Gene expression during preimplantation mouse development

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To develop a resource for the identification and isolation of genes expressed in the early mammalian embryo, large and representative cDNA libraries were constructed from unfertilized eggs, and two-cell, eight-cell, and blastocyst-stage mouse embryos. Using these libraries, we now report the first stages at which the cytokines interleukin (IL)-6, IL-1β, and interferon (IFN)-γ are transcribed in the developing embryo and the presence of IL-7 transcripts in the unfertilized egg. Transcripts for IL-1α, -2, -3, -4, or -5 were not detected at these stages. To identify novel genes expressed on activation of the embryonic genome, the egg and eight-cell stage-specific cDNA libraries were subtracted from the two-cell library, yielding a specialized cDNA library enriched for transcripts expressed at the two-cell stage. Sequence and Southern blot analysis of several of these cDNAs expressed predominantly at the two-cell stage of embryogenesis revealed them to be from novel genes, thereby providing the first molecular tools with which to approach the study of gene expression in the early mammalian embryo.

[Key Words: cDNA libraries; cytokines; interleukins; IFN-γ; PCR; preimplantation embryos; subtractive hybridization]

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The molecular control of mammalian preimplantation embryogenesis remains largely unexplored, due mainly to the difficulty of obtaining sufficient quantities of timed embryos for experimentation. Nonetheless, knowledge about the changes in gene expression that underlie this period is essential to understanding mammalian development. Several lines of evidence, most notably that inhibition of transcription at the one-cell stage blocks protein synthesis and all subsequent development after the first cleavage division, and that initiation of synthesis of all classes of RNA occurs at the two-cell stage, point to the early activation of the embryonic genome (for review, see Telford et al. 1990a). No resources existed that allowed investigation of whether activation leads to generalized gene transcription or synthesis of independent stage-specific transcripts. Temporal changes in transcription are also likely to herald the completion of cleavage and the formation of the first differentiated cells, those of the trophectoderm (for review, see Schultz 1986), whose origin and fate have been difficult to study without probes to lineage-specific markers. One approach to identifying genes relevant to mammalian development has focused on the sequence homology with genes of developmental importance in other vertebrate or invertebrate organisms. However, considering that early development in mammals results in an implantation-competent embryo, it is likely that a unique combination of genes controlling this process will be utilized in the mammalian embryo. In an effort to identify genes expressed in these early mammalian developmental stages, several investigators have applied the polymerase chain reaction (PCR) technique, thus circumventing the problem of obtaining sufficient embryonic material for study (Rappolee et al. 1988). However, only transcripts of known genes can be readily identified with this technique. Classically, cDNA libraries have provided a useful resource for identifying novel genes transcribed in specific cell types or tissues. Yet for technical reasons, cDNA libraries prepared from unfertilized eggs or single stages of the preimplantation embryo (Taylor and Pikó 1987; Weng et al. 1989; Ko et al. 1990; Welsh et al. 1990) have not provided reliable sources for the comprehensive analysis of genes differentially expressed during early embryonic development.

Here, we describe the use of large and representative cDNA libraries constructed from poly[A]+ mRNA of preimplantation mouse embryos to demonstrate stage-specific transcription of several cytokines. Subtractive hybridization of these libraries served to identify cDNAs representing novel genes expressed in the two-cell-stage embryo. These libraries provide the first resource for molecular information about genes expressed in the egg and early embryonic stages and a tool to access novel genes.
expressed at the two-cell stage when the mouse embryonic genome is first activated.

Results

Library and insert size of egg and embryonic cDNA libraries

A single mouse egg or mouse embryo at any stage of preimplantation development contains no more than 50 pg of poly[A]+ mRNA (Clegg and Pikó 1983). We therefore optimized a cDNA cloning strategy to permit efficient library construction using 10–100 ng mRNA (J.L. Rothstein, D. Johnson, J. Jessee, J. Skowronski, D. Solter, and B. Knowles, in prep.). Plasmid vectors, which can accommodate the directional cloning of cDNA, were employed so that T7 and T3 RNA polymerase promoter sequences could be used to generate sense and antisense transcripts for subtractive hybridization. Libraries containing $1 \times 10^6$ to $2 \times 10^6$ independent cDNA clones were constructed from the 50–175 ng of poly[A]+ RNA isolated from unfertilized eggs, and two-cell, eight-cell, and blastocyst-stage embryos (Table 1). Because a library of $10^6$ clones has a $>99\%$ probability of including rare transcripts (<10 copies per cell) at a detectable frequency (Sambrook et al. 1989), these egg- and embryonic-stage libraries are not only likely to contain representatives of abundant but also of medium- and low-abundance transcripts in the egg or embryonic stage. Each of these four libraries contains at least $10^6$ independent cDNA clones (Table 1). The insert size of 25–50 randomly picked independent clones per library was determined by PCR, and blastocyst-stage embryos were probed with a mouse β-actin cDNA. Between 200 and 355 of the 250,000 clones screened in each library hybridized with the β-actin probe (Table 2). These values, when converted (see Materials and methods), suggest that 18,700 actin mRNA molecules are present in the mouse egg and 5600 are present in the late two-cell stage. We find that the levels of actin correspond well to those reported previously for the egg and two-cell stages, that is, 21,000 copies of actin mRNA in the egg and 3700 in the two-cell stage (Taylor and Pikó 1990), and are close to those of Bachvarova et al. (1989). We observe an increase in the number of actin transcripts in the eight-cell stage (18,460) and blastocyst stages (41,480), a pattern similar to previous reports (Taylor and Pikó 1990). The levels of actin in the libraries corroborate those of Taylor and Pikó and Bachvarova et al. and are substantially lower than the figures from comparable stages reported previously (Giebelhaus et al. 1983, 1985).

Transcripts of tissue-type plasminogen activator (t-PA) previously have been shown to decrease in maturing oocytes until they become nearly undetectable at ovulation (Huarte et al. 1985, 1987; Strickland et al. 1988). Expression of t-PA in the oocyte has been estimated at <0.05% of the total mRNA (Huarte et al. 1987). To determine the representation of t-PA in the unfertilized ovulated egg library, we hybridized a mouse t-PA cDNA clone (Rickles et al. 1988) to replica filters containing 250,000 clones and found that 60, or 0.024%, of the clones in the egg library were positive (Table 3). Three representative t-PA clones were partially sequenced from the 3' end and found to be homologous to the 3'-untranslated region of the mouse t-PA gene cloned from the F9 teratocarcinoma-derived cell line (Rickles et al. 1988; data not shown). As expected, no t-PA cDNAs were detected in the two-cell, eight-cell, or blastocyst-

<table>
<thead>
<tr>
<th>Embryonic stage</th>
<th>Number of females</th>
<th>Number of embryos</th>
<th>RNA [ng]</th>
<th>Library size [cfu]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unfertilized egg</td>
<td>200</td>
<td>5,000</td>
<td>1750²</td>
<td>2 x 10⁶</td>
</tr>
<tr>
<td>Two-cell</td>
<td>665</td>
<td>13,500</td>
<td>910³</td>
<td>1 x 10⁶</td>
</tr>
<tr>
<td>Eight-cell</td>
<td>300</td>
<td>2,778</td>
<td>1740⁴</td>
<td>2 x 10⁶</td>
</tr>
<tr>
<td>Blastocyst</td>
<td>100</td>
<td>600</td>
<td>500⁵</td>
<td>1 x 10⁶</td>
</tr>
</tbody>
</table>

²Estimate based on 5% poly[A]+ RNA.
³Total number of independent cDNA clones (colony forming units) plated on primary filters.
⁴Amount of RNA calculated based on values of Pikó and Clegg (1982).
⁵Values determined by use of Northern analysis of 18S/28S rRNA in the embryonic samples compared with rRNA from a standard amount of cellular RNA.
Table 2. Analysis of gene expression in cDNA libraries

<table>
<thead>
<tr>
<th>cDNA library</th>
<th>Probesa</th>
<th>cytochrome-c oxidase</th>
<th>β-actin</th>
</tr>
</thead>
<tbody>
<tr>
<td>28S/18S rRNA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unfertilized egg</td>
<td>0</td>
<td>0</td>
<td>275 (0.110)</td>
</tr>
<tr>
<td>Two-cell</td>
<td>0</td>
<td>20 (0.008)</td>
<td>200 (0.080)</td>
</tr>
<tr>
<td>Eight-cell</td>
<td>1 (0.0004)b</td>
<td>0</td>
<td>355 (0.142)</td>
</tr>
<tr>
<td>Blastocyst</td>
<td>0</td>
<td>0</td>
<td>305 (0.122)</td>
</tr>
</tbody>
</table>

Probes used for hybridization are pA-28S/18S rRNA; cytochrome-c oxidase I and II, and mouse β-actin isolated from a blastocyst-stage library [see Materials and methods].

Probes used were pTAM, t-PA; clone 11, IAP, and pB1/B2 (B1 and B2 repeats).

Number of positive colonies of 250,000 independent cDNA clones screened (%).

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Probes used were pTAM, t-PA; clone 11, IAP, and pB1/B2 (B1 and B2 repeats).

Number of positive colonies of 250,000 independent cDNA clones screened (%).

Detection of cytokines in the cDNA libraries

As a first approach to identifying genes expressed during early embryogenesis that may have regulatory functions, we investigated the representation of several cytokines in the libraries. Studies of polypeptide growth factors in early mammalian development have focused on expres-
sion of epidermal growth factor (EGF), transformation growth factor (TGF-α), platelet-derived growth factor (PDGF), TGF-β, insulin-like growth factor (IGF)-I, and IGF-II at specific times during preimplantation development (Rappolee et al. 1988; Lee et al. 1990; Telford et al. 1990b). Although some of these growth factors are known to play a role in differentiation, their major function appears to involve regulation of the cell cycle. On the other hand, the cytokines not only regulate proliferation but also appear to induce differentiated functions.

To investigate whether interleukins IL-1α, IL-1β, IL-2-7, or γ-interferon (IFN-γ) are transcribed during preimplantation embryogenesis, the cDNA libraries from the unfertilized egg, eight-cell, and blastocyst stages were screened. Cytokine expression, identified initially by PCR analysis of pooled inserts from each cDNA library, was confirmed by direct screening of the cDNA libraries with authentic probes. PCR analysis of the libraries revealed expression of IL-1β, IL-6, IL-7, and IFN-γ but not IL-1α, or IL-2–IL-5 (Table 4). As expected, β₂ microglobulin (β₂M) was present at all stages tested (Sawicki et al. 1981). Southern hybridization of the PCR gels using probes to IL-1β, IL-6, IL-7, IFN-γ, and β₂M verified the presence of these transcripts in the libraries (Fig. 1).

To quantify cytokine expression in the embryonic libraries, we screened each stage with a PCR-generated gene-specific probe. Screening 250,000 clones of each library with an IL-7 probe indicated 8 positive clones in the unfertilized egg library, whereas no colony hybridization was seen with the same number of clones from eight-cell and blastocyst libraries. Thus, IL-7 transcripts appear to be rare in the mouse egg (0.003% of the independent cDNA clones in the library) and undetectable in the early embryo. Sequence analysis of two of the IL-7 hybridizing clones from the unfertilized egg library confirmed these to be the mouse IL-7 gene (Namen et al. 1988; data not shown). IL-6, a cytokine with effects on many cell types (Hirano et al. 1990; Sehgal 1990), was shown previously to be expressed at the blastocyst stage (Murray et al. 1990). Here, we report that IL-6 is transcribed as early as the eight-cell stage, persisting into the blastocyst stage (Table 4; Fig. 1). IL-1β, a pleiotropic cytokine expressed by multiple cell types with an important role in the inflammatory response (Oppenheim et al. 1986; Dinarello 1989), is expressed by mammalian placental tissue and cultured trophoblast-derived cell lines (Taniguchi et al. 1991). The function of IL-1β in the developing embryo is not known, and there have been no reports of its synthesis during early embryonic development. Although IL-1β expression at the blastocyst stage was indicated by our PCR analysis of the libraries (Fig. 1) and by direct analysis of freshly isolated blastocysts by

### Table 4. PCR analysis of embryonic cytokine and β₂M gene expression

<table>
<thead>
<tr>
<th>Target gene</th>
<th>Embryonic stage a</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>egg</td>
</tr>
<tr>
<td>IL-1α</td>
<td>–</td>
</tr>
<tr>
<td>IL-1β</td>
<td>–</td>
</tr>
<tr>
<td>IL-2</td>
<td>–</td>
</tr>
<tr>
<td>IL-3</td>
<td>–</td>
</tr>
<tr>
<td>IL-4</td>
<td>–</td>
</tr>
<tr>
<td>IL-5</td>
<td>–</td>
</tr>
<tr>
<td>IL-6</td>
<td>–</td>
</tr>
<tr>
<td>IL-7</td>
<td>+</td>
</tr>
<tr>
<td>IFN-γ</td>
<td>–</td>
</tr>
<tr>
<td>β₂M</td>
<td>+</td>
</tr>
</tbody>
</table>

aPlated cDNA libraries were scraped, plasmid DNA was prepared by alkaline lysis, and cesium chloride-purified DNA was digested with MluI and SalI. Insert cDNA was isolated by agarose gel electrophoresis, and 10–25 ng was amplified by PCR by use of the indicated cytokine-specific oligonucleotide primers for 45 cycles (94°C for 30 sec, 50°C for 30 sec, 72°C for 1 min). (+) The presence of a specific signal for the indicated cytokine by Southern hybridization (see Fig. 1). All cytokine-specific primers were tested in RT–PCR reactions with total RNA derived from mouse peritoneal exudate and spleen cells, and were shown to give the appropriate size bands on EtBr-stained agarose gels (not shown).
reverse transcriptase [RT]–PCR [data not shown], no hybridi-
zing colonies were detected in the $5 \times 10^6$ colonies
screened using a probe homologous to the 5′ end of IL-
1β. However, 20-fold more cDNA was screened by PCR
than by filter hybridization. These data suggest that IL-
1β is either expressed as a rare message in each cell or by
a small number of specialized cells in the mouse blasto-
cyst, or is actually present in the blastocyst library at a
higher level but not detectable using the 5′ IL-1β probe.
Analysis of IL-1β expression in the inner cell mass and
the trophectoderm, as well as screening the blastocyst
library by use of a 3′ IL-1β probe, will likely resolve this
issue. IFN-γ is also expressed in the mouse blastocyst
(Table 4, Fig. 1.), consistent with the observation that the
mouse blastocyst secretes a factor with IFN-like antivi-
ral activity in vitro [Cross et al. 1990; Nieder 1990]. A
member of the IFN-α gene family was identified previ-
ously as one of the major proteins expressed by bovine,
ovine, caprine, and porcine blastocysts [Imakawa et al.
1987, 1989; Hansen et al. 1988; Cross and Roberts 1989;
Roberts et al. 1989; Baumbach et al. 1990]. Our data pro-
vide the first evidence for expression of any interferon in
the murine blastocyst. Thus, the screening of these li-
braries with probes of known cytokines demonstrates
the transcription of genes whose products are often ex-
pressed in differentiated cell types and mediate changes
in gene expression.

**Isolation of novel stage-specific genes**

by subtractive hybridization

To identify genes whose expression changes during pre-
implantation development, we generated specialized li-
braries by subtractive techniques. Directional cloning in
the Bluescript vector allowed us to use a modification of
the biotin–streptavidin method [Sive and St. John 1988;
J.L. Rothstein, D. Johnson, J. Jessee, J. Skowronski, D.
Solter, and B. Knowles, in prep.] to obtain unique mRNA
molecules. T3-initiated, biotinylated, antisense single-
stranded, and hybrid RNA molecules were separated from
T7-initiated, sense single-stranded molecules after bind-
ing to streptavidin [Fig. 2]. Using this approach, we
generated a two-cell-specific subtraction library by hybridi-
zation of a fivefold excess of biotinylated RNA from the
egg library to that of the two-cell library. The resulting
two-cell-specific single-stranded RNA was separated from
biotinylated RNA bound to streptavidin, and hybridi-
zation to a 10-fold excess of biotinylated RNA from the
eight-cell library. Following a second streptavidin treat-
ment, the remaining single-stranded RNA was reverse-
transcribed and cloned into plasmid vectors. The average
insert size of the cDNAs in the two-cell subtraction li-
brary (2CSL-I), which contains $2 \times 10^6$ clones, was 1.0
kb [data not shown]. Repeating the procedure schema-
tized in Figure 2 with the 2CSL-I library as starting ma-
terial resulted in a second two-cell-specific subtraction
library (2CSL-II) of $2 \times 10^6$ clones with an average insert
size of 300–400 bp [data not shown]. The smaller size of
the cDNAs in the 2CSL-II library is consistent with
RNA degradation during the multiple and long incuba-
tion periods of double-stranded RNA hybrids at high
temperatures.

To determine whether these subtraction libraries are
reduced in complexity, we hybridized both subtracted
libraries with probes to IAP, β-actin, and B1/B2 repeat
sequences; no positive hybridization was detected in ei-
ther subtraction library (250,000 clones screened; data
not shown). Because cDNAs in the 2CSL-I and -II librar-
ies should be highly enriched for transcripts expressed in
greatest abundance at the two-cell stage of embryogene-
sis, we randomly selected clones for further analysis.
Twenty such clones with inserts >500 bp in length were
partially sequenced and compared with those listed in the
GenBank/EMBL data bases. Of these 20 clones, 14
did not match any sequence listed in the data bases. The
other six clones proved to be bacterial cDNAs, most
likely from the *Escherichia coli* tRNA used as a carrier in
the preparation of the subtraction libraries. Following
sequence analysis, the 14 unique cDNAs were hybridi-
zied to the original two-cell cDNA library and any clones
that showed specific hybridization were analyzed further
for stage-specific expression by hybridization to the egg
and eight-cell libraries. In this manner, four stage-spe-
cific cDNAs were identified that were expressed pre-
dominantly or exclusively at the two-cell stage of pre-
implantation development (Table 5). One clone, stage-
specific embryonic clone-3 (SSEC-3), appears to be
expressed predominantly at the two-cell stage, with a
low number of hybridizing clones [3 hybridizing clones/
250,000 colonies screened] in the egg library. SSEC-D
is a highly expressed message; 0.16% of the clones in the
two-cell library hybridized with this cDNA, approxi-
mately fivefold more than in the egg library. SSEC-C is
two-cell specific, but only 0.002% of the colonies
screened hybridized with this cDNA. There are more
colones hybridizing with the SSEC-P probe in the two-
cell-stage library [0.02%] than in the egg [0.01%], sug-
gesting that this gene is either newly transcribed at the
two-cell stage or that its message is somehow protected
during the generalized RNA degradation that occurs af-
fter fertilization [Clegg and Pikó 1983]. Each of these
clones represents an authentic single-copy mouse gene
determined by Southern analysis [data not shown]. In
addition, expression of SSEC-C, SSEC-D, and SSEC-P
was confirmed by direct RT–PCR analysis of freshly iso-
lated two-cell embryos using SSEC-specific primers [data
not shown]. Since SSEC-3, SSEC-C, SSEC-D, and SSEC-P
are small cDNAs without apparent open reading frames,
isolation of full-length cDNAs corresponding to these
clones is a necessary and an ongoing effort. All of the
remaining 14 cDNAs were confirmed by Southern blot
hybridization to be of mouse origin, but were not de-
ected in the two-cell library after screening 250,000
clones and, therefore, are likely to represent extremely
rare transcripts.

**Discussion**

The unfertilized egg and embryonic stage-specific cDNA
libraries we have described provide a unique resource to
Figure 2. Scheme for the generation of primary subtractive cDNA libraries (2CSL-I). For secondary subtractive libraries, the entire procedure was repeated with sense transcripts from 2CSL-I to yield 2CSL-II. Phenol extraction of the hybridized streptavidin-treated RNA leaves the two-cell-specific RNA in the aqueous phase for use in cDNA cloning. Sense RNA was synthesized from the linearized two-cell library and hybridized to a fivefold excess of antisense biotinylated RNA from the egg library. The RNA remaining after streptavidin treatment was subsequently hybridized to a 10-fold excess of RNA transcribed from the eight-cell library. RNA left in the aqueous phase was used to make a cDNA library (2CSL-I). This subtraction library was used to make a second subtraction library by hybridizing the sense RNA transcribed from this subtraction library to a 10-fold excess of the biotinylated antisense RNA transcribed from the egg library. The RNA isolated in the aqueous phase was hybridized further to a 10-fold excess of the biotinylated antisense RNA transcribed from the eight-cell library, and a cDNA library was constructed from the remaining two-cell-specific RNA as described previously (2CSL-II).

study genes expressed in the early mammalian embryo. The results obtained by probing these libraries with single-copy genes suggest that they are representative of the genes transcribed at these stages. One aspect of this analysis is that our estimate of actin levels will serve to resolve the controversy over the quantity of actin message in the preimplantation embryo. Previous estimates of actin mRNA abundance were made by comparing the level of embryonic actin mRNA to that in mRNA from a nonembryonic standard source, a technique subject to variation. Quantitation of independent actin clones in cDNA libraries overcomes this limitation. Another interesting aspect arises from the difference between B1 and B2 transcript levels in the unfertilized egg and two-cell-stage libraries on the one hand, and the eight-cell- and blastocyst-stage libraries on the other. Previous studies of total unfractionated RNA revealed an increase in B1 and B2 repeat-containing transcripts throughout preimplantation development (Taylor and Pikó 1987; Pozanski and Calarco 1991), whereas the frequency of B1 and B2 repeat-containing cDNAs decreases in the libraries after the two-cell stage. These data suggest that there may be changes in the RNA polymerase II and III activity in the embryo after the activation of the embryonic genome at the two-cell stage. Previous studies have suggested that changes occur in the relative amounts of RNA polymerase II and III activity between the eight-cell stage and blastocyst, the earliest embryonic stages investigated (Warner 1977).

The expression of polypeptide growth factors was investigated because the interactions of these factors with their receptors mediate changes in gene expression, re-
Ten of these clones represent rare transcripts, and the genes. The first of these libraries, enriched for genes expressed at the two-cell stage of embryogenesis, has been generated. These embryonic cDNA libraries will now serve as a source of novel cDNA probes for genes whose transcription changes during development, and in isolating novel cDNA clones of relatively rare transcripts from a specific embryonic stage.

Because each cDNA library described in this report is representative, it should contain at least one cDNA clone of most of the genes transcribed in the corresponding stage in the mouse. Thus, by using probes derived from known genes and new probes isolated by such techniques as subtraction, the libraries provide the much needed instrument to determine whether the genes transcribed at the two-cell stage are activated independently to perform a stage-specific function or whether most of the embryonic genome is transcriptionally activated at the two-cell stage and, on differentiation, enhanced expression or specific repression of specific gene subsets occurs. Although few studies have directly addressed either notion, the generalized decrease in methylation during preimplantation mouse development (Monk et al. 1987) suggests a global activation of the embryonic genome. Similarly, the low-level constitutive expression of cell-lineage-specific genes, such as myoD and other mesoderm-associated genes (Rupp and Weintraub 1991), at the time when the frog embryonic genome is first activated, also supports the generalized activation hypothesis. The availability of cDNA libraries representing serial stages of early mouse development may now allow this basic issue to be experimentally addressed in the earliest embryonic stages.

### Materials and methods

#### Mice and embryo recovery

Unfertilized eggs, two-cell, eight-cell, and blastocyst embryos were collected from immature B6D2F1 mice (Jackson Laboratories, Bar Harbor, ME or Harlan-Spague Dawley, Indianapolis, IN) after superovulation (Hogan et al. 1986) and mating to B6D2F1 males, where appropriate. Unfertilized eggs were treated with hyaluronidase and subsequently with Pronase, whereas cleavage-stage embryos and blastocysts were treated with Pronase alone (Hogan et al. 1986). Eggs and embryos from all stages were washed repeatedly in modified Whitten’s medium (Abramczuk et al. 1977), and pools of 500–1000 were

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**Table 5. cDNA clones obtained from two-cell subtraction library**

<table>
<thead>
<tr>
<th>Clone</th>
<th>cDNA insert size (bp)</th>
<th>Positive clones sequence information</th>
<th>Sequence length (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSEC-3</td>
<td>500</td>
<td>3 [0.0011]</td>
<td>10 [0.004]</td>
</tr>
<tr>
<td>SSEC-C</td>
<td>600</td>
<td>0</td>
<td>5 [0.002]</td>
</tr>
<tr>
<td>SSEC-D</td>
<td>600</td>
<td>75 [0.030]</td>
<td>400 [0.160]</td>
</tr>
<tr>
<td>SSEC-P</td>
<td>900</td>
<td>25 [0.0010]</td>
<td>50 [0.020]</td>
</tr>
</tbody>
</table>

*Approximate size based on agarose gel (6X174 standard).

aNumber of positive colonies detected of 250,000 independent cDNA clones screened.

Obtained from combining partial 3' and 5' sequence of each clone. Nucleotide sequences were found to be novel when compared with

those listed in GenBank/EMBL by use of WORDSEARCH and FASTA commands of the GCG software program (Devereux et al. 1984).
placed in 200 µl of embryo lysis buffer [ELB: 100 mM NaCl, 50 mM Tris-HCl (pH 7.5), 5 mM EDTA, 0.5% SDS, 5 µg of E. coli tRNA (Boehringer Mannheim)], which had been initially incubated with 0.5 mg/ml of protease K (Boehringer Mannheim) for 30 min at 37°C to remove any contaminating RNase.

**Embryo RNA isolation**

The embryo/ELB solution was incubated for 1 hr at 37°C and extracted twice with phenol–chloroform, and nucleic acids were collected by ethanol precipitation and stored at −70°C in absolute ethanol (Sambrook et al. 1989). Aliquots were removed and microcentrifuged for 60 min. The 70% ethanol-washed pellet was air-dried, redissolved in 80 µl of RNase-free water and 20 µl of 5× DNase buffer (250 mM Tris-HCl [pH 7.5], 1 mM NaCl, 50 mM MgCl₂, 25 mM CaCl₂), and 1.5 µg of DNase I (worthington Biochemicals), which was initially incubated with 0.5 mg/ml of protease K (Boehringer Mannheim) at 37°C for 30 min to remove contaminating RNase, was added, and the sample was incubated at 37°C for 30 min. DNase digestion was terminated by adding 10 µl of 0.25 M trans-1,2-diaminocyclohexane-

**RNA quantification**

RNA was quantified by visually comparing the amount of RNA in 1% of each embryo sample to serial dilutions of cellular RNA standards on Northern blots (Sambrook et al. 1989) after hybridization to a random-primed (Feinberg and Vogelstein 1983) radiolabeled probe. The embryo/ELB solution was incubated for 1 hr at 37°C and extracted twice with phenol–chloroform, and nucleic acids were collected by ethanol precipitation and stored at −70°C in absolute ethanol (Sambrook et al. 1989). Aliquots were removed and microcentrifuged for 60 min. The 70% ethanol-washed pellet was air-dried, redissolved in 80 µl of RNase-free water and 20 µl of 5× DNase buffer (250 mM Tris-HCl [pH 7.5], 1 mM NaCl, 50 mM MgCl₂, 25 mM CaCl₂), and 1.5 µg of DNase I (Worthington Biochemicals), which was initially incubated with 0.5 mg/ml of protease K (Boehringer Mannheim) at 37°C for 30 min to remove contaminating RNase, was added, and the sample was incubated at 37°C for 30 min. DNase digestion was terminated by adding 10 µl of 0.25 M trans-1,2-diaminocyclohexane-

**cDNA synthesis**

First- and second-strand cDNA syntheses were performed by modification of the method described (Gubler and Hoffman 1983). RNA in 8.3 µl of H₂O was heated to 65°C for 15 min to remove secondary structure and placed immediately on ice. All first-strand reaction mixtures were assembled on ice in a total volume of 33 µl. Each reaction contained 6.6 µl of 5× reverse transcriptase buffer [GIBCO/BRL], 3.3 µl of 10 mM dNTPs [Pharmacia], 500 ng of oligo(dT)₁₇ linker primer [Wistar Institute Nucleotide Synthesis Facility, 5'-CGGTGCAGGCCT-GACGG[T₃₋₅]₃', 1.3 µl of BSA (2.4 mg/ml; Boehringer Mannheim), 1 unit of human placental RNase inhibitor (Boehringer Mannheim), 50 µCi of [³²P]dCTP (3000 mCi/mM, Amersham), and 200 units of Superscript RNase H⁻. Moloney murine leukemia virus [MLV] RT [GIBCO/BRL]. The sample was then incubated at 37°C for 60 min. The amount of RNA converted to cDNA was quantified as described (Sambrook et al. 1989) and was consistently 30 ± 8%. Second-strand synthesis was performed in the same tube in a total volume of 200 µl. Briefly, 30 µl of first-strand reaction, 20 µl of 10× second-strand buffer [Sambrook et al. 1989], 2 units of RNase H [Pharmacia], 70 units of DNA polymerase I holoenzyme (Boehringer Mannheim), 20 µl of dNTPs (10 mM each), and sterile, nuclease-free H₂O were incubated for 1 hr at 15°C, followed by 1 hr at room temperature. The reaction was terminated by the addition of 2.5 µl of 0.25 M CTDA, 5 µl of glycerol (1 mg/ml; Boehringer Mannheim), 7 µl of 10% SDS, and 50 µg of proteinase K, followed by a 15-min incubation at 56°C. Following phenol–chloroform extraction, addition of an equal volume of 5 M Na₂OAc, and ethanol precipitation, S1 nuclease treatment of double-stranded cDNA was performed (Gubler and Hoffman 1983) in a total volume of 100 µl with 200 units of S1 nuclease (Boehringer Mannheim) in 1× S1 buffer (0.1 M NaOAc, 0.8 M NaCl, 2 mM ZnCl₂), for 20 min at 37°C. Nuclease reactions were terminated by adding 20 µl Tris-HCl (pH 8.3), followed by phenol extraction and ethanol precipitation. Nuclease-treated cDNA was end repaired by resuspension in 11 µl of nuclease-free H₂O, 4 µl of 5× T4 polymerase buffer [0.2 M Tris-HCl [pH 7.5], 50 mM MgCl₂, 10 mM EDTA, 40 mM DTT, 1 mg/ml of BSA], 4 µl of dNTPs (10 mM each), 1 unit of T4 polymerase (Boehringer Mannheim), and incubation at 37°C for 15 min (Sambrook et al. 1989). End-repaired cDNA was phenol/chloroform-extracted and ethanol-precipitated. 5'-Phosphorylated MluI linkers (3 µg, Pharmacia LKB Biotechnology) were ligated to blunt-ended cDNA by use of 1 Weiss unit of T4 ligase in 30 µl for 16–18 hr at 15°C (Sambrook et al. 1989). Ligase was inactivated (65°C for 10 min), and the cDNA was double-digested with the restriction enzymes Sall and MluI [New England Biolabs] in a total volume of 400 µl for 5–6 hr at 37°C, under conditions suggested by the manufacturer. Digestion reactions were terminated, phenol/chloroform-extracted, and ethanol-precipitated as described above. Digested cDNA was resuspended in 15 µl of nuclease-free H₂O, 10 µl of saturated urea, and 1 µl of bromphenol blue tracer dye (1 mg/ml) and loaded onto a 1-ml Sepharose CL4B column [Pharmacia] initially washed in column buffer [20 mM Tris-HCl [pH 7.5], 0.2 mM NaOAc, 4 mM EDTA, 0.1% SDS)]. Fractions of 100–200 µl were collected; cDNA >500 bp eluted in the first radioactive peaks. After counting, the peak fractions were pooled and cDNA was precipitated with ethanol using 15 µg of glycerol as carrier. Precipitated cDNA was resuspended in water to a concentration of 0.25–1 ng/µl and ligated into excess of linearized pBS vector for 18 hr at 15°C (Stratagene, modified so that the EcoRI site was converted to an MluI site, and the HindIII site was converted to a Sall site). Ligation reactions were phenol/chloroform-extracted, ethanol-precipitated, and resuspended in 10 µl of TE [10 mM Tris-HCl [pH 7.5], 0.1 mM EDTA] prior to bacterial electroporation.

**Bacterial electroporation, plating, and composition of cDNA libraries**

E. coli strain DH10B was kindly provided by Joel Jessee

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**Preimplantation mouse development**

**E. coli** strain DH10B was kindly provided by Joel Jessee
Rothstein et al.

Bacteria used for electroporation were grown and made electrocompetent as described [Hanahan et al. 1991]. All electroporations were performed by use of a Cell Porator (GIBCO/BRL) set at 400 V and 4000 ohms (resulting in line voltages of 2.4-2.5 kV). Electrotransformation efficiencies of 3 x 10^10 to 6 x 10^10 cfu/μg of plasmid were routinely obtained with 10 pg of pUC19. Electroporation of cDNA libraries resulted in transformation efficiencies of 2 x 10^6 to 3 x 10^6 cfu/μg of cDNA. cDNA was electroporated into independent aliquots containing 1 μl of cDNA and 25 μl of electrocompetent bacteria. Aliquots were electroporated, grown in 1 ml of SOC medium [Hanahan et al. 1991] at 37°C for 60 min, pooled, and spun at 400g for 10 min. Cell pellets were resuspended in 1 ml of SOC for every 250,000 estimated transformants, and each milliliter was spread onto 8.5 x 8.5-inch MSi nylon membranes that were placed on top of LB/Amp agar plates [Sambrook et al. 1989] and incubated overnight at 37°C. The following day, filters were replica-plated and prepared for hybridization, and master filters were stored at −70°C [Rothstein et al. 1992]. For all libraries, additional plates were scraped with 20 ml of LB/Amp, and 0.1- to 0.5-ml aliquots were stored at −70°C.

Genetic probes and reagents

Probes used in this study were: pTAM (full-length t-PA cDNA), a kind gift from S. Strickland [State University of New York at Stony Brook], clone 11 (genomic clone containing the 5' LTR and coding regions of a mouse IAP gene), and mitochondrial cytochrome-c oxidase I and II cDNA clone [Pikó and Taylor 1981], kindly provided by P. Pikó [Veterans Administration Hospital, Sepulveda, CA]; murine B1/B2, cDNA probe [Larín et al. 1991], a kind gift from M. Bucán [University of Pennsylvania, Philadelphia, PA]. We isolated the murine β-actin cDNA clone from a mouse blastocyst library using a chicken β-actin cDNA [Alonso et al. 1986]. The full-length murine IL-7 cDNA probe was kindly provided by S. Gillis and L. Park [Immunex, Seattle, WA]. PCR primer sets specific for the central and 5' and 3' regions of mouse IL-7 and IFN-γ genes were purchased from Clonetech Laboratories. Radioactive probes were obtained by isolating inserts from plasmids by appropriate restriction enzyme digestion, agarose gel purification, and 32P-labeling by use of the random primer method [Feinberg and Vogelstein et al. 1983]. Probes were hybridized to library filters (1 x 10^6 to 2 x 10^8 cpm/ml) in Church buffer [7% SDS, 1 mM EDTA, 0.5 M sodium phosphate buffer (pH 7.2)] for 18–20 hr at 65°C.

cDNA library screening

Only colonies hybridizing with a given probe on two replicated library filters were considered positive. Subsequent secondary screening was performed to verify positive signals and to isolate clones for sequencing. The frequency of a given transcript in a library was determined by calculating the number of positive colonies per total cDNA colonies screened. For example, the frequency of β-actin transcripts in the egg library is 0.0011 (275 positive colonies/250,000 independent cDNA colonies screened). To estimate the total number of transcripts of a given gene in a single egg or embryo, the frequency of its occurrence in the egg or embryonic libraries was multiplied by the total number of poly(A) RNA molecules previously estimated to be present in a single egg or embryo, that is, 1.7 x 10^8 poly(A) mRNA molecules in the mouse egg, 7 x 10^8 in the two-cell embryo, 1.3 x 10^8 in the eight-cell-stage embryo, and 3.4 x 10^8 in the early blastocyst [Clegg and Pikó 1983]. Therefore, from the frequency of β-actin expression in the egg library of 0.0011, the total number of actin molecules calculated in the mouse egg is estimated to be (0.0011) 1.7 x 10^8 or 18,700 mRNA molecules.

PCR analysis of cDNA libraries and Southern blotting

Aliquots from each library were plated at high density [2 x 10^8 cfu] on Nuncleone 8.5 x 8.5-inch LB plates containing 70 μg/ml of ampicillin, incubated at 37°C for 16 hr, scraped into 50-ml centrifuge tubes, and spun at 2500g. Plasmid DNA was obtained by use of a standard alkaline lysis method [Birnboim and Doly 1979], purified by CsCl gradient centrifugation [Sambrook et al. 1989], and digested with MluI and SalI or Pvull as described by the manufacturer [New England Biolabs]. Insert cDNA was purified by gel electrophoresis and separated from agarose with spin columns [Bio-Rad]. For each PCR reaction, 10–50 ng of gel-purified insert cDNA was used. Primers for T7 and T3 polymerase promoters were synthesized in the Wistar Institute Nucleotide Synthesis Facility with the sequence published by Stratagene, and PCR was performed. Briefly, DNA template was denatured at 100°C for 15 min in 30 μl of autoclaved 1 x PCR buffer [10 mM Tris-HCl (pH 8.3), 50 mM KCl, 2.5 mM MgCl2, 0.1 mg/ml of gelatin, 0.45% NP-40, 0.45% Tween 20], followed by the addition of PCR mix [0.1 μg/ml of T7/T3 primer, 0.2 μM dNTPs, 2 units of Thermalse, [IBI/Kodak]], and placed in a thermal cycler (Perkin-Elmer) for 35–45 cycles at 94°C for 30 sec, 50°C for 30 sec, and 72°C for 1.0 min. PCR reactions for cytokine primers were performed as recommended by Clonetech. PCR products were analyzed on 1–3% agarose gels and, when necessary, blotted onto nylon membrane [Sambrook et al. 1989], exposed to 1200 J of UV light using a Stratalinker 2400, and hybridized with appropriate probes.

RNA transcription and subtractive hybridization

The template for sense RNA was generated by digesting CsCl-purified plasmid DNA with SalI for 18 hr at 37°C, followed by treatment with 5 μg/ml of proteinase K [Boehringer Mannheim] for 15 min at 56°C, phenol–chloroform extraction, and ethanol precipitation. Template for antisense RNA was prepared as described above, except that MluI was used instead of SalI. RNA synthesis was performed with T7 or T3 RNA polymerase, and the reaction buffer was supplied by the manufacturer [Promega]. For a single transcription reaction, 5–10 μg of template DNA was mixed with 1 μm each ATP, GTP, CTP, and either UTP or, for antisense RNA, biotin–UTP and UTP [10: 1, respectively], and 100 units of polymerase in reaction buffer. Tracer [32P]UTP was added at 1–2 μCi/reaction. Following transcription, template was removed by DNase treatment [50 μg/ml at 37°C for 30 min], and the synthesized RNA, purified by phenol–chloroform extraction and ethanol precipitation, was quantified either spectrophotometrically or by calculating [32P]UTP incorporation. Hybridization reactions between egg and two-cell-stage RNA were as described [Sive and St. John 1988]. Briefly, 200 ng of two-cell library-derived RNA was coprecipitated with 1 μg of biotinylated egg library-derived RNA, resuspended in 4.5 μl of hybridization buffer [250 mM HEPES (pH 7.5), 10 mM EDTA, 1% SDS] and 0.5 μl of 5 M NaCl, and hybridization was carried out for 48 hr at 65°C under oil. Hybridization buffer without SDS was added [50 μl], followed by 5 μl of streptavidin [1 mg/ml, Bethesda Research Labs], and the reaction mixture was incubated for 5 min at room temperature, followed by phenol–chloroform extraction. The organic phase was extracted twice with 25 μl of hybridization buffer without SDS, and the aqueous phases were pooled, phenol/chloroform-extracted three more
times, ethanol-precipitated, and washed. The two-cell library-derived sense RNA remaining after hybridization with egg library-derived RNA was hybridized to a 10-fold excess of eight-cell antisense RNA and treated as described above. The two-cell library-derived sense RNA remaining after hybridization and phenol–chloroform subtraction was reverse-transcribed and cloned into the pBS cloning vector as described above.

Sequencing

All cDNA clones were sequenced by use of the Sequenase kit (U.S. Biochemical) from the 5′ end with the T7 primer and the 3′ end with [35S]dATP as described by the manufacturer. Sequencing reactions were run on a 10% polyacrylamide/6% urea gel at 2000 V for 6–8 hr and exposed to X-ray film (X-Omat, Kodak) overnight at room temperature. All sequences were compared with those listed in the GenBank/EMBL data bases by use of the WORDSEARCH and FASTA commands of the GCG sequence analysis program (Devereux et al. 1984).

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