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Molecular Characterization of a cDNA Encoding Putative Vitellogenin from the Pacific Oyster *Crassostrea gigas*

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ABSTRACT—To elucidate the molecular mechanisms involved in oogenesis, we applied a differential display method to identify genes whose expression was detected only in ovaries containing oocytes. One of the cDNA fragments isolated by mRNA differential display was similar in structure to vitellogenin. Using this fragment, a full-length cDNA encoding putative vitellogenin in the Pacific oyster *Crassostrea gigas* was cloned by RACE (rapid amplification of cDNA ends), and its amino acid sequence was deduced. The open reading frame predicted 1583 amino acid residues. The deduced primary structure of putative vitellogenin in *C. gigas* was shown to be similar to vitellogenins of various other mollusk, fish, crustacean and nematode species, especially in the N-terminal region. Reverse transcription-mediated PCR revealed that mRNA encoding putative vitellogenin was expressed only in the ovary. *In situ* hybridization analysis revealed that the follicle cells are the site of putative vitellogenin synthesis.

Key word: follicle cells, mRNA differential display, oogenesis, oyster, vitellogenin

INTRODUCTION

In marine bivalves as in most oviparous animals, a large amount of yolk protein (vitellin) is accumulated in oocytes during ovarian maturation. In vertebrates such as fish, amphibians and birds, yolk protein is synthesized from a precursor, vitellogenin (Vg), produced by the liver, and is transported to the oocytes via the blood circulation system. The relationship between volk protein and Vg has been extensively studied in Xenopus laevis and in the chicken, Gallus gallus, and the amino acid sequences of these species have been determined (Gerber-Hunber et al., 1987; Van het Schip et al., 1987). Vgs have been shown to be present in almost all species of oviparous animals ranging from nematodes to vertebrates, and extensive sequence conservation is seen among these groups (Chen et al., 1997). Recently, complete cDNA sequences of Vg have been reported from several fish and crustacean species (LaFleur et al., 1995; Mouchel et al., 1996; Tsutsui et al., 2000; Okuno et al., 2002).

In bivalve molluscs, a few biochemical studies of yolk proteins have been carried out (Osada *et al.*, 1992; Suzuki

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et al., 1992). In the oyster, vitellin has been isolated and characterized (Suzuki *et al.*, 1992; Li *et al.*, 1998), but the characterization of Vg has not been reported, and information on bivalve vitellogenesis is still limited. To identify the genes associated with vitellogenesis, we cloned and sequenced more than 100 cDNA fragments using mRNA differential display, which is a method for studying differential gene expression from different sources (Liang and Pardee, 1992), and found that one of the isolated genes was similar to Vg. The objective of the present study was to clone the full-length of the Pacific oyster putative Vg cDNA on the basis of the cDNA fragment obtained by mRNA differential display, to deduce the complete primary amino acid sequence, and to examine the expression of its mRNA.

MATERIALS AND METHODS

Animals and isolation of RNA

Cultured Pacific oysters were collected monthly from March to September 1999 in Gokasho Bay, Mie Prefecture. The gonad was homogenized using ISOGEN (Nippon Gene, Toyama, Japan), and the total RNA was extracted according to the manufacturer's instructions (Chomzynski and Sacchi, 1987).

mRNA differential display

First strand cDNA was synthesized by annealing $pd(N)_6$ Primer per 5 µg of total RNA using a First Strand cDNA Synthesis kit

(Amersham Biosciences Corp., NJ), according to the manufacturer's protocol. The resulting cDNAs were amplified with seven different 10-nucleotide random primers (RA1, 3, 5, 7, 13, 14 and 16) selected from the list of Monna *et al.* (1994). The cycling parameters were as follows : one cycle of 92°C for 1 min, and 40 cycles of 92°C for 1 min, 45°C for 1 min, 72°C for 1 min and lastly one cycle of 72°C for 5 min. The amplified cDNA was then separated on an 8% polyacrylamide gel and stained with ethidium bromide. Complementary DNA fragments whose expression was detected only in the spawning season (July and August) were recovered directly by cutting out the gel slices, and reamplified using the same primer and PCR conditions except an annealing temperature at 55°C. Reamplified PCR fragments were subcloned and sequenced.

Oligonucleotides

The oligonucleotides used as PCR primers are listed here and shown in Fig. 2. Primers for β -actin were based on the nucleotide sequences of sea scallop and oyster (Patwary *et al.*, 1996; Cadoret *et al.*, 1999).

Primer 1. 5'- ACG GGT AAG TAC TGT TGG AC -3' Primer 2. 5'- GAC GAA TGT TCC GAC GTG GG -3' Primer 3. 5'- CCC GAT GTC AAT GGC TTA TGG -3' Primer 4. 5'- GCA GAT GGA AGG ATG TCC ATC AG -3' Primer 5. 5'- TTC ACA GTC ATG GAG CCC AGC AT -3' Primer 6. 5'- GAC CCG TCA GAA TTG TTG TCA GAC -3' Primer 7. 5'- GAC TTC GAA CAA GAG ATG -3' Primer 8. 5'- GAT ATC GAC ATC ACA TTT C -3'

Rapid amplification of cDNA ends (RACE)

5' RACE was performed using a 5' RACE System (Invitrogen Corp., Carlsbad, CA) according to the manufacturer's instructions (Frohman *et al.*, 1988). First-strand cDNA was synthesized in a 25 μ l solution containing 5 μ g of total RNA, 2.5 pM of primer 1, and 200 U of Superscript II. PCR with primer 2 and 5' RACE abridged anchor primer was initiated at 94°C for 2 min, followed by 30 cycles of 94°C for 0.5 min, 55°C for 0.5 min, 72°C for 2 min and lastly one cycle of 72°C for 7 min. For 3'RACE, first strand cDNA was synthesized by annealing Not I-d(T)₁₈Primer using a First strand cDNA Synthesis kit (Amersham Biosciences) as described above. PCR was conducted with primer 3 and adapter primer using the same conditions but 35 cycles of amplification. The PCR products were subcloned and sequenced. Since these fragments were too long to sequence, PCR walking was performed with the designed primers (primers 4, 5 and 6).

Nucleotide sequence analysis

The cDNA fragments amplified by PCR were subcloned into a pCR2.1 plasmid using a TA Cloning kit (Invitrogen). Both strands of the insert in the plasmid were sequenced on a model 377 DNA sequencer using a BigDye Terminator Cycle Sequencing kit (Applied Biosystems, Foster City, CA).

Reverse transcription-mediated PCR

Total RNA was prepared from ovary, gill, muscle, mantle tissues of female oysters collected in April, and testis and ovary of the oysters collected in March, May, July and September, and first strand cDNA was synthesized as described above. Twenty-five (Fig. 5A) and 20 (Fig. 5B) cycles of amplification for vitellogenin (primers 4 and 5) and β -actin (primers 7 and 8) using AmpliTaq Gold (Applied Biosystems) were carried out under the following conditions : denaturation at 94°C for 0.5 min, annealing at 54°C for 0.5 min, and extension at 72°C for 0.5 min. At completion of the PCR, fragments were separated on a 1.5% agarose gel and stained with ethidium bromide.

In situ hybridization

For in situ hybridization, the digoxigenin (DIG)-labeled sense

and antisense RNA probes were transcribed in vitro from 807 bp cDNA fragment of oyster putative Vg (region between primers 4 and 5) using a DIG RNA Labeling kit SP6/T7 (Roche Diagnostics, Mannheim, Germany). The ovarian fragments were fixed in 4% paraformaldehyde in 0.1 M phosphate buffer at pH 7.4 for 16 hr, embedded in paraffin, and cut into 5 µm serial sections. Deparaffinized sections were treated with proteinase K (10 µg/ml) in PBS at 37°C for 15 min, and then incubated at 42°C for 16 hr in a hybridization buffer (50% formamide, 2 x SSC, 1 µg/µl tRNA, 1 µg/µl Salmon sperm DNA, 1 µg/µl BSA, 10% Dextran sulfate) and 1 mg/ ml DIG-labeled probe. After hybridization, the sections were washed three times with 2 x SSC/50% formamide at 42°C for 1 hr and twice with 2 x SSC at room temperature for 15 min. The sections were incubated with the Fab fragment of an anti-DIG alkaline phosphatase-conjugated antibody (1:500 dilution, Roche Diagnostics), and visualized with nitroblue tetrazolium and 5-bromo-4chloro-3-indolyl phosphate (Roche Diagnostics).

RESULTS AND DISCUSSION

Isolation and characterization of cDNA encoding putative Vg

In order to identify the expressed genes in the ovary, transcripts derived from the ovaries during the spawning season and the spent gonads containing no oocytes were examined by mRNA differential display using seven different primers. During this procedure, we identified a cDNA of approximately 1.2 kb in length, which was detected in the ovary during the spawning season (Fig. 1). Sequence analysis of the cDNA fragment followed by a database search revealed a similarity to Vg (Fig. 2: C1184).

Subsequently the 5' and 3' ends of this cDNA were obtained by 5' and 3' RACE with the gene-specific primers, respectively (Fig. 2). The deduced amino acid sequence of

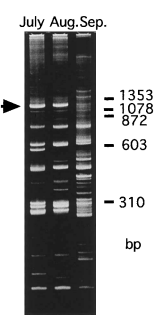


Fig. 1. Differential display of mRNAs from ovary (July and August) and spent gonad (September) of oyster. RNA was reverse transcribed with random primer. Resulting cDNA was amplified with an arbitrary primer (RA16, CCGACAGCTT). A differentially expressed fragment (arrow) was recovered, eluted from the gel slice, and reamplified. Numbers to the right represent molecular size markers.

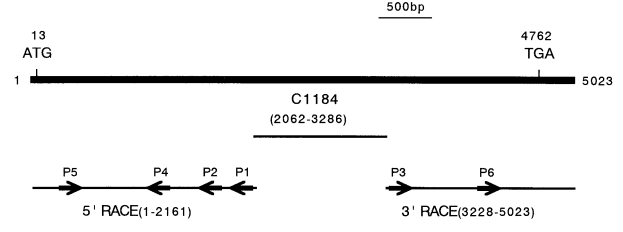


Fig. 2. Cloning strategy and schematic view of *C. gigas* putative vitellogenin cDNA. Location of the specific primers is shown corresponding to oligonucleotides listed in the Materials and Methods. The obtained cDNA is delineated, showing positions of the start codon (13 ATG) and stop codon (4762 TGA). C1184 indicates the PCR product obtained by mRNA differential display.

C. gigas putative Vg cDNA is shown in Fig. 3. The cDNA consisted of 5023 bp, comprising a 5' untranslated region (12 bp), and open reading frame (4749 bp), a stop codon (TGA), and a 3' untranslated region (259 bp). The 3' untranslated region contained a polyadenylation signal (AATAAA) (Fig. 3).

Characterization of the deduced amino acid sequence

The cDNA had an ORF (open reading frame) that encoded 1583 amino acid residues with a predicted molecular mass of 179,191 Da. The deduced amino acid sequence contained a consensus cleavage site, R-X-R-R (Arg⁸²⁸ to Arg⁸³¹), capable of undergoing processing by endoproteases of the subtilisin family (Barr, 1991) that have also been reported in Vgs of insects and crustaceans (Chen et al., 1994; Yano et al., 1994; Tsutsui et al., 2000; Okuno et al., 2002). If this processing site is cleaved by an endoprotease, the resultant two subunits would have calculated molecular masses of 94,000 Da and 85,209 Da. Previous studies have shown that in the Pacific oyster C. gigas, two major bands (179 and 110 kD) and several minor bands (Li et al., 1998), and seven protein bands with relative molecular masses of 105, 85, 66, 64, 60, 45 and 41 kD (Suzuki et al., 1992) were recognized in SDS-immunoblot analysis of the vitellin. The cDNA isolated in the present study may correspond to one or some of these proteins. The overall serine content of the oyster putative Vg (11.9%) was higher than in the mosquito (10.1%) and the silkworm Vg (9.8%), which possess polyserine clusters. The deduced amino acid sequence contained 56.2% of serine residues in the region between Ser¹¹⁴⁹ and Ser¹¹⁸⁰, and this region may be the polyserine domain in C. gigas putative Vg. This protein possessed six potential N-linked glycosylation sites, which is conserved in vertebrate and invertebrate Vgs.

The deduced amino acid sequence was compared to those of Vgs of the scallop *Patinopecten yessoensis* (partiallength, accession number AB055960), *Caenorhabditis elegans* (Spieth *et al.*, 1991), *Fundulus heteroclitus* (LaFleur *et al.*, 1995), rainbow trout (Mouchel *et al.*, 1996), and kuruma prawn (Tsutsui *et al.*, 2000). The BLAST algorithm indicated

homologies of 35% to scallop Vg (residues 24-519), 21% to C. elegans vit-5 (residues 24-954), 19% and 24% to Fundulus Vg1 (residues 22-285 and 454-953, respectively), 25% and 22% to rainbow trout Vg (residues 24-266 and 567–932, respectively) and 28% to kuruma prawn Vg (residues 559–940). As shown in Fig. 4, the region of *C. gigas* putative Vg which showed similarity to other Vg was limited to the N-terminal and the central region. The C-terminal region showed no significant homology to any of the other reported species. Thus this region appears to possess unique properties compared with Vg of the other known species. Best conserved amino acid motif between invertebrate and vertebrate Vgs (Mouchel et al., 1996) is found in the sequence of C. gigas putative Vg (KTIGNAG: positions 603-609). This sequence is speculated to play a highly conserved role in vitellogenesis, such as specific recognition by oocytes (Spieth et al., 1991).

Tissue distribution of oyster putative Vg mRNA

The levels of putative Vg mRNA in various tissues from female oyster and stage-specific expression were measured by reverse transcription-mediated PCR. In April, the oyster contained fully grown oocytes and putative Vg mRNA expression was detected only in the ovary (Fig. 5A). As Fig. 5B shows, putative Vg mRNA expression was detected only in the ovary, and indicated maximum level in March (the early stage of maturation). To determine the distribution of oyster putative Vg mRNA expression in ovary, we performed *in situ* hybridization using DIG-labeled sense and antisense RNA probes, respectively. A strong signal was detected in the follicle cells using an antisense RNA probe (Fig. 6A). In contrast, hybridization with a sense control probe produced no significant signal (Fig. 6B).

The synthesis, secretion and processing of Vgs differ among phyla. Vgs are synthesized by extraovarian tissues such as the liver in vertebrates and the fat body in insects (Sappington and Raikhel, 1998), secreted into the circulatory system, and transported into the ovary. In crustaceans, three tissues have been reported as sites of Vg synthesis:

1	1 TTACACGGCAGAATGTTGCTTCTGCTTGCGGTCTCAACAGCCCTGGTGGCGGCAGCCGCCCAGACTCCCAGTCAAGCCTATGAAAGCAAT														Г										
		М	L	L	L	LA	v	S	Т	А	ΓV	A	А	А	А	Q	т	Ρ	s	Q	A	Y	E	S N	26
	KEYLYEYETQA	LTGI	PMG	STI	YSG	IKMKS	DVR	IQFF	RSRS	SAT	LKMDF	LTF7	YKIN	DPII	ESV	DPT	QQQV	PAE	MFG	2PL1	「GRD	AE	QML'	TDLSR	P 116
	INFRYVRGNVK	DIHH	EADI	DPE	wsvi	IVKKG	LLSN	1L EM	INLE	KRK	ELRKS	GVII	PQVL	RPQS	SDI	EGSN	1FTV	MEP	SIV	/GEC	ETL	YRI	ISPV	TSTS	206
	NPWMHITKVRN	YQHC	LDRI	PKY	FGSI	1FHLR	QCAI	ECVF	REQS	EPL	RSASÇ	IRYI	LRG	NMR(QFQ	IQSZ	AIAE	SQH	VF	PYS	AKG	GH	VAT	LNQT	296
	NLVKEEDVKTS	LSEP	QS P	TKV	KSG	LQYTS	RELI	EMRI	SSS	ILQ	QSSDS	LSTI	KSS	PESI	PQH	SSS	SSNH	HPS	PQI	PPSE	PQPK	TT	SNQ	ADRI	R 386
	KLLKMLEVFMK	PTIQ	PEA	GPL	LMS	LEEI	RTAI	OSES	SLRA	VYR	ECSNG	ETA/	ΔTVQ	DADO	SHP	SICO	FTTS	ATE	VLV	/DSI	LND	QLI	QSE	avva	476
	GILSLSARPDV	LIAK:	KLLI	DLT	RSS	QVSKD	RYLH	(RAA	AFLC	LGS	VAGMI	REEG	WRR	SREI	ISR	QEEI	IVKT	LIS	QES	SRSÇ)TRN	ELF	KSKI	RELE	566
	LKKREHS IKTK	IKQE	IVKI	ELQI	MLMI	RSTVF	DDKI	LSF	FKTI	GNA	GLWEM	IPTI	RTY	IQDI	KSQI	PQVI	LRTQ	AIY	SLF	RKLA	RHY	PDI	DIQI	TLLP	656
	YFDQSEKEEVR	IGSY	LVM	r FTI	EPSI	QLLE	MVAÇ	QSLH	IRER	NPH	VGTFV	YTHI	EQM	зизт	TYP	CLMS	WAK	NAS	FAM	IRFA	KKF	SPN	IYHY	SRFMI	1746
	LSGFNEMHKMG	AAAE:	LGLV	VTTI	PEEB	TIPRA	GAVN	ILHI	YVF	GRS	VNFAE	IGFN	ITEG	LQTI	JVSI	KLVG	PLG	ELTI	KGK	SLV	DVL	ĸQ	RVRF	SAES	836
	QPSDPISQIHK	QLKV	SPRI	rsp:	грко	HMYM	KMMG	SNEI	JQYI	TLD	GTLVD	TLLF	EGK	LLSC	svsi	EQDI	LKTG	LNV	EVH	IRST	MPL	EAE	IMI	PSEC	, 926
	LPLRLKLHGTA	AIKV	IGK\	VGV!	FGMI	SIFE	INRE	?GKÇ	AKE	LSF	NFELR	PSVI	FQL	RGEM	1EMI	DAES	FKM	GVA	LKI	MAH	MET	PLS	SLTA	SANLI	2 1016
	KAKFYTKFNIE	KLSEI	KIA	RLEV	VSPY	TYFK	EDPS	SEIT	KYP	LPR	ETQEI	SVAK	NAK	VFPN	ISM	AYGI	QTL	GLE	IKI	SGQ	AVV	RDE	EMI	VPYYI	9 1106
	MIGKQEMIVTL	SPGTI	DPQI	KYVI	EIQI	QLMK	RMSÇ	QSKÇ	QPT	SEE	SSSST	GLLS	wvs	SLIG	SDÇ	QSDÇ	QSSP	SSS	sss	STS	PPK	DIF	RSKI	TSLKI	1196
	LLEHLKESQSVI	HPDG	svvi	FRM	NSMO	LIFN	VIGI	DQS	RSI	KRH	FHISM	AAGM	NPS	PKTI	SII	LLRM	<u>in</u> rs	PIP	TME	TKP	WNM	DVA	VNV	NLPSH	1286
	LADPSELLSDA	YQRE:	LEQÇ	QIE	VYVI	RHGE	SDYF	IRWI	MLE	EPR	WESSV	LQRI	LSL	SDTÇ	2EVI	I DE \	/MNS	ELSI	KSE	ESG	TPE	DRN	III	LIRK	0 1376
	KQILKEARKIW	QRSYI	DSL	RVQI	DIKA	KLSS	LIKF	RSES	VRK	DLE	KDSAS	VSKS	EKS	DALF	RKY	CALI	LQQL	IDQI	RLD	EAV	RSSI	RRS	LSS	KERQI	1466
	IAKSLESSKES:	LRKI	LQQÇ	2HR.	SPSI	VDSS	інні	LALF	rksi	LHR	SVKVI	TKKI	KNL	RTQN	IKKI	INPS	NNN.	SLLI	NSP	RTP	CLAI	INL	PKC	LVVRI	1556
4.604																									
4681	TGTCAAAGTCA	GTGT	CAG/ O	ACC!	FTT	GCAA C K	AAAC N	R R	AAG K	ATG'	rggtt w f		ATA' T	TTTC F	CAC P	GGTC					.GGG2 R I		'GAA *	ATCTO	; 1583
4771	TTGATGAGAAA		~	-	-								-	-	_	-	~						TCC	TCAAZ	
4861	ATATCGAGAGG																								-

Fig. 3. Deduced amino acid sequence and 5' and 3' untranslated regions of putative Vg cDNA. Positions of nucleotides and amino acids are indicated by numbers on the left and right sides of the figure, respectively. Putative *N*-glycosylation sites are underlined and consensus cleavage sequence for processing by endoproteases of the subtilisin family is framed. The polyadenylation signal is shown in white lettering with black background. The position of DIG-labeled RNA probe used for *in situ* hybridization is indicated by a dotted underline. The nucleotide sequence has been submitted to DDBJ/EMBL/GenBank with the accession number AB084783.

the ovary (Lee and Watson, 1995; Tsutsui *et al.*, 2000), adipose tissue (Okuno *et al.*, 2000) and hepatopancreas (Soroka *et al.*, 2000; Tsutsui *et al.*, 2000; Okuno *et al.*, 2002). Autosynthetic yolk formation is thought to be the main type of vitellogenesis in bivalves and gastropods (Jong-Brink *et al.*, 1983). In marine bivalves, autosynthesis of yolk proteins in the ovary has been postulated, based on the morphological evidence (Pipe, 1987; Dorange and Le Pennec, 1989; Eckelbarger and Davis, 1996). In oyster and other bivalve species, the ovarian acinus is a simple structure containing only developing oocytes and associated follicle cells within a thin germinal epithelium. The function(s) of follicle cells in the bivalve ovary are not well understood, though they are suspected of playing some role in oocyte nutrition. In the present study putative Vg mRNA expression

was in the follicle cells in the ovary of *C. gigas*. This suggests that yolk synthesis in *C. gigas* may take place in the follicle cells.

5023

In teleosts, as in other oviparous vertebrates, it is clearly established that the Vg gene expression is regulated by estradiol- 17β (E₂), produced by the maturing female ovarian follicles, in the liver (Vaillant *et al.*, 1988; Pakdel *et al.*, 1991; Okumura *et al.*, 2002). In the Pacific oyster, E₂ has been detected in the ovary, and its content shows a synchronous profile with gonadal maturity and decreased in the spawning season (Matsumoto *et al.*, 1997). This is correspondent to the change of putative Vg mRNA levels from early vitellogenic stage to spawning stage, suggesting that E₂ is at least one of the factors which promotes vitellogenesis in ovsters. However, further investigation is required in

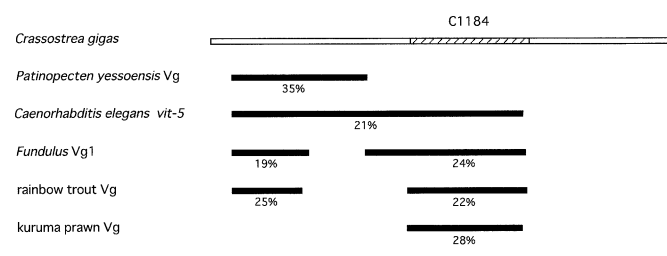


Fig. 4. Alignment of *C. gigas* putative Vg with other animal Vgs: scallop *Patinopecten yessoensis* (partial-length, accession number AB055960); *Caenorhabditis elegans* (Spieth *et al.*, 1991); *Fundulus heteroclitus* (LaFleur *et al.*, 1995); rainbow trout (Mouchel *et al.*, 1996); and kuruma prawn (Tsutsui *et al.*, 2000). The open bar represents the deduced amino acid sequence of *C. gigas* putative Vg. The solid bars indicate the regions of Vg which have homology to those of other animals. Percentage of homology is also indicated.

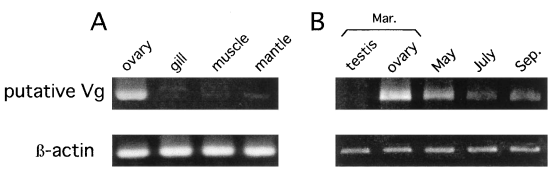


Fig. 5. Distribution of putative Vg mRNA in tissues of oyster. RNA was prepared from ovary, gill, muscle, mantle of female oysters collected in April (A), testis and ovary in March, and ovary in May, July and September (B). β-actin was used as a positive control.

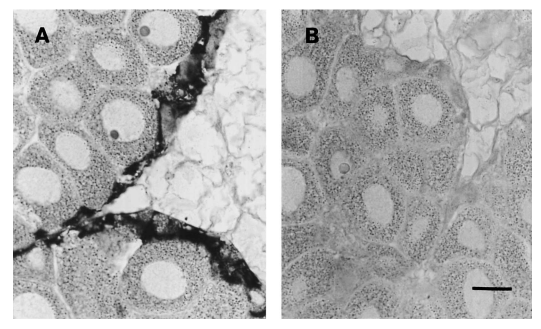


Fig. 6. Localization of putative Vg mRNA in the ovary by *in situ* hybridization. Strong expression of putative Vg was seen in the follicle cells with an antisense probe (A) in the ovary. Hybridization with a sense probe (B) produced no significant signal in the ovary. Bar=25 µm.

order to clarify the estrogen regulatory mechanisms in the oyster.

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