

Development of a murine hematopoietic progenitor complementary DNA microarray using a subtracted complementary DNA library

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With the goal of creating a resource for in-depth study of myelopoiesis, we have executed a 2-pronged strategy to obtain a complementary DNA (cDNA) clone set enriched in hematopoietic genes. One aspect is a library subtraction to enrich for underrepresented transcripts present at early stages of hematopoiesis. For this, a hematopoietic cDNA library from primary murine bone marrow cells enriched for primitive progenitors was used as tester. The subtraction used 10 000 known genes and expressed sequence tags (ESTs) as driver. The 2304 randomly picked clones from the subtracted cDNA libraries represent 1255 distinct genes, of which 622 (50%) are named genes, 386

(30%) match uncharacterized ESTs, and 247 (20%) are novel. The second aspect of our strategy was to complement this subtracted library with genes known to be involved in myeloid cell differentiation and function. The resulting cDNAs were arrayed on polylysine-coated glass slides. The microarrays were used to analyze gene expression in primary and cultured murine bone marrow-derived progenitors. We found expression of various types of genes, including regulatory cytokines and their receptors, signal transduction genes, and transcription factors. To assess gene expression during myeloid differentiation, we examined patterns of change during induced differentiation of

EML cells. Several hundred of the genes underwent fluctuations in expression level during myeloid cell differentiation. The complete database, accessible on the World Wide Web at <http://yale130132115135.med.yale.edu/>, allows for retrieval of information regarding these genes. Our microarray allows for genomewide expression analysis of myeloid stem cells, which will help in defining the regulatory mechanisms of stem cell differentiation. (Blood. 2002; 100:833-844)

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Introduction

Acute myeloid leukemia (AML) remains a highly lethal malignancy requiring novel therapeutic strategies.^{1,2} An integral component of the AML phenotype is the loss of the capacity to differentiate into mature myeloid cells. Consequently, a major focus of research in this area has been on the molecular mechanisms controlling normal myeloid differentiation. One conclusion of this work is that differential expression of key regulatory genes in hematopoietic stem cells controls their differentiation into mature cell types, including erythrocytes, platelets, neutrophils, monocytes, eosinophils, and basophils. A detailed understanding of the gene expression patterns throughout hematopoietic differentiation obtained by means of messenger RNA (mRNA)–expression profiling and bioinformatics can provide valuable insights into this complex process and will perhaps lead to novel treatment approaches for AML. We are interested in the patterns of gene expression at early stages of myeloid commitment and differentiation. Previous studies have identified a small but diverse group of genes that are down-regulated during this process, including *CD34*,³ *ckit*,⁴ *Jagged2*,⁵ *mpl*,⁶ *sca-1*,⁷ *SCL*,⁸ *GATA-1* and *GATA-2*,⁸ *Flt-1*,⁹ *Notch*,¹⁰ *Ap-1*,¹¹ *Mzf-1*,¹² *C/ebp*,¹³ and *STATs*.¹⁴ Up-regulated genes include *Pu.1*¹⁵ and others.¹⁶ However, there are likely to be many more genes, some known and some yet to be identified, involved in the molecular events of differentiation.^{17,18}

To better understand the interacting pathways and networks involved in hematopoiesis, we decided to employ the genomewide strategy of gene expression profiling using complementary DNA (cDNA) microarrays.¹⁹ A requisite component of this technology is inclusion of potentially critical genes on the array. With the aim of developing a cDNA microarray for use in the study of gene expression during early myelopoiesis, we constructed a subtracted cDNA library derived from sorted hematopoietic progenitor cells, and we complemented this set of genes with a set of available clones known to be important to myelopoiesis. This clone set was sequenced, characterized, and then spotted onto glass slides to create a microarray for analyzing the profile of gene expression during early steps in myelopoiesis. We have employed this microarray to assess expression in primary hematopoietic precursors and to analyze changes in gene expression during induced differentiation of the myeloid progenitor cell line EML.

Materials and methods

Reagents

The α -[³²P]deoxycytidine 5′triphosphate (α -[³²P]dCTP) (3000 Ci/mmol [111 TBq/mmol]) was purchased from Amersham Pharmacia Biotech

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(Buckinghamshire, United Kingdom), and restriction enzymes were purchased from New England Biolab (Beverly, MA). Iscoves modified Dulbecco medium (IMDM) (Life Technologies, Rockville, MD) supplemented with 20% horse serum was the culture medium used throughout. Recombinant human stem cell factor (SCF) was purchased from Peprotech (Rocky Hills, NJ). Rhodamine123 and Hoechst 33342 dyes were from Molecular Probes (Eugene, OR). Plasmid vectors λ ZipLox and pSport were from Life Technologies (Rockville, MD).

Cell lines

The EML and EPRO cell lines²⁰ were generously contributed by S. Tsai. EML cells were maintained in IMDM supplemented with 20% horse serum, 15% BHK/MKL-conditioned medium (containing SCF), 1% L-glutamine, 1% penicillin/streptomycin/amphotericin (P/S/A), and 1% nonessential amino acids. EPRO cells were maintained in IMDM supplemented with 20% horse serum, 10% HM-5-conditioned medium (containing granulocyte-macrophage colony-stimulating factor), 1% L-glutamine, 1% P/S/A. Cell lines were cultured at 37°C in 5% CO₂. EML cells were induced to differentiate into myeloid cells with 10 μ M all *trans*-retinoic acid (ATRA) and 5% WEHI-conditioned medium as a source of interleukin-3 (IL-3).

cDNA library for subtraction

The lineage⁻rhodamine^{Low}Hoechst^{Low} (Lin⁻rhodamine^{Low}Hoechst^{Low}) (LRH) library was derived by means of previously described techniques,^{21,22} and its construction has been described.²³ Briefly, primary bone marrow cells were depleted of lineage-committed cells and then further enriched for primitive cells by fluorescent-activated cell sorting for cells with low-level staining with rhodamine123 and Hoechst 33342 dyes. From 30 mice, 5000 cells were obtained, from which a directionally cloned cDNA library was created in the lambda vector λ ZipLox (Gibco BRL) as *SalI*-*EagI* fragments, in such a way that the 5' end of the cDNA was adjacent to the *SalI* site and the 3' end was adjacent to the *EagI* site. The original library had an initial plating complexity of 1.44×10^7 clones.²³

The LRH library was converted to single-stranded DNA (ssDNA) by *in vivo* excision via cre-mediated excision and filamentous phage rescue as described.²⁴ Briefly, we electroporated 50 ng library DNA into competent *Escherichia coli* DH5 alpha F' bacteria, F' ϕ 80 Δ lacZ Δ M15 Δ (lacZYA-argF) U169deoR recA1 endA1 hsdR17 (r_K⁻, m_K⁺) phoA supE44 λ -thi-1 gyrA96relA1. We incubated the transformed bacteria in 100 mL 2 \times YT broth at 30°C on an orbital shaker for 1 hour and then added 150 μ L ampicillin (50 mg/mL) and 1 mL 20% glucose. Bacteria were grown overnight until OD₆₀₀ = 0.1 (OD indicates optical density), followed by incubation at 37°C for 1 hour until OD₆₀₀ = 0.2. The culture was superinfected with M13 KO7 helper phage and then cultured an additional 2 hours. After eliminating bacteria by centrifugation filamentous phage particles were precipitated with the addition of 4 g polyethylene glycol and 2.92 g NaCl into 100 mL solution and incubation at 4°C for 16 hours. Particles were collected by centrifugation; the phage DNA was purified with phenol-chloroform extraction; and the final product was dissolved in TE (10 mM Tris-HCl pH 7.9, 1 mM EDTA). The ssDNA was confirmed by digestion with mung-bean nuclease. This yielded approximately 25 mg single-stranded phage DNA composing the entire LRH library of cDNAs.

Prior to hybridization, the single-stranded library DNA was purified by means of hydroxyapatite (HAP) column chromatography to eliminate double-stranded DNAs (dsDNAs). First, 10 μ g library ssDNA was digested with *PvuII*, and then it was applied to a 10-mL HAP column. The flowthrough (7 to 8 mL), representing the ssDNA, was collected and concentrated by means of Qiagen (Valencia, CA) spin columns following the manufacturer's protocol. DNA eluted from the Qiagen spin column was precipitated and resuspended in 5 μ L double-distilled water (ddW). The ssDNA was confirmed by digestion with mung-bean nuclease.

Preparation of driver DNA

The DNA driver pool was prepared with 10 000 mouse cDNA clones that were a gift from Research Genetics (Huntsville, AL). These clones were derived from mouse testes, kidney, diaphragm, skin, lung, brain, heart, and whole embryonic

fetus; mouse melanoma; embryonic carcinoma; and mouse macrophages. The inserts were amplified by polymerase chain reaction (PCR) with Expand high-fidelity PCR system (Invitrogen, Carlsbad, CA) under the following conditions: 94°C, 7 min for 1 cycle; 20 cycles at 94°C for 1 minute, 55°C for 2 minutes, and 72°C for 3 minutes; and a final extension of 7 minutes at 72°C. The PCR products were purified by phenol-chloroform extraction and checked by ethidium bromide-stained agarose gel electrophoresis. All inserts were combined to make the driver pool by transfer of 2 μ L from each PCR product.

Subtraction of cDNA library

Subtraction of the LRH library was performed essentially as described by Bonaldo et al²⁴ with minor modifications. The hybridizations of library ssDNA and pooled driver DNA were performed in 20 μ L volume hybridization buffer (50% formamide, 0.12M NaCl, and 1% sodium dodecyl sulfate [SDS]) with 2.5 μ g driver DNA and 50 ng tracer ssDNA from cDNA library at 30°C for 110.4 hours (C₀t = 50; here, C₀ is substrate concentration of total DNA in solution, and t is hybridization time at 30°C). To block hybridization via the vector and poly(adenylic acid) (poly[A]) tail sequences, blocking oligonucleotides were designed (Table 1) and were included in the hybridization at a concentration of 2 μ g/ μ L for blocking vector homology sequence and 0.5 μ g/ μ L for blocking poly(A) tail sequence. Following hybridization, DNA molecules remaining single stranded were purified by HAP chromatography and were concentrated to 11 μ L volume as described above. The ssDNA was converted into dsDNA *in vitro* by transferring the ssDNA into premixed reaction solution, (5 μ L sequenase buffer [5 \times] and 1 μ L M13 forward primer [1 μ g/mL]), heating at 65°C for 5 minutes, then 37°C for 3 minutes, adding 2 μ L deoxynucleoside 5'-triphosphates (10 mM each), 1 μ L dithiothreitol (DTT) (0.1 M), and 1 μ L sequenase (5 U/ μ L) into reaction solution, incubating at 37°C for 30 minutes, and then purifying the dsDNA with phenol-chloroform extraction.

The resultant dsDNA was transformed into *E coli* DH10 α , which was then plated on Luria-Bertani broth/ampicillin agar plates. The total number of clones were calculated. An aliquot of the subtracted LRH library was submitted to Lawrence Livermore National Laboratory (Livermore, CA) for transformation, plating, and robotic picking of colonies into 96-well plates as part of the Cancer Genome Anatomy Project (National Institutes of Health, Bethesda, MD). A separate aliquot of the libraries was amplified as a population and used to prepare DNA.

DNA preparation and sequencing

Plasmid DNA was prepared in 96-well plates. The clones derived from the subtracted cDNA libraries were grown in 96-well plates, and plasmids were isolated by the alkaline-lysis method. The final plasmid was dissolved in 100 μ L ddW. (Note: for sequencing purposes, we used 20 μ L plasmid directly, and for arraying, we further purified the plasmid DNA with a 96-well filter plate. The picked clones were sequenced with single-pass automated sequences by the W. M. Keck Facility at Yale University (New

Table 1. Blocker sequences for library subtraction for LRH library

Vector	Blocker sequences
pSport	
3' end	5' gtg agc gga taa caa ttt cac aca gga aac agc tat gac cat gat tac gc 3'
	5' caa gct cta ata cga ctc act ata ggg aaa gct ggt acg cct gca ggt ac 3'
5' end	5' ggc cgc tct aga gga tcc ggg ctt acg tac gcg tgc atg cga cgt cat ag 3'
	5' ctc ttc tat agt gtc acc taa att caa ttc act ggc cgt cgt ttt aca ac 3'
	5' gtc gtc act ggg aaa acc ctg gcg tta ccc aac tta atc gcc ttg cag 3'
NA	5' ttt ttt ttt ttt ttt 3'

NA indicates not applicable.

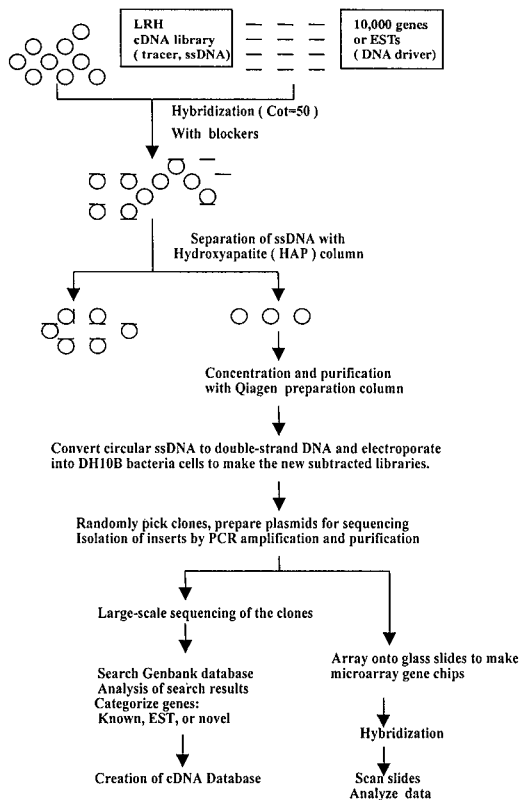


Figure 1. Schematic for production of the normalized hematopoietic progenitor cDNA library and construction of microarray gene chips. At top, single-strand circles represent the starting cDNA library produced by means of filamentous phage rescue. The ssDNA was hybridized with driver (10 000 IMAGE consortium cDNAs) with appropriate blocking oligonucleotides. The fraction that remains single stranded (flowthrough from HAP column) was converted to double-strand circles, electroporated into DH10B, and propagated under ampicillin selection to generate an amplified normalized cDNA library. Large-scale sequencing of clones was performed with the use of the M13AEK forward primer. To make the myeloid-specific gene chips, the sequenced cDNA clones were amplified by PCR, purified, and printed onto polylysine-coated glass slides.

Haven, CT) and/or the Genome Sequencing Center at Washington University Medical School (St Louis, MO) with the use of an M13AEK forward primer (5' CAA AAG GGT CAG TGC TG 3'), which primes synthesis at the 3' of clones. Some clones were also sequenced from the 5' end with the use of the T7 promoter primer (5' TAA TAC GAC TCA CTA TAG GG 3'). The M13/pUC reverse primer (AGC GGA TAA CAA TTT CAC ACA GGA) for 5' termini was used to confirm LRH novel sequences.

Sequence editing and analysis

Because some sequencing primers contained common vector sequence, we first removed vector sequences from the sequences with CodonCode-Cross_Match software (<http://www.codoncode.com>). FASTA formatted DNA sequences were compared with known nucleotide sequences with the use of the Blast algorithm in batches of 3228 sequences and the use of the blastall program (BLASTN and BLASTX programs²⁵) installed in a Dell Workstation with a Linux operating system. Three publicly accessible databases were searched: Genbank nonredundant (nr) nucleotide, database for expressed sequence tags (dbESTs), and Genbank nr protein. Internal redundancy within our clone set was determined by comparison of each sequence against our own database. Categorization of sequence homology was based on the following criteria: exact match to known named mouse genes (threshold score exceeding 200) or protein, or near-identity to a known gene or protein from a species other than mouse (usually either human or rat); EST only (no extensive homology to any published or characterized protein, but identity to ESTs from mouse, rat, or human); or novel (no

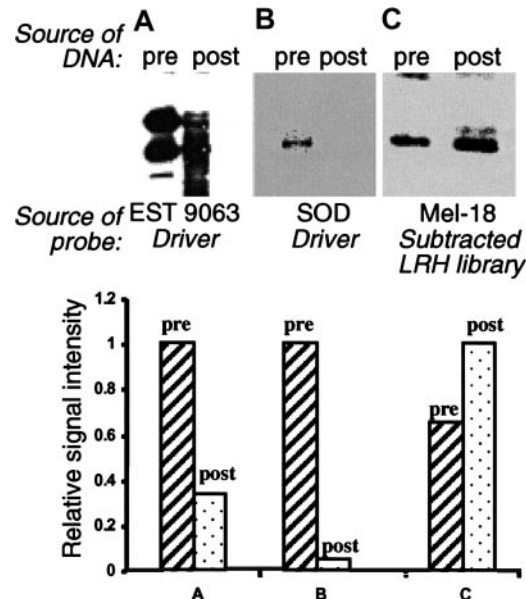


Figure 2. Southern blot analysis of subtracted LRH cDNA library. Pooled plasmids from the presubtracted (pre) and postsubtracted (post) libraries were double-digested with *Bam*HI/*Eco*R1 and analyzed by Southern blot. The probes were as follows: EST 9063, derived from the driver pool (panel A); superoxide dismutase (SOD), derived from the driver pool (panel B); and Mel-18, derived from subtracted LRH cDNA library (panel C). The bar graphs present quantitation of the data, revealing the effective elimination of the representative driver sequences and enrichment of the selected clones not present in the driver.

extensive homology to any nucleotide or protein sequence in these public databases). Sequence data from 5' and 3' sequence reads were assembled with the use of the PHRAP software package (<http://www.phrap.org/>) kindly provided by Phil Green (Washington University). Protein motifs within the assembled sequences were identified by converting the DNA sequence to open reading frame using the ORF analysis program (<http://curagen.com/>) (CuraGen, New Haven, CT) and then performing domain searches with Pfam, ProDom, Prosite, and Prints software programs (<http://curagen.com/>) (CuraGen). Cutoff parameters for match selection were *P* < .05; identities exceeded 40%, and positives exceeded 50%.

Southern hybridization

Five micrograms of library DNA was double-digested with restriction enzymes *Bam*HI and *Eco*R1, fractionated on 0.8% agarose gel, and transferred to nylon membranes.²⁹ Hybridization probe DNAs were cut with restriction enzyme, gel-purified, and labeled with random primer DNA labeling. The labeled probes were purified with Sephadex G-50 Quick Spin Column (Boehringer Mannheim, Germany), and Southern blot analysis was performed according to standard methods.

Table 2. Summary of subtracted cDNA library from Blast search of 2304 LRH clones

Type of clone	No.
Nonredundant clones*	
Novel	247
ESTs	386
Known	622
Redundant	599
Ribosomal	10
Bad sequences	80
Vector	309
Mitochondrial	35
Bacterial	16

*There were 1255 clones in this category.

Table 3. Examples of known genes in subtracted cDNA library placed into functional categories

Category/gene name	Description	Log ₂ (EML24/EML0)	Log ₂ (LH ^{LowR} ^{Bright} /LH ^{LowR} ^{Low})
Transcription factors/DNA binding proteins			
Cbfb	Core binding factor β	—	- 0.9 ± 0.2
Cebpa-rs1	Transcription regulator related to C/ebp α	- 0.9 ± 0.3	—
Cebpb	Transcription regulator C/ebp β	—	—
Cebpg	Controls the activity of IL-4 promoter activity	- 1.9 ± 0.68	- 2.5 ± 0.29
Creml	Transcriptional modulator of cAMP responsive genes	—	- 1.5 ± 0.7
Elk4	Member of ETS oncogene family	- 1.2 ± 0.5	- 0.9 ± 0.08
Fos	Part of AP1; propagation of mitogenic signals	- 1.2 ± 0.4	- 2.0 ± 0.05
Hmga1	High-mobility group AT-hook 1	—	—
Hoxa5	Homeobox A5	—	—
Jun	Jun oncogene, part of AP1 transcriptional complex with Fos	—	—
Klf9	Erythroid-specific transcription	1.8 ± 0.53	- 2.5 ± 0.15
Lrf	BTB-POZ transcription factor, dimerizes with Bcl6	—	- 2.7 ± 0.1
Mint	Organizes transcriptional complexes	2.9 ± 0.3	2.5 ± 0.7
Mlf2	Related to Mlf1, a partner in t(3;5) with Npm in leukemia	- 2.9 ± 0.45	- 2.4 ± 0.3
Nkx2-3	Regulates emigration of circulating leukocytes	- 1.6 ± 1.0	—
Sox4	Important for very early B-cell differentiation	- 2.1 ± 1.1	- 2.7 ± 0.26
Sox5	SRY-box containing gene 5	—	- 3.1 ± 0.22
Tal1	T-cell acute lymphocytic leukemia 1	—	—
Xbp1	Essential for plasma cell differentiation, liver development	1.6 ± 0.4	- 1.1 ± 0.5
Signaling proteins			
Arhgef1	Rho guanine nucleotide exchange factor (GEF) 1	- 0.9 ± 0.58	—
Calm	Calmodulin, component of calcium signaling pathway	—	- 1.2 ± 0.13
Calr	Calreticulin, major ER calcium-binding protein	—	- 2.2 ± 0.3
Camkk2	CaM-kinase kinase β, regulates calcium signaling	—	- 3.1 ± 0.2
Ccnd1	Cyclin D1, cell cycle regulation	- 1.7 ± 1.0	- 2.7 ± 0.2
Csnk1a	Casein kinase i, α isoform	—	- 2.9 ± 0.45
Hus1	Cell cycle checkpoint protein	- 1.4 ± 0.07	- 3.1 ± 0.32
Iqgap	IQ motif containing GTPase-activating protein 1	—	—
Itpr3	Inositol 1, 4, 5-triphosphate receptor 3	—	—
Lats1	Large tumor suppressor, modulates CDC2 activity	—	—
Map2k3	Mitogen-activated protein kinase kinase 3	—	—
Nsp3	Novel SH2-protein; couples Eph receptors to R-Ras	—	- 2.1 ± 0.06
Parg1	PTPL1-associated RhoGAP 1; activates Rho	ND	- 2.7 ± 0.55
Pde4b	Phosphodiesterase 4B, cAMP specific	—	—
Pde8a	Phosphodiesterase 8A, cAMP specific	—	—
Rala	Ras-related GTPase	—	- 0.74 ± 0.22
Rangap1	RAN GTPase-activating protein 1	—	—
Tax1bp1	Antiapoptotic; involved in IL-1 signaling pathway	0.6 ± 0.35	—
Tpd52	Regulator of cell proliferation, gene transcription	- 2.4 ± 0.2	- 0.9 ± 0.02
Traf1	TNF receptor-associated factor 1	—	—
Traf6	TNF receptor-associated factor 6	—	—
Tyk2	Plays a restricted role in IFNα signaling	- 2.1 ± 0.2	- 2.6 ± 0.2
Vav3	Activates Rho GTP-binding proteins	—	—
Wnk1	Serine/threonine protein kinase	- 1.9 ± 0.5	- 2.5 ± 0.4
Growth factors			
Efnb1	Ephrin B1, cek5 receptor ligand	—	—
Hdgf	Hepatoma-derived growth factor; a mitogen for fibroblasts	—	- 1.4 ± 0.04
Hegfl	Heparin-binding EGF-like, potent mitogen	—	- 3.2 ± 0.12
Scgf	Controls the proliferation of hematopoietic stem cells	- 1.4 ± 0.17	0.93 ± 0.04
Surface molecules/membrane proteins			
Fgfr1	Basic fibroblast growth factor receptor 1 precursor	—	- 3.1 ± 0.1
Gpr21	G protein-coupled receptor 21 (GPR21)	—	- 0.65 ± 0.1
H2-K	Class I antigen of MHC, mediates transplant rejection	—	- 1.5 ± 0.08
Hegfl	Heparin-binding EGF-like growth factor	—	—
Itn1	Integral membrane protein 1; unknown function	—	- 1.5 ± 0.08
Ncam2	Neural cell adhesion molecule 2	—	—
Oprs1	Opioid receptor, σ1	—	—
Tnfrsf1b	TNF receptor superfamily, member 1b	—	—

Preparation of DNA samples for arraying

Bacterial cultures were grown overnight in 96-well culture plates (Qiagen), and plasmid DNA was prepared as described above. The cDNA inserts were amplified by means of PCR (96-Well GeneAmp PCR System 9700) (Perkin

Elmer—Applied Biosystems, Foster City, CA) in 96-well plates with the use of M13 AEK forward and reverse primers (1 μM) for amplification. The PCR reaction was carried out in 100 μL solution of 1 mM deoxyadenosine 5'-triphosphate (dATP), dCTP, deoxyguanosine 5'-triphosphate (dGTP),

Table 3. Examples of known genes in subtracted cDNA library placed into functional categories (continued)

Category/gene name	Description	Log ₂ (EML24/EML0)	Log ₂ (LH ^{LowR} Bright/LR ^{LowH} Low)
Related to blood cell function			
Gpx1	Glutathione peroxidase; red cell antioxidant defense	1.2 ± 1.0	—
H2-Aa	Histocompatibility 2, class II antigen A, α	—	—
Ncf1	NADPH oxidase subunit (47 kd)	—	0.14 ± 0.1
Prg2	Eosinophil major basic protein	—	—
Scya19	Small inducible cytokine A19	—	- 2.8 ± 0.3

The Log₂ ratios of EML24/EML0 and LH^{LowR}Bright/LR^{LowH}Low were the average values of 3 experiments and normalized the signal intensity of each hybridization to internal control glyceraldehyde-3-phosphate dehydrogenase. A minus sign in front of a Log₂ ratio indicates downregulation (and the absence of a minus sign indicates upregulation) of (1) gene expression when EML cells were treated with all ATRA/IL-3 for 24 hours, or (2) in primary sorted bone marrow cells, LH^{LowR}Low relative to LH^{LowR}Bright. A data cell containing only a minus sign indicates no change.

LR^{LowH}Low indicates lineage-minus (Lin⁻)rhodamine^{Low}Hoechst^{Low}; LH^{LowR}Bright, Lin⁻Hoechst^{Low}rhodamine^{Bright}; cAMP, cyclic adenosine monophosphate; GTPase, guanosine 5'-triphosphatase; Eph, ephrin; TNF, tumor necrosis factor; IFN, interferon; GTP, guanosine 5'-triphosphate; EGF, epidermal growth factor; MHC, major histocompatibility complex; and NADPH, nicotinamide adenine dinucleotide phosphate.

Table 4. List of domains identified in novel genes by means of multiple domain search tools

Clone ID	Gene Ontology category	Description	Searching tool
2405	Cell cycle regulator	Cell division control protein 54	ProDom
2073	Chaperone	Heat shock 70-kd protein. ATP-binding; heat shock;	ProDom
2247	Enzyme	Prenyltransferase and squalene oxidase	Pfam
2656	Enzyme	E-class p450 group i signature (motif source owl:s22339)	Prints
2100	Enzyme	Acetyl esterase	ProDom
2923	Enzyme	Cyclase adenylate type ATP	ProDom
2632	Enzyme	Sulfite reductase (NADPH) flavoprotein	ProDom
3121	Enzyme	NOS-type oxidoreductase	ProDom
3332	Enzyme	Nitrate reductase (NAD[P]H)	ProDom
2363	Enzyme	Nitrate reductase	ProDom
2941	Enzyme	Acyl-coA oxidase	Pfam
2255	Enzyme	Dehydrogenase γ-glutamyl	ProDom
1860	Enzyme	3-isopropylmalate dehydrogenase precursor	ProDom
2247	Enzyme	Farnesyltransferase β subunit	ProDom
1667	Enzyme	DNA-directed DNA polymerase	ProDom
2148	Enzyme	Preprotein translocase SECA subunit	ProDom
2821	Enzyme	Glutathione S-transferases	Pfam
3063	Enzyme inhibitor	Trypsin inhibitor, serine protease inhibitor, 41 aa	ProDom
2940	Ligand binding	Phosphoribulokinase/recF protein	Pfam
2839	Ligand binding	Pyridine nucleotide disulfide reductase class-II signature	Prints
2199	Nucleic acid binding	Ribosomal protein 40S S28 S33 repeat acetylation	ProDom
1248	Nucleic acid binding	U1 small nuclear ribonucleoprotein	ProDom
2097	Nucleic acid binding	Chromosome segregation protein SMC2	ProDom
2097	Nucleic acid binding	λ and other repressor helix-turn-helix signature	Prints
2625	Nucleic acid binding	Protein ATP-binding	ProDom
2526	Nucleic acid binding	Transmembrane protein HWLF3 late glycoprotein	ProDom
2524	Nucleic acid binding	Elongation factor Tu mitochondrial precursor protein	ProDom
1998	Nucleic acid binding	Ribosomal protein L20 signature	Prints
1497	Nucleic acid binding	Protein homeobox DNA-binding	ProDom
3103	Nucleic acid binding	Transcriptional regulator YCF30 protein transcription	ProDom
3132	Nucleic acid binding	Double-stranded RNA-specific editase 1	ProDom
2952	Nucleic acid binding	Homeobox protein GSH-1	ProDom
2940	Nucleic acid binding	Exonuclease ABC subunit a SOS response excision	ProDom
2704	Signal transducer	Dopamine receptor. G-protein coupled	ProDom
1984	Signal transducer	Receptor polypeptide gastric inhibitory precursor (GIP-R)	ProDom
3023	Signal transducer	Receptor precursor repeat glycoprotein	ProDom
1981	Signal transducer	SL cytokine precursor (FLT3 ligand)	ProDom
3001	Signal transducer	Tyrosine-protein kinase JAK3	ProDom
2995	Signal transducer	mapk/erk kinase kinase 1	ProDom
2366	Signal transducer	Gap junction cx32.7	ProDom
2316	Signal transducer	Chemotaxis protein transducer	ProDom
2316	Signal transducer	HAMP domain/methyl-accepting chemotaxis protein (mcp)	Pfam
1925	Signal transducer	Insulinlike receptor precursor	ProDom
2858	Signal transducer	Tyrosine-protein kinase JAK3	ProDom
3091	Transporter	Transport protein precursor signal periplasmic	ProDom
1939	Transporter	Calcium calcium-transporting ATPase plasma membrane	ProDom

NOS indicates nitric-oxide synthase.

and deoxyribothymidine 5'-triphosphate (dTTP); 1.5 mM MgCl₂; and 2.5 U Taq polymerase in 96-well plate with the following cycles: 5 cycles of 94°C for 50 seconds, 55°C for 1 minute, and 72°C for 1.5 minutes; followed by 30 cycles of 94°C for 30 seconds, 56°C for 1 minute, and 72°C for 1.5 minutes; and then 1 cycle of 72°C for 10 minutes. Resulting PCR products were purified with the use of a 96-well glass-fiber filter (MAFB NOB) (Millipore, Bedford, MS) according to the manufacturer's user manual. The purity and yield were approximated by running the purified PCR products on a 0.8% agarose gel. The DNAs were prepared for arraying by transferring 5 μ L to 384-well plates and adding SSC to a final concentration of 3 \times . Glass slides were prepared for printing and arrayed by the Yale Microarray Facility (http://info.med.yale.edu/wmkeck/dna_arrays.htm) with the use of a GeneMachines (San Carlos, CA) Omnigene Arrayer. After printing, the slides were postprocessed as described by P. Brown and J. DeRisi (<http://www.microarrays.org.protocols.html>).

RNA preparation, probe labeling, and hybridization

Total RNA was prepared by means of Trizol reagent (Life Technologies). Microarray slide hybridization was performed as follows. The cDNA probes were synthesized by reverse transcription with oligo deoxythymidine (dT) as primer, incorporating allyl amine-deoxyuridine triphosphate (aa-dUTP) (Sigma, St Louis, MO) into synthesized cDNA. Reactions were performed in 100 μ L reaction with the following final concentrations: 85.8 μ g/mL oligo dT primer; 0.5 mM each dATP, dCTP, and dGTP; 0.2 mM aa-dUTP; 0.3 mM dTTP; 10 mM DTT; and 1280 U MLV reverse transcriptase (SuperscriptII) per milliliter. Coupling of cyanine-3 (cy-3) or cy-5 dyes to aa-dU-modified cDNAs was done with the use of NHS-ester cy-3 or cy-5 dye (Pharmacia, Piscataway, NJ) by incubation at 25°C for 1 hour in subdued light. Hybridization of fluorescently labeled probes to glass slides was performed with hybridization buffer (50% deionized formamide, 12.5% SSPE, 0.625% SDS, 1.5 \times Denhardt reagent with blockers [0.5 μ g/ μ L mouse Cot 1 DNA, 0.1 μ g/ μ L poly(A) (15A), and 0.2 μ g/ μ L yeast transfer RNA] at 42°C for 18 to 24 hours. After hybridization, the slides were washed first with 1 \times SSC, 0.1% SDS, at 25°C for 15 minutes, then with 0.2 \times SSC, 0.1% SDS, and finally with 0.2 \times SSC. The slides were scanned with a GSI Lumonics (Packard, Billerica, MA) or Axon (Axon Instruments, Union City, CA) laser scanner. Analysis of the fluorescent hybridization signal of microarray slide was performed with Quantarray (Packard) or Genepix software (Axon Instruments, Union City, CA), and the data were analyzed by means of Microsoft Excel. Further clustering of data was performed with the use of Genespring software (Silicon Genetics, Redwood City, CA).

Northern blot analysis

Northern hybridization was carried out following standard methods. Total RNA (10 μ g) was electrophoresed on a 1% agarose/formaldehyde gel and was blotted onto Hybond-N nylon membranes (Amersham Pharmacia Biotech) followed by UV cross-linking. DNA probes were labeled with random primers, and the hybridization was performed at 65°C for 16 hours. Signals on the washed filter were visualized by autoradiography.

Results

Creation of a subtracted myeloid cDNA library that is enriched for low-abundance transcripts

With the long-term goal of fully characterizing changes in gene expression during the early stages of myelopoiesis, we wanted to develop a cDNA microarray that was enriched in genes expressed in primitive hematopoietic cells and early committed myeloid cells. We took a 2-pronged approach to achieve this: one prong was to create a subtracted cDNA library from an early hematopoietic library; the second was to complement this set

with available genes known to be involved in myelopoiesis. As starting points for the library subtractions, we used cDNA library LRH, derived from primary bone marrow samples that were sorted for early progenitors by flow cytometry (Degar et al²³). The initial complexity of the LRH cDNA library was 1.44×10^7 clones. Subtraction of the libraries was performed by using as driver a pool of 10 000 mouse Integrated Molecular Analysis of Genomes and Their Expression (IMAGE) Consortium cDNA clones that were derived from several different mouse organ cDNA libraries (Figure 1). Following subtraction, we obtained 1×10^6 total clones; the complexity within this population is not known.

To assess the efficacy of the subtraction process, we performed Southern blot analysis of library-derived cDNA populations derived before and after normalization. As hybridization probes, we used 3 different sequences known to be present in both the driver and the tracer populations. The results (Figure 2) clearly indicate that the subtraction

A WEE1 kinase alignment with Clone 1317

```

C1. 1317  EEEEEEEEG--EAPIG---THRFPDQCPSPDIPPEEPESHLYQVYTSQ--LPLGLSHQI
WEE1 (Hs)  EEEEEEEEGSHSTGEDSAFQPPDPLPSARSAAEAQRRRRSPGAPSSPGELEDDI
WEE1 (Mm)  EEEEEEEEGSHSTGEDSAFQPPDPLPSARSAAEAQRRRRSPGAPSSPGELEDDI
WEE1 (Rn)  EEEEEEEEGSHSTGEDSAFQPPDPLPSARSAAEAQRRRRSPGAPSSPGELEDDI

```

B Zinc finger protein CNBP alignment with Clone 1508

```

C1. 1508  KVARCGVLEEHRVAVGG--SGQLE---LFCISLEDAELKKPLQVHPADQWMEDE
CNBP (Mm)  KCRSSHWL--ECITSGCGGNMRSRSGQFQVSSSLDPIIYCCGESHAKLQCDLQED
CNBP (Hs)  KCRSSHWL--CPGGGRKCRGFRKCTTSRRCQVSSSPDLCRCGESHAKLQCDLQED
CNBP (Rn)  KCRSSHWL--CPGGGRKCRGFRKCTTSRRCQVSSSPDLCRCGESHAKLQCDLQED
CNBP-Ho1  KCRVGHILRDCRNGGGSGRFRGGHRSNMNCTACGSYGHAADCTNIVKCYCGKIGH
CNBP-Ho2  KCRSSHWL--CPGGGRKCRGFRKCTTSRRCQVSSSLDPIIYCCGESHAKLQCDLQED

```

C DNA binding protein GLI alignment with Clone 2265

```

C1. 2265  HSEQLLITQVFF--CRNGICSEQLSPFSSPY
GLI1 (Mm)  GERRFVHWQEFVCHWGGCSRRLRPFKAGY
GLI (Hs)  NSEHHEERREVFCHWGGCSRRLRPFKAGY
GLI4 (Hs)  KCKQDPHHIYENINHWDCSKLFDNDQQLV
GLI2 (Hs)  CKQEAELVYENINHWDCRKYDNDQQLV
GLI (Dm)  NNDHTQTNKRAFVCRWDCTRGEKPFKAGY
GLI3 (Ga)  HSEQLPVVYVYENINHWDCSRRLRPFKAGY

```

D C-ETA protein alignment with Clone 1405

```

C1. 1405  PKLRMSGPFVRSVSNYIKGQALSPISLSSLELVS
PKCL (Hs)  PRINSRD--VSNFDPDFIKERFVLTPID--EGHLPIN
PKCL (Mm)  PRINSRD--VSNFDPDFIKERFVLTPID--EGHLPIN

```

E IGF growth factor alignment with Clone 1797

```

C1. 1797  GTEQIALDRRRNTETAVASLEHSLIPASGGLEGL
INS (Mm)  GERGFYTFMSRREVEHPVQALELGGPAGDQLTL
INS1 (Mm)  GERGFYTFMSRREVEHPVQALELGGSP--DQLQTL
INS2 (Mm)  GERGFYTFMSRREVEHPVQALELGGPAGDQLTL
INS (Hs)  GERGFYTFMSRREVEHPVQALELGGPAGDQLTL
INS (Sh)  GERGFYTFMSRREVEHPVQALELGGPAGDQLTL

```

F GPR1/FUN34/yaaH. GTW motif alignment with Clone 2131

```

C1. 2131  EVWEVLEFLEDFLIVYENAIKRAFYSSALSLA
GPR1 (Y1)  LFGVCCVQAQMWVYQENTFGAAALTSYGFVW
Y215 METH  FANGGILQILASVMEFRGNTFFGTVAAGSYGLFW
YAAH ECOLI  IFPGGILQIAGILEKKNFTFLAATSYGFVW
YD84 YEAST  HFGGAILQIAGLLCVYGDYAMVTSYGFVW
YF34 YEAST  HFGGILQIAGIMEALENTFGGIALCSYGFVW
YCQ0 YEAST  HFGGILQIAGIMEALENTFGGIALCSYGFVW

```

G MEKK1 protein motif alignment with Clone 2995

```

C1. 2995  SAQREPARGAARQEPGEG--RG
MEKK1 (Mm)  GDGGEMSNLAELC---GEQAG--
M3K1 (Rn)  GDGGEMSNLAELC---GEQAG--
MEKK1 (Hs)  GDGGEMSNLAELC---GEQAG--

```

H JAK3 protein motif alignment with Clone 3001

```

C1. 3001  HSCSPGCSGARSVEQLASNSQSETCLLISAPITNK
JAK3 (Mm)  FSYSSGCPPTHFVRPGWPRTKSAACLLISLVAIKG

```

I JAK3 Protein motif alignment with Clone 2858

```

C1. 2858  SSFFFLWFLRWGISGCGPTWFLYSPGWF
JAK3 (Mm)  NGFAKQGFYSY----SGCPGTH--FVRPGWF

```

Figure 3. Alignments of important motifs for 9 proteins identified in hematopoietic stem cells. (A) Clone 1317 with 3 closely related WEE1 kinase protein family members, which play an important role in mitosis. (B) Clone 1508 with 5 closely related members of the Zinc-finger protein CNBP DNA-binding family, which play a role in gene transcription. (C) Clone 2265 with 6 GLI protein family members, all of which are Zinc-finger DNA-binding proteins. (D) Clone 1405 with 2 C-ETA type protein kinases. (E) Clone 1797 with 5 closed relative insulin/IGF/relaxin family members, which play a role in insulin expression and distribution. (F) Clone 2131 with 6 GPR1/FUN34/yaaH family members. (G) Clone 2995 with a region of homology to MEKK1 proteins. (H) (I) Regions of similarity of 2 clones (3001 and 2858) and the Jak3 protein.

Table 5. Clone sources and description of myeloid complementary DNA microarray

Sources	No. clones	Description
LRH clones: Flow-sorted HSCs from mouse bone marrow	1288	Subtracted myeloid cDNA library; represent early myeloid stem/progenitor cells
EEE clones: EML cells and differentiated derivatives	310	Subtracted myeloid cDNA library, composed of EML(0), EML(6) and EPRO cDNA libraries; represent late-stage myeloid cells; and clones derived from RDA
cDNAs from assorted mouse tissues (Research Genetics)	692	Assortment of regulatory and housekeeping genes expressed in a variety of cell types
Putative Evi-1 target genes	96	cDNAs identified by in vitro selection for Evi-1-binding sites
T-cell RDA clones*	576	T cell-specific genes
Total	2962	

HSCs indicate hematopoietic stem cells; RDA, representation difference analysis.

*From B. Lu, S. Kim, and R. A. Flavell, unpublished data, 2001.

was effective in greatly reducing the abundance of these clones, but the degree of reduction for these genes was variable. For example, clone ID9063 (IMAGE: 421622) was reduced around 3-fold with subtraction, but superoxide dismutase precursor was reduced more than 20-fold

(Figure 2). We also tested for enrichment of genes present only in the tracer population, not in the driver. In this instance, some low-copy genes were enriched more than 1.5- to 5-fold through hybridization (eg, Mel-18; Figure 2C).

Table 6. Cytokines, lineage-specific receptors, and transcription factors on the myeloid cDNA microarray

Clone ID	UniGene No.	Locus link	Symbol	Name
Cytokines				
4 345	795	12 977	Csf1	CSF 1 (M-CSF)
4 154	35 814	16 156	Il11	IL-11
4 284	371	16 189	Il4	IL-4
22	28 830	59 027	Pbef	Pre-B-cell colony-enhancing factor
Cytokine receptors				
4 594	22 574	12 978	Csf1r	Colony stimulating factor 1 receptor
1 105	156 264	12 982	Csf2ra	GM-CSFR- α
4 400	1 940	12 984	Csf2rb2	CSF 2 receptor, beta 2, low-affinity (GM)
4 377	22 673	14 127	Fcer1g	Fc receptor, IgE, high affinity I, gamma polypeptide
4 269	10 809	14 130	Fcgr2b	Fc receptor, IgG, low affinity IIb
4 165	3 303	14 132	Fcgrt	Fc receptor, IgG, alpha chain transporter
4 307	4 154	16 155	Il10rb	IL-10 receptor, beta
4 233	731	16 161	Il12rb1	IL-12 receptor, beta 1
3 963	896	16 177	Il1r1	IL-1 receptor, type I (Il1r1)
4 158	1 349	16 178	Il1r2	IL-1 receptor, type II
4 157	24 771	16 180	Il1rap	IL-1 receptor accessory protein
4 142	4 944	16 188	Il3ra	IL-3 receptor, alpha chain
4 224	389	16 197	Il7r	IL-7 receptor alpha chain precursor
Transcription factors				
3 604	1 318	11 350	Abl1	v-abl Abelson murine leukemia oncogene 1
1 113	34 537	12 606	Cebpa	C/EBP, alpha
1 067		110 794	Cebpe	C/EBP, epsilon
4 521	131	15 399	Hoxa2	Homeobox A2
4 557	173	15 402	Hoxa5	Homeobox A5
4 565	3 546	15 412	Hoxb4	Homeobox B4
4 553	207	15 413	Hoxb5	Homeobox B5
4 578	215	15 414	Hoxb6	Homeobox B6
4 432	255	15 417	Hoxb9	Homeobox B9
4 331	1 351	15 423	Hoxc4	Homeobox C4
4 134	20 305	15 424	Hoxc5	Homeobox C5
4 556	4 444	15 425	Hoxc6	Homeobox C6
4 602	4 765	15 427	Hoxc9	Homeobox C9
1 057	2 444	17 869	Myc	Myelocytomatosis oncogene
1 037	88 061	20 185	Ncor1	Nuclear receptor co-repressor 1
4 197	3 107	18 021	Nfatc3	Nuclear factor of activated T-cells, cytoplasmic 3
4 486	31 255	18 128	Notch1	Notch gene homolog 1
4 187	4 173	18 130	Notch2l	Notch2-like
4 494	4 945	18 131	Notch3	Notch gene homolog 3
4 146	173 813	18 132	Notch4	Notch gene homolog 4
3 865	22 675	18 514	Pbx1	Pre-B-cell leukemia transcription factor 1
4 588	4 788	19 401	Rara	Retinoic acid receptor, alpha
4 585	1 273	19 411	Rarg	Retinoic acid receptor, gamma
3 496	4 869	19 696	Rel	Reticuloendotheliosis oncogene
4 203	455	21 405	Tcf1	Transcription factor 1

CSF, colony-stimulating factor; M, macrophage; GM, granulocyte-macrophage; C/EBP, CCAAT/enhancer binding protein.

The subtracted LRH cDNA library contains a high percentage of novel sequences

To determine the identity of the cDNA clones derived from the library subtraction, we subjected 2304 LRH clones to partial sequence determination and analysis (Table 2). Of the clones, 54% (1255) were nonredundant cDNAs, representing protein-encoding mRNAs. Of these, 247 (20%) were novel sequences; 386 (31%) ESTs; and 622 (50%) known genes. Of the LRH sequences, 46% were not useful, the majority because of being redundant, ribosomal, or empty vector. Sequence data for all of the novel genes have been submitted to GenBank (dbEST).

To facilitate the analysis, retrieval, and further accrual of information concerning these genes, we created a database that is accessible via the World Wide Web (<http://yale130132115135.med.yale.edu/>).

Examples of the known genes derived from the subtracted cDNA library are shown in Table 3, categorized by the functional criteria. Of interest is the presence of 3 members of the *C/ebp* family of transcription factors, as well as *Cbfb*, *Klf9*, *Lrf*, *Sox4*, *Tall1*, and *Xbp1*, each of which is an important regulator of cell differentiation of blood cell lineages and/or other organs.²⁶ It is also notable that while genes for growth factors (eg, *Hdgf*, *Hegfl*, *Efnb1*) and growth factor receptors (eg, *Fgfr1*, *Tnfrsf1b*, *Oprsl*) are present, none of the classical hematopoietic-specific cytokines or their receptors is present in our subtracted library. Components of apoptotic pathways are represented by *Tnfrsf1b*, *Traf1*, *Traf6*, *Prg2*, *Tax1bp1*, *Snip3l*, and *Casp6*. Calcium signaling transducers are included, eg, *Calm*, *Calr*, *Cmkk2*, and *Itp3*. Also present are regulators of the cell cycle, including *Ccnd1*, *Hus1*, and *Lats*.

Protein structure analysis of novel genes

We identified 247 novel sequences among the subtracted LRH clones. These clones are considered novel because of our inability to find any matching sequence in available databases. For each of the potentially novel genes, we subjected clones to additional sequencing from both the 5' and 3' ends of the clones. After compiling the 5' and 3' sequence data, we derived potential open reading frames from these sequences and analyzed them for domains and/or functional motifs (Table 4). This revealed that our novel sequences contained 13 potential nucleic acid-binding proteins, including 4 transcription factors, 11 signal transducers including 2 with similarity to Jak3, 1 with homology to Flt3 ligand, 1 bearing resemblance to the insulin receptor, and 1 mapk/erk kinase kinase-like protein. Sixteen proteins with similarities to known enzymes or enzyme inhibitors were identified, including some potential drug targets (eg, farnesyltransferase, prenyltransferase, and adenylate cyclase). What is notable is the relative paucity of more structural proteins (Table 4). In Figure 3, we show detailed analyses of 9 potentially important novel genes and their homology to known proteins.

Development of a cDNA microarray for analysis of early hematopoiesis

A major goal of this endeavor was to create a cDNA microarray for evaluating gene expression changes during hematopoietic differentiation with specific interest in the myeloid lineage. Thus, the second prong of our approach was to supplement the subtracted library with genes known to be expressed in myeloid cells as well as genes encoding proteins that regulate cell cycle, apoptosis, differentiation, and cell signaling. Thus, we added

587 cDNAs for known genes from an IMAGE Consortium clone set, 310 genes from EML cells isolated by 2 separate subtractive cloning procedures,³⁶ 96 putative Evi-1 target genes,²⁷ and 576 T-cell-expressed genes (B. Lu, S. Kim, and R. A. Flavell, unpublished data, 2001) (Table 5). A significant number of cytokines, hematopoietic transcription factors, growth factors, and growth factor receptors are also on the array (Table 6). A detailed description of these cDNAs and their sources can be found on the Web-accessible database mentioned earlier (<http://yale130132115135.med.yale.edu/>). Purified PCR-amplified cDNA inserts from this collection of plasmids were robotically spotted on polylysine-coated glass slides.

Myeloid cell differentiation is accompanied by abundant fluctuations in gene expression

We tested the cDNA microarray by employing it to analyze patterns of gene expression during induced differentiation of EML cells, a myeloid progenitor cell line.²⁰ We compared the spectrum of gene expression in uninduced EML cells to that of EML cells induced to differentiate for 6, 24, and 72 hours, as

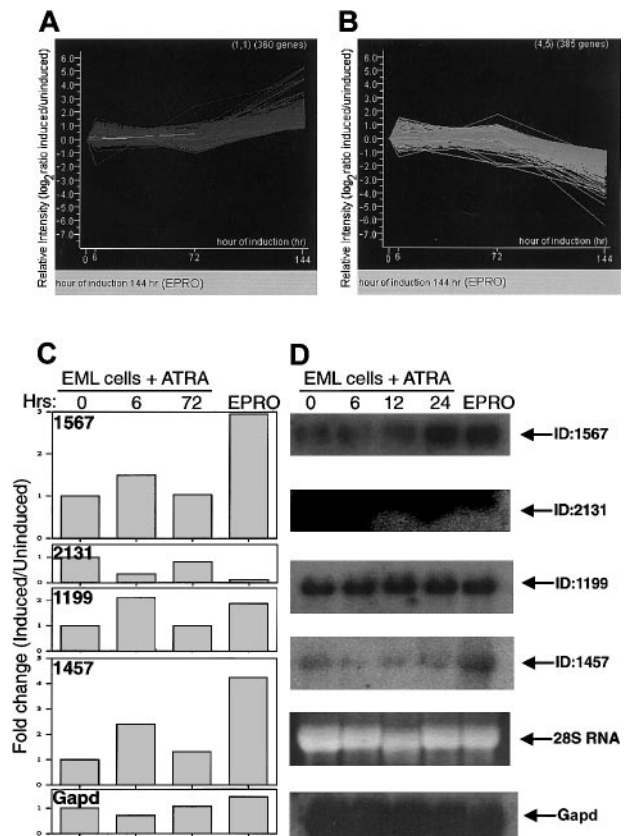


Figure 4. Graphical representation of the change in gene expression over time in EML cells treated with ATRA/IL-3. (A) (B) Each line represents a different gene. The time points (x-axis) represent 0, 6, 24, and 72 hours. The last point on this axis represents EPRO cells, a stage in differentiation beyond 72 hours. The y-axis shows the relative change in gene expression level as \log_2 (induced/uninduced). Thus, a value of +1 represents a 2-fold increase, and -1 equals a decrease to 50% of the value at time zero. (C) Quantitation of fold change in gene expression (as a ratio of induced/uninduced), for 4 novel genes and (*Gapd*) derived from microarray data, for 0 hours, 6 hours, and 72 hours of induction, and for EPRO cells. (D) Northern blot to confirm microarray data obtained with RNA from EML cells. Presented are hybridizations with 4 novel clones from our subtracted myeloid cDNA library, as well as *Gapd* for normalization. Also shown is the ethidium bromide staining of 28S RNA. RNAs are from EML cells treated with ATRA plus IL-3 for 0, 6, 12, and 24 hours to induce myeloid differentiation as well as from EPRO cells.

Table 7. Genes up-regulated during ATRA/IL-3–induced EML cell differentiation

Symbol	Name	Gene Ontology
Gzmb	C11 encoding T-cell–specific protein CCPI CTLA-1, granzyme B	Apoptosis
Prg2	Proteoglycan 2, esoinophil major basic protein	Apoptosis
Dad1	Defender against cell death 1	Apoptosis
Bnip3l	BCL2/adenovirus E1B 19-kd interacting protein 3-like	Apoptosis
Tial1	Tial1 cytotoxic granule-associated RNA-binding protein-like 1	Apoptosis
Calm	Calmodulin	Calcium binding
Calr	Calreticulin	Calcium binding
Cdkn1a	Cyclin-dependent kinase inhibitor 1A (P21)	Cell cycle
Ppm1g	Protein phosphatase 1G, magnesium-dependent, γ isoform	Cell cycle
LATS1	Large tumor suppressor homolog 1	Cell cycle control
Ran	GTP-binding nuclear protein RAN, member RAS oncogene family	Cell cycle regulator
Mdm2	Transformed mouse 3T3 cell double minute 2	Cell growth and maintenance
Cct5	Chaperonin subunit 5 ϵ	Chaperone
C1qa	Complement component 1, q subcomponent, α polypeptide	Complement component
Myo1c	Myosin 1c	Cytoskeleton organization and biogenesis
Ncl	Nucleolin	DNA, RNA binding
Madh5	MAD homolog 5	Embryogenesis and morphogenesis
Mt1	Metallothionein1	Heavy metal binding
Elavl3	ELAV-like 3 (Hu antigen C)	Histogenesis and organogenesis
Igj	Immunoglobulin joining chain	Humoral defense mechanism
Ppia	Peptidylprolyl isomerase A	Immune response
Scya19	Small inducible cytokine A19	Inflammatory response
Irgb7	Integrin β 7	Integrin receptor signal signaling pathway
Emp3	Epithelial membrane protein 3	Membrane fraction
Vasp	Vasodilator-stimulated phosphoprotein	Mineralocorticoid metabolism
Ada	Adenosine deaminase	Nucleic acid metabolism
Ubb	Ubiquitin B	Nucleus
Gnas	Guanine nucleotide binding protein, α stimulating	Protein ADP-ribosylation
DHPS	Deoxyhypusine synthase	Protein biosynthesis
Eef1a1	Elongation factor 1-alpha (EF 1- α)	Protein biosynthesis
Ptpn11	Protein tyrosine phosphatase, nonreceptor type 11	Protein tyrosine phosphatase
Impdh2	IMP dehydrogenase	Purine nucleotide biosynthesis
Various	Ribosomal proteins L3, L4, L5, L6, L7, L10, L11, L13, L23a, L29	Ribosome biogenesis
Various	Ribosomal proteins S3, S4, S6, S7, S12, S14, S16, S17, S19, S20	Ribosome biogenesis
Fgfr4	Fibroblast growth factor receptor 4	Signal transduction
Nsp3	SH2-containing protein 3	Signal transduction
Rgl2	Ral guanine nucleotide dissociation stimulator-like 2	Signal transduction
Mint	Msx2-interacting nuclear target protein	Transcription factor
Mtf2	Metal response element binding transcription factor 2	Transcription factor
Bteb3	Basic transcription element binding protein 3	Transcription from Pol II promoter
Klf9	Kruppel-like factor 9	Transcription regulation
Creg	Cellular repressor of E1A-stimulated genes	Transcription regulation
Rb1	Retinoblastoma 1	Transcription regulation

ELAV indicates embryonic lethal, abnormal vision; IMP, inosine 5'-monophosphate. Other abbreviations are explained in the Table 3 footnote.

well as to that of EPRO cells, which represent a promyelocyte-like stage derived from the EML cells.²⁸ In each experiment, a competitive hybridization was performed between labeled cDNA from uninduced EML cell and from induced EML or EPRO cells, except for the 24-hour time point samples, for which some hybridizations were not competitive. Fluorescently labeled cDNA samples (with either cy-3 or cy-5) (“Materials and methods”) were hybridized. Following washing of the slide, the amount of hybridized probe was quantitated as pixel intensity of fluorescence; low-intensity signals were discarded; and the normalized data were expressed as a \log_2 of the ratio of signal from induced RNA to uninduced RNA cells. These values for EML cells induced for 6, 24, and 72 hours, and for EPRO cells were subjected to clustering by means of a self-organizing map algorithm. This yielded 20 different sets of genes, each of which contained genes that varied in expression level in a similar manner across the samples. Figure 4A-B shows composite graphs for the sets containing genes that increased the most

(Figure 4A) or decreased the most (Figure 4B) over the time points. Tables 7 and 8 list the named genes in each of these sets, respectively.

The major class of genes expressed at higher level in RA-induced EML cells and EPRO cells relative to EML cells was the class encoding ribosomal proteins (Table 7) and included proteins in both the large- and small-ribosomal subunit. These data, together with the increase observed for elongation factors 1 α 1 and Tu-binding and polyA-binding protein, are consistent with a generalized increase in protein synthesis. The calcium signaling pathway also appeared up-regulated: calmodulin, calreticulin, and annexin A1 were all higher. I κ B α , an inhibitor of nuclear factor- κ B (NF- κ B), was also induced, suggesting down-regulation of the NF- κ B–signaling pathway. In addition, there were 74 uncharacterized genes (ESTs) and 37 novel genes that increased in expression during EML cell differentiation. The latter were derived from our library subtractions, and this demonstration that they are

Table 8. Genes down-regulated during ATRA/IL-3-induced EML cell differentiation

Symbol	Name	Gene Ontology
Gzma	Granzyme A	Apoptosis
Pde1b	Phosphodiesterase 1B, Ca ²⁺ -calmodulin dependent, 63 kd	Calmodulin binding
Vcam1	Vascular cell adhesion molecule 1	Cell adhesion
Ppm1g	Fibroblast growth factor inducible 13 (FIN13)	Cell cycle
Ccng	Cyclin G	Cell cycle control
Yes	Yamaguchi sarcoma viral (v-yes) oncogene homolog	Cell cycle regulator
Vav2	Vav2 oncogene	Cell cycle regulator
Fos	FBJ osteosarcoma oncogene	Cell cycle regulator
Hsp70-2	Heat shock protein, 70 kd 2	Chaperone
Fosl2	fos-like antigen 2	DNA binding
Hox1111	Homeobox 11-like 1	DNA binding
Rbpsuh	Recombining binding protein suppressor of hairless (Drosophila)	DNA recombination
Madh4	MAD homolog 4	Morphogenesis
Gna13	Guanine nucleotide binding protein, α 13	GTP binding
Gnao	Guanine nucleotide binding protein, α o	GTP binding
Sh3bp1	SH3 domain-binding protein 1	GTPase activator
Notch1	Notch gene homolog 1	Organogenesis
Eif2ak2	Eukaryotic translation initiation factor 2 α kinase 2	Immune response
Scya2	Small inducible cytokine A2	Inflammatory response
Pla2g7	Phospholipase A2 group VII (platelet-activating factor acetylhydrolase)	Inflammatory response
Apba2	Amyloid β (A4) precursor protein-binding, family A, member 2	Intracellular protein traffic
Hegfl	Heparin-binding epidermal growth factor-like growth factor	Membrane fraction
Kif1b	Kinesin heavy chain member 1B	Microtubule motor
Cited2	Cbp/p300-interacting transactivator, with Glu/Asp-rich C-terminal domain, 2	Nucleus
Psmc4	Proteasome (prosome, macropain) 26S subunit, ATPase, 4	Nucleus
Crat	Carnitine acetyltransferase	Peroxisome
Acox	Acyl-Coenzyme A oxidase	Peroxisome
Cd63	Cd63 antigen	Plasma membrane
Cd152	CD152 antigen	Plasma membrane
Cd14	Cd14	Posttranslational targeting
Ptpn8	Protein tyrosine phosphatase, nonreceptor type 8	Protein tyrosine phosphatase
Expi	Extracellular proteinase inhibitor	Proteinase inhibitor
Es1	Esterase 1	Serine esterase
Serpinb2	Serine (or cysteine) proteinase inhibitor, clade B (ovalbumin), member 2	Serine protease inhibitor
Foxd3	Forkheadbox D3	Transcription regulation
Hnf4	Hepatic nuclear factor 4	Transcription regulation
Hoxb4	Homeobox B4	Transcription regulation
Klf1	Kruppel-like factor 1 (erythroid)	Transcription regulation
Mid1	Midline 1	Transcription regulation
Nfatc3	Nuclear factor of activated T cells, cytoplasmic 3	Transcription regulation
Stat4	Signal transducer and activator of transcription 4	Transcription regulation
Tcf1	Transcription factor 1	Transcription regulation
Xbp1	X box binding protein	Transcription regulation
Ybx1	Y box protein 1	Transcription regulation
Zfp038	Zinc finger protein 38	Transcription regulation
Zfp100	Zinc finger protein 100	Transcription regulation
Zfp216	Zinc finger protein 216	Transcription regulation
ZNF183	Zinc finger protein 183 (RING finger, C3HC4 type)	Transcription regulation
Hoxd12	Homeobox D12	Transcription regulation
LATS2	Serine/threonine kinase KPM	Tumor suppressor

Some abbreviations are explained in the Table 3 footnote.

differentially expressed and thus likely to be of interest, shows the utility of this undertaking for investigating myelopoiesis.

Genes that were down-regulated in this differentiation pathway were more varied (Table 8), but also included 13 novel genes derived from our cloning effort, as well as 68 uncharacterized genes. Several key transcription factors were down during ATRA/IL-3-induced differentiation of EML cells (Table 8). Some of these, such as *Klf1*,²⁹ *Hoxb4*,³⁰ and *Xbp1*,³¹ have known regulatory roles in hematopoietic cells. These data provide an important starting point for further analyses aimed at understanding myelopoiesis at the molecular level, studies that are ongoing in the laboratory.

Northern blot analysis was performed as confirmation of the microarray results obtained for several novel genes (ID1567, ID2131, ID1199, and ID1457). For example, ID2131 (which contains a GTW motif of G-protein receptor [GPR1]/FUN34/yaaH family proteins) is dramatically down-regulated during differentiation, and ID1567 (containing a KGR motif) and ID1457 are up-regulated during EML cell differentiation (Figure 5). Novel gene ID1199 showed similar expression levels before and after induction of differentiation.

While the EML cell culture system has proved useful in identifying changes in gene expression during hematopoietic differentiation, it is nonetheless an immortalized cell line and

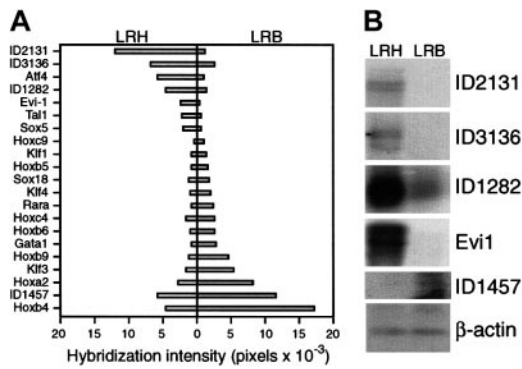


Figure 5. (A) Graphic representation of hybridization intensity in pixels of 21 genes to the cDNA microarray. Values for LRH are depicted by bars to the left; for Lin⁻Hoechst^{Low}rhodamine^{Bright} (LRB), by bars to the right. Gene name or novel gene ID number is indicated to the left. (B) Southern blot of amplified cDNAs derived by reverse-transcription PCR of entire mRNA populations from 2 cell preparations from primary sorted mouse bone marrow cells. The samples in each lane are as indicated on the figure. The probes used for hybridization are listed to the right of each panel. Hybridization with β -actin serves as a control for loading.

thus may not accurately represent normal hematopoietic cells. To compare the changes in gene expression observed in EML cells with normal hematopoiesis, we analyzed RNA from sorted primary bone marrow cells. We obtained 2 pools of cDNAs from sorted primary mouse bone marrow cells: lin⁻Hoechst^{Low} rhodamine^{Bright} (LRB), representing late-stage progenitor cells; and LRH, representing more primitive progenitors.²³ These cDNAs were amplified by PCR with the use of primers specific to adaptor sequences and concomitantly labeled with fluorescent dyes. The LRH and LRB pools were competitively hybridized to the cDNA microarray. Hybridizations were performed in triplicate, and data were normalized to internal control (*Gapd*). This analysis revealed differences in expression between the LRH and LRB preparations of a number of key regulatory transcription factors, including Hox, Klf, and Sox family genes, Evi-1, Tal-1, GATA-1, and Rara (Figure 5A). In addition, a number of novel genes (designated by ID number) were differentially expressed, including ID2131 and ID1457, which were up-regulated and down-regulated, respectively, during EML cell differentiation (Figure 4). To confirm these microarray results, samples of amplified cDNA from LRH and LRB cells were fractionated by gel electrophoresis and then subjected to Southern blot analysis with specific cDNAs used as probes. These data (Figure 5B) support the microarray data in that they demonstrate differential expression between LRH and LRB preparations.

Discussion

We have described the creation of a resource for the in-depth study of gene expression in early hematopoietic cells that should be useful in the study of the molecular regulation of myeloid cell differentiation. Several features of this work are notable and essentially novel. First, the library that we exploited for the creation of the subtracted library represents an early stage of hematopoietic differentiation distinct from that used previously.³² Second, we undertook a library subtraction step that successfully removed most commonly expressed genes, leaving a residual that was relatively enriched in regulatory genes and novel genes. In aggregate, we netted 1255 different gene sequences from the subtraction effort. Given that more than 50% of the clones picked were nonredundant,

it is likely that further sequencing of clones from this subtracted library will allow isolation of additional interesting genes.

Third, we have been successful in creating a glass slide-based microarray from the gene sequences we have isolated. To complement the clones from the subtracted library, we have added genes from a variety of other sources (Table 5). We have tested the utility of this array in 2 initial hybridization experiments. The first identifies genes that are up-regulated or down-regulated during ATRA/IL-3-induced differentiation of EML cells. The second documents transcriptional differences between sorted primary bone marrow cells. This investigation is continuing. However, some of our initial results with the microarray have been confirmed by Northern blot analysis (Figure 4D) or by Southern blot analysis of cDNA populations (Figure 5B), which attests to the validity of the microarray-based quantitation of mRNA or cDNA copies.

Fourth, we have created a Web-accessible database for the genes on the microarray. Using this database, one can download a list of genes present on the array and can query to obtain information regarding specific genes. This Web site represents the starting point for a variety of features, including posting of downloadable microarray data and accrual of information on genes important to hematopoietic progenitor and myeloid cell biology.

A remarkable feature of our sequence analysis was the high number of novel gene sequences present in the subtracted library. This should prove to be an important resource for the isolation of genes that play regulatory roles in early hematopoiesis. Initial protein motif analysis reveals the presence of numerous interesting motifs (Table 4; Figure 3) within these genes. Also remarkable is the paucity of growth factor receptors or cytokines among the known genes in the subtracted library. This is likely to be due to their being present in the driver or to their lack of expression in the LRH library. Our finding of Ephrin-B1 in LRH is novel. Previous studies have shown that a related transmembrane ligand, Ephrin B2, is expressed in certain leukemias and lymphomas.³³ It has also been shown that the receptor for Ephrin B2, EphB4 (hepatoma transmembrane kinase), is expressed on human erythroid progenitors cord blood cells and that it was regulated by SCF.^{34,35} However, no report of expression of EphB1, the receptor for Ephrin B1, in hematopoietic cells has been made. The role of this signaling system in hematopoietic cells is unknown. Interestingly, in the subtracted library, we also identified Nsp3, which encodes a protein that couples Eph receptors to Ras, further suggesting that this is an important pathway in early hematopoietic cells.

Our studies complement and extend data reported by Phillips et al,³² who reported on 2119 nonredundant gene products and the creation of a Stem Cell Database as a repository for these sequences. In aggregate, our effort, combined with theirs, provide an abundance of cloned sequences from early hematopoietic progenitors that allow for investigation into the molecular control of hematopoiesis.

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