

# Spemann organizer transcriptome induction by early beta-catenin, Wnt, Nodal, and Siamois signals in Xenopus laevis

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The earliest event in Xenopus development is the dorsal accumulation of nuclear β-catenin under the influence of cytoplasmic determinants displaced by fertilization. In this study, a genome-wide approach was used to examine transcription of the 43,673 genes annotated in the Xenopus laevis genome under a variety of conditions that inhibit or promote formation of the Spemann organizer signaling center. Loss of function of β-catenin with antisense morpholinos reproducibly reduced the expression of 247 mRNAs at gastrula stage. Interestingly, only 123 β-catenin targets were enriched on the dorsal side and defined an early dorsal β-catenin gene signature. These genes included several previously unrecognized Spemann organizer components. Surprisingly, only 3 of these 123 genes overlapped with the late Wnt signature recently defined by two other groups using inhibition by Dkk1 mRNA or Wnt8 morpholinos, which indicates that the effects of β-catenin/ Wnt signaling in early development are exquisitely regulated by stage-dependent mechanisms. We analyzed transcriptome responses to a number of treatments in a total of 46 RNA-seg libraries. These treatments included, in addition to β-catenin depletion, regenerating dorsal and ventral half-embryos, lithium chloride treatment, and the overexpression of Wnt8, Siamois, and Cerberus mRNAs. Only some of the early dorsal β-catenin signature genes were activated at blastula whereas others required the induction of endomesoderm, as indicated by their inhibition by Cerberus overexpression. These comprehensive data provide a rich resource for analyzing how the dorsal and ventral regions of the embryo communicate with each other in a self-organizing vertebrate model embryo.

embryonic induction  $\mid$  Wnt signaling  $\mid$  embryonic patterning  $\mid$  Cerberus  $\mid$  Spemann organizer

With the recent completion of the *Xenopus laevis* genome (1), it has now become practical to study global transcriptional changes at the earliest stages of development in this classical model organism: for example, by analyzing the transcriptomes of the dorsal and ventral regions of the gastrula embryo (2). Recent studies analyzed the zygotic Wnt pathway in *X. laevis* and *Xenopus tropicalis* embryos, defining late gene signatures transcriptionally activated by endogenous Wnt8 expressed in the ventro-lateral marginal zone (3, 4). This zygotic Wnt signal occurs during mid- to late-gastrulation, promotes the formation of ventral-posterior tissues, and inhibits head development. For example, when Wnt8 DNA constructs that are expressed only at gastrula are injected into embryos (5), head structures are reduced. Reciprocally, overexpression of Wnt antagonists such as Dickkopf1 (Dkk1) (6) and Frzb1 (7) results in enlarged head structures.

The opposite phenotype is elicited by microinjection of *Wnt* mRNAs during early cleavage, generating embryos with spectacular secondary axes containing complete heads when injected into ventral blastomeres (8). When injected radially into embryos,

Wnt8 mRNA leads to a dorsalized phenotype consisting entirely of head structures without trunks and a radial Spemann organizer (9–11). Similar dorsalizing effects are obtained by incubating embryos in lithium chloride (LiCl) solution at the 32-cell stage (12). LiCl mimics the early Wnt signal by inhibiting the enzymatic activity of glycogen synthase kinase 3 (GSK3) (13), an enzyme down-regulated by the canonical Wnt signal (14, 15). The molecular nature of the endogenous dorsal signal in *Xenopus* eggs remains elusive, but it is clear that its main effect is to cause the stabilization of  $\beta$ -catenin during cleavage on the dorsal side of the embryo (16, 17). A  $\beta$ -catenin antisense morpholino ( $\beta$ -CatMO) is one of the most potent and reliable reagents in *X. laevis* embryology, and its microinjection results in embryos lacking Spemann organizer tissue and dorsal development (18).

Dorsal β-catenin stabilization requires a cortical rotation of the egg cortex toward the animal pole and the displacement of maternal cytoplasmic determinants containing Dishevelled (Dvl) toward the animal pole (19, 20). Interestingly, the endogenous cytoplasmic determinants are refractory to inhibition by Dkk1and Frzb mRNAs, which, however, readily inhibit microinjected

#### **Significance**

We present a genome-wide study of the signals responsible for the early induction of the body axis in the following experimental conditions:  $\beta$ -catenin morpholino;  $Wnt,\ Siamois,\$ and  $Cerberus\$ mRNAs; LiCl treatment; and dorsal-ventral regenerating half-embryos bisected at gastrula. Comparing 46 RNA-seq libraries, we uncovered the genetic networks that initiate dorsal-ventral patterning and Spemann's organizer formation. We defined an early  $\beta$ -catenin signature that has only minor overlap with recently published late zygotic Wnt signatures. The relation of these early steps of development to endome-sodermal germ layer induction was studied by overexpressing the growth factor antagonist Cerberus. This study offers a rich resource for understanding the earliest inductive events in the body plan of a model vertebrate embryo.

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Data deposition: The data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database, www.ncbi.nlm.nih.gov/geo (accession no. GSE93195).

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Wnt mRNA (21, 22). The combination of maternal β-catenin, which induces the homeobox transcriptional activator Siamois/Twin (10), with high zygotic Nodal/TGF-β (called Xnr1/2/5/6 in Xenopus) signaling on the dorsal side induces formation of Spemann organizer tissue (23). The organizer in turn secretes BMP antagonists such as Chordin and Noggin, Wnt antagonists such as Dkk1, Frzb1 and Crescent, and multivalent inhibitors such as Cerberus (24). These proteins shape dorsal-ventral (D-V) and anterior-posterior (A-P) patterning (25, 26). Therefore, understanding the signaling pathways that induce the transcriptional activation of Spemann organizer tissue is of fundamental importance for understanding vertebrate morphogenesis.

In the present study, we applied genome-wide transcriptome analyses to investigate the induction of the Spemann organizer by the maternal Wnt/ $\beta$ -catenin signal. We used  $\beta$ -catenin MO-, Wnt8 mRNA-, Siamois mRNA-, and Cerberus mRNA-injected embryos and LiCl-treated embryos. We also performed RNA-seq of uninjected embryos bisected into dorsal and ventral fragments at midblastula and allowed to develop for 5 h before isolating RNA at early gastrula. This procedure maximizes dorsal-ventral (D-V) differences and, in addition, tests the regenerative potential of the embryo (26). A total of 46 RNA-seq libraries were examined, comparing levels of the expression of 43,673 transcripts at late blastula (stage 9) and early gastrula (stage 10.5) of X. laevis. The results define an early  $\beta$ -catenin

signature that has very little overlap with the zygotic Wnt signature and generate a comprehensive genome-wide catalog of the earliest developmental transcriptional changes in a vertebrate embryo.

#### **Results**

**Defining Spemann Organizer Induction by β-Catenin.** As shown in Fig. 1*A*, the dorsal maternal determinant signal can be blocked by microinjection of β-CatMO at the two- to four-cell stage. This signal can be mimicked by microinjection of *Wnt8* mRNA, LiCl treatment, or microinjection of mRNA for the downstream effector *Siamois*. The early β-catenin signal results in the induction of the TGF-β-related molecules Xnr5/6, and a mixture of organizer growth factor inhibitors on the dorsal side that oppose signals provided by BMP and the zygotic (late) Wnt signaling pathways on the ventral side of the embryo (24, 25) (Fig. 1*A*).

As indicated in Fig. 1B, the effect of  $\beta$ -catenin loss of function or *Wnt8* mRNA gain of function was analyzed at early gastrula by RNA-seq in three independent experiments. In addition, inhibition of GSK3 by LiCl at the 32-cell stage and overexpression of *Siamois* mRNA were analyzed in duplicate, and the effect of blocking endomesoderm induction with *Cerberus* mRNA was analyzed in triplicate (Fig. 1B). As controls, we used uninjected embryos because previous work indicates that they behave like mock-injected embryos (27). We also analyzed transcripts from

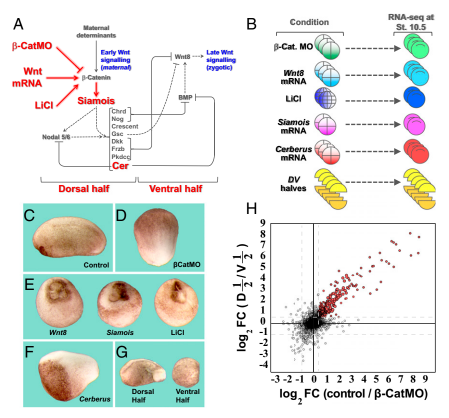


Fig. 1. RNA-seq libraries for transcriptome analysis of early β-catenin targets. (A) Diagram showing the early β-catenin-dependent gene network on the dorsal side and the zygotic Wnt8 signal that arises later on the ventral side. The embryonic manipulations that were subjected to RNA-seq in the present study are indicated in red. (B) For RNA-seq, embryos were injected with β-catenin MO, Wnt8, Siamois, or Cerberus mRNA, treated with LiCl, or cut into dorsal and ventral halves at stage 8. All embryos were harvested at stage 10.5 for RNA extraction and RNA-seq. Some embryos were also analyzed at blastula stage 9. (C-G) Phenotypes of embryos subjected to manipulations in this study. Note that β-CatMO-injected embryos lacked all traces of a dorsal axis whereas Wnt8, Siamois mRNA microinjected, and LiCl-treated embryos were all radially dorsalized. Cerberus mRNA-microinjected embryos lacked endomesoderm and consisted mostly of anterior neural tissue and cement gland. Ventral half-embryos lacked dorsal axes whereas dorsal half-embryos developed a complete axis. (Magnification: 8x.) (H) Scatter plot comparing the  $\log_2$  of fold change (FC) in RPKM upon β-catenin knockdown with that of dorsal halves/ventral halves. Shown here are 40,157 transcripts; red dots indicate transcripts decreased by β-catenin MO (by at least 1.4-fold in average of three experiments) and enriched in dorsal halves compared with ventral halves (by at least 1.5-fold in average of five experiments). Full datasets are provided in Dataset S1. Cer, Cerberus; D, dorsal half; St., stage; V, ventral half.

dorsal and ventral half-embryos (bisected at stage 8 and allowed to regenerate for 5 h until stage 10.5) in quintuplicate (Fig. 1*B*). In all experiments, sibling embryos were allowed to develop until tailbud stage, and gastrula stage 10.5 RNA samples were used for RNA-seq only when completely dorsalized or ventralized phenotypes, such as those in Fig. 1 *C–G*, were obtained. The RPKM (reads per kilobase per million mapped reads) for all annotated *X. laevis* genes are provided in Dataset S1, and the raw data for all 46 libraries are available at the GEO repository (accession no. GSE93195).

β-Catenin binds to thousands of gene promoters during early *Xenopus* development (3, 4). However, the initial accumulation of β-catenin is dorsal (16). In agreement with this finding, a scatter plot of the fold change (FC) of transcripts inhibited by β-CatMO correlated strongly with dorsal/ventral enrichment at early gastrula (Fig. 1*H*, red dots). Overall, many more transcripts were inhibited by β-catenin loss of function than were activated (Fig. S1*A*). We conclude that β-catenin knockdown provides an excellent method for D-V gene discovery.

**Defining the Early Dorsal** β-Catenin Transcriptional Signature. Among the genes repressed by β-CatMO, we found that 247 were reproducibly inhibited in each of all three experiments at stage 10.5 (Fig. S1B). About half of those genes were equally expressed in dorsal and ventral regenerating half-embryos, indicating that, although they required β-catenin, they were not dependent on the early dorsal maternal determinant signal (Fig. S1B, dashed box). Once genes without dorsal enrichment were removed, we were left with a set of 123 transcripts (Fig. S1C) that not only were decreased by β-CatMO in three independent replicates, but also were reproducibly enriched dorsally in four of five D-V experiments. The D-V quintuplicate samples identified most of the known Spemann organizer genes, as well as the classical ventrally enriched genes (Fig. S1D).

The early dorsal β-catenin signature identified by these experiments is listed in Fig. 2A. Of these 123 dorsal genes, 86 were reported to be bound by  $\beta$ -catenin in X. tropicalis (4) (Dataset S2). There were also 124 genes that were decreased by  $\beta$ -CatMO but not dorsally enriched, of which 34 bound β-catenin in ChIP experiments (Dataset S3) (4). The early dorsal signature includes many secreted factors and transcription factors known to be expressed in Spemann's organizer (2, 24), as well as many previously undescribed dorsal genes. As shown in Fig. 3, nine of these genes were enriched dorsally in a statistically significant way, and included PLEKHG5 (a RhoA GEF also known as SYX); TMEM150B (a transmembrane protein); BHLHE22 (a class E basic helix-loop-helix protein); FAM89A (Family with Sequence Similarity 89 Member A); Xlaev18002038m (unknown); CRNRIP1 (cannabinoid receptor interacting protein); NAT8B (probable N-acetyl transferase 8B); and LOC100170590.L (unknown Xenopus ORF MGC115585). In situ hybridization for LOC100170590.L confirmed that it was indeed expressed in the Spemann organizer at stage 10.5 whereas, at tailbud (stage 24), it was localized in the eye, brain, somites, and proctodeum (Fig. 3 J and K). The requirement for  $\beta$ -catenin and dorsal enrichment for LOC100170590.L were validated through qRT-PCR (Fig. S24). This gene has a possible homolog in coelacanth fishes, but not in mammals (Fig. S2B).

In generating our early β-catenin signature, several arbitrary cutoff criteria were applied (RPKM above 1, decrease by 1.4-fold in three β-CatMO experiments, and dorsal enrichment above 1.5-fold in four libraries). We therefore sought an independent method to validate this signature. To this end, we applied Weighted Gene Coexpression Network Analysis (WGCNA) to 29 libraries sequenced at stage 10.5 (28). This method, which detects modules of highly correlated genes, generated a cluster of 233 genes that contained 106 of the 123 transcripts (86%) in the early β-catenin signature (Fig. S3 and Dataset S4), which

suggests that our early  $\beta$ -catenin signature is independent of subjective cutoff points.

The signature presented here defines 123 X. laevis dorsal transcripts that require the early  $\beta$ -catenin signal. These results should be compared with those recently obtained by Kjolby and Harland (3), who used embryos microinjected with the Wnt antagonist Dkk1 to identify a late Wnt signature of 82 genes that constitute zygotic Wnt/ $\beta$ -catenin transcriptional targets at midgastrula (stage 11.5). Remarkably, only 3 genes (IRX1, MYF5, and SEBOX) overlapped between the early and late Wnt signatures (Fig. 2B). We conclude that a very specific early dorsal  $\beta$ -catenin gene signature can be defined by combining a requirement for  $\beta$ -catenin and dorsal expression.

The Early  $\beta$ -Catenin Gene Signature Is Activated by LiCl and by *Wnt8* and *Siamois* mRNAs. The repression by  $\beta$ -CatMO and dorsal enrichment of the 123 transcripts that constitute the early  $\beta$ -catenin signature can be visualized in Fig. 4A. Unsupervised hierarchical grouping clustered the classical Spemann organizer genes together, such as *Chordin*, *Xnr3*, *Crescent*, *Siamois*, *Goosecoid*, *PKDCC*, *OTX2*, *Cerberus*, *Noggin*, *Dkk1*, *ADMP*, and others (Fig. 4A, sidebar).

To investigate the relationship between the early β-catenin signature and the Wnt pathway, we microinjected embryos radially with Wnt8 mRNA in triplicate. Heat map analysis indicated that most early dorsal β-catenin signature genes were activated by Wnt8 mRNA (Fig. 4B). Similarly, LiCl treatment and Siamois mRNA microinjection (in duplicate) induced most genes in this signature (Fig. 4B). The expression of the early β-catenin signature across the different conditions sequenced was reproducible, as indicated by Pearson correlation matrix analysis (Fig. 4C). In addition, principal component analysis (PCA) showed that replicates and Spemann organizer-inducing conditions clustered together, confirming reproducible expression of the early β-catenin signature (Fig. S4). Dataset S2 provides the RPKM levels for the early β-catenin signature genes in each library.

Analysis of the early  $\beta$ -catenin signature enrichment among *Wnt8* mRNA-, LiCl-, *Siamois* mRNA-, and *Cerberus* mRNA-induced genes was performed using Gene Set Enrichment Analysis (GSEA) (29). All genes with RPKM values above 1 (in corresponding control embryos, ~16,000 genes) were ranked according to their average fold change. The early  $\beta$ -catenin gene set was greatly enriched within the genes ranked according to their levels of induction by *Wnt8* mRNA (indicated as vertical black lines in Fig. 4D). Similarly, LiCl- and Siamois-induced gene ranking showed enrichment of this gene set at the top of the list (Fig. 4 E and F). We conclude that the dorsal  $\beta$ -catenin gene set closely mirrors early Wnt, LiCl, and Siamois signaling in the early embryo.

Cerberus Inhibits Most Endomesodermal Organizer Genes. Cerberus is a protein secreted by the head organizer that functions as a multivalent growth factor antagonist by inhibiting the Nodal/ TGFβ, BMP, and Wnt pathways (30, 31). To investigate the convergence of these pathways with the early  $\beta$ -catenin signature, we overexpressed Cerberus mRNA in X. laevis embryos, and RNA-seq libraries were prepared at early gastrula in three independent experiments. Most genes within the early dorsal β-catenin signature were inhibited, presumably indicating a requirement for endomesodermal differentiation (Fig. 4B, in blue). However, a few genes in the signature were increased by Cerberus mRNA, including Xnr5, Xnr6, and many neural genes such as Hes1, Zic1, Zic4, Otx2, Irx1, Nkx6-2, and Tiki/Trabd2a (Fig. 4B). GSEA analysis confirmed that many components of the  $\beta$ -catenin gene set were inhibited by Cerberus mRNA although some transcripts still ranked highly in the list (Fig. 4G).

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## A Early dorsal β-Catenin Signature

#### **Secreted Factors** ADMP bone morphogenetic protein 2 preproprotein AGR2 PREDICTED: anterior gradient protein 2 homolog isoform X1 BMP2 bone morphogenetic protein 2 preproprotein CER1 cerberus precursor CHRD chordin precursor CPE carboxypeptidase E preproprotein CRESCENT Crescent / secreted frizzled-related protein 5 precursor DKK1 dickkopf-related protein 1 precursor FGF20 fibroblast growth factor 20 secreted frizzled-related protein 3 precursor **FRZB** KAZALD1 Mig30 / kazal-type serine protease inhibitor domain-containing NOG noggin precursor protein kinase domain-containing protein, cytoplasmic precursor **PKDCC** PRSS27 serine protease 27 precursor SHISA2 protein shisa-2 homolog precursor Xnr3 / Nodal homolog precursor XNR3 XNR5 Xnr5 / Nodal homolog precursor XNR6 Xnr6 / Nodal homolog precursor

#### Transcription Factors and Regulators

Transcription ractors and Regulators		
BHLHE22	class E basic helix-loop-helix protein 22	
CRX	homeobox protein OTX2 isoform b	
DMBX1	Otx3 / diencephalon/mesencephalon homeobox protein 1	
EGR1	early growth response protein 1	
FOXB1	forkhead box protein B1	
FOXD3	forkhead box protein D3	
GATA4	transcription factor GATA-4	
GATA5	PREDICTED: transcription factor GATA-5 isoform X1	
GSC	homeobox protein goosecoid	
HES1	transcription factor HES-1	
HHEX	hematopoietically-expressed homeobox protein HHEX	
HLX	H2.0-like homeobox protein	
IRX1	iroquois-class homeodomain protein IRX-1	
LEF1	lymphoid enhancer-binding factor 1 isoform 2	
LHX1	LIM/homeobox protein Lhx1	
LMO1	rhombotin-1 isoform a	
LMO4	PREDICTED: LIM domain transcription factor LMO4 isoform X1	
MYF5	myogenic factor 5	

OSR1 PREDICTED: protein odd-skipped-related 1 isoform X1
OTX2 homeobox protein OTX2 isoform b
homeobox protein SEBOX
SIAMOIS Siamois (Sia) homeobox protein
SKOR1 SKI family transcriptional corepressor 1
T brachyury protein isoform 1

homeobox protein Nkx-6.2

oligodendrocyte transcription factor 3

ZIC1 zinc finger protein ZIC 1
ZIC2 zinc finger protein ZIC 2
ZIC3 zinc finger protein ZIC 3
ZIC4 zinc finger protein ZIC 1

NKX6-2

OLIG4

#### **Transmembrane Proteins and Receptors**

ADAM19	disintegrin and metalloproteinase domain-containing protein 19
CHRNA4	neuronal acetylcholine receptor subunit alpha-4 isoform 1 precursor
CNRIP1	CB1 cannabinoid receptor-interacting protein 1 isoform X2

**EFNB2** ephrin-B2 precursor **FZD8** frizzled-8 precursor

LPAR6 lysophosphatidic acid receptor 6

LRIG3 leucine-rich repeats and immunoglobulin-like domains protein 3 isoform 2

PCDH9 PREDICTED: protocadherin-9 isoform X2

PDGFRA PREDICTED: platelet-derived growth factor receptor alpha isoform X1

PRRT1 proline-rich transmembrane protein 1

SLC2A1 solute carrier family 2, facilitated glucose transporter member 1 SLC34A2 sodium-dependent phosphate transport protein 2B isoform a

TIKI metalloprotease TIKI1 isoform 1 precursor

TMCC1 transmembrane and coiled-coil domains protein 1 isoform X7

TMEM150B transmembrane protein 150B isoform X3

#### Enzyme/Kinases

ALDH1A2	retinal dehydrogenase 2 isoform 1
APOBEC2	probable CU-editing enzyme APOBEC-2
ARHGAP39	rho GTPase-activating protein 39
DENND2C	DENN domain-containing protein 2C isoform 1
DNASE1L3	deoxyribonuclease gamma isoform 1 precursor
HS6ST1	heparan-sulfate 6-O-sulfotransferase 1
NAT8B	probable N-acetyltransferase 8B
DI E1/110 F	and a structure to a second a second and a second at the s

PLEKHG5 pleckstrin homology domain-containing family G member 5 ST3GAL5 lactosylceramide alpha-2,3-sialyltransferase isoform 2

#### Others and Unknowns

#N/A #1	Xelaev18042095m
#N/A #2	Xelaev18003555m
#N/A #3	Xelaev18000592m
#N/A #4	Xelaev18004073m
#N/A #5	MGC115585 protein Xenopus laevis
#N/A #6	Xelaev18002038m
#N/A #7	LOC101733854.1
FAM110C	PREDICTED: protein FAM110C isoform X1
FAM89A	protein FAM89A
KRT12	keratin, type I cytoskeletal 12

keratin, type I cytoskeletal 12

LZTS1 PREDICTED: leucine zipper putative tumor suppressor 1 isoform X1

NCK2 PREDICTED: cytoplasmic protein NCK2 isoform X1

RAB20 ras-related protein Rab-20
RGS1 regulator of G-protein signaling 1
SNCG gamma-synuclein
SPNS2 protein spinster homolog 2

SRR1-like protein

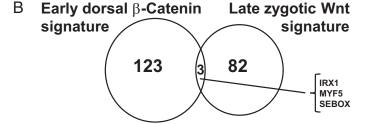


Fig. 2. The early dorsal β-catenin gene signature. (A) Genes were categorized as follows: secreted factors, transcriptional factors and regulators, transmembrane protein and receptors, enzymes/kinases, and others and unknowns. Human gene symbols and full names are shown (2), except when *Xenopus*-specific names exist. Because the subtetraploid *X. laevis* has long and short homologs for many genes, duplicate names have been removed from this list. For unknown genes, the gene IDs from the JGI9 *X. laevis* genome are indicated. (B) Venn diagram showing that the early β-catenin gene signature is distinct from the late zygotic Wnt signature (3), with only three genes in common.

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When the entire transcriptome of *Cerberus* mRNA-injected embryos was examined, many more genes were significantly repressed than were induced (Fig. 5A). This observation was

consistent with the inhibition of mesodermal and endodermal differentiation marked by *Xbra* and  $Sox17\beta$ , respectively (Fig. 5*B*). It is noteworthy that expression of *xWnt8*, the major Wnt

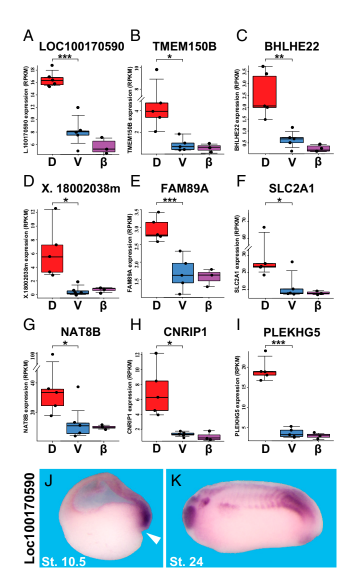


Fig. 3. The early  $\beta$ -catenin signature contains previously undescribed dorsal transcripts. (A-I) Box plots indicate dorsal and ventral gene expression levels (in RPKM) in five independent regenerating half-embryo experiments. These genes have either not been previously characterized or have been only superficially investigated. Box plots illustrate the reproducibility of replicate experiments. Each dot represents the expression in RPKM of the corresponding gene in a distinct experiment. Genes above the whisker are considered outliers according to the interquartile range (IQR) calculated for each box plot. The difference between dorsal and ventral expression was statistically significant for all nine genes, with all genes surpassing the 95% confidence interval mark at a minimum. The  $\beta$ -CatMO RPKMs ( $\beta$ ) were similar to those of ventral halves. Statistical significance was determined through two-tailed pairwise P values. Significant differences in expression values (RPKM) are indicated as  $\leq 0.05$ ,  $\leq 0.01$ , and  $\leq 0.005$ . (J) In situ hybridization validating that unknown Loc100170590 is indeed expressed in Spemann organizer mesoderm at gastrula. White arrowhead indicates the dorsal blastopore lip. (Magnification: 16x.) (K) At tailbud stage 24, Loc1001170590 is expressed in eye, central nervous system, somites, and proctodeum. (Magnification: 14x.) D, dorsal; St., stage; V, ventral.

that signals from ventrolateral mesoderm during *Xenopus* gastrulation, is essentially eliminated in *Cerberus* mRNA-injected embryos (Fig. 5B and Dataset S1). *Cerberus* overexpression represses, and induces, a wide range of genes of both dorsal and ventral origin by stage 10.5 (Fig. 5C and Datasets S5 and S6). Previous RNA-seq studies used the Nodal/TGF- $\beta$  receptor inhibitor SB431542 to define Nodal target genes (27). We observed

that *Cerberus*, a Nodal antagonist, strongly repressed 107 genes, of which 49 overlapped with genes repressed by SB431542 (Dataset S5). In GSEA analyses, a set of 107 genes repressed by *Cerberus* were found predominantly at the bottom of the correlation of those inhibited by  $\beta$ -CatMO (Fig. 5D). Cerberus-repressed genes also correlated with genes arranged according to their D-V enrichment (Fig. 5E). When the Cerberus-repressed gene set was probed against rankings of transcripts induced by *Wnt8* or *Siamois*, they appeared at either the top or bottom of the ranked list, presumably reflecting the dual inhibition of Nodal and BMP pathways by this multivalent antagonist (Fig. 5 F and G). We conclude that many, but not all, of the early  $\beta$ -catenin–induced genes require endomesoderm germ layer specification by stage 10.5.

The Early β-Catenin Signature Is Established at Gastrula. In this study, we concentrated on the analysis of genes expressed at early gastrula because it is at this stage that Spemann organizer tissue is fully functional. We also analyzed gene expression at the earlier stage 9 (blastula, about 3.5 h before harvesting RNA from siblings used for gastrula studies) in some experiments and provide the complete RPKM data for stage 9 libraries in Dataset S7. As shown in Fig. 6A, most of the 123 genes in the early β-catenin signature were not activated until gastrula (shown as blue at stage 9 in the RPKM heat map). However, a subset of β-catenin targets had higher expression at blastula (in red) relative to gastrula stage 10.5 (Fig. 6A, boxed on top). Accordingly, most genes in the early  $\beta$ -catenin signature were unaffected by β-CatMO at stage 9 (Fig. 6B). Results from all experimental conditions tested at blastula and gastrula (control, β-CatMO, Wnt8 mRNA, Cerberus mRNA, LiCl, and Siamois mRNA) were allowed to cluster hierarchically by row according to RPKM expression as shown in Fig. 6C. At blastula stage 9, most genes in the signature were unaffected by β-catenin depletion or dorsalizing treatments (Fig. 6C, in blue). However, a group of genes did respond to Wnt-mimicking agents and were inhibited by β-CatMO at stage 9 (boxed at top of Fig. 6C and listed in Dataset S8), which included the classical Wnt targets Siamois (10) and Xnr3 (32). In addition, many organizer-specific genes were induced at stage 9, such as Xnr5, Xnr6, Admp, Cerberus, Noggin, and Zic2. The relative delay in expression of the majority of the early β-catenin signature genes expressed at gastrula is likely explained by the need of the early  $\beta$ -catenin signal to activate *Siamois* as well as *Xnr5/6* expression to achieve full organizer expression (33, 34), as indicated in the pathway outlined earlier in Fig. 1A.

#### **Discussion**

In Xenopus, fertilization triggers a cortical rotation that transports Dvl-containing cytoplasmic organelles, called maternal dorsal determinants, along microtubules toward prospective dorsal regions (19, 35). The rotation brings cytoplasmic determinants from vegetal cytoplasm in contact with the marginal zone of the egg before first embryonic cleavage (36, 37). This process determines the site at which the dorsal side of the embryo is formed by stabilizing maternal β-catenin protein, which accumulates in dorsal nuclei as early as the two- to four-cell stage (16, 17). The full molecular composition of the cytoplasmic determinant and the mechanism by which β-catenin is stabilized remain unknown although the participation of maternal Wnt11 and Dvl proteins, perhaps in combination with the formation of multivesicular bodies, has been proposed (22, 38). It is likely that the elusive cytoplasmic determinant is protein in nature because recent RNA-seq analyses of early cleavage stages (eight-cell) of X. laevis or X. tropicalis embryos have not detected any asymmetrically expressed mRNAs along the D-V axis (39, 40). The early β-catenin stabilization, in combination with zygotic Siamois and Nodal signaling, induces Spemann organizer formation at the gastrula stage (11). There has been considerable progress

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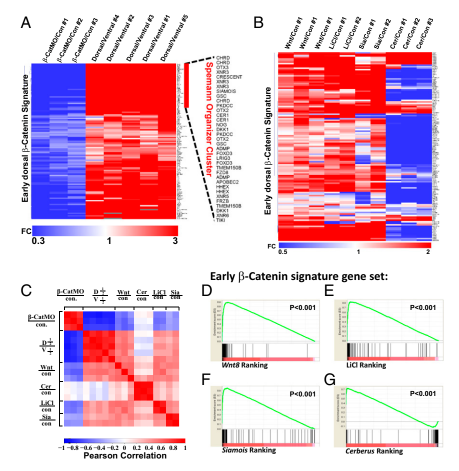


Fig. 4. Correlation of the early dorsal β-catenin gene signature with Wnt8, LiCl, Siamois-induced genes, and Cerberus-repressed genes. (A) Heat map showing the fold change of early β-catenin signature genes in three β-catenin MO and five dorsal halves with respect to control and ventral half-embryo libraries, respectively. Fold changes (FC) over controls were used as inputs, and unsupervised hierarchical clustering of columns and rows was performed. Note that hierarchical clustering of rows identified most classic known Spemann organizer genes on the top of the heat map. (β) Heat map displaying regulation of genes composing the early β-catenin signature by Wnt8 mRNA, LiCl, and Siamois or Cerberus mRNAs. (C) Correlation matrix of β-CatMO-, Wnt8-, Wnt8-

in identifying the components of the Spemann organizer (2, 41-43). However, a global examination of the transcriptomic landscapes elicited by the early nuclear  $\beta$ -catenin signal and associated pathways that induce formation of Spemann organizer tissue was lacking.

The Early  $\beta$ -Catenin Dorsal Gene Set. In this study, we set out to determine the transcriptional output of the early maternal β-catenin signal. To this end, we sequenced the transcriptome of β-CatMO-injected embryos at gastrula (stage 10.5) in three independent experiments and compared them with their uninjected controls. We used β-CatMO as a loss-of-function reagent because it blocks dorsal development with 100% penetrance (18). Secreted Wnt antagonists (such as Dkk1 and Frzb) are ineffective at blocking the endogenous early β-catenin cytoplasmic determinant, yet inhibit the zygotic Wnt8 ventral-lateral signal that mediates A-P patterning (24). We found 247 transcripts that reproducibly required  $\beta$ -catenin, but, of these, to our surprise, only about half were dorsal genes, which suggested that many β-catenin transcriptional targets were not dependent on the dorsal cytoplasmic determinant signal. In fact, recent ChIP-seq studies indicate that endogenous β-catenin binds to

5,000 genes in the *X. tropicalis* genome, but only a small fraction of these are regulated by Wnt at any one stage (4).

To specifically focus on the genetic program activated by  $\beta$ -catenin on the dorsal side, we sequenced libraries from regenerating dorsal and ventral half-embryos bisected at midblastula and cultured until early gastrula in five independent experiments. This approach resulted in much higher D-V differences than those observed in previous studies using dissected dorsal and ventral blastopore lips (2) because the BMP and late Wnt8 signals in the ventral half were unopposed by diffusing organizer-secreted antagonists (26). Transcripts were filtered so that only those decreased by  $\beta$ -catenin knockdown and dorsally enriched resulted in a set of 123 genes, which were named the early dorsal  $\beta$ -catenin signature (Fig. 24).

This filtering approach was validated by WGCNA (28), a conventional clustering method that grouped 86% of the early  $\beta$ -catenin signature within a single cluster. The early  $\beta$ -catenin signature included many Spemann organizer genes. In addition, it identified many other genes that have been previously undescribed (such as LOC100170590.L) (Fig. 3 A, J, and K), unknown to be dorsally enriched, or, in one case, only superficially described

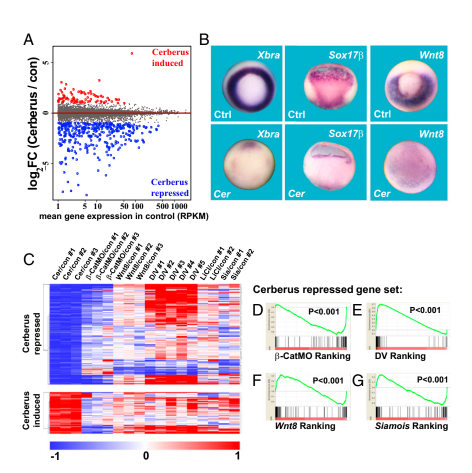


Fig. 5. The Cerberus gene signature identifies early dorsal β-catenin genes that require endomesodermal induction. (A) MA-plot comparing gene expression of 16,729 mRNAs between control and Cerberus mRNA-injected embryos. The average log<sub>2</sub> fold change in expression of transcripts in Cerberus mRNA-injected embryos over control embryos is plotted on the y axis; the average gene expression in RPKM of uninjected control embryos is on the x axis. This MA-plot confirms the proper normalization of the RNA-seq data because the horizontal nature of the line across the zero baseline shows that there is no systematic bias. (B) Cerberus overexpression greatly inhibits Xbra, xSox17β, and xWnt8 mRNA expression when examined by in situ hybridization. (Magnification: 10x.) (C) Heat map showing how gene signatures repressed or induced by Cerberus mRNA were differentially regulated by various dorsalizing or ventralizing conditions. Note that a large block of Spemann organizer genes strongly activated in D/V regenerating half-embryos (in red, top of the heat map) are repressed by Cerberus-mediated endomesodermal inhibition. (D–G) GSEA analysis of the Cerberus-repressed gene set. Approximately 16,000 transcripts were ranked according to β-CatMO, D/V expression, and Wnt8 and Siamois mRNA induction and were examined for the ranking of Cerberus-repressed genes (vertical lines). Cer, Cerberus; con, control; Ctrl, control; D/V, dorsal/ventral ratio; Sia, Siamois.

(PLEKHG5) (Fig. 3*I*) (43). Studies are needed to dissect the function of these dorsal genes in D-V patterning. A likely explanation for why these genes had been missed in our previous studies (2) is that dorsal and ventral half-embryos were allowed to regenerate, maximizing differences as a result of unopposed BMP and Wnt8 signaling in ventral halves.

A point of interest is that the *Wnt8* gene itself is highly increased in ventral half-embryos and, conversely, has one of the highest fold decreases in *Siamois*-injected and LiCl-treated embryos (Dataset S1). Wnt8 is responsible for most of the late Wnt signature in *Xenopus* and is known to be induced by BMP4 (4, 44). The genes inhibited by  $\beta$ -catenin but not dorsally enriched also constitute an interesting group, in particular in light of the recent finding that  $\beta$ -catenin is bound to thousands of gene loci in *Xenopus* (4). Of these 124 genes, 34 bound  $\beta$ -catenin in ChIP assays (4) (Dataset S3).

A Rich Genome-Wide Embryonic Resource. We have compiled the RNA-seq results in complete RPKM files of expression levels for the 43,673 genes annotated in the *Xenopus* genome (1) comprising a total of 46 RNA-seq libraries. This rich resource is now accessible to the *Xenopus* community or anyone interested in the complete transcriptional repertoire of Spemann organizer-inducing signals

in vertebrate embryos. The embryonic transcriptional response should be considered globally because, for each action in the organizer, there is a reaction in the ventral side (45).

The widespread dorsal gene activation by overexpression of the homeobox protein Siamois was particularly interesting because it implied that most of the early maternal  $\beta$ -catenin signal is mediated through this Xenopus-specific transcription factor (10, 46, 47). The X. laevis genome contains six active Siamois/Twin genes, which ensure a rapid response to the maternal β-catenin signal (48). We note that Siamois overexpression tended to generate more robust RPKM inductions than Wnt8 or LiCl, presumably because this transcription factor reached higher concentration levels after direct injection (Dataset S1), even though the dorsalized embryonic phenotypes were comparable (Fig. 1E). Microinjection of Cerberus mRNA repressed genes that require endoderm and mesoderm formation and increased neural genes within the early dorsal β-catenin signature. The Cerberus data also defined global gene signatures repressed and induced by this mRNA that may help decipher the physiological role of this growth factor antagonist.

We also examined the timing of the activation of the early β-catenin signature. By comparing the transcriptomes of Spemann organizer activating or inhibiting treatments at stage 9 (blastula) versus 10.5 (about 3.5 h later), we confirmed that,

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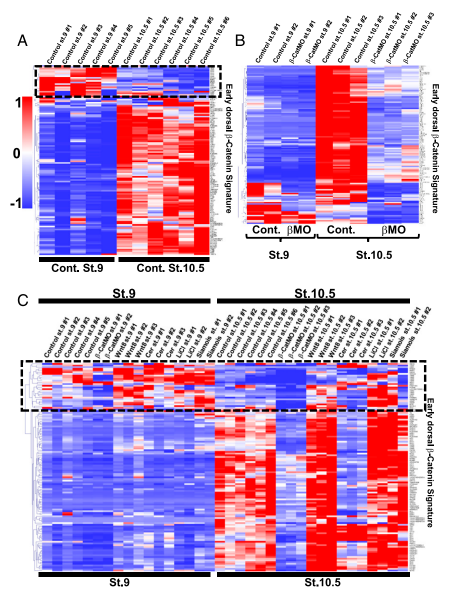


Fig. 6. Comparison of the expression of the early dorsal β-catenin gene signature at blastula and gastrula stages. (A) Heat map showing that the majority of the early dorsal β-catenin gene signature had higher expression levels (in red) in WT control (Cont.) embryos at stage 10.5 rather than at blastula stage 9. RPKMs of the genes constituting the early β-catenin signature genes at blastula stage 9 or gastrula stage 10.5 (results from five and six sequenced libraries, respectively) were normalized by row. (B) The majority of genes in the early β-catenin signature were unaffected by β-CatMO at stage 9; the effect of β-catenin MO becomes evident at stage 10.5. (C) Heat map comparing RPKM levels of the early dorsal β-catenin signature in 46 libraries at blastula and gastrula stages. Only a minority of the β-catenin signature were inhibited by β-CatMO or activated by Wnt-like signals at blastula stage 9. RPKM values of all libraries were allowed to cluster automatically according to rows. The cluster delimited by a box at the top indicates transcripts that are capable of being induced or repressed by dorsalizing or ventralizing treatments at blastula stage 9. A list of these genes is provided in Dataset S8 and includes many known early Wnt targets, such as Siamois, Xnr3, Xnr5, Xnr6, Noggin, and ADMP. The scale bar at the bottom indicates normalized RPKM values. In conclusion, most genes in the early β-catenin signature become sensitive to β-catenin depletion only at gastrula stage, presumably indicating the requirement of the endodermal and mesodermal germ layer induction for Spemann organizer development. Cer, Cerberus; Cont, control; St., stage.

whereas all genes within the dorsal β-catenin signature are activated by stage 10.5, a small cluster of these genes were expressed at higher levels at stage 9 (Fig. 6). It is known that in *Xenopus* β-catenin bound to TCF-3 plays an important role in recruiting epigenetic regulators even before zygotic transcription (49). Nuclear β-catenin recruits Prmt2 (a protein-histone arginine methyltransferase) that causes an epigenetic mark called H3R8Me2 (dimethylation of histone H3 at Arginine 8), which deposits an epigenetic memory that poises promoters for transcription up to six cell cycles before Wnt target genes are first transcribed (49). The maternal β-catenin/Wnt signal, followed by zygotic Siamois and Nodal signaling, provides the initial step in the

process that gives rise to the formation of the Spemann organizer in the dorsal marginal zone at the gastrula stage (11, 23). Spemann organizer tissue then orchestrates dorsal ventral patterning by secreting a mixture of growth factor antagonists (24).

Comparison of the Early and Late Wnt Signatures. The cellular responses to Wnt ligands in early development are stage-dependent, which represents a difficulty when trying to assign a general transcriptional signature for the pathway. There are two distinct waves of Wnt signaling in *Xenopus* early development. Maternal Wnt/ $\beta$ -catenin signaling drives the formation of the Spemann organizer in dorsal regions whereas zygotic Wnt8,

expressed at mid- to late-gastrula in ventro-lateral mesoderm, promotes the development of the posterior region of the embryo.

Recently, two studies have defined the late zygotic Wnt8 gene set using a combination of RNA-seq and ChIP-sEq (3, 4). Kjolby and Harland (3) used the Wnt inhibitor Dkk1 to define the transcriptome activated by Wnt signal at mid- to late-gastrula in X. laevis. A total of 82 genes constituted their late Wnt signature. Nakamura et al. (4) used X. tropicalis to screen for target genes repressed by a morpholino specifically targeting Wnt8. As controls, they rescued these effects with Wnt8 DNA and also selected only those genes bound by endogenous  $\beta$ -catenin at TCF/LEF sites. These more stringent criteria identified fewer genes (4), but all were included in the late signature from the other laboratory (3). Strikingly, it was found that  $\beta$ -catenin bound to 5,193 genes at gastrula, most of which are not regulated by Wnt at this stage;  $\beta$ -catenin recruitment does not imply transcriptional regulation, which must depend on additional developmental contexts (4).

That the early and late Wnt signatures would differ was expected (11). However, the extreme degree of divergence we found was surprising. We identified only 3 genes (Irx, Myf5, and Sebox) that overlapped between our early β-catenin signature of 123 genes and the late Wnt signature of 82 genes (3) (Fig. 2B). It is currently unknown what provides this "switch" from maternal to late  $\beta$ -catenin/Wnt signaling although epigenetic mechanisms or stage-specific TCF proteins may be involved (49, 50). It has been proposed that β-catenin recruitment may occur during early embryonic stages to DNA regulatory elements responsible for Wnt-regulated expression in other tissues at much later stages (4). In this regard, cross-talk between signaling pathways such as BMP, Nodal, and FGF may be important. In addition, as yet unknown lineage master regulators, as occurs with the recruitment of β-catenin in hematopoietic progenitor cells (51), might be involved in triggering differential transcriptional programs. It is generally accepted that the main role of  $\beta$ -catenin at gastrula is to signal dorsally and establish the Spemann organizer. However, in the present study it was found that many other genes require β-catenin for transcription but are not dorsally enriched, perhaps indicating as yet unknown transcriptional roles for β-catenin in early development. Studies are needed to investigate the effects of  $\beta$ -catenin/Wnt signaling at later embryonic stages: for example, to examine the deployment of the Hox complexes that are not yet active at the stages studied here.

#### **Conclusion**

This work presents an early dorsal  $\beta$ -catenin signature of 123 genes and provides the complete transcriptional landscape of treatments that massively stimulate or inhibit formation of the Spemann organizer in X. laevis. The genome-wide transcriptomic signatures associated with  $\beta$ -catenin loss of function, with dorsal and ventral half-embryos, and with Wnt8, LiCl, Siamois, and Cerberus phenotypes and the specific differences and similarities among them provide a rich substratum for investigations into the self-organizing nature of the earliest gene regulation events during development of a vertebrate embryo.

### Materials and Methods

Embryo Manipulations, mRNA Injections, and Whole-Mount in Situ Hybridization. *X. laevis* were purchased from Nasco. Embryos were generated through in vitro fertilization and cultured in 0.1× Marc's modified Ringer's (MMR) and staged according to Nieuwkoop and Faber (52). A total of 24 ng of morpholino oligonucleotide against *Xenopus* β-catenin (18) was injected four times marginally at two-cell stage. For in vitro mRNA synthesis, pCS2-*xWnt8*, pCS2-*xCerberus*, and pCS2-*xSiamois* were linearized with NotI and transcribed with SP6 RNA polymerase using the Ambion mMessage mMachine kit. Synthetic mRNAs were injected into the marginal zone of each blastomere at the four-cell stage as follows: 3 pg of *xWnt8*; 100 pg of *xCerberus*; and 20 pg of *xSiamois*. All of the injected embryos were cultured until stage 9 or 10.5. For LiCI treatment, 32-cell stage embryos were treated with 0.3 M LiCl in 0.1× MMR for 7 min, washed three times with 0.1× MMR, and cultured

until stage 9 or 10.5. For dorsal and ventral half-embryos, regularly cleaving stage 8 midblastula embryos were bisected into dorsal and ventral halves (44) and cultured until gastrula stage 10.5. For hybridization probe synthesis, pCS2-xWnt8 was linearized with BamH1 and transcribed with T3 RNA polymerase; pCS2-xbra with Sal1 and transcribed with SP6 RNA polymerase; pCS2-xSox17 $\beta$  with Apa1 and transcribed with SP6 RNA polymerase. Wholemount in situ hybridizations were performed as described at www.hhmi.ucla. edu/derobertis.

cDNA Library Preparation, RNA Sequencing, and Data Analysis. Total RNA was isolated from two whole embryos or six half-embryos (dorsal or ventral) with the Absolutely RNA Miniprep Kit (Agilent). Libraries were constructed with the Illumina TruSeq RNA Library Preparation Kit V2 according to the manufacturer's protocol. One microgram of total RNA was used as input RNA to make each library, each of which was sequenced on an Illumina Hi-SEq. 2000 using standard methods to generate 100-bp single-end reads at the Broad Stem Cell Research Center at the University of California, Los Angeles (UCLA). RNA sequencing data were processed as described previously (2, 53, 54). All RNA-seq data reported in this paper have been deposited in the Gene Expression Omnibus (GEO) database (accession No. GSE93195). Gene set enrichment analysis (GSEA) was performed using the GSEA software from the Broad Institute (software.broadinstitute.org/gsea/index.jsp) (29, 55). Statistical significance was measured with a permutation based Kolmogorov-Smirnoff nonparametric rank test (1000 permutations). For Weighted Gene Coexpression Network Analysis (WGCNA) (28), a gene coexpression network was constructed using the 2,000 transcripts that showed greatest variations upon microinjection of  $\beta$ -CatMO and their expression values throughout the 29 libraries sequenced at stage 10.5 (Dataset S1). To find modules of highly correlated genes, average linkage hierarchical clustering was performed, and modules were assigned by color codes.

**Heat Maps.** Heat maps were generated in Multiexperiment Viewer (MeV) (56). For Fig. 6, RPKMs were used as inputs, and rows/genes were normalized using the mean and SD of the row to which the value belongs, using the following formula: Value = [(Value) – Mean(Row)]/[SD(Row)]; hierarchical clustering using Manhattan distance was performed. The heat map in Fig. 5 used fold change as input and was similarly normalized.

Mean Log Ratios-Average Plot and PCA Analysis. Mean log ratios-average plots (MA-plots) were generated in R. For Fig. S1A, the  $\log_2$  fold change (FC) in expression of transcripts between control embryos corresponding to β-CatMO embryos was plotted versus the average RPKM expression of transcripts in control embryos. In Fig. 4A, the  $\log_2$  FC in expression of transcripts between Cerberus embryos to control embryos was plotted against the average RPKM levels of expression in control embryos. Transcripts with an average RPKM below 1 across all conditions were eliminated. Principal component analysis (PCA) in Fig. S4 was generated in R by comparing  $\log_2$  FC in all libraries for transcripts identified in our early β-catenin signature.

Scatter and Box Plots. The scatter plot matrix in Fig. 1H contains 40,157 transcripts from dorsal and ventral half RNA-seq libraries. The  $\log_2$  FC in average expression of transcripts between dorsal and ventral half-embryos (five experiments) was plotted versus the  $\log_2$  FC of control embryos compared with β-CatMO embryos (three experiments). Transcripts highlighted in red are those with fold changes higher than 1.5 and 1.4 in the dorsal/ventral (D/V) and con/β-CatMO conditions, respectively. Box plots in Fig. S1 and Fig. 3 were generated in R. The middle line in the box indicates the median, the box edges indicate the 25th/75th percentiles, and the whiskers indicate min and max values. The statistical significance of differences in gene expression levels between pairwise sets of genes was tested using the Mann–Whitney test and indicated as follows:  $\leq$ 0.05, \*\* $\leq$ 0.01, and \*\*\* $\leq$ 0.005.

Cloning and qRT-PCR. To clone full-length *Loc100170590*, forward and reverse PCR primers were designed according to the genomic sequence deposited in the Xenbase database (www.xenbase.org/entry/). The oligos also contained upstream sequences for Gateway-mediated cloning. PCR was performed on cDNA obtained from gastrula stage *X. laevis* embryos, resulting in an amplification product migrating at the expected size (about 1.5 kb). The PCR product was purified, cloned in a pDonr221 vector and subsequently in a home-made Gateway-compatible pCS2 vector suitable for antisense probe and in vitro mRNA synthesis. Primers for cloning were as follows: Fwd, GGGGACAAGTITGTACAAAAAAGCTGGGTCATGATGTTCCGATGGGCCCTGTC; Rev, GGGGACCACTITGTACAAGAAAGCTGGGTCATGAAGCTTATTCAACCCTTTTTGAC. Primers for qRT-PCR were as follows: Fwd, GCCCTTGAGTCTTCTTAT; Rev, CAGACATGAGAGAGAGTGG.

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