LYAR, a novel nucleolar protein with zinc finger DNA-binding motifs, is involved in cell growth regulation

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A cDNA encoding a novel zinc finger protein has been isolated from a mouse T-cell leukemia line on the basis of its expression of a Ly-1 epitope in a λgt11 library. The putative gene was mapped on mouse chromosome 1, closely linked to Idh-1, but not linked to the Ly-1 (CD5) gene. The cDNA is therefore named Ly-1 antibody reactive clone (LYAR). The putative polypeptide encoded by the cDNA consists of 388 amino acids with a zinc finger motif and three copies of nuclear localization signals. Antibodies raised against a LYAR fusion protein reacted with a protein of 45 kD on Western blots and by immunoprecipitation. Immunolocalization indicated that LYAR was present predominantly in the nucleoli. The LYAR mRNA was not detected in brain, thymus, bone marrow, liver, heart, and muscle. Low levels of LYAR mRNA were detected in kidney and spleen. However, the LYAR gene was expressed at very high levels in immature spermatocytes in testis. The LYAR mRNA is present at high levels in early embryos and preferentially in fetal liver and fetal thymus. A number of B- and T-cell leukemic lines expressed LYAR at high levels, although it was not detectable in bone marrow and thymus. During radiation-induced T-cell leukemogenesis, high levels of LYAR were expressed in preleukemic thymocytes and in acute T leukemia cells. Fibroblast cells overexpressing the LYAR cDNA from a retrovirus vector, though not phenotypically transformed in vitro, had increased ability to form tumors in nu/nu mice. Therefore, LYAR may function as a novel nucleolar oncoprotein to regulate cell growth.

[Key Words: Zinc finger protein, nucleolus, radiation-induced thymic leukemogenesis]

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Oncogenic transformation of cells leads to a number of changes in cellular metabolism, physiology, and morphology. One of the major changes is the pleomorphism and hyperactivity of the nucleolus (Yeoman et al. 1976; Busch et al. 1979, 1986). Nucleolar proteins have been identified that are specifically expressed in actively dividing cells and tumor cells (Reddy et al. 1989). The functions of these proteins are not clear, and it has been speculated that they may be involved in cell growth regulation (Tan 1982, Mathews et al. 1983) and in transcriptional control (Dyman and Tjian 1983).

The precise molecular pathways and secondary changes leading to malignant transformation for most cell types are not clear. A number of cases have been reported in which altered expression or activity of some proteins with putative DNA-binding motifs are found in some tumors or leukemias. For example, chromosomal translocations involving lyl-1 [a protein with a putative DNA-binding motif (Mellentin et al. 1989)] and the fusion proteins between RARα and PML [a zinc finger-encoding gene (de The et al. 1991; Goddard et al. 1991)] are detected in some leukemic cell lines. In two recent reports, a zinc finger protein has been identified which, when overexpressed, appears to cooperate with the myc oncogene to induce B-cell leukemias in transgenic mice [Haupt et al. 1991, van Lohuizen et al. 1991].

The zinc finger domain is a sequence-specific DNA-binding motif in a family of proteins that was first identified in the Xenopus laevis transcription factor IIIA (Miller et al. 1985). Substantial evidence has established these proteins as a class of trans-acting molecules with regulatory roles in cellular growth and differentiation. Here, we report a cDNA isolated from a T-cell leukemia line which encodes a putative nuclear protein with zinc finger motifs. We present evidence that the protein is localized in the nucleolus of cells. Expression of the gene is induced during radiation-induced thymic leukemogenesis in mice. Over-expression of the cDNA in NIH-3T3 cells increases their tumorigenic potential in nu/nu mice.

Results

Isolation of a cDNA encoding a zinc finger protein from a mouse T-cell leukemia line

In an attempt to clone the T-cell surface marker Ly-1 (CD5) several years ago (Hershberger 1989), a λgt11 li-
The Ly-1 antibody reactive clone (LYAR) was characterized and sequenced. It was ruled out as encoding Ly-1 by chromosomal mapping (see below). Comparison of LYAR to the Ly-1 sequences, subsequently elucidated by others (Huang et al. 1987), showed that they encoded polypeptides with no significant similarity.

The LYAR clone encodes a 1164-bp open reading frame consisting of 388 amino acids (Fig. 1A). A sequence similarity search of the gene bank data bases revealed no significant similarity to any known genes either at the DNA or at the amino acid levels (data not shown). Further inspection of the encoded polypeptide revealed that two putative zinc finger domains are present in the first 60 amino acids (Fig. 1A, underlined). Both of the zinc finger motifs are similar to the standard zinc finger consensus sequence (Berg 1986), although the space between the last two zinc-binding residues (Cys or His) is two instead of three amino acids.

Recently, a new family of zinc finger proteins has been identified whose spacing between the last two zinc-binding residues consists of two instead of three amino acids (Fig. 1B, Goddard et al. 1991; Haupt et al. 1991, van Lohuizen et al. 1991). These proteins all have eight zinc-binding residues (Cys or His) in a cluster (C4HC3) at their amino termini. The putative zinc finger domains at the amino terminus of LYAR fit the consensus of this new class of zinc finger motifs (Fig. 1B), although the LYAR domain has the His residue in a different position (C4HC). Overall, this Cys/His motif seems to characterize proteins that interact with DNA. Most of the proteins are implicated in cell proliferation and tumorigenesis, and other regulatory functions.

Three copies of nuclear localization signals [NLS] are also detected in the protein (Fig. 1A,C). One is almost identical to the first NLS identified in the SV40 T antigen (Kalderon et al. 1984a,b). The other two are bipartite nuclear localization signals [BNLS] similar to the ones identified in nucleoplasm and thyroid β-receptor (Robbins et al. 1991). Thus, the LYAR cDNA appears to encode a nuclear protein with a putative DNA-binding motif.

A sequence search for conserved motifs also revealed that LYAR has a potential site for tyrosine phosphorylation immediately following the second zinc finger (double underlined in Fig. 1A). A potential Ser residue for cAMP-dependent protein kinase is also detected at the carboxyl terminus (Fig. 1A, carboxyl terminus underline). In addition, a stretch of amino acids is found to be mostly charged residues (32 of 47) with a positive net charge (11 negative, 21 positive residues, Fig. 1A, wavy underline).

Chromosomal location of the LYAR gene

The chromosomal location of the LYAR gene in mice was established using the BXD recombinant inbred strains (Bailey 1971). A restriction fragment length polymorphism [RFLP] for LYAR between the BXD parental strains C57Bl/6 and DBA was identified, and this was used to determine the strain distribution pattern of the LYAR locus in 21 BXD mice (Table 1). The strain distribution pattern for LYAR exactly matched that of isocitrate dehydrogenase-1 [Idh-1] on chromosome 1, implying that these loci are tightly linked (P>0.99). Adjacent loci on chromosome 1 also showed significant linkage, which diminished with increasing distance from Idh-1. The proximal chromosome 1 markers gave the best match to LYAR of 144 loci on 16 chromosomes (Taylor 1989).

LYAR, a 45-kD protein, is localized predominantly in the nucleolus

Antibodies were raised against a LYAR fusion protein produced in bacteria. The purified antibodies specifically recognized a protein of 45 kD from EL4 cells, which was not detected in a bone marrow-derived stromal cell line (AC6) on Western blot (Fig. 2A, B).

By immunoprecipitation of labeled cell extracts, it was shown that the antibody reacted with a protein of ~45 kD, which was not detected in AC6 cells, and the preimmune antiserum did not react with this protein (Fig. 2C). The mobility of the protein on SDS-PAGE did not change under reducing or nonreducing conditions (data not shown).

Indirect immunofluorescence staining was performed to localize the LYAR protein in mouse cell lines and in COS-7 cells transfected with the LYAR cDNA. The LYAR protein was localized predominantly in some subcompartments of the nuclei of T or B leukemia cells (data not shown). Using mouse embryonic fibroblast (EF) cells (Fig. 3A), it was shown clearly that the LYAR protein was localized in the nucleoli of the nucleus, which was visualized by staining with DNA with Hoechst 33342.

In addition, COS-7 cells transiently transfected with the LYAR cDNA also showed predominantly nucleolar staining (Fig. 3B). COS-7 cells transfected with the vector plasmid DNA did not give any detectable staining (data not shown). This indicates that the antibody was specific to the protein encoded by the LYAR cDNA. A few of the LYAR-expressing cells, however, seemed to express LYAR at higher levels, and LYAR was localized both in the nucleolus and in the nucleoplasm. Interestingly, the nucleoli of these cells appeared to be more organized along the nuclear membrane (see arrows in Fig. 3B), perhaps as a result of overexpression of LYAR in these cells.

In actively dividing mouse EF cells, the LYAR protein is present mostly in the nucleolus. However, in some cells undergoing mitosis with condensed chromosomal DNA, LYAR protein is localized throughout the cells. The LYAR protein appears to be excluded from the condensed chromosomal DNA (Fig. 3C, c). The rhodamine immunofluorescence signal [red] of LYAR appears to be excluded from nuclear compartments where high levels of DNA [blue or white] are detected (Fig. 3C, c). This indicates that LYAR is probably not an integral part of the chromosomes.
A novel nucleolar oncoptotein

Figure 1. Molecular structure of the LYAR gene. (A) Nucleotide and deduced amino acid sequences of the LYAR cDNA. The zinc finger region is underlined. The double underline indicates a potential Tyr phosphorylation site. The region underscored by a wavy line consists of 47 residues with 32 charged ones (11 negative, 21 positive). The three putative NLS are marked with bold letters ([BNLS1 bipartite nuclear localization signal], [NLS] SV40 T-type nuclear localization signal]). A putative phosphorylation site for cAMP-dependent kinase is marked at the carboxyl terminus. (B) The zinc finger motifs of LYAR and a new family of proteins. The residues conserved in all proteins are boxed and defined as the consensus, except Pro [P], which is not conserved in all the proteins. (X) Any amino acid. Residues conserved in LYAR are in bold letters. Sequences not cited in the text are the baculovirus early proteins CG30 [Thiem and Miller 1989] and PE-38 [Kraupa and Knebel-Morsdorf 1991], a trypanosome protein with a leucine repeat motif, T-LR [Smiley et al. 1990], and the SS-A/Ro protein [Chan et al. 1991], which is part of a nuclear ribonucleoprotein particle. (C) The putative NLS in the LYAR protein. As marked in Fig. 1A, NLS is aligned with that of the SV40 T. BNLS1 and BNLS2 are aligned with the NLS of thyroid β-receptor and nucleoplasmin, respectively. (X) Any amino acid.
**Table 1.** LYAR pattern distribution in BXD and linkage analysis

| BXD | LYAR | Chromosome 1 locus | Distance to Idh-1 (cm) | Number of differences | Probability of linkage |
|-----|------|--------------------|------------------------|-----------------------|-----------------------|
| 1   | B    | Aox-1              | 6.43 ± 1.55            | 1/21                  | >0.99                 |
| 5   | B    | Idh-1              | 0                      | 0/21                  | >0.99                 |
| 6   | B    | Lsh                | 11.46 ± 3.59           | 6/21                  | >0.95                 |
| 8   | D    | Pep-3              | 23.31 ± 1.81           | 7/21                  | <0.90                 |
| 9   | B    | Mls                | 31.92 ± 6.01           | 9/21                  | <0.90                 |

Strain distribution pattern of the LYAR RFLP among the BXD recombinant inbred strains C57Bl/6 (B) and DBA (D). The summary of the strain distribution comparisons to loci on chromosome 1, listed in proximal to distal order, is based on data given in Taylor (1989). The genetic distances to Idh-1 are from published results (Davidson et al. 1989), using the combined male and female recombination data. Probability of linkage was calculated by the \( \chi^2 \) distribution.

The LYAR mRNA is detected in a number of T and B leukemia cells but not in most adult mouse tissues

To investigate the expression pattern of LYAR in T- and B-cell leukemias, we first examined its expression in a number of cell lines by Northern blot analyses (Fig. 4A). High levels of LYAR mRNA were detected in all leukemic T (BW, EL4, and VL3.1) and leukemic B (L1-2, 200-2 and RAW112) lineage cell lines tested but not in a bone marrow-derived stromal cell line (AC6) or in an interleukin-2 (IL-2)-dependent CTL line (AR1). Thus, LYAR seemed to be expressed at high levels in the transformed leukemia cells tested.

Expression of LYAR in different tissues of adult mice was also examined (Fig. 4B). By Northern blot analyses, LYAR mRNA was not detected in brain, thymus, liver, bone marrow, heart, or muscle. Low levels of expression were detected in kidney and spleen, and high levels of LYAR mRNA were detected in testis. In situ hybridization with the LYAR cDNA probes confirmed the expression patterns in mouse primary tissues (see below and data not shown). This indicated that the T- and B-cell leukemia lines tested were induced to express the LYAR gene sometime during or after their transformation.

LYAR expression during radiation-induced T-cell leukemogenesis

As mentioned above, LYAR mRNA was not detectable in thymus or bone marrow but was highly expressed in...
Figure 3. (See following page for legend.)
all of the T and B lineage leukemia cells tested [Fig. 4A,B]. We therefore decided to test whether the expression of LYAR occurred in the preneoplastic or neoplastic stages following irradiation (12-16 weeks after irradiation for the preleukemic thymocytes) was blotted and probed with a LYAR-specific probe [Fig. 4C]. High levels of LYAR mRNA were detected in thymic populations containing preleukemic thymocytes, as well as in acute leukemic thymocytes.

The increased LYAR mRNA detected could be the result of induction of the LYAR gene in preleukemic or leukemic thymocytes. Alternatively, it could also be attributable to expansion of a minor thymocyte population expressing high levels of LYAR in the normal thymus. We investigated the latter possibility, in situ hybridization was performed with thymus sections from normal mice [Fig. 4D]. No cells in the thymus were found to express high levels of the LYAR message. Therefore, expression of the LYAR gene appeared to be highly induced early during the thymic leukemogenesis process.

Expression of the LYAR gene during embryonic development

The LYAR gene was expressed at high levels in mouse embryonic stem (ES) cells [Fig. 5A, lane 1]. Analyses of RNA isolated from day 9 and day 13 embryos also detected high levels of LYAR mRNA [Fig. 5A, lanes 2,3]. Low levels of LYAR mRNA were detected in adult spleen, as shown above in Figure 4B and in Figure 5A [lane 4].

To localize specific tissues expressing LYAR, early embryos were studied for LYAR expression by in situ hybridization. The LYAR mRNA was detected in day 10 fetal liver [Fig. 5B] and day 13 fetal thymus [Fig. 5C]. No significant levels of hybridization were detected in fetal heart at either stage. Expression of LYAR was also detected in fetal livers and fetal thymuses of other fetal developmental stages, and in some other tissues such as vertebrae and lung [L. Su and I.L. Weissman, in prep.]. Therefore, the LYAR gene is expressed in early embryos and preferentially in some tissues, including fetal liver and fetal thymus, where fetal hematopoiesis is occurring.

The LYAR gene is expressed at very high levels in immature spermatocytes

In adult testis the LYAR mRNA was present at a very high level [Fig. 4B]. In situ hybridization was performed to examine what cell types in the testis express the LYAR gene. As shown in Figure 6, LYAR-positive cells appeared to reside at the basal layer of the seminiferous tubules, where the immature spermatocytes are located [Fig. 6A]. The Sertoli cells, interstitial cells (Leydig cells), the mature, and maturing spermatocytes did not express detectable levels of LYAR [Fig. 6A, a,b]. The sense probe, which served as negative control for specific hybridization, showed no hybridization signal in the testis [Fig. 6A, c,d].

To confirm that the LYAR protein is expressed in immature spermatocytes, immunofluorescence staining of testis sections with the anti-LYAR antibodies was performed [Fig. 6B]. The primary and secondary spermatocytes are specifically labeled, consistent with the in situ hybridization results.

To confirm that only the germ cells in testis expressed the LYAR gene, in situ hybridization was performed with testis sections from the germ cell-deficient mouse [W/W*]. It is known that W/W* mice are sterile because they lack germ cells, and the nongerm cells of the testis are all present [Coulombre and Russell 1954]. As expected, no LYAR-positive cells were detected in testis from W/W* mice [Fig. 6C].

LYAR is a novel nucleolar protein with oncogenic potential

We have demonstrated that LYAR is a nucleolar protein with a putative DNA-binding domain. Expression of the LYAR gene seems to correlate with abnormal cell growth. The oncogenic potential of LYAR was tested by overexpressing the LYAR cDNA in a mouse fibroblast cell line [NIH-3T3] via a retrovirus vector. Figure 6 shows that two independent clones stably expressing the LYAR cDNA, Vly12 and Vly36 [Fig. 7A, lanes 1,2], expressed much higher levels of LYAR proteins than the control cells expressing the neomycin resistance gene [Fig. 6A, lane 3] or the parental NIH-3T3 cells [data not shown].

The cells expressing high levels of LYAR [Vly12, Vly22, Vly36], although not morphologically transformed in vitro [data not shown], showed increased ability (i.e., higher frequency and shorter latent period) to form tumors at the sites of injection than the control.

Figure 3. Subcellular localization of LYAR. [A] LYAR is localized in the nucleolus. Mouse EF cells were stained with anti-LYAR antisera [a,c,e] or preimmune serum [b,d,f]. [a,b] Phase-contrast pictures show the cellular morphology; [c,d] fluorescence signals of anti-LYAR staining. [e,f] DNA (Hoechst dye) staining. The arrows in a,c, and e indicate one of the nucleoli stained with the anti-LYAR antibody. [B] The LYAR cDNA encodes the nucleolar protein reactive with the anti-LYAR antibody. COS-7 cells transiently expressing the LYAR cDNA, Vly12 and Vly36 [Fig. 7A, lanes 1,2], expressed much higher levels of LYAR proteins than the control cells expressing the neomycin resistance gene [Fig. 6A, lane 3] or the parental NIH-3T3 cells [data not shown]. [C] LYAR localization during cell mitosis. Actively dividing mouse EF cells were stained with polyclonal antisera. [a] Anti-LYAR signals (rhodamine), [b] DNA (Hoechst dye) staining, [c] double exposure of both the anti-LYAR signals [red] and the DNA-staining signals [blue and white].
cells [Neo1, Vneo22, Vneo45], when their tumor-forming potential was tested in nu/nu mice (Table 2). Tumors derived from Vly12 were shown to express the LYAR cDNA at high levels (data not shown) and were typical fibrosarcomas (Fig. 7B). The tumor cells could be retransferred to form tumors efficiently in other nu/nu mice (Table 2).

The cells expressing the neo gene from the retroviral vector (Vneo22 and Vneo45) or from an expression plasmid (Neo1) did not give rise to any visible tumors until the fourteenth week after injection. By that time, some of the mice with tumors from the LYAR cDNA-expressing cells had died from tumors that were usually 3-4 cm in diameter. Therefore, overexpression of the LYAR cDNA in NIH-3T3 fibroblast cells rendered them more tumorigenic, indicating that LYAR is a novel nuclear protein that can be involved in cell growth regulation.

**Discussion**

**LYAR** is a cDNA isolated from a T-cell leukemia line encoding a novel nuclear zinc finger protein

A cDNA was isolated and characterized from a T-cell leukemia, which encoded a polypeptide cross-reactive...
Figure 5. Detection of LYAR mRNA during early fetal development. (A) Northern blot analysis of LYAR mRNA in early embryos. RNA isolated from ES cells [lane 1], day 9 whole embryo [E9, lane 2], day 13 whole embryo [E13, lane 3] or adult spleen [Sp, lane 4] was hybridized with a LYAR-specific probe [LY, top] or a human actin probe [AC, bottom]. (B) Detection of LYAR mRNA in day 10 fetal liver by in situ hybridization. The bright-field picture demonstrated that high levels of LYAR RNA were detected [black grains] in fetal liver (Lv) but not in fetal heart (Ht). [a] Day 10 embryo section hybridized with a LYAR sense probe as a background control; [b] day 10 embryo section hybridized with a LYAR antisense probe. Magnification, 19×. (C) Detection of LYAR mRNA in day 13 fetal thymus by in situ hybridization. The bright-field picture showed high levels of LYAR RNA present in day 13 fetal thymus [Th] but not in fetal heart [Ht]. [a] Day 13 embryo section hybridized with a LYAR sense probe as a background control; [b] day 13 embryo section hybridized with a LYAR antisense probe. Magnification, 94×.

with the anti-Ly-1 monoclonal antibody 53.7. The putative gene was mapped by RFLP on mouse chromosome 1 and was not the Ly-1 gene. The LYAR clone encoded a protein that has a unique zinc finger motif within the first 60 amino acid residues and three copies of NLS [Fig. 1].

A new family of zinc finger proteins has been identified recently [Goddard et al. 1991]. Like LYAR, these proteins all have eight zinc-binding residues in a cluster at the amino terminus [Fig. 1B]. Most of the proteins in this family have been implicated in regulating cell growth or gene expression. For example, PML [dc The et al. 1991; Goddard et al. 1991], bim-1 [Goebl 1991], Haufl et al. 1991], van Lohuizen et al. 1991], MEL-18 [Tagawa et al. 1990; Goebl 1991], and rfp [Takahashi et al. 1988] are probably involved in cell transformation, IE110 [Perry et al. 1986], VZV61 [Davison and Scott 1986], and Rpt-1 [Pataraca et al. 1988] are believed to be involved in regulating gene expression, and RAG-1 [Schatz et al. 1989] and RAD-18 [Jones et al. 1988] are involved in site-specific DNA recombination and DNA repair, respectively.

The three copies of putative NLS present in LYAR suggest that its nuclear localization may be of interest (Fig. 1C). The SV40 T antigen signal, which is sufficient to direct most, but not all, proteins into the nucleus, requires other NLS motifs to transport some proteins [Richardson et al. 1986; Greenspan et al. 1988; Morin et al. 1989]. Thus, LYAR may use these multiple copies of NLS to direct its nuclear and/or subnuclear localization. Multiple copies of NLS may also play a role in regulated nuclear localization. For example, two copies of NLS have been reported in the glucocorticoid receptor, whose nuclear localization is regulated by binding to steroid hormones [Picard and Yamamoto 1987; Picard et al. 1990].

Nucleolar localization and the putative DNA-binding domain of LYAR

Immunolocalization of LYAR in different cells clearly showed that LYAR is predominantly localized in the nucleolus [Fig. 3]. A putative NLS consisting of mostly positively charged residues has been identified [Siomi et al.
The charged region in LYAR (Fig. 1A) has a net positive charge that might serve as a NLS.

The subcellular location of a protein usually implies its potential function in the cell. According to our current understanding of the nucleolus, LYAR may be involved in the following processes: (1) ribosomal gene

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**Figure 6.** Localization of LYAR-expressing cells in mouse testis. (A) In situ hybridization detection of the LYAR mRNA in mouse testis. (a,b) LYAR antisense probe; (c,d) LYAR sense probe. a and c are bright-field pictures showing the morphology; b and d are dark-field pictures showing the LYAR mRNA hybridization signals (white grains). (st) Sertoli cells; (s) spermatozoa; (ms) maturing spermatozoa. (B) Detection of the LYAR protein in mouse testis by immunofluorescence staining. (a) Testis section was stained with preimmune serum, (magnification, 22×); (b) testis section stained with anti-LYAR specific antiserum, (magnification, 22×); (c) anti-LYAR specific antiserum, (magnification, 112×). (st) Sertoli cells; (s) spermatozoa; (ms) maturing spermatozoa. (C) Analysis of LYAR mRNA in W/Wv mouse testis by in situ hybridization. (a) Phase contrast of the testis section; (b) dark-field picture of testis section hybridized with the LYAR antisense probe; (c) dark-field picture of testis section hybridized with the LYAR sense probe.
transcription; [2] rRNA processing; [3] ribosome assembly; or [4] protein transport between the nucleus and cytoplasm [Meier and Blobel 1990, 1992].

The zinc finger motifs in LYAR and the fact that the glutathione s-transferase (GST)-LYAR fusion protein can bind to double-stranded DNA with high affinity (L. Su and I.L. Weissman, unpubl.) strongly suggest that LYAR is a DNA-binding protein. It is conceivable that LYAR may be involved in regulating ribosomal gene transcription. On the other hand, LYAR may also function in other cell compartments either during different stages of development or different stages of the cell cycle. Identification of the DNA sequences interacting with LYAR should be informative about the target genes for LYAR action. Ribosomal genes are candidate targets for LYAR action because of its subcellular location. In addition, general methods can be employed that have been developed to identify the DNA-binding sites of any putative DNA-binding protein (Kinzler and Vogelstein 1989; Mavrothalassitis et al. 1990).

**Induction of LYAR during oncogenic transformation**

The LYAR gene is highly expressed in all T and B leukemia cells tested but is not detected in normal thymus and bone marrow cells. During the radiation induction of thymic leukemias, the LYAR gene was induced in preleukemic thymocytes and remained active in acute leukemic thymocytes (Fig. 5). Therefore, LYAR expression seemed to be associated with the early stages of oncogenic transformation.

The results described above raised the possibility that LYAR may be associated with cell proliferation in general. However, several lines of evidence indicated that the LYAR gene was not expressed during proliferation of some normal cells. First, actively proliferating cells are present in bone marrow (>20%), yet no or very low levels of LYAR expression were detected in bone marrow.

### Table 2. Oncogenicity test of LYAR in NIH-3T3 cells

| Cells | 8 weeks | 14 weeks |
|-------|---------|----------|
| Neo1  | 0/5     | 1/5<sup>a</sup> |
| Vly12 | 4/5     | 5/5<sup>3</sup> |
| Vneo45| 0/5     | 0/5      |
| Vly22 | 2/5     | 3/5<sup>1</sup> |
| Vly36 | 3/5     | 3/5<sup>2</sup> |
| Vly12T<sup>e</sup> | 10/10 | 10/10<sup>8</sup> |

<sup>a</sup>NIH-3T3 cells were transfected with a neomycin resistance gene [Neo1], with a LYAR cDNA in a retroviral vector [Vly12, Vly22 and Vly36], or with the same retroviral vector expressing the neo gene [Vneo22 and Vneo45], and G418-resistant cells were cloned and analyzed.

<sup>b</sup>About 2 × 10<sup>5</sup> cells were injected subcutaneously into each nu/nu mouse. Tumors at the sites of injection were scored weekly.

<sup>c</sup>Small tumors (<0.5 cm in diam.) appeared around the fourteenth week postinjection, when tumors observed at the eighth week were bigger than 3 cm in diam.

<sup>d</sup>Number of mice that died from tumors.

<sup>e</sup>Tumor cells derived from [experiment 1] Vly12-injected nu/nu mice.
cells [Fig. 5]. Second, AR1 (an IL-2-dependent CTL line) can proliferate efficiently in the presence of IL-2, but no LYAR expression was detected in AR1 cells (Fig. 5). Therefore, LYAR expression appeared to be associated with abnormal cell growth.

Interestingly, the LYAR gene was expressed at high levels in ES cells and in a number of fetal tissues including fetal liver and fetal thymus [Fig. 6; L. Su and I.L. Weissman, unpubl.] This suggests that the LYAR function may be important during normal fetal development. The expression of LYAR in immature spermatocyte cells also suggests that LYAR may be involved in growth of undifferentiated cells or early cell differentiation.

LYAR: a novel nucleolar oncoprotein

We demonstrated that NIH-3T3 cells overexpressing the LYAR cDNA from a retrovirus vector, when injected into nu/nu mice, had higher activity (higher frequency and shorter latent period) for tumor formation. Therefore, LYAR behaved like an oncoprotein. However, the cells overexpressing LYAR did not grow like completely transformed cells in vitro, that is, no focus formation or anchorage-independent growth in agarose colony assay was detected (data not shown).

Although it has been shown that the nucleoli of tumor cells may be distinguishable from those of normal cells [Busch et al. 1979, 1986] very little is known about the significance of these differences. Proliferating cells and tumor cell-specific nucleolar antigens have been identified. Their precise role in cell proliferation, in transformation, or in the maintenance of tumorigenicity is not clear [Yeoman et al. 1976; Tan 1982; Mathews et al. 1983].

Whereas different types of genetic alterations may all lead to altered expression or function of cell-growth regulatory genes and to abnormal growth, it is generally believed that more than one event is required to lead to the neoplastic transformation of a normal cell to a malignant one [Land et al. 1983; Weinberg 1989]. It is conceivable that LYAR may partly transform cells and render them more sensitive to other transformation events that lead to malignant transformation. Alternatively, LYAR may complement a pre-existing transformation event present in the NIH-3T3 fibroblast cell line to cause malignant transformation. Experiments testing the oncogenic potential of LYAR in vivo or in primary cells are in progress.

A new zinc finger-encoding oncogene, mdm-2, has been reported recently [Fakharzadeh et al. 1991]. The mdm-2 protein has two zinc fingers of CXXC/CXXC type. The mdm-2 gene is found to be overexpressed in a number of tumor cells. Like LYAR, overexpression of mdm-2 in NIH-3T3 cells does not lead to morphologic transformation in vitro, but it can increase tumorigenicity of the cells in nu/nu mice. Two recent studies have reported that mdm-2 can physically interact with and modulate the function of the p53 protein to affect cell growth [Momand et al. 1992; Oliner et al. 1992].

Materials and methods

Cell lines

NIH-3T3 cells, COS-7 cells, and EF cells are maintained in Dulbecco's modified Eagle medium supplemented with 10% fetal bovine serum [FBS] [Sigma]. The T leukemia cells EL4, VL3-1, and BW5147, B leukemia cells L2-1, 200-2, RAW112, and the bone marrow-derived stromal cell line AC-6 are maintained in RPMI-1640 supplemented with 10% FBS. The CTL line AR-1 cells are cultured in RPMI-1640 supplemented with 10% FBS and 5% RAT T-stim with concanavalin A [Collaborative Research, Inc.].

Molecular cloning and DNA sequence analyses

A lig11 cDNA library of EL4 cells [a gift from T. St. John, ICOS, Seattle, WA] was screened with a monoclonal antibody against the mouse Ly-1 antigen [57.3.1. Ledbetter and Herzenberg 1979], as described previously [St. John et al. 1986]. LYAR clones were used to screen the library to get the full-length LYAR cDNA. DNA sequence was obtained by subcloning the cDNA into M13mp18 and M13mp19. Both strands were sequenced by chain termination using the Sequenasc 2.0 kit [U.S. Biochemicals].

The DNA sequence was analyzed on the Stanford Cell Biology VAX computer, with the Wisconsin GCG programs, and on the CMGM computer at Stanford, with the IntelliGenetics programs.

Northern blot analyses

Total RNAs from cell lines or fresh tissues were purified, and 15 µg of total RNA was used for each Northern blot analysis [Ausubel et al. 1989]. A human β-actin cDNA [a kind gift from M. Hu, Stanford University, CA] was used as probe to standardize the amount of RNA from each sample.

In situ hybridization

In situ hybridization was performed essentially as described [Wilkinson et al. 1987] except that the LYAR cDNA in pBlue-script KS or SK [Stratagene] was used as template for T7 RNA polymerase transcription to generate a sense or antisense RNA probe labeled with α-35S-labeled UTP.

LYAR fusion proteins and antibody production

The entire coding region of LYAR was fused with the bacterial GST gene, and the fusion protein was overexpressed and purified as described [Smith and Johnson 1988]. The purified fusion protein was used to immunize rabbits or rats for polyclonal or monoclonal antibody production, respectively. Purified LYAR protein fused with the bacterial maltose binding protein [MBP, New England Biolabs] was used for screening anti-LYAR-positive antibodies by ELISA [Harlow and Lane 1988]. The antibodies were purified on protein A or protein G affinity column [Harlow and Lane 1988].

Protein analysis

Total cell extract from 10⁶ cells was analyzed by Western blot as described [Harlow and Lane 1988]. The rabbit anti-LYAR serum or preimmune serum was used at different dilutions. Protein A conjugated with horseradish peroxidase [HRP, Sigma] was used as the second-stage reagent, and an ECL Western blot detecting kit [Amersham] was used.

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Immunoprecipitation was performed essentially as described [Harlow and Lane 1988]. Briefly, cell extract labeled with [35S]methionine (5 x 10^6 cpm) was used for each reaction with the antisera, and protein A-Sepharose (Sigma) was used to pellet the rabbit IgG.

For indirect immunofluorescence staining, it was adapted from the protocol described [Harlow and Lane 1988]. Cells grown on chamber slides (at ~70% confluence) were washed with PBS and fixed in 3.7% formaldehyde for 10 min. The cells were then washed with PBS again and treated with acetone for 2 min. After washing with H2O and PBS, the anti-LYAR antibody or control antibody was diluted in PBS and incubated with the cells for 60 min at room temperature. The cells were then washed three times with PBS and incubated with rhodamine-labeled goat anti-rabbit IgG antibody for 60 min. After the second antibody staining, the cells were incubated with Hoechst dye 33342 (CalBiochem) at 400 nM for 15 min at room temperature, followed by three washes with PBS. A Nikon (Nikon, Inc.) microscope was used for imaging and photography.

Radiation induction of T-cell leukemia in C57Bl/Ka mice

The radiation protocol was as described [Kaplan and Brown 1954]. Cells from preleukemic thymuses were isolated from mice 12–16 weeks after the initial irradiation. Thymocytes at this stage contain a subpopulation that has acquired leukemic potential but is not fully transformed. Leukemic thymocytes were isolated from mice 20 weeks after the initial irradiation, when the mice had visible lymphoma in the thymus.

Transient expression of LYAR cDNA in COS-7 cells

LYAR cDNA was cloned into pSDD2 vector (a gift from D. Denny, Stanford University, CA) and transfected into COS-7 cells by the DEAE–dextran method as described [Ausubel et al. 1989].

Stable transfection of LYAR cDNA in NIH-3T3 cells

LYAR cDNA cloned in a retrovirus vector (a gift from C. Shih, Systemix, Palo Alto, CA) was cotransfected with pGKNeo (a gift from E. Li, MIT, Cambridge, MA) into NIH-3T3 cells, and G418-resistant cells were selected. Cells transfected with pGKNeo alone or with the same retrovirus vector expressing the neomycin resistance gene were also selected and cloned.

Tumorigenicity test in nu/nu mice

Stable transfectants were expanded, and 2 x 10^3 cells were injected subcutaneously into each nu/nu mouse. Tumors at the injection sites were monitored weekly for at least 14 weeks.

Mapping the chromosomal location of the LYAR gene with recombinant inbred mice

A RFLP was identified for the LYAR probe between the inbred mouse strains C57Bl/6 and DBA using the restriction enzymes KpnI and ApaI. Segregation of this RFLP marker was examined in 21 BXD recombinant inbred strains, kindly provided by Benjamin Taylor (Jackson Laboratory, Bar Harbour, ME). The resultant pattern was compared with 144 other markers on 16 chromosomes from the BXD strains [Taylor 1989]. Linkage was determined by X^2 analysis.

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