikaros influences transcription elongation in a similar manner at other transcriptionally active Ikaros target genes and in other hematopoietic cells.

In conclusion, we show that Ikaros contributes to efficient transcription initiation and elongation of the hγ-genes in primitive EryC. As such, this study contributes to a better understanding of the molecular mechanisms of Ikaros action to control gene expression in hematopoietic cells. Therefore, results presented herein should provide a basis for understanding malignant hematopoietic processes characterized by Ikaros- or GATA-1-dependent gene dysregulation.
SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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