Lin28b promotes fetal B lymphopoiesis through the transcription factor Arid3a

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Abbreviations used: BCR, B cell antigen receptor; ELP, early lymphocyte progenitor; FDR, false discovery rate; FL, fetal liver; miRNA, microRNA; Q-PCR, quantitative PCR; TdT, terminal deoxynucleotidyl transferase; UTR, untranslated region.

Mouse B cell precursors from fetal liver and adult bone marrow (BM) generate distinctive B cell progeny when transplanted into immunodeficient recipients, supporting a two-pathway model for B lymphopoiesis, fetal “B-1” and adult “B-2.” Recently, Lin28b was shown to be important for the switch between fetal and adult pathways; however, neither the mechanism of Lin28b action nor the importance of B cell antigen receptor (BCR) signaling in this process was addressed. Here, we report key advances in our understanding of the regulation of B-1/B-2 development. First, modulation of Let-7 in fetal pro-B cells is sufficient to alter fetal B-1 development to produce B cells resembling the progeny of adult B-2 development. Second, intact BCR signaling is required for the generation of B1a B cells from Lin28b–transduced BM progenitors, supporting a requirement for ligand-dependent selection, as is the case for normal B1a B cells. Third, the $V_H$ repertoire of Lin28b–induced BM B1a B cells differs from that of normal B1a, suggesting persisting differences from fetal progenitors. Finally, we identify the Arid3a transcription factor as a key target of Let-7, whose ectopic expression is sufficient to induce B-1 development in adult pro-B cells and whose silencing by knockdown blocks B-1 development in fetal pro-B cells.

B cells, a key arm of the immune system responsible for humoral immunity, are generated through a tightly regulated sequence of developmental stages, in the liver before birth and in the BM of adults. During B cell development, Ig heavy and light chains are rearranged and selected, yielding a diverse antigen receptor repertoire that is largely purged of high-affinity pathogenic self-reactivity (Nemazee, 2006; Goodnow, 2007). Importantly, mature B cells in mice are not completely homogenous across anatomical sites. In particular, certain functionally distinct subsets, such as the CD5$^+$ B cell (“B1a”) subset, show some degree of self-reactivity (Hayakawa et al., 1984). A key unresolved issue is how these self-reactive cell types diverge from the primary B cell development pathway that generates follicular B cells. Study of the Ig heavy and light chains rearranged in these cells has shown that they constitute a biased set of B cell antigen receptors (BCRs; Förster et al., 1988; Pennell et al., 1989; Carmack et al., 1990), some of which have been shown to be selected by interaction with self-determinants (Hayakawa et al., 1999), suggesting an instructive antigen-dependent model for CD5$^+$ B cell generation.

Early BM transfer experiments revealed poor generation of CD5$^+$ B cells (B1a) in adult hosts (Hayakawa et al., 1985). Later experiments, using more defined populations of B cell progenitors from fetal and adult sources, showed that fetal precursors supported efficient production of B1a B cells, but repopulation of typical follicular B cells was inefficient (Hardy and Hayakawa, 2001). These results led us to propose a switch in B cell lymphopoiesis during ontogeny, similar to the well-known switch from fetal to adult hemoglobin in erythropoiesis (Groudine et al., 1983). Specifically, we suggested that the fetal pathway of development (termed “B-1”) is responsible for generating most of the CD5$^+$ B cell pool, whereas an adult pathway (termed “B-2”) generates most of the CD5$^-$ B cells that populate the adult (Hardy and Hayakawa, 2001). The latter cells are often
Figure 1. Fetal and adult pro-B cells generate B cells with distinct phenotype and show differential expression of mRNAs and miRNAs. (A) Pro-B cells were sorted from FL and BM and then transferred i.v. into immunodeficient SCID mice. Peritoneal cavity washout cells (PerC) were analyzed 3 wk later for the indicated surface proteins by flow cytometry. Data are representative of five independent transfer experiments with two mice per group. (B) Heat map displays of mRNA fold change comparing fetal and adult pre-pro-B (Fr. A) and pro-B (Fr. BC) stage cell samples by microarray of genes higher in FL or in BM using both the Affymetrix (C57BL/6J) and Agilent Technologies (BALB/c ICR) platforms. This analysis identifies 219 differentially expressed genes. (C) Heat map display of microarray data (top) and Q-PCR analysis (bottom) of miRNA species differentially expressed between adult BM and FL pro-B cells (Fr. BC). miRNAs in the heat map that are also analyzed by Q-PCR are indicated by a red asterisk. Q-PCR data were normalized to SN0202 expression. All expression analyses were performed in duplicate or triplicate, with each sample representing RNA or miRNA from at least four mice. Error bars represent ±SEM.
identified as follicular or "B2" B cells. We hypothesized that a distinctive gene program operating in B cell progenitors is responsible for the fetal-biased B–1 generation of CD5+ B cells.

Therefore we analyzed fetal- and adult-origin B cell precursors for mRNA and microRNA (miRNA) expression differences by microarray to rigorously identify potential regulators that might play a role in the B–1/B–2 developmental switch. Cell fractions where initial Ig heavy chain rearrangement takes place, pre–pro-B (Fr. A) and pro-B (Fr. BC), were analyzed because these populations span the stage at which B lineage commitment occurs (Rumfelt et al., 2006). Based on this idea of distinctive fetal and adult lymphopoiesis, Yuan et al. (2012) performed and published a similar analysis of pro-B stage cells, comparing gene expression and miRNA expression in such cells purified from fetal liver (FL) and adult BM of mice. They found that retroviral expression of Lin28b in BM stem cells generated innate-type B and T cells in transfer recipients (Yuan et al., 2012). Considering the differential expression of Lin28b and Let-7 that they identified, they hypothesized that this “axis” functions to promote fetal development of B1a cells and innate-type T cells.

Here we have asked whether perturbation of this regulatory axis can reprogram cells later than stem cells, at the committed pro-B cell stage. We have also asked whether reprogrammed cells still depend on BCR signaling for B1a cell generation and identified genes altered by this axis to understand the mechanism of reprogramming. We found that ectopic expression of Lin28b as late as the pro-B stage in adults can redirect cells to the B–1 developmental pathway. Conversely, expression of Let-7b miRNA in fetal pro-B cells can switch fetal development to B–2. Transduction of Btk-deficient precursor cells from Xid mouse BM (Hayakawa et al., 1983; Rawlings et al., 1993; Khan et al., 1995) reveals that normal BCR signaling is required for accumulation of CD5+ B1 B cells, even when they are generated from Lin28b-directed BM pro-B cells in the adult. Finally, we report a critical new role for the Lin28b–Let-7 axis can reprogram cells later than stem cells, at the committed pro-B cell stage.

Redirecting fetal and adult B cell development
at the B cell–committed pro–B stage

To assess the developmental stage at which these regulators determine development, fetal and adult cells were retrovirally transduced to express Let-7b and Lin28b. Quantitative PCR (Q-PCR) validated expression of Let-7b in FL pro–B (not depicted); cytoplasmic staining and Western blotting verified Lin28b expression in a pro–B cell line (Fig. 2 A). The developmental consequences of Lin28b expression in BM pro–B or early lymphoid progenitors were assessed by culturing transduced cells for an extended period. Many of the IgM+ cells produced in these cultures were CD5−, whereas cells transduced with empty vector control retrovirus were not (Fig. 2 B). Mature B cells are not generated efficiently in vitro, so we next performed cell transfer experiments with pro–B cells transduced to inappropriately express these regulators, i.e., Let-7 in FL and Lin28b in BM. Results, shown in Fig. 2 C, were very clear. SCID mice receiving empty vector BM pro–B cells generated GFP+ B cells that were IgM1IgD2CD5−CD23−CD21med (follicular type); in contrast, Lin28b BM pro–B cells generated GFP+ B cells that were IgM1IgD+CD5+CD232CD21med and many CD5+. In contrast, SCID mice receiving empty vector FL pro–B cells generated GFP+ B cells that were IgM2IgD+/−CD23− and many CD5−; in contrast, animals receiving Lin28b FL pro–B cells generated GFP+ B cells that were IgM1IgD+CD5+CD23+. These findings indicate that alterations in the Lin28–Let-7 axis are both necessary and sufficient for fetal and adult CD19+ B cell precursors. Microarray analyses were performed to identify genes whose expression distinguished B–1 and B–2 development at pre–pro–B and pro–B stage cells, using two different platforms, selecting genes that showed concordant results in both stages and platforms. As shown previously by RT-PCR analysis, expression of several genes key for B cell development, including Rag1, Rag2, Lambda5/Igl1, and VpreB1, does not differ significantly between these two cell types, whereas terminal deoxynucleotidyl transferase (TdT; encoded by the Dntt gene) is strikingly abundant in adult precursors but absent from those in FL (Li et al., 1993). These criteria filtered 567 Fr. A genes and 592 Fr. BC genes to a final set of 219, shown in the heat map in Fig. 1 B. Gene lists and measurements used to derive this set are provided in Datasets S1–S7.

miRNA expression was analyzed using the Agilent Technologies miRNA microarray platform, identifying a set of miRNAs that were differentially expressed, including many Let-7 family miRs that were up-regulated in BM and miR-451 that was up-regulated in FL (Fig. 1 C, top; and all data in Dataset S8). This pattern of expression was confirmed using miRNA TaqMan assays, showing the most significant differences among those tested to be miR−10a, miR−125b, miR−146, and Let−7 members in BM and miR−451 in FL (Fig. 1 C, bottom). Importantly, Lin28b and Let−7 can function as a molecular switch, in that Let-7 targets Lin28b mRNA for degradation and Lin28b functions to sequester pre–Let-7 from processing to functional miRNA (Mayr et al., 2012).
to specify fetal and adult lymphopoietic programs even after progenitors have adopted the B cell fate. These differential B cell phenotypes are shown for spleen B cells in Fig. 3 and for peritoneal cavity–resident B cells in Fig. 4. Quantitation of cell numbers engrafted in SCID mice 3 wk after transfer, shown in Table 1, reveals somewhat increased numbers in Lin28b-transduced recipients and reduced numbers in Let-7–transduced recipients. These differences likely arise from changes in cell proliferation at the pre-B stage resulting from expression of these regulators.

**Lin28b–induced BM B1a B cells: Requirement for intact BCR signaling**

The generation of normal CD5+ B cells depends on the presence of antigen (Hayakawa et al., 1999). Furthermore, mouse mutants that weaken BCR signaling have a deficit in CD5+ B cells (Tarakhovsky et al., 1995; Inaoki et al., 1997; Suzuki et al., 1999). For example, the Btk-deficient Xid mouse has fewer B cells in spleen, accumulates B cells with a distinctive phenotype, has very low levels of serum IgM, and completely lacks CD5+ B cells (Hayakawa et al., 1983; Khan et al., 1995). However, the ability of ectopic expression of Lin28b to divert cells to the B-1 fate in progenitors with defective BCR signaling has never been directly tested. Therefore, we asked whether BCR signaling is required for production of Lin28b-induced BM-derived B1a B cells. For this purpose, transduced pro-B cell isolated from BM of Xid mice were transferred into SCID mice, and then engrafted cells were examined for generation of CD5+ B cells. As shown in Fig. 5 A, even provision of Lin28b did not induce production of...
We then asked whether the failure to generate a typical B1a fetal-type repertoire was the result of relatively weak down-regulation of Dntt, the gene which codes for TdT, by Lin28b. Because increased N-addition in Ig heavy chains sequenced from B1a B cells generated by Lin28b-transduced BM pro-B cells was observed, similar transfer experiments were performed using pro-B cells from TdT−/− mice. Results from single cell sequence analysis still showed a repertoire quite distinct from that of normal B1a B cells (Fig. 5 C), indicating that N-addition in Lin28b-transduced pro-B cell development is not the reason for the different VH gene distribution.

Genes regulated by the Lin28b–Let-7 axis: Role of Arid3a
To gain insight into genes that may be influencing the development of fetal versus adult B cells as a consequence of expression of Lin28b or Let-7, fetal and adult pro-B cells were transduced as above, and then, 4 d after transduction, RNA was prepared from sorted GFP+ cells. Microarray analysis identified 219 genes as differentially expressed between normal fetal and adult pro-B cells (data in Dataset S7). Of these, 40 genes showed altered expression in Lin28b-transduced BM pro-B cells and 50 genes in Let-7–transduced FL pro-B cells (Fig. 6 A and Dataset S10). The reciprocal overlap

Figure 3. Phenotype of Lin28b–transduced BM and Let-7–transduced FL generates B cells in recipient spleen that resemble B1 and FO/B2. CD19+GFP+ mature (CD93−) B cells in spleen were generated from transfer of the indicated pro-B–transduced cells, 3 wk after transfer in SCID mice, and analyzed by flow cytometry using the indicated markers. The red boxes indicate an adult-type and the green boxes indicate a fetal-type phenotype. The blue gates represent IgM+CD21hi cells, a marginal zone B cell type. Data are representative of five independent transfer experiments with two mice per group.
Figure 4. Phenotype of Lin28b-transduced BM and Let-7–transduced FL generates B cells in recipient peritoneal cavity that resemble B1 and B2. CD19^GFP^ cells in the peritoneal cavity were generated from transfer of the indicated pro-B–transduced cells, 3 wk after transfer in SCID mice, and analyzed by flow cytometry using the indicated markers. The red boxes indicate an adult-type and the green boxes indicate a fetal-type phenotype. Data are representative of five independent transfer experiments with two mice per group.

(Fig. 6 B) identified 16 genes, in addition to Lin28b, showing an expression pattern that might be relevant to the fetal–adult developmental switch. One particularly interesting gene in the list is a transcription factor, Arid3a, also known as Bright, that has been previously implicated in enhancing Ig heavy chain expression and altering B cell responses (Schmidt et al., 2009). Importantly, the 3’ untranslated region (UTR) of the Arid3a mRNA contains several Let-7 target sites (shown in Fig. 6 C), indicating that this transcription factor could be regulated by the miR Let-7. Another miRNA differentially expressed in fetal and adult pro-B stage cells is miR–125b, previously shown to regulate Arid3a function in human and mouse B cell progenitors (Puisségur et al., 2012). Target sites for this miRNA are also shown in Fig. 6 C.

Transduction of BM pro-B cells with Arid3a-expressing retrovirus showed that this transcription factor was sufficient to support the generation of B1a B cells (Fig. 7 A). Thus, Arid3a is a key transcription factor induced by Lin28b and targeted by Let-7b for reprogramming adult B cell development to resemble that ongoing in FL. The phenotype of peritoneal cavity B cells induced by retroviral provision of Arid3a is shown in more detail in Fig. 7 B.

Table 1. Cell numbers engrafted in SCID mice by transfer of transduced pro-B cells

| Cell type (x10⁴) | BM pro-B | FL pro-B |
|------------------|----------|----------|
|                  | pMIG SE | Lin28b SE | MSCV SE | Let-7b SE |
| Spleen B1a       | 4.74 1.59 | 34.81 8.44 | 10.56 0.74 | 1.08 0.25 |
| Spleen F0        | 32.81 8.71 | 10.78 2.45 | 4.99 0.96 | 6.12 1.25 |
| Spleen MZ        | 29.57 8.01 | 12.22 3.11 | 12.83 1.12 | 4.09 0.47 |
| PerC B1a         | 7.02 2.94 | 51.04 10.67 | 40.12 10.78 | 3.14 0.71 |

M2, marginal zone. CD19^GFP^ cells of the indicated cell type were engrafted in SCID mice 3 wk after cell transfer of transduced pro-B cells. Mean and standard error (SE) are shown; for each transfer sample type, n = 4. pMIG and MSCV are control empty vectors.
functions to regulate the switch from fetal-type to adult-type lymphopoiesis, generating innate-like T and B cells. However, based on this work, it remained unclear whether altering Lin28B–Let-7 could reprogram development after commitment to the B cell fate had occurred. Here we show that B cell development can be efficiently reprogrammed as late as the CD19+ pro-B stage. We also show that retroviral provision of Let-7 is sufficient to reprogram fetal pro-B cells to generate adult-type (B2) B cell progeny. Thus, we confirm and extend the earlier study, showing that the Lin28b–Let-7 axis mediates the B cell developmental switch that we proposed many years ago (Hardy and Hayakawa, 1991, 2001).

In contrast, although we find that generation of CD5+ B cells from Lin28b-transduced adult pro-B cells requires Btk signaling, as with fetal B1a generation, it does not fully recapitulate the fetal B1a BCR repertoire. Specifically, we observe a deficit of anti-PtC B cells that use V_H11 and VH12 BCRs. One possible reason for lack of VH11 B cells, where the prototypic VDJs lack TdT-mediated N-addition at the junctions, is relatively modest regulation of TdT by Lin28b. However, even in BM pro-B cells lacking TdT, transduction of Lin28b did not recapitulate the normal fetal B1a repertoire. Considering that the typical B1a fetal-type repertoire requires appropriate initial BCR repertoire generation, selection, and accumulation, the failure of Lin28b–induced adult pro-B cells to generate a

Retroviral knockdown of Arid3a in fetal pro-B cells reprograms development

We performed a loss-of-function experiment, asking how B cell development would be affected if we reduced the amount of Arid3a in fetal pro-B cells by retroviral transduction of shRNA specific for this gene. We screened five shRNA hairpins specific for Arid3a from the Broad Institute Public TRC portal, expressing them in the Open Biosystems LMP retroviral vector, and found three that decreased the level of Arid3a 5–10-fold, as tested by Q-PCR (Fig. 8 A). We isolated pro-B stage cells from FL, cultured cells overnight, transduced them the next day with a pool of the shRNAs, and then transferred cells to SCID mice the following day. 3 wk later, recipients were sacrificed and analyzed for engraftment of GFP+ B cells. As is clear from Fig. 8 B, knockdown of Arid3a by this procedure reduced the frequency of CD5+ B cells generated from FL pro-B cells, reprogramming development to generate cells with a B2 B cell phenotype.

DISCUSSION

Yuan et al. (2012) recently showed that retroviral transduction of BM hematopoietic progenitors and stem cells could generate large numbers of fetal-type T and B lymphocytes. Based on differential expression of Lin28b and Let-7 in FL and BM B cell progenitors, they proposed that the Lin28b–Let-7 axis functions to regulate the switch from fetal-type to adult-type lymphopoiesis, generating innate-like T and B cells. However, based on this work, it remained unclear whether altering Lin28B–Let-7 could reprogram development after commitment to the B cell fate had occurred. Here we show that B cell development can be efficiently reprogrammed as late as the CD19+ pro-B stage. We also show that retroviral provision of Let-7 is sufficient to reprogram fetal pro-B cells to generate adult-type (B2) B cell progeny. Thus, we confirm and extend the earlier study, showing that the Lin28b–Let-7 axis mediates the B cell developmental switch that we proposed many years ago (Hardy and Hayakawa, 1991, 2001).

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In SCID recipients, similar to results obtained by Lin28b transduction. Finally, loss-of-function shRNA knockdown showed that diminishing Arid3a mRNA levels in fetal pro-B cells could reprogram B-1 development, generating B cells with an adult-type phenotype. Thus, we conclude that a key consequence of altering the Lin28b–Let-7 axis is changing expression levels of Arid3a.

How might Arid3a be inducing the generation of B1a B cells from BM pro-B cells? This transcription factor was originally identified by its capacity to bind to IgH V segments, increasing their expression level in a cell line treated with antigen and IL-5 (Herrscher et al., 1995). Later work showed that Arid3a was not restricted to activated B cells, but was also expressed in B cell progenitors (Webb et al., 1998). Further

Normal B1a repertoire may indicate further requirements for the generation of an appropriate initial BCR repertoire and/or fetal antigens lacking in the adult microenvironment.

Global analysis of genes reciprocally perturbed by misexpression of these regulators generated a relatively short list of candidate fetal/adult regulators. We considered the Arid3a transcription factor to be particularly interesting, based on its known functions of altering Ig heavy chain expression and inducing autoantibody production when overexpressed as a transgene (Oldham et al., 2011). Moreover, analysis of Arid3a on the TargetScan website revealed conserved Let-7 and miR-125b miRNA target sites in its 3’-UTR, both miRs which show relatively elevated expression in BM. Transduction of Arid3a into BM pro-B cells induced CD5+ B1 B cells in SCID recipients, similar to results obtained by Lin28b transduction. Finally, loss-of-function shRNA knockdown showed that diminishing Arid3a mRNA levels in fetal pro-B cells could reprogram B-1 development, generating B cells with an adult-type phenotype. Thus, we conclude that a key consequence of altering the Lin28b–Let-7 axis is changing expression levels of Arid3a.
lymphoid progenitors present in embryonic day 12.5 (E12.5) FL showed reduced numbers of both cell types, and colony-forming assays revealed reduced function of hematopoietic progenitor cells. Rare surviving progeny (<1%) were significantly smaller in size and had perturbed B lineage development, with fewer pro-B, pre-B, and immature B cells in BM and decreased numbers of transitional B cells in spleen. The follicular compartment was nearly normal, likely because of homeostatic compensation, but marginal zones were decreased and B1a B cells were essentially lacking. Thus, expression of Arid3a is required for normal early hematopoiesis and B lineage development. The difference that we describe between fetal and adult pro-B is therefore not all or none, but a difference in expression level, higher in fetal and lower in

analysis showed that Arid3a-binding site motifs were common in the promoter regions of both human and mouse V_{H} genes (Goebel et al., 2002). We speculate that Arid3a binding could alter the accessibility of V_{H} genes to rearrangement, biasing the expression of these genes, selecting for B1a “preferred” BCRs. In addition, Arid3a overexpression has been shown to perturb B cell subset generation (Oldham et al., 2011), and a dominant-negative transgene diminishes B1 B cell function (Nixon et al., 2008). Thus, there is clear precedent for BRIGHT/Arid3a altering B cell development.

Figure 7. **Arid3a induces fetal-type B cell generation from BM pro-B cells.** (A) BM pro-B cells were transduced with an Arid3a retrovirus or pMIG control and then transferred to SCID recipients. 3 wk later, spleen or PerC B cells, gated as CD19-GFP+, were analyzed by flow cytometry. The green regions indicate a fetal phenotype. Data are representative of three independent experiments using two mice per group. (B) The expression of other markers on PerC B cells generated from Arid3a-reprogrammed BM pro-B cells was analyzed by flow cytometry. Cells were gated as CD19-GFP+. The red regions indicate an adult-type and the green regions indicate a fetal-type phenotype. Data are representative of three independent experiments using two mice per group.

Figure 8. **shRNA knockdown of Arid3a inhibits FL pro-B generation of B1a B cells.** (A) FL pro-B cells or the N38 Abelson mouse line was transduced with empty vector (LMP) or vector containing each of five different shRNAs. 3 d after transduction, GFP+ cells were sorted for RNA. ARID expression was quantified by Q-PCR. Arid3a expression was first normalized to β-actin, and then the level in control vector (LMP) was set to 100. Error bars represent ±SEM. (B) Phenotype of CD19-GFP+ B cells generated in SCID mice 3 wk after transfer of fetal pro-B cells, transduced with empty vector or a pool of three shRNA constructs. The red regions indicate an adult-type and the green boxes indicate a fetal-type phenotype. Data are representative of three independent experiments using two mice per group.
adult, that may serve to tune the response of the BCR, as described below, promoting B1 generation from FL and B2 production from adult BM.

Arid3a consists of an acidic N-terminal domain, an A–T–rich ARID DNA-binding domain, a transactivation domain, a helix-loop-helix (HLH) protein interaction domain, and a short C-terminal domain. Interestingly, Arid3a has been found to shuttle between the cytoplasm and the nucleus, under the control of its HLH domain (Kim and Tucker, 2006). The transcriptional activation function in mature B cells was found to be dependent on this domain, and specific residues, mapped by mutational analysis, were shown to be critical for nuclear or cytoplasmic localization. Arid3a has been shown to interact with the Tec kinase Btk (Webb et al., 2000), and in fact, functional Btk is required for Arid3a activity (Rajaiya et al., 2005), by phosphorylation of TFII-I which then forms a tripartite complex with Arid3a and Btk. Thus, Arid3a is not functional in B lineage cells from Btk-deficient Xid mice, known to lack B1a B cells (Hayakawa et al., 1983).

Importantly, Arid3a has also been found to alter BCR signaling because of its association with BCR-containing lipid rafts (Schmidt et al., 2009). Arid3a can be palmitoylated, redirecting its localization to lipid rafts where it associates with the BCR signalsome and interacts with sUmoylation enzymes, blocking calcium flux and phosphorylation of Btk and TFII-I. Thus, higher levels of Arid3a decrease B cell responsiveness to BCR cross-linking and so function to tune BCR responsiveness. We speculate that the threshold for selection into the B1a B cell pool is altered in Lin28b/Arid3a-expressing early B lineage cells, facilitating selection of cells with BCRs that normally would be eliminated in BM (B-2) development because of their recognition of self-ligands. The requirement for selection by self-reactivity (Hayakawa et al., 1999) accounts for the observation that many B cells developing from pro-B that express Lin28b (or Arid3a) do not enter the CD5+ B cell pool. Manipulating Arid3a will be key in advancing our understanding of differences in BCR selection during fetal and adult B cell development.

MATERIALS AND METHODS

mRNA expression analysis using Agilent Technologies whole genome arrays. Total RNA, isolated as described previously (Rumfelt et al., 2006), was used for production of fluorescent-labeled probe and then hybridized to the array. RNA purity and integrity were evaluated using the 2100 Bioanalyzer (Agilent Technologies) and NanoDrop 1000 (Thermo Fisher Scientific) before probe generation. For mouse BM and FL gene expression microarray analysis, a two-color array was used. Experimental samples, BM and FL cells, were labeled with Cy3, and a common reference RNA, a pool of total RNAs from several mouse organs generated in the laboratory, was Cy5 labeled. A single color (Cy3) array format was used for the Lin28b and Let-7 transduction experiment. Hybridized slides were scanned on an Agilent Technologies scanner, and fluorescent intensities of hybridization signals were extracted using Agilent Technologies Feature Extraction software. Statistical analysis was performed in the Fox Chase Biostatistics Facility.

miRNA expression analysis using Agilent Technologies miRNA arrays. Total RNA was isolated using TRIzol reagent (Invitrogen) and washed with 80% ethanol to better retain small RNAs. The RNA quality and integrity were evaluated using the 2100 Bioanalyzer and NanoDrop 1000. 100 ng total RNA was dephosphorylated with calf intestinal phosphatase and then end-labeled with Cyanine-3-pCp using T4 RNA ligase. Labeled RNA was hybridized to the Agilent Technologies mouse miRNA microarray at 55°C for 20 h. Slides were washed according to the Agilent Technologies miRNA protocol and scanned as above, and signals were extracted as above.

Real-time PCR. miRNAs were analyzed by TaqMan miRNA assays (Life Technologies) according to the manufacturer’s protocol. In brief, cDNA was reverse transcribed from total RNA samples using specific target miRNA primers from the TaqMan MicroRNA Assay and reagents from the TaqMan MicroRNA Reverse Transcription kit. PCR was then performed using the TaqMan assay together with the TaqMan Universal PCR Master Mix using an ABI 7500 real-time PCR system (Applied Biosystems). Data were normalized using SN0202 expression. Q-PCR. Applied Biosystems IDs are as follows: miR-10a, 000387; miR-125b, 002508; miR-146, 000468; Let-7b, 000378; Let-7c, 000379; miR-451, 001141; and Sno202, 001232.

Production of Lin28B pMIG-eGFP/mCherry and Let-7b pmSCV-CFP constructs. The mouse full-length Lin28b cDNA flanked by NotI and EcoRI sites in pQCXIP retroviral vector was obtained from D. West (Fox Chase Cancer Center). Lin28b cDNA was subcloned into pMIG-EFGP retroviral vector by digesting the original construct with NotI, followed by filling in the cohesive end with T4 polynucleotide. This construct was then digested with EcoRI, releasing the Lin28b cDNA fragment. The pMIG-EFGP vector was first digested with BglII and filled in to generate a blunt end, then followed by digestion with EcoRI. The Lin28b fragment (EcoRI–blunt) was then ligated into pMIG-EFGP (EcoRI–blunt) yielding the Lin28b retrovirus. A similar procedure was used to generate an mCherry retrovirus containing Lin28b. For the miRNA expression construct, mouse Let-7b miRNA stem loop sequence identified from miRBase, along with its flanking genomic sequence of ~200 bp to preserve the putative hairpin structure and proper endogenous processing, was amplified by PCR with primers that incorporated BglII and XhoI sites. The PCR fragment was then cloned into the retroviral vector pMSCV-CFP at BglII and XhoI sites. Both retroviral constructs were verified by DNA sequencing and then used to generate retroviral supernatant by calcium phosphate–mediated transfection of Phoenix-E cells.

shRNA constructs for knockdown of Arid3a. The retroviral vector LMP from GE Healthcare was used to express the shRNAmir construct from RNA Polymerase II (Pol II) promoters (Dickins et al., 2005, 2007). These vectors produce highly efficient knockdown even when present at single copy. Hairpin sequences specific for Arid3a were from the Broad Institute Public TRC portal. Oligos for hairpins were synthesized and cloned into pMSCV-CFP retroviral vector. Retroviral constructs were verified by DNA sequencing and then used to generate retroviral supernatant by calcium phosphate–mediated transfection of Phoenix-E cells.

Animals. BALB/c, BALB/c XID, and BALB.TdT− female mice, 6–12 wk old, bred in our animal facility were used as sources of BM cells for retroviral transduction. Embryos generated from timed mating of BALB/c were used as sources of day 16 FL. BM from BALB/c Rag1−/−GFP reporter mice was used for isolation of early lymphoid progenitors. C.B-17 SCID female mice, 6–12 wk old, were used as recipients in transfer experiments. All experiments were conducted under an animal protocol approved by the Fox Chase Institutional Animal Care and Use Committee.

Retroviral transduction of pro-B cells for cell transfer and microarray analysis. OP9 cells were grown in a 37°C humidified, 10% CO2 gassed incubator in 16% FBSS/αMEM (supplemented with 1× glutamine and sodium pyruvate, 10 mM Hepes, 5 × 10–3 M 2-ME, and 0.5 mg/ml gentamycin). Cells were passaged into 24-well plates in the same medium 1–2 d before use, so as to avoid confluence. Immediately before precursor cell deposition, medium was replaced with fresh medium containing cytokine...
Culture of retrovirally transduced B cell precursors for B cell differentiation. Early B lineage precursors were purified as “early lymphocyte progenitors” (ELPs) from BALB/c Rag1-GFP transgenic BM by flow cytometry, depositing them onto OP9 cells prepared as described above. Medium was supplemented with 10 ng/ml stem cell factor (SCF), 10 ng/ml Flt3L, and 100/μl IL-7, all obtained from R&D Systems. Retroviral transduction was performed as above, and cells were cultured for an additional 3 d. Then cells were passaged onto FLST2 stromal cells in 24-well plates, growing in 5% FBS/RPMI1640 medium supplemented with 1× glutamine, 10 mM Hepes, 5 x 10⁻⁵ M 2-ME, 0.5 mg/ml gentamicin, and 100/μl IL-7. After a further 4 d, cells were again passaged into RPMI medium, but with 100-fold reduced IL-7 to promote B cell differentiation. Cells were split once more into low–IL-7 medium and then analyzed 3 d later by flow cytometry (day 15). In other experiments, pro-B cells from BALB/c BM were deposited onto FLST2 stromal cells in 24-well plates (100/μl IL-7), transduced the following day, and then replated onto FLST2 cells in 100-fold reduced IL-7. Cells were split 3 d later, split again after 3 d, and finally analyzed by flow cytometry after 4-d further culture (day 11).

Flow cytometry sorting and analysis. Cells were sorted using a BD FACSVantage SE/DIVA equipped with lasers enabling 12-color detection or a FACSArray equipped with lasers enabling 15-color detection. For microarray analysis, pre-pro-B cells and pro-B cells were isolated: for pre-pro-B as (Ter119,Ly6c,CD3,CD11b,GR1,IgM)CD93⁺CD43⁺Cd117⁺IL7Ra⁺B220⁺CD24⁻CD19⁻ and for pro-B as (Ter119,Ly6c,CD3,CD11b,GR1,IgM⁺)CD93⁺CD43⁺B220⁺CD24⁺CD19⁻. For cell culture and transfer experiments, yield was maximized by sorting pre-pro-B cells from BM and FL as (Ter119,Ly6c,CD3,CD11b,GR1,IgM⁻)CD43⁻B220⁺CD24⁺CD19⁻. ELPs were purified from BM of Rag1-GFP reporter mice as (Ter119,Ly6c,CD3,CD11b,GR1,IgM⁺)CD93⁺CD43⁺Cd117⁺IL7Ra⁺B220⁺CD24⁻CD19⁻. GFP⁺ cells were sorted into RNA lysis buffer (“Solution D”).

Bioinformatics analysis. Raw expression data obtained from Agilent Technologies two-channel microarrays were background corrected and normalized (Quantile) across experimental conditions. The LIMMA (linear models for microarray data) methodology was applied to the log2-transformed expression data to identify differentially expressed genes in each comparison. The LIMMA module in the Open Source R/Bioconductor package was used in the computations. Differentially expressed genes were identified based on statistical significance as well as biological significance. Statistical significance was measured by the false discovery rate (FDR) to account for multiple testing. For Agilent Technologies two-channel microarrays, a cutoff of FDR ≤5% and fold change ≥2 were used; for Affymetrix microarrays, a cutoff of FDR ≤20% and fold change ≥1.5 were used. Gene expression heat maps were generated using Java TreeView (version 1.1.5r2). The gene interaction networks of significant genes were generated through the use of IPA (Ingenuity Systems). All edges are supported by at least one reference from the literature, from a textbook, or from canonical information stored in the Ingenuity Knowledge Base. The intensity of the node color indicates the degree of up (red) or down (green) regulation. Nodes are displayed using various shapes that represent the functional class of the gene product. Raw microarray data are available under GEO accession no. GSE65536.

Online supplemental material. Datasets S1–S7 present microarray expression data. Dataset S8 shows miRNA expression microarray data. Dataset S9 shows heavy chain sequence data used to produce Fig. 5. Dataset S10 contains gene expression perturbated by the Lin28b–Let-7 axis for 219 genes differentially expressed in fetal and adult B cell progenitors. All datasets are included in separate Excel files. Online supplemental material is available at http://www.jem.org/cgi/content/full/jem.20141510/DC1.

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