

Construction of a cDNA Microarray Derived from the Ascidian *Ciona intestinalis*

Authors: Azumi, Kaoru, Takahashi, Hiroki, Miki, Yasufumi, Fujie, Manabu, Usami, Takeshi, et al.

Source: Zoological Science, 20(10) : 1223-1229

Published By: Zoological Society of Japan

URL: <https://doi.org/10.2108/zsj.20.1223>

BioOne Complete (complete.BioOne.org) is a full-text database of 200 subscribed and open-access titles in the biological, ecological, and environmental sciences published by nonprofit societies, associations, museums, institutions, and presses.

Your use of this PDF, the BioOne Complete website, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at www.bioone.org/terms-of-use.

Usage of BioOne Complete content is strictly limited to personal, educational, and non - commercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.

Construction of a cDNA Microarray Derived from the Ascidian *Ciona intestinalis*

Kaoru Azumi^{1,4*}, Hiroki Takahashi^{2,4}, Yasufumi Miki⁴, Manabu Fujie³,
Takeshi Usami¹, Hisayoshi Ishikawa³, Atsusi Kitayama²,
Yutaka Satou³, Naoto Ueno², and Nori Satoh^{3,4}

¹Department of Biochemistry, Graduate School of Pharmaceutical Sciences, Hokkaido University, Sapporo 060-0812, Japan

²Laboratory of Developmental Biology, National Institute for Basic Biology, Okazaki 444-8585, Japan

³Department of Zoology, Graduate School of Science, Kyoto University, Kyoto 606-8502, Japan

⁴CREST, Japan Science and Technology Corporation, Tokyo 103-0027, Japan

ABSTRACT—A cDNA microarray was constructed from a basal chordate, the ascidian *Ciona intestinalis*. The draft genome of *Ciona* has been read and inferred to contain ~16,000 protein-coding genes, and cDNAs for transcripts of 13,464 genes have been characterized and compiled as the “*Ciona intestinalis* Gene Collection Release I”. In the present study, we constructed a cDNA microarray of these 13,464 *Ciona* genes. A preliminary experiment with Cy3- and Cy5-labeled probes showed extensive differential gene expression between fertilized eggs and larvae. In addition, there was a good correlation between results obtained by the present microarray analysis and those from previous EST analyses. This first microarray of a large collection of *Ciona intestinalis* cDNA clones should facilitate the analysis of global gene expression and gene networks during the embryogenesis of basal chordates.

Key words: ascidian, *Ciona intestinalis*, cDNA microarray, gene expression, development

INTRODUCTION

Ascidians or tunicates (the subphylum Urochordata) are basal chordates, and they share a common ancestor(s) with vertebrates. Ascidians provide an appealingly simple experimental system for investigating the molecular mechanisms that underlie cell-fate specification during development (Satoh, 1994; Corbo *et al.*, 2001; Satoh, 2001; Satoh *et al.*, 2003). First, the ascidian tadpole is composed of only ~2600 cells, which constitute a small number of organs including epidermis, central nervous system (CNS), endoderm and mesenchyme in the trunk, and notochord and muscle in the tail. Extensive information on the cell lineages of most major tissues and organs of ascidians is available (Conklin, 1905; Nishida, 1987). Second, the blastomeres of early ascidian embryos are large and easy to manipulate (Nishida, 2002), and permit detailed visualization of changes in gene expres-

sion during development (Satoh, 2001). Third, *Ciona* embryogenesis is rapid (about 18 hr elapses from fertilization to a free-swimming tadpole at 18°C) and the entire life cycle takes less than 3 months, facilitating genetic analyses (Nakatani *et al.*, 1999; Sordino *et al.*, 2000). Fourth, novel functions of developmental genes can be determined by misexpressing or overexpressing a variety of regulatory genes that encode transcription factors (Takahashi *et al.*, 1999) or signaling molecules (Imai *et al.*, 2002) and by functional suppression of genes with morpholino oligonucleotides (Satou *et al.*, 2001a). Finally, transgenic DNAs can be introduced into developing embryos using simple electroporation methods (Corbo *et al.*, 1997) that permit simultaneous transformation of hundreds of synchronously developing embryos, facilitating studies of transcriptional regulation.

Ciona intestinalis is a cosmopolitan species that spawns all year round and is used by researchers worldwide. Very recently, the draft genome of *Ciona intestinalis* has been determined, and its ~160 Mbp genome is estimated to contain ~16,000 protein-coding genes (Dehal *et al.*, 2002), similar to the number in other invertebrates, but

* Corresponding author: Tel. +81-11-706-3917;
FAX. +81-11-706-3917.
E-mail: kaoru@pharm.hokudai.ac.jp

only half that found in vertebrates. It has been proposed that large-scale gene duplications occurred in the vertebrate lineage after the divergence from cephalochordates and urochordates (Holland *et al.*, 1994). The ascidian, however, has a basic, non-duplicated set of the chordate-type genome (Dehal *et al.*, 2002).

In addition, we are now performing an in-depth cDNA analysis of *Ciona intestinalis* that includes the generation of expressed sequence tags (ESTs), the collection and sequencing of a non-redundant set of cDNAs, and the clarification of their expression profiles (Satoh *et al.*, 2003; Satou *et al.*, 2002a). Six cDNA libraries were made from fertilized eggs (Nishikata *et al.*, 2001), cleaving embryos (Fujiwara *et al.*, 2002), tailbud embryos (Satou *et al.*, 2001b), larvae (Kusakabe *et al.*, 2002), young adults (Ogasawara *et al.*, 2002), and the testes of mature adults (Inaba *et al.*, 2002). We then generated ESTs from both the 5' and 3' ends from ~90,000 cDNA clones. The total number of 3' ESTs was 81,637, and they were grouped into 13,464 independent cDNA clusters. The 13,464 clones were re-arrayed in 36 384-well plastic dishes for release as "*Ciona intestinalis* Gene Collection Release I (CiGCR1)" (Satou *et al.*, 2002b). This number of cDNAs corresponds to ~80% of all protein-coding genes of this ascidian. The 5' ESTs were searched against the DDBJ/Genbank/EMBL database to obtain information about the proteins the genes encode. In addition, ~10,000 cDNAs have been sequenced to determine their full-length insert. All of the *Ciona intestinalis* cDNA information can be seen at our website <http://ghost.zool.kyoto-u.ac.jp/indexrl.html>.

Given the present research circumstances described above, application of DNA microarray strategies may facilitate the identification of developmental genes with novel functions and characterization of gene networks in *Ciona intestinalis*. DNA microarrays have been used to analyze gene expression profiles during developmental stages in *Caenorhabditis elegans* (Hill *et al.*, 2000), *Drosophila melanogaster* (Furlong *et al.*, 2001; Arbeitman *et al.*, 2002), the zebrafish (Ton *et al.*, 2002), and the mouse (Miki *et al.*, 2001). To date, however, no DNA microarray containing an array of cDNAs or oligo DNAs corresponding to all the protein-coding genes of a vertebrate species is available. Our present study was therefore an attempt to construct a microarray of ascidian cDNAs containing a large number of cDNAs corresponding to ~80% of all protein-coding genes of this ascidian.

MATERIALS AND METHODS

Biological materials

Ciona intestinalis were cultivated at the Maizuru Fisheries Research Station of Kyoto University, Maizuru, Kyoto, Japan. Adults were maintained under constant light to induce oocyte maturation. Eggs and sperm were obtained surgically from the gonoduct. After insemination, eggs were washed and maintained in seawater at room temperature (18–20°C). Larvae hatched at about 17 hr of development.

Construction of cDNA microarray

CiGCR1, which contains a set of 13,464 unique *Ciona* genes, was used. Clones were obtained from the 384-well plates and cDNA inserts were amplified by PCR using Accu Taq LA DNA polymerase (Sigma-Aldrich, USA) in a 20- μ l reaction mixture. PCR products were purified using Corning FilterEXTM 384-well filter plates (Corning, USA). The quality and quantity of purified PCR products were confirmed using agarose gel electrophoresis. More than 90% of the clones gave a single band. Purified PCR products were dried and resuspended in 10 μ l of distilled water, and then mixed with 10 μ l of spotting solution (Microarray Crosslinking Reagent D, Amersham Biosciences, USA). cDNA solutions were spotted onto silanized microarray slides (type 7 Star, Amersham Biosciences) using a capillary pen type arrayer (Gene III, Amersham Biosciences). cDNA spotted slides were then exposed to 50mJ of 254nm light to crosslink DNA on slides. Each slide was spotted with duplicate sets of about 4,500 unique cDNAs on the left- and right-hand sides of the array; in total 13,396 cDNAs were spotted on four slides in duplicate. Lambda phage DNA (Takara, Japan) was used as an external control gene and was spotted onto each array block (32 $\times$ 12) with more than one spot. Spotting solution without DNA was spotted as a negative control on each block with at least 14 spots. This cDNA microarray was termed "*Ciona intestinalis* cDNA chip version 1 (Ci cDNA chip ver. 1)".

Preparation of probes and hybridization to cDNA microarray

Total RNAs were isolated from fertilized eggs and larvae by ultracentrifugation in a cesium trifluoroacetate gradient, and mRNA was extracted using an mRNA purification kit (Amersham Biosciences). The RNA quality was verified by electrophoresis using an Agilent 2100 bioanalyzer (Agilent Technologies, USA). Labeling and hybridization were performed as follows: first-strand cDNA probes were generated by incorporation of 50 μ M of either Cy3-dCTP or Cy5-dCTP (Amersham Biosciences) during reverse transcription with 400 units of SuperScript II (Gibco-BRL, USA), each 1 μ g of mRNA, 200 pg of lambda polyA⁺ RNA-A (Takara) as external control, 4 μ g of random hexamer (Invitrogen, USA), and 1 μ g of oligo-dT in a total volume of 20 μ l reaction mixture at 42°C. After the reaction, residual RNAs were degraded and neutralized, and the resulting cDNA probes purified using a Cyscribe GFX purification kit (Amersham Biosciences), according to the manufacturers protocol, and then concentrated using a vacuum centrifuge dryer. The labeling reaction efficiency of each cDNA probe was confirmed by spectrophotometry, measuring the absorbance at 260nm, 550nm and 650nm. One-fourth of the labeled probe mixture was used for hybridization of one arrayed slide, and the four slides of one set were hybridized with the same probes and under the same conditions. Hybridization reactions were performed using a manufacturer's hybridization buffer (Amersham Biosciences) containing formamide for 16 hr at 55°C in an air phase incubator. After hybridization, the slides were washed in 2X SSC/0.1% SDS for 10 min, followed by washing in 1X SSC/0.1% SDS for 5 min each at 55°C, then in 0.1XSSC at room temperature to remove SDS. Spinning at low speed in a centrifuge for 1 min dried the slides.

Scanning and data processing

The slides were scanned using a GenePix4000B DNA microarray scanner (Axon Instruments, USA) at ~635 nm (Cy5) and then at ~532 nm (Cy3). The voltage of both channels of the photo multiplier tubes were adjusted to obtain a ratio value of 1.0 for the external control spots. The resulting fluorescent intensities for each spot were quantified by an algorithm for adaptive circle segmentation of an image, and signal intensities calculated from the value of the median of the background intensities subtracted from the median of the intensities of each pixel ([F532 Median] - [B532] or [F635 Median] - [B635]) using GenePix Pro 4.0 microarray analysis software (Axon Instruments). Two independent hybridization exper-

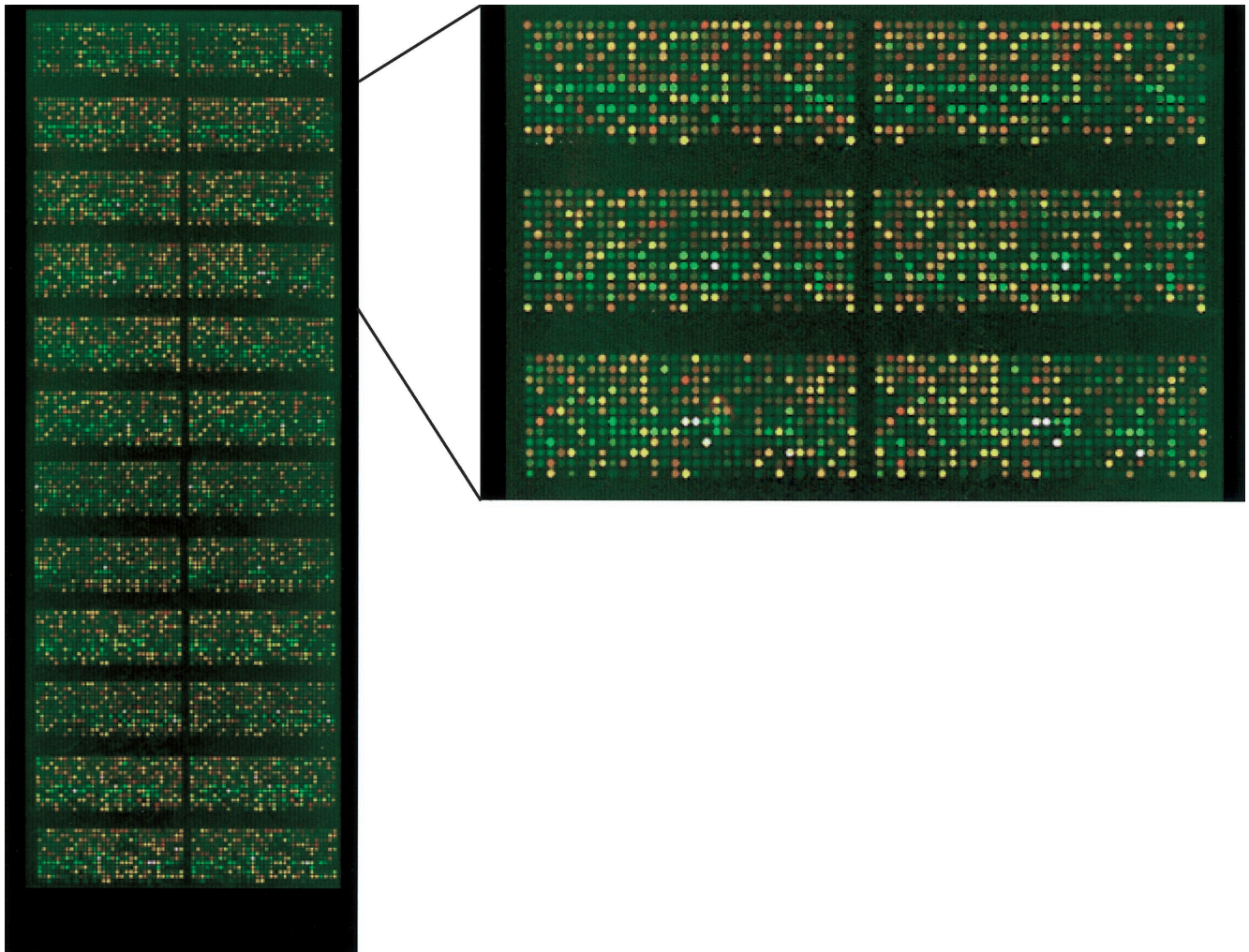


Fig. 1. The “*Ciona intestinalis* cDNA chip version 1 (Ci cDNA chip ver. 1)”: a first-generation cDNA microarray with 13,394 PCR products derived from “*Ciona intestinalis* Gene Collection Release I”. In total 13,396 kinds of cDNA were divided and spotted on four slides. A color-image of a slide of the four Ci cDNA chip ver. 1 are shown in Fig. 1, one slide was spotted with duplicate sets of about 4,500 unique cDNAs on the left- and right-hand sides of the array. Colors indicate the hybridization to each probe: green spots indicate up-regulated in larvae, red up-regulated in fertilized eggs, and yellow almost equally expressed at both stages.

iments, each using one set of slides, were performed. An intensity-dependent normalization (non-linear of LOWESS normalization) method (Cleveland and Devlin, 1988) was applied to normalize the data from replicate experiments using Genespring software version 5.0 (Silicon Genetics, USA). The fluorescence units of each duplicate spot on each slide were averaged and the ratio value then calculated; so that the signal intensity of each tested sample (larvae sample) was divided by that of the control sample (fertilized egg sample). We labeled the same RNA samples with different dyes to evaluate the variation of the results depending on the labeled dye bias; and also to determine cutoff values for up- or down-regulated genes. Under our experimental conditions, we set the significance values at >2.0 for up-regulated genes and <0.5 for down-regulated genes, respectively. The processed overlay image of a hybridized slide is shown in Fig. 1.

RESULTS AND DISCUSSION

DNA microarray analysis is a powerful technology for global gene expression profiling. The recent accumulation of

huge amounts of data on *Ciona intestinalis*, in the form of EST analyses and cDNA clone collections, led us to successfully construct an ascidian microarray containing 13,396 cDNAs. To assess the qualities of the *Ciona* cDNA microarray, we used it to investigate transcriptional differences between fertilized eggs and larvae. The mRNAs obtained from larvae and fertilized eggs were used to prepare cDNA probes differentially labeled with Cy3 and Cy5, and hybridized to the Ci cDNA chip ver. 1. In total 13,396 cDNAs were divided and spotted on four slides. Each slide of four of Ci cDNA chip ver. 1 was spotted with duplicate sets of about 4,500 unique cDNAs on the left- and right-hand sides of the array. As we have shown in a color-image of the slide of the four in Fig. 1, the signal profiling of both sides exhibited the same patterns, and the fluorescent intensities derived from duplicate spots were also almost the same. An average of ratio value of lambda phage DNA controls which were spotted elsewhere onto four slides was 1.01 ± 0.1 ($n=152$). These

data indicated that the *Ciona* cDNA microarray provides reproducible results.

We spotted cDNA onto mirror-type slides (type 7 Star, Amersham); this type of slide gave a stronger signal intensity than those of the usual clear glass because the fluorescence derived from each spot was reflected on the surface of the mirror and with high efficiency detected by the scanner. Also, this slide shows the lower background signals. We detected fluorescent signals derived from around 70% of the 13,396 spots of the *Ciona* arrays, even though we used probes prepared from 0.25 µg of each mRNA for hybridization to one slide. To determine the significant cutoffs for up-regulated genes in fertilized eggs and in larvae, we first labeled mRNAs from the fertilized eggs with both dyes and then hybridized them to the arrays. The results of hybridization yielded a linear scatter plot, with almost all of the spots lying within the range of a twofold difference (Fig. 2A). On the basis of this result, we chose >2.0 as the cutoff value for up-regulated genes in larvae and <0.5 for genes specifically expressed in fertilized eggs. As shown in Fig. 2B, experiments with fertilized egg samples and larvae samples showed variations. A total of 2,400 genes exhibited signal intensity >2.0 (up-regulated in larvae), and 2,100 genes exhibited intensity <0.5 (up-regulated in fertilized eggs).

Next, we compared the microarray data to data from EST analyses that have already been included in the *Ciona* cDNA database (<http://ghost.zool.kyoto-u.ac.jp/indexr1.html>). The cDNA libraries of *Ciona* fertilized eggs (Nishikata *et al.*, 2001) and larvae (Kusakabe *et al.*, 2002) are neither normalized nor amplified. Therefore, the number of ESTs (hereafter we call this the EST count) corresponds to the quantity

of at least the majority group of the mRNAs in the library. In other words, we can determine whether the gene is expressed either in eggs or in larvae or in both. Table 1 summarizes the 15 cDNAs with highest expression ratios as determined by the microarray analysis. Table 1 includes, for each cDNA, information about the EST count of the corresponding gene at the larval stage and fertilized egg stage, and the gene product predicted by BLASTX search. As is evident in Table 1, all of the cDNAs had large EST counts at the larval stage, but no or very small EST counts at the fertilized egg stage. For example, the ID00617 gene, which encodes a protein with sequence similarity to a mouse alpha tectorin, scored the ratio (larvae/eggs) of 240 in the array data, and its EST count was 119 in larvae and 2 in fertilized eggs. Another example is the ID00244 gene, which encodes a protein with sequence similarity to a mouse polydomain protein, and had a ratio (larvae/eggs) of 223 in the array data, while its EST count was 118 in larvae and 0 in fertilized eggs. The other 13 cDNAs showed similar results. Therefore, the quantitative expression of genes assessed by the array analysis corresponds to that assessed by EST analysis. Strictly speaking, the microarray ratio does not always show one-to-one relationship with the ratio determined by EST analysis, but in general the array ratio shows fairly good accordance with EST counts.

On the other hand, Table 2 summarizes the 15 cDNAs with the lowest array ratios (larvae/eggs). The occurrence of large EST counts in eggs clearly indicates that all of the genes are expressed maternally. The data shown in Table 2 also show a distinct co-relationship between the array ratio and EST count as to the qualitative expression of genes. For

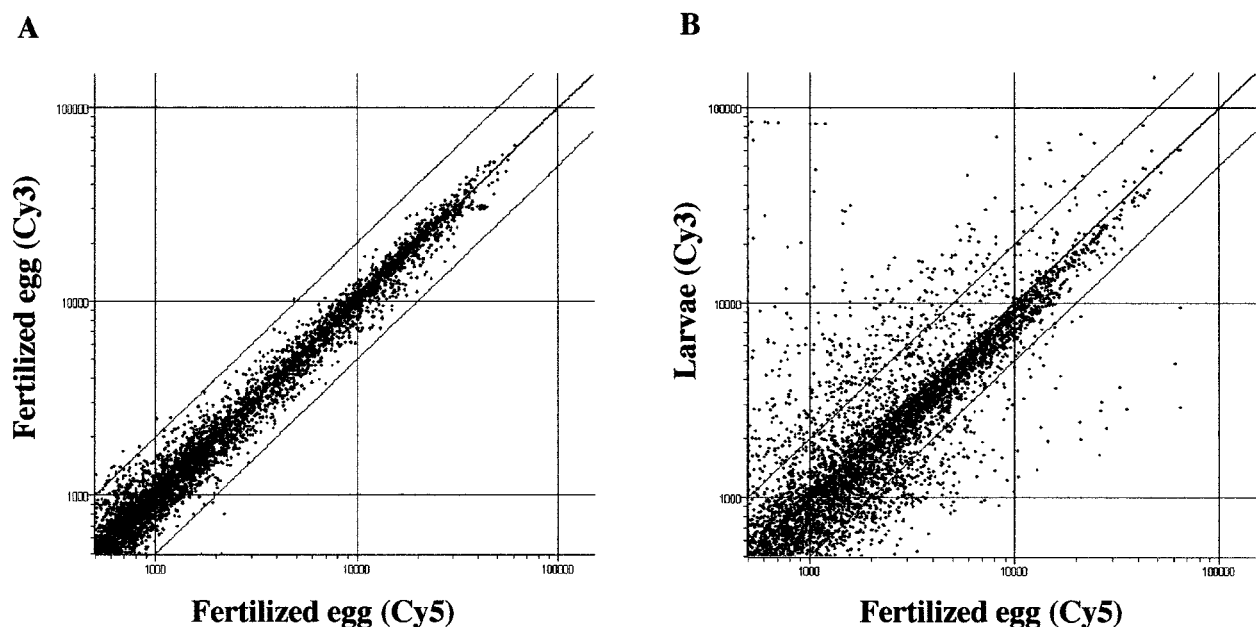


Fig. 2. (A) Scatter plot of fluorescent signal intensity from a hybridization of fertilized eggs in both the Cy3 and Cy5 channels to the *Ci* cDNA chip ver. 1. Lines show two-fold different expression in both channels. (B) Scatter plot of fluorescent signal intensity from a hybridization to the *Ci* cDNA chip ver. 1 of larvae and fertilized eggs with Cy3- and Cy5-labeled probes, respectively. Lines show twofold different expression in both channels. Spots with background-subtracted intensity lower than 500 in either the Cy3 or Cy5 channel were filtered out.

Table 1. List of genes of which expression is up-regulated in larvae.

Clutster ID	clone ID	Ratio LV/FE	Larvae(LV) /24532 EST clones	Fertilized eggs (FE) /29444 EST clones	Accession No	Data Base entry name	Organism	p
00617	cicl05f19	240	119	2	CAA68138	alpha tectorin	<i>Mus musculus</i>	2e-16
00244	cilv06e24	223	118	0	BC008135	polydomain protein	<i>Mus musculus</i>	4e-13
00056	cilv12n18	207	135	0	AAA60229	Pur protein	<i>Homo sapiens</i>	1e-37
00171	cilv30p03	194	75	0	BAA23597	CsEpi-1	<i>Ciona savignyi</i>	9e-74
00419	cilv22b11	143	363	0	BAA23647	BAI 1	<i>Homo sapiens</i>	1e-18
00035	cilv15p02	119	313	0	AAA18911	ribosomal protein L8	<i>Xenopus laevis</i>	1e-53
00093	ciad01j05	81.7	44	0	BAA36973	alpha 1 type I collagen	<i>Cynops pyrrhogaster</i>	1e-26
00065	cilv10c11	75.4	1488	0	BAA08112	nonmuscle actin	<i>Halocynthia roretzi</i>	2e-53
02124	cilv14a06	60.5	51	1	AAB33312	Tamm-Horsfall protein (THP)	<i>Bos taurus</i>	1e-18
00087	cicl22b06	59.6	395	1	AAA50025	elongation factor 1-alpha	<i>Danio rerio</i>	4e-34
00578	cilv14e05	59.2	70	2	AAA48675	HMW/LMW collagen subunit precursor	<i>Gallus gallus</i>	3e-22
00837	cilv34n07	47.6	336	1	CAB90289	alpha 5 type IV collagen	<i>Homo sapiens</i>	1e-51
01325	cilv04i10	39.0	42	1	—			
00605	cilv02p13	37.0	34	0	CAB88872	serine protease	<i>Anopheles gambiae</i>	4e-16
00031	cilv01j06	31.7	55	0	AAB30787	superfast myosin light chain-2 (MyLC2)	<i>Felis catus</i>	2e-51

Table 2. List of genes of which expression is up-regulated in fertilized eggs.

Clutster ID	clone ID	Ratio LV/FE	Larvae(LV) /24532 EST clones	Fertilized eggs (FE) /29444 EST clones	Accession No	Data Base entry name	Organism	p
00585	cieg07e03	0.08	3	112	AAH02983	similar to gene rich cluster, C9 gene	<i>Homo sapiens</i>	1e-24
00032	cicl10b06	0.11	16	136	AAD22962	claudin-1	<i>Homo sapiens</i>	1e-16
06444	cieg28d21	0.11	0	19	BAA91002	unnamed protein product	<i>Homo sapiens</i>	2e-16
02605	ciad79i24	0.12	2	39	AAF46189	CG3184-PA	<i>Drosophila melanogaster</i>	1e-63
02257	citb10p18	0.12	0	20	BAB33089	hypothetical protein	<i>Macaca fascicularis</i>	1e-24
01544	cicl20p10	0.18	2	33	BAA11926	posterior end mark	<i>Ciona savignyi</i>	4e-19
01137	cieg01m07	0.19	0	78	1D8J	GENERAL TRANSCRIPTION FACTOR TFIIIE-BETA	<i>Homo sapiens</i>	1e-16
00734	cicl01d09	0.19	0	77	CAA45968	cyclin B1	<i>Mus musculus</i>	1e-18
03719	cieg22g14	0.20	0	21	—			
04422	cicl08a24	0.23	0	25	AAC18425	NAD+-dependent isocitrate dehydrogenase	<i>Bos taurus</i>	1e-40
06573	cieg09n14	0.24	0	38	BAB93517	OK/SW-CL.48	<i>Homo sapiens</i>	1e-85
05917	cicl15d13	0.27	2	29	AAB84279	OmpR	<i>Proteus vulgaris</i>	1e-61
04248	cicl52k02	0.30	0	20	AAD09339	NAD+-specific isocitrate dehydrogenase beta subunit isoform A	<i>Homo sapiens</i>	2e-41
03061	cieg08k24	0.32	1	37	BAB28333	unnamed protein product	<i>Mus musculus</i>	1e-19
01271	cieg0402	0.32	2	20	CAA18425	cell division control protein 18	<i>Schizosaccharomyces pombe</i>	1e-19

example, the ID00585 gene scored a ratio (larvae/eggs) of 0.08 in the array data, while the EST count was 112 in fertilized eggs and 3 in larvae. The array ratio of the ID00320 gene was 0.11, while the EST count was 136 in fertilized eggs and 16 in larvae. Therefore, as in the case of genes that are up-regulated at the larval stage, the quantitative expression of down-regulated genes assessed by the microarray analysis coincided with that assessed by EST analyses. Interestingly, five genes listed in Table 2 encode proteins with sequence similarity to vertebrate or *Drosophila* counterparts, but whose function is not known. Because these genes show dramatic changes in the quantity of their transcripts during embryogenesis, their roles should be elu-

cidated in further studies.

Finally, Table 3 lists 15 cDNAs whose array ratio fell in the range of 2.0-0.5. Namely, these are cDNAs for genes whose expression was at the same level at both the fertilized egg stage and larval stage. As is evident in this Table, most of the genes showed similar EST counts at both the fertilized egg stage and larval stage.

All of the data described above clearly demonstrate that the quantitative expression of *Ciona* developmental genes can be assessed using cDNA microarrays and that the expression measured by the array analysis corresponds well with that deduced from EST analyses. Therefore, it can be concluded that the *Ciona intestinalis* cDNA microarray

Table 3. List of genes which show non-differential expression.

Cluster ID	clone ID	Ratio LV/FE	Larvae(LV) /24532 EST clones	Fertilized eggs (FE) /29444 EST clones	Accession No	Data Base entry name	Organism	p
01501	cicl20d13	1.35	80	86	AAD24490	phosphate transporter precursor	<i>Drosophila melanogaster</i>	1e-63
00555	ciad37a13	1.29	25	22	AAC46585	ribosomal protein DL11	<i>Drosophila melanogaster</i>	1e-72
00774	ciég54d17	1.26	19	14	AAA93261	zinc finger protein C2H2-25	<i>Homo sapiens</i>	2e-17
00289	cicl17o21	1.14	23	17	CAB96129	putative claudin	<i>Halocynthia roretzi</i>	5e-20
01777	ciad02b22	1.13	16	23	AAB48626	ribosomal protein L27 homolog	<i>Caenorhabditis elegans</i>	1e-37
00051	ciad06k02	1.08	18	14	AAA42291	tropomyosin TM4	<i>Rattus norvegicus</i>	1e-16
01786	ciég01c10	1.05	28	24	AAF50751	CG10576-PA	<i>Drosophila melanogaster</i>	1e-19
01131	cilv10i18	1.01	119	94	AAF52454	CG10203-PA	<i>Drosophila melanogaster</i>	1e-25
00086	ciég57m19	0.88	84	81	AAG15316	beta tubulin	<i>Notothenia coriiceps</i>	1e-71
00754	ciég34k18	0.86	32	33	BAB23963	unnamed protein product	<i>Mus musculus</i>	1e-41
00264	ciég14o23	0.78	15	14	BAA22933	proteasome p45/SUG	<i>Rattus norvegicus</i>	8e-93
03297	cicl46l24	0.78	15	14	P30153	serine/threonine protein phosphatase 2A	<i>Homo sapiens</i>	1e-70
01175	citb07m06	0.73	22	21	BAB02430	nucleolar protein	<i>Arabidopsis thaliana</i>	1e-30
02686	ciad20i18	0.72	27	19	CAB56042	glycine rich RNA binding protein	<i>Ciona intestinalis</i>	3e-40
03319	ciég49o23	0.71	18	12	AAA40884	catalase (EC 1.11.1.6)	<i>Rattus norvegicus</i>	1e-63

prepared in the present study is of a high enough quality so that it can be used for further analysis of the gene expression profiles.

PERSPECTIVE

As described above, a cDNA microarray based on CiGCR1 was constructed. EST projects as well as cDNA projects of *Ciona intestinalis* are progressing. By adding new data for transcripts expressed in gastrulae/neurulae and those expressed in the gonad (ovary), endostyle (the primordium of vertebrate thyroid gland), neural complex, heart, and blood (coelomic) cells of the adult, the present data set consists of more than 480,000 ESTs (Sato *et al.*, 2003). The ESTs are categorized into 17,834 independent cDNA clusters. An additional 4,370 cDNA clones identified from the second series are ready for release as “the supplement of CiGCR1”. Therefore, a total of 17,834 independent cDNA clones are now available as a *Ciona* gene resource. BLAST search suggests that this cDNA resource covers ~85% of the transcripts expressed in this ascidian. Along with these advances in *Ciona* cDNA projects, we are now making a “*Ciona intestinalis* cDNA chip version 2”, which includes all cDNAs including “the supplement of CiGCR1”, on a single slide. Furthermore, “*Ciona intestinalis* oligo chip version 1” can also now be prepared custom-made (Agilent Technologies). Despite their relatively high position in animal phylogeny, ascidians show several characteristic phenomena (Sawada *et al.*, 2001). Some live as colonial ascidians, formed by budding. They synthesize cellulose-like carbohydrates in the tunic. Some ascidians extensively accumulate heavy metals such as vanadium from seawater. It is to be expected that complex gene networks are involved in each of the biological activities mentioned above. Therefore, it is intriguing to ask what kinds of gene networks are responsi-

ble for various biological phenomena in the basal chordates. Furthermore, reading of the *Ciona intestinalis* draft genome has been completed (Dehal *et al.*, 2002). Thus, the results of analyses using the cDNA microarray can be interpreted using various sources of genetic information. The present construction of a cDNA microarray may therefore enormously facilitate research on the genes and gene networks of *Ciona intestinalis*.

ACKNOWLEDGMENTS

This work was a part of a CREST project “Development, Differentiation and Regeneration”, CREST, JST, Japan (N.S., K.A. and H.T.). The production of “*Ciona intestinalis* Gene Collection Release I” was supported in part by a Grant-in-Aid from the MEXT, Japan (12202001) to N.S. Kazuko Hirayama, Chikako Imaizumi, and Asako Adachi are acknowledged for their technical support.

REFERENCES

- Arbeitman MN, Furlong EE, Imam F, Johnson E, Null BH, Baker BS, Krasnow MA, Scott MP, Davis RW, White KP (2002) Gene expression during the life cycle of *Drosophila melanogaster*. *Science* 297: 2270–2275
- Cleaveland WS, Devlin SJ (1988) Locally-weighted regression: an approach to regression analysis by local fitting. *J Amer Statist Assoc* 83: 596–610
- Corbo JC, Levine M, Zeller RW (1997) Characterization of a notochord-specific enhancer from the Brachyury promoter region of the ascidian, *Ciona intestinalis*. *Development* 124: 589–602
- Corbo JC, Di Gregorio A, Levine M (2001) The ascidian as a model organism in developmental and evolutionary biology. *Cell* 106: 535–538
- Conklin EG (1905) The organization and cell lineage of the ascidian egg. *J Acad Nat Sci (Philadelphia)* 13: 1–119
- Dehal P *et al.* (2002) The draft genome of *Ciona intestinalis*: insights into chordate and vertebrate origins. *Science* 298: 2157–2167
- Fujiwara S, Maeda Y, Shin-i T, Kohara Y, Takatori N, Sato N (2002) Gene expression profiles in *Ciona intestinalis* cleavage-

- stage embryos. *Mech Dev* 112: 115–127
- Furlong EEM, Andersen EC, Null B, White KP, Scott MP (2001) Patterns of gene expression during *Drosophila* mesoderm development. *Science* 293: 1629–1633
- Hill AA, Hunter CP, Tsung BT, Tucker-Kellogg G, Brown EL (2000) Genomic analysis of gene expression in *C. elegans*. *Science* 290: 809–812
- Holland PWH, Garcia-Fernandez J, Williams NA, Sidow A (1994) Gene duplications and the origins of vertebrate development. *Development Suppl*: 125–133
- Imai KS, Satoh N, Satou Y (2002) Early embryonic expression of *FGF4/6/9* gene and its role in the induction of mesenchyme and notochord in *Ciona savignyi* embryos. *Development* 129: 1729–1738
- Inaba K, Padma P, Satou Y, Shin-i T, Kohara Y, Satoh N, Satou Y (2002) EST analysis of gene expression in testis of the ascidian *Ciona intestinalis*. *Mol Reprod Dev* 62: 431–445
- Kusakabe T, Yoshida R, Kawakami I, Kusakabe R, Mochizuki Y, Yamada L, Shin-i T, Kohara Y, Satoh N, Tsuda M, Satou Y (2002) Gene expression profiles in tadpole larvae of *Ciona intestinalis*. *Dev Biol* 242: 188–203
- Miki R *et al.* (2001) Delineating developmental and metabolic pathways *in vivo* by expression profiling using RIKEN set of 18,816 full-length enriched mouse cDNA arrays. *Proc Natl Acad Sci USA* 98: 2199–2204
- Nakatani Y, Moody R, Smith WC (1999) Mutations affecting tail and notochord development in the ascidian *Ciona savignyi*. *Development* 126: 3293–3301
- Nishida H (1987) Cell lineage analysis in ascidian embryos by intracellular injection of a tracer enzyme. III. Up to the tissue restricted stage. *Dev Biol* 121: 526–541
- Nishida H (2002) Patterning the marginal zone of early ascidian embryos: localized maternal mRNA and inductive interactions. *BioEssays* 24: 613–624
- Nishikata T, Yamada L, Mochizuki Y, Satou Y, Shin-i T, Kohara Y, Satoh N (2001) Profiles of maternally expressed genes in fertilized eggs of *Ciona intestinalis*. *Dev Biol* 238: 315–331
- Ogasawara M, Sasaki A, Metoki H, Shin-i T, Kohara Y, Satoh N, Satou Y (2002) Gene expression profiles in young adult *Ciona intestinalis*. *Dev Genes Evol* 212: 173–185
- Satoh N (1994) *Developmental Biology of Ascidians*, Cambridge Univ Press, New York
- Satoh N (2001) Ascidian embryos as a model system to analyze expression and function of developmental genes. *Differentiation* 68: 1–12
- Satoh N, Satou Y, Davidson B, Levine M (2003) *Ciona intestinalis*: An emerging model for whole-genome analyses. *Trends Genetics* 19: 376–381
- Satou Y, Imai KS, Satoh N (2001a) Action of morpholinos in *Ciona* embryos. *Genesis* 30: 103–106
- Satou Y *et al.* (2001b) Gene expression profiles in *Ciona intestinalis* tailbud embryos. *Development* 128: 2893–2904
- Satou Y, Takatori N, Fujiwara S, Nishikata T, Saiga H, Kusakabe T, Shin-i T, Kohara Y, Satoh N (2002a) *Ciona intestinalis* cDNA projects: expressed sequence tag analyses and gene expression profiles during embryogenesis. *Gene* 287: 83–96
- Satou Y, Yamada L, Mochizuki Y, Takatori N, Kawashima T, Sasaki A, Hamaguchi M, Awazu S, Yagi K, Sasakura Y, Nakayama A, Ishikawa H, Inaba K, Satoh N (2002b) A cDNA resource from the basal chordate *Ciona intestinalis*. *Genesis* 33: 153–154
- Sawada H, Yokosawa H, Lambert CC (Eds) (2001) *The Biology of Ascidians*. Springer-verlag, Tokyo
- Sordino P, Heisenberg C-P, Cirino P, Toscano A, Giuliano P, Marino R, Pinto MR, De Santis R (2000) A mutational approach to the study of development of the protochordate *Ciona intestinalis* (Tunicata, Chordata). *Sarsia* 85: 173–176
- Takahashi H, Hotta K, Erives A, Di Gregorio A, Zeller RW, Levine M, Satoh N (1999) *Brachyury* downstream notochord differentiation in the ascidian embryo. *Genes Dev* 13: 1519–1523
- Ton C, Stamatiou D, Dzau VJ, Liew C-C (2002) Construction of zebrafish cDNA microarray: gene expression profiling of the zebrafish during development. *Biochem Biophys Res Commun* 296: 1134–1142

(Received June 2, 2003 / Accepted June 28, 2003)