Interference with gene regulation in living sea urchin embryos: Transcription factor Knock Out (TKO), a genetically controlled vector for blockade of specific transcription factors

LEONARD D. BOGARAD*, MARIA I. ARNONE†‡, CHIEH CHANG*, AND ERIC H. DAVIDSON*§

*Division of Biology, California Institute of Technology, Pasadena, CA 91125; and †Stowers Institute for Medical Research, Kansas City, MO 64110

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ABSTRACT “TKO” is an expression vector that knocks out the activity of a transcription factor in vivo under genetic control. We describe a successful test of this concept that used a sea urchin transcription factor of known function, P3A2, as the target. The TKO cassette employs modular cis-regulatory elements to express an encoded single-chain antibody that prevents the P3A2 protein from binding DNA in vivo. In normal development, one of the functions of the P3A2 transcription factor is to repress directly the expression of the CyIIIa cytoskeletal actin gene outside the aboral ectoderm of the embryo. Ectopic expression in oral ectoderm occurs if P3A2 sites are deleted from CyIIIa expression constructs, and we show here that introduction of an αP3A2-TKO expression cassette causes exactly the same ectopic oral expression of a coinjected wild-type CyIIIa construct. Furthermore, the αP3A2-TKO cassette derepresses the endogenous CyIIIa gene in the oral ectoderm and in the endoderm. αP3A2-TKO thus abrogates the function of the endogenous SpP3A2 transcription factor with respect to spatial repression of the CyIIIa gene. Widespread expression of αP3A2-TKO in the endoderm has the additional lethal effect of disrupting morphogenesis of the archenteron, revealing a previously unsuspected function of SpP3A2 in endoderm development. In principle, TKO technology could be utilized for spatially and temporally controlled blockade of any transcription factor in any biological system amenable to gene transfer.

In this communication we describe a means of blocking a specific gene regulatory interaction in living sea urchin embryos. Sea urchins have a relatively simple process of embryogenesis, and an efficient and straightforward method of gene transfer has been developed, affording the opportunity to introduce expression constructs into thousands of eggs per day. Sea urchin eggs and embryos have thus emerged as a major experimental system for analysis of developmental cis-regulatory functions (1–3). Quantitative temporal and spatial patterns of reporter-gene expression can be conveniently assessed (e.g., refs. 3–5). Many relevant sea urchin transcription factors have been purified by using affinity chromatography after identification of their cis-regulatory target sites and subsequently have been cloned (6). Antisense oligonucleotides can be used to destroy maternal mRNAs encoding transcription factors; however (7), there has been no general or direct way to examine the function of specific transcription factors in sea urchin embryos by blocking their activity in vivo. This would afford the opportunity to compare the effects of canceling transcription factor activity with the effects of mutations of the relevant target sites in an expression vector (8). More generally, it would provide a means of determining downstream functions of the targeted factor.

Here we demonstrate the functional blockade of a transcription factor that had previously been shown to be responsible for spatial control of a developmentally regulated sea urchin embryo gene. This was accomplished by introducing into fertilized eggs an expression construct that encodes a single-chain antibody that binds to the factor and prevents it from forming complexes with its DNA target sites. We term this a “TKO” (Transcription factor Knock Out) vector.

The TKO vector described herein was designed to attack the SpP3A2 transcription factor (9), the initial member of a small family of transcription factors that now includes Drosophila erect wing (10), chicken IBR (11), and human NRF-1 (12). P3A2 was cloned and characterized (9, 13) after identification of two of its target sites in the cis-regulatory element of the CyIIIa cytoskeletal actin gene, which were found to be required for correct spatial expression of this gene. This gene is normally expressed only in aboral ectoderm lineages beginning early in development. CyIIIα-CAT expression constructs reproduce the aboral expression of the parent gene, but if either of the P3A2 sites in the wild-type construct is destroyed, expression spreads dramatically to the oral ectoderm (14). Similarly, if the endogenous P3A2 factor is titrated away from the CyIIIα-CAT expression construct by coinroduction of excess target-site, ectopic oral ectoderm expression is also observed (15). Our objectives in this work were threefold: first, to develop a TKO vector that would effectively sequester endogenous P3A2 transcription factor (αP3A2-TKO), the functionality of which could be assayed by determining its effects on spatial expression of CyIIIα-CAT; second, to determine whether we could affect the expression of the endogenous CyIIIa gene during embryonic development; and third, to look for any other phenotypes in αP3A2-TKO embryos that may indicate additional embryonic functions of the P3A2 transcription factor.

MATERIALS AND METHODS

DNA-Binding Inhibition Assays. P3A2 DNA-binding electrophoretic mobility-shift assays (EMSA) were performed as previously described by Calzone et al. (9) using the P3A2 binding site (top strand: 5′-GATCTTTTCGGCTTCTGCG-3′ and bottom strand: 5′-CACACCCCACGCGCATGGGC-3′) and crude nuclear extracts. Inhibition assays were performed by incubating P3A2 DNA binding reactions with serial dilutions of supernatant from hybridoma-producing αP3A2 mAbs. The αP3A2 mAbs

Abbreviations: EMSA, electrophoretic mobility-shift assay; WMISH, whole-mount in situ hybridization; scFv, single-chain antibody; GFP, green fluorescent protein; H2b, histone 2b.

†Present address: Stazione Zoologica, Anton Dohrn, Villa Comunale-Napoli 80121, Italy.
§To whom reprint requests should be addressed at: Division of Biology 156-29, California Institute of Technology, Pasadena, CA 91125. e-mail: davidson@mirsky.caltech.edu.
were created in the Caltech Monoclonal Antibody Facility (16). Hybridoma ascites fluid and bacterial extracts containing the αP3A2 sites were similarly assayed. In these assays, inhibition of P3A2–DNA complex formation was estimated quantitatively as described in Fig. 1.

**Production of the Single-Chain Anti-P3A2 Antibody.** Hybridoma poly(A) RNA was purified by using the Stratagene RNA isolation kit and Dynal (Great Neck, NY) oligo(dT) beads as described by the manufacturers. Heavy- and light-chain cDNAs were isolated from hybridoma cDNA libraries constructed by using the Stratagene Uni-Zap vector system. Subsequently, the variable heavy- and light-chain coding regions of these cDNAs were fused in-frame (separated by an encoded flexible Gly/Ser linker) into the pcDNA3.1 vector by using the expression module of the Pharmacia single-chain antibody (scFv) system.

**Construction of TKO and Green Fluorescent Protein (GFP) Expression Cassettes.** The TKO cassette was constructed in steps. First, the simian virus (SV)40 large T 3′ untranslated intron, transcriptional attenuation, and polyadenylation signals isolated from the pNL vector (17) were cloned into the EcoRI site of pBluescript II KS− [Bspoly(A) construct]. Second, the coding region of the αP3A2 scFv was recovered from the positive pcDNA3.1 phagemid by PCR, using either the Ab1 5′ primer to make the TKO expression cassette or the Ab3 5′ primer to make a missense control cassette. Primers were as follows: Ab1, 5′-GGGCGGCGCCGCCACATGGGCGGCC-ACCGCCCGCAAAAGAGAGCGTAAGGCCCCCTAGC-GA-3′; Ab3, 5′-GGGCGGAGCTCAGTGGATGTTGATTCCTCA-AGAATGC-3′. The italicized base is deleted in primer Ab3 5′; otherwise, primers Ab1 5′ and Ab3 5′ are identical.

Sequences encoding a nuclear localization site following the ATG site: Ab-reverse, 5′-GGGACTAGTCTTTGCGGTGCTG-CTTCGTTGTACCCCGCACCTGCGGC-3′. Standard PCR was carried out in 100-μl volumes containing 50 pmol of each primer, 1× Deep Vent buffer (New England Biolabs), 2 μM of dNTPs, ∼10 ng of target sequence, and sterile H2O. Reactions were heated to 94°C for 2 min, 5 units of Deep Vent polymerase was added (New England Biolabs), and the mixture was cycled 25 times (30 sec, 94°C; 30 sec, 60°C; 60 sec, 72°C) in a Perkin–Elmer 9600 thermocycler. The PCR products were digested with NotI, purified, and ligated into the Bspoly(A) I vector. The final TKO and missense cassettes were made by ligating either the cis-regulatory element (18) of the SpH2b early histone gene into the SacII site or the SpHE hatchwing enzyme gene cis-regulatory element (a gift from L. Angerer, University of Rochester, Rochester, NY; see ref. 19) into the SacI site of the respective TKO and missense constructs. The histone gene regulatory sequences were amplified from the SpH2b clone (18) with the following oligonucleotide primers: H2b5, 5′-CCCGCGGCCGGTGCTCCTAAATATG-ATTGGCAGCTTAAATTGG-3′ and H2b3, 5′-CCCCCGCGGAGTAGATTGTGATTTCTCACGAATGC-3′. The hatchwing enzyme gene regulatory sequence was amplified from the parent SpH2b-linear probe by using HE5: 5′-CCCCCGCGGA-GCTTTTTGATTGTTTGTTTTGG-3′ and HE3: 5′-CCCCCGGGGATGAAAGAAGTGATAATTGATTTC-3′. The pCANTAB5 anti-P3A2 scFv and TKO inserts were sequenced in an Applied Biosystems 373 sequencer. TKO and αP3A2-TKO mis (missense) plasmids were linearized with Clal for microinjection. A pHE/GFP expression vector was constructed by amplifying the SpHE regulatory element with primers HES5, 5′-CCCCAGGCTCAAGCTTGGTTTGTGTTTGTGGG-3′ and HE3, 5′-CCCCGGGTAGATGAAGAAGTGATAATTGATTTC-3′ and ligating the product into the GFP expression vector pGL3-Basic (5). The resulting plasmid was linearized with KpnI for microinjection.

**Microinjection, Whole-Mount In Situ Hybridization, and Fluorescence Microscopy.** Microinjection and culture of *Strongylus purpuratus* embryos were performed essentially as described (14). However, when multiple constructs were coinfected, we maintained the total amount of DNA injected at a constant level by proportionately lowering the amount of carrier DNA. Detection of endogenous *CyIIIA* expression by whole-mount in situ hybridization (WMISH) was carried out following the protocol of Ransick et al. (24) with the following modifications. The fixation step was extended to 20 min and the probe concentration was raised to 0.4 ng/μl. The *CyIIIA* antisense probe was generated by *in vitro* transcription from the T7 promoter of the *CyIIIA*–*HpaI* vector that had been digested with *PvuII*. This generates a digoxigenin-labeled probe containing 147 bases of the *CyIIIA*, 3′ untranslated region, and an 873-bp tail of pbBlueScript II SK− (procedure of David G.-W. Wang, personal communication). Fluorescence microscopy, sorting, and WMISH of GFP-injected embryos were performed as described (5).

**RESULTS**

αP3A2 mAb and αP3A2-TKO Construct. The TKO expression construct was designed to express, translate, and transport an anti-P3A2 reagent into the nucleus so as to inhibit P3A2 from carrying out its developmental regulatory functions *in vivo*. A set of αP3A2 mAbs was generated against the full-length P3A2 protein. These were screened for their ability to inhibit this protein from binding to its target DNA sequence by adding them to gel-shift reactions at increasing dilutions. Inhibitory activity was measured quantitatively as described in Fig. 1. Inhibition constants were determined for 12 different hybridoma subclones. mAb 7B12/2E7 displayed the strongest inhibitory effects on P3A2 DNA binding and was chosen as the
basis for the αP3A2-TKO vector. The same mAb was used for an extensive series of measurements of P3A2 prevalence throughout embryological development (16). The specificity of this antibody was confirmed by the observation that it reacted with only a single molecular species of the known size of the P3A2 transcription factor in total nuclear extracts from all stages of development as well as in total egg lysate (Fig. 1 of ref. 16).

To build the αP3A2-TKO vector, the mRNA was extracted from 7B12/2E7 hybridoma cells, and by using the universal IgG primers amplified DNAs encoding the heavy-chain and light-chain V-regions were cloned in-frame into the Pharmacia pCANTAB vector surrounding a flexible Gly/Ser linker. Thus, a construct encoding a single-chain antibody was produced. Gel-shift inhibition assays were repeated to test the inhibitory activity of the αP3A2 scFv, which was extracted for this purpose from bacteria expressing the pCANTAB vector. Its activity was found to be quantitatively equivalent to that of the starting mAb. As described in Materials and Methods, the αP3A2 scFv was then incorporated into an expression cassette, the organization of which is summarized in Fig. 2. In addition to the scFv itself, the major features of the αP3A2-TKO construct are: (i) a nuclear localization site intended to increase the concentration of the scFv in the nucleus relative to that of the endogenous nuclear P3A2 factor; (ii) simian virus (SV)40 intron, 3′ trailer, termination, and poly(A) addition sites, features that are conventionally used in most sea urchin gene-transfer vectors; and (iii) a cis-regulatory control element. This last design feature, in principle, permits expression of the TKO vector to be targeted to any domain of the embryo for which a cis-regulatory element is available. However, in the present experiments we utilized instead ubiquitously, or nearly ubiquitously, active cis-regulatory elements derived from a histone H2b gene (18) and from a hatching enzyme gene (19).

Stable incorporation of exogenous DNA in sea urchin embryo blastomeres is mosaic, although entirely random with respect to lineage (20), and therefore embryos are obtained that have incorporated the exogenous DNA into every possible spatial domain. In some of the experiments that follow, the αP3A2-TKO vector was coinjected with a GFP reporter under control of the same cis-regulatory element, so that the particular cells incorporating the αP3A2-TKO construct could be identified visually by GFP expression following the procedures of Arnone et al. (5). Thus, we could examine embryos in which the construct was present in endoderm, aboral ectoderm, or oral ectoderm, as desired.

**αP3A2-TKO Effects on Spatial Expression of a Wild-Type CyIIα-CAT Construct.** When CyIIα-CAT constructs lacking the two known P3A2 target sites are injected into eggs, ectopic oral ectoderm expression is observed in an average of 34% of stained embryos (14). Because this is statistically equivalent to the percentage of embryos that incorporate exogenous DNA into oral ectoderm founder cells, it follows that essentially all oral ectoderm cells will express the CyIIα-CAT construct if interactions with the P3A2 repressor are obviated by deletion. In the CyIIα-CAT experiments summarized in Table 1, αP3A2-TKO or a missense frame shift control differing from αP3A2-TKO by deletion of only 1 bp were coinjected with wild-type CyIIα-CAT (14). Wild-type CyIIα-CAT and ΔP3A2-HF, the CyIIα-CAT construct lacking the two P3A2 sites, were injected at the same time as the TKO constructs and performed as reported (14). When coinjected with wild-type CyIIα-CAT, the missense αP3A2-TKO had no effect (2% oral expression). However, a remarkable result was obtained when wild-type CyIIα-CAT was co-injected with αP3A2-TKO: 26% of stained embryos now displayed oral ectoderm expression, as illustrated in the example reproduced in Fig. 3c and d. The 26% value could represent a slight underestimate of αP3A2-TKO effects or CyIIα-CAT expression because in these experiments only morphologically normal stained embryos were counted, and as we discuss below, a significant fraction of embryos receiving αP3A2-TKO develop abnormally. In any case, the results are those predicted if the αP3A2-TKO vector effectively sequesters the endogenous P3A2 factor, thus preventing its interaction with the CyIIα-CAT regulatory target sites. That is, as Table 1 shows, the fraction of ectopically expressing embryos is about the same as when P3A2-CyIIα-CAT interactions are instead prevented by deletion of these target sites.

We note here that coinjection of the anti-P3A2 antibody protein into eggs together with CyIIα-CAT also led to ectopic oral expression of the CyIIα-CAT reporter (C. Kirchhamer and E.H.D., unpublished results). These effects were not observed when an irrelevant antibody was injected. However, this approach can be applied only to early embryos; furthermore, the αP3A2 and the control IgG caused lethal arrest of development in an unacceptably large fraction of embryos.

**αP3A2-TKO Effects on Spatial Expression of the Endogenous CyIIα Gene.** The foregoing results confirm the role of the P3A2 transcription factor as a spatial repressor of CyIIα-CAT expression and raise the question whether the endogenous gene could be similarly affected by αP3A2-TKO. This was not obvious a priori, because we had found earlier that the stability of endogenous CyIIα regulatory complexes exceeds that of the equivalent CyIIα-CAT complexes: thus, in vivo competition with excess target sites for a number of CyIIα factors stoichiometrically reduced expression of exogenous CyIIα-CAT, whereas expression of the endogenous CyIIα gene was unaffected in the same embryos (22).

The normal aboral ectoderm expression of the endogenous CyIIα gene is displayed by WMISH in Fig. 3a and b. When eggs were injected with αP3A2-TKO and endogenous CyIIα expression was similarly monitored, ectopic expression of the

![Fig. 2. The TKO expression cassette. Histone 2b or hatching enzyme cis-regulatory elements (A) employed in the cassette are modular and can easily be exchanged with any other regulatory sequence. The start of transcription (B) is set to be at least 20 bp upstream of a canonical Kozak ATG (C) and a nuclear localization sequence (D) that are best fit to sea urchin initiators and codon usage. The heavy- (E) and light-chain (G) variable regions of the anti-P3A2 scFv are separated by an encoded flexible Gly/Ser linker (F). To process, stabilize, and terminate the TKO transcripts, we inserted the SV40 large T 3′ intron (I), polyadenylation (J), and transcriptional termination (K) sequences after the stop codon (H).](image-url)
endogenous gene was observed. As summarized in Table 1, 27% of these embryos display clear expression in clones of oral ectoderm, and frequently in limited clones of gut cells as well. Examples are reproduced in Fig. 3e and f. Three embryos that displayed ectopic expression in the missense control were also found (Table 1). Because the endogenous CyIIIa gene is never expressed erroneously, the single base mutation that distinguishes aP3A2zTKOmis from aP3A2zTKO might not have entirely prevented synthesis of a functional TKO product. For example, a partially functional protein might have been generated by use of a downstream ATG. In any case, the percentage of ectopic oral expression observed in the aP3A2zTKO sample is very similar to that seen with either the mutated CyIIIa reporter lacking P3A2 sites (14) or in the coinjections of aP3A2zTKO and wild-type CyIIIa reporters (Table 1). This is as expected, because ectopic oral expression depends on clonal incorporation of the aP3A2zTKO construct in oral ectoderm cells in all three experiments.

**A Specific aP3A2zTKO Embryonic Phenotype.** Embryos developing from eggs injected with aP3A2zTKO display a specific lethal phenotype. This occurs in a relatively high fraction of embryos, 44% in the experiment tabulated in Table 2, although in other experiments somewhat more modest frequencies of 20–30% were recorded. The aP3A2zTKOmis control resulted in only 5% abnormal embryos (Table 2). This phenotype is best described as a failure to form a complete archenteron, caused by a disorganization of the endoderm. Fig. 3g shows a normal 72-hr prism-stage embryo that developed from an egg that had been injected with the missense control construct, whereas the embryo in Fig. 3h displays the
Results are listed in Table 3. Significantly, all of the embryos endodermal clones. This result again supports a weak expres- gastrulation defects (Table 3), and all of these contained large embryos in the missense control experiment also showed marker (2, 23, 24), as shown, for example, in Fig. 3 of specification; that is, they continue to express the endoderm cells in these embryos retain their endodermal state endoderm are illustrated in Fig. 3 gastrulate and that contained the exogenous DNA in the ectoderm clones displayed ectopic expression of 19), whereas 9 of 10 embryos that contained large oral- large endoderm clones again were gastrulation-defective (19 of

Table 2. *aP3A2-TKO* effect on embryo development

<table>
<thead>
<tr>
<th>Construct</th>
<th>Embryos, no. tested</th>
<th>Gastrulation-defective, %</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>aP3A2-TKO</em></td>
<td>419</td>
<td>44</td>
</tr>
<tr>
<td><em>aP3A2-TKOMis</em></td>
<td>345</td>
<td>5</td>
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*aP3A2-TKO* phenotype. This embryo has not completed gas- trulation. Its endoderm consists of a pile of nonadherent cells, and the form of the embryo is round rather than triangular. The secondary mesoderm appears unaffected, however. Such embryos appear to develop normally up to gastrulation. The implication is that *P3A2* transcription factor is necessary for morphogenesis of the archenteron, a previously unknown function.

To further investigate the role of the *P3A2* transcription factor in archenteron formation, we performed a series of coinjection experiments making use of GFP as a marker to determine location of the *aP3A2-TKO* construct in the experimental embryos. GFP expression constructs can be used to determine the mosaic expression domain of a coinjected construct if both constructs are driven by the same cis-regulatory elements, because shortly after injection the exogenous DNAs are all ligated together into a concatenate and are then incorporated into the same blastomeres (5). In these experiments, the hatching enzyme cis-regulatory element was used to drive both the GFP and *aP3A2-TKO* constructs. Results are listed in Table 3. Significantly, all of the embryos bearing *aP3A2-TKO* that failed to gastrulate express the GFP marker in the endoderm (89 of 89), and less than 10% of the set of embryos in which GFP is expressed in the endoderm develop normally. All of those that do are distinguished by small GFP clone sizes (not shown). On the other hand, most of the 48% of the embryos injected with the missense control that contained endodermal clones developed normally, even though these clones were often quite large. An example is shown in Fig. 3j. Embryos bearing *aP3A2-TKO* that failed to gastrulate and that contained the exogenous DNA in the endoderm are illustrated in Fig. 3 k and l. The disorganized endoderm cells in these embryos retain their endodermal state of specification; that is, they continue to express the *Endo16* marker (2, 23, 24), as shown, for example, in Fig. 3i. A few embryos in the missense control experiment also showed gastrulation defects (Table 3), and all of these contained large endodermal clones. This result again supports a weak expres- sion of an *aP3A2-TKO* activity because of override of the frameshift, as considered above.

To explore further the relation between location of clonal incorporation and occurrence of the gastrulation phenotype, we carried out the experiment summarized in Table 4. Em- bryos bearing *aP3A2-TKO* plus GFP constructs were sorted according to the locus of incorporation of the expression constructs, and *CyIIIa* expression was then monitored by WMISH in the sorted samples. Table 4 shows that the locus of incorporation correlates perfectly with the effect of *aP3A2-TKO* on *CyIIIa* expression. All embryos selected for large endoderm clones again were gastrulation-defective (19 of 19), whereas 9 of 10 embryos that contained large oral-ectoderm clones displayed ectopic expression of *CyIIIa* in the oral ectoderm. Conversely, all embryos bearing large aboral ectoderm clones displayed normal *CyIIIa* expression.

It may be concluded from these experiments that the function of *P3A2* in archenteron morphogenesis is cell-autonomous, i.e., that the factor is needed in the archenteron cells themselves. Similarly, we confirm that the function of *P3A2* in oral ectoderm as a spatial repressor of *CyIIIa* trans- scription is also autonomous, as proposed earlier (14).

**DISCUSSION**

We describe here the use of a genetic expression vector that abrogates the activity of a transcription factor in a living embryo. Three main results were obtained. First, the intracel- lular efficacy of a scFv for this purpose was established by demonstration of the same ectopic expression of the *CyIIIa-CAT* vector as is caused by deletion of the target sites for this transcription factor. Second, we confirmed that the *P3A2* factor indeed serves as a spatial repressor for the *CyIIIa* gene and extend this observation from *CyIIIa-CAT* expression constructs to the endogenous *CyIIIa* gene. Third, we discov- ered a hitherto unknown function of *P3A2* in archenteron morphogenesis. Both this function and the effects of the *P3A2* factor on *CyIIIa* expression are cell autonomous.

It is striking that *CyIIIa-CAT* and *CyIIIa* gene expression are so efficiently expanded to the oral ectoderm in embryos bearing *aP3A2-TKO* in oral ectoderm clones. *P3A2* is initially a relatively prevalent maternal factor, present in about 2 × 10^6 molecules per embryo (16) and is later zygotically transcribed; there are a few thousand to a few hundred molecules per nucleus throughout development (16, 25). The affinity of the scFv encoded by the *aP3A2-TKO* vector for the *P3A2* factor is significantly higher than that of the factor for its DNA target site (9, 13) as calculated from experiments such as that shown in Fig. 1. Thus, the scFv apparently acts as a very effective intracellular sequestering agent. Nor is any nonspecific early embryonic lethality observed, in contrast to the general dele- terious effect of injecting the mAb per se into eggs (unpublished results). Furthermore, use of a genetic vehicle for producing the *aP3A2* scFv ensures that it will continue to function through development (depending of course on the cis-regulatory element driving its expression), which is never reliably the case for either protein or mRNA introduced directly into the egg. The results shown in this paper confirm that *aP3A2-TKO* continues to function through embryogenesis (Fig. 3).

An unexpected aspect of the spatial derepression of the endogenous *CyIIIa* gene caused by introduction of *aP3A2-TKO* is that ectopic expression was observed not only in oral ectoderm but also in gut (Fig. 3 e and f). Recent studies (ref. 14, and earlier observations reviewed therein) proved that the two *P3A2* sites deleted in the experiments of Kirchhamer and Davidson (14) are required to prevent ectopic oral-ectoderm expression of *CyIIIa-CAT* constructs, but no ectopic gut ex- pression was observed in these experiments. However, there are additional potential *P3A2* sites in the *CyIIIa* cis-regulatory element, and these may be responsible for controlling expres- sion in gut.
A new role for P3A2 in archenteron formation was revealed when we examined the postgastrular lethal phenotype produced by aP3A2-TKO injection into fertilized eggs. This effect occurs only when the construct is present in large clones of archenteron cells. These cells fail to adhere to one another so that the archenteron disintegrates into a wholly or partially disorganized pile of endoderm cells. Yet these cells retain their state of specification, as monitored by expression of an endoderm-specific gene, Endo16. The implication is that genes encoding some cell surface proteins are controlled by P3A2 either directly or indirectly. If directly, it is possible that P3A2 acts positively in this context, although its function is clearly negative in regulation of the CyIIa gene.

There are many obvious possible extensions of the TKO technology described here, both in research and potentially for gene therapy (26). We are now extending the TKO approach to several other sea urchin embryo transcription factors. This may become a general method for examination of transcriptional functions. One powerful potential advantage of the TKO method is the possibility of exactly controlling the time and place of TKO expression. This depends directly on the cis-regulatory system employed to drive the transcription of the TKO vector. Thus, for example, TKO vectors could be built and inserted into the mouse genome that would blockade given the TKO vector. Thus, for example, TKO vectors could be built and inserted into the mouse genome that would blockade given the TKO vector.

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