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# XOL-1, primary determinant of sexual fate in *C. elegans*, is a GHMP kinase family member and a structural prototype for a class of developmental regulators

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In *Caenorhabditis elegans*, an X chromosome-counting mechanism specifies sexual fate. Specific genes termed X-signal elements, which are present on the X chromosome, act in a concerted dose-dependent fashion to regulate levels of the developmental switch gene *xol-1*. In turn, *xol-1* levels determine sexual fate and the activation state of the dosage compensation mechanism. The crystal structure of the XOL-1 protein at 1.55 Å resolution unexpectedly reveals that *xol-1* encodes a GHMP kinase family member, despite sequence identity of 10% or less. Because GHMP kinases, thus far, have only been characterized as small molecule kinases involved in metabolic pathways, for example, amino acid and cholesterol synthesis, XOL-1 is the first member that controls nonmetabolic processes. Biochemical investigations demonstrated that XOL-1 does not bind ATP under standard conditions, suggesting that XOL-1 acts by a mechanism distinct from that of other GHMP kinases. In addition, we have cloned a XOL-1 ortholog from *Caenorhabditis briggsae*, a related nematode that diverged from *C. elegans* ~50–100 million years ago. These findings demonstrate an unanticipated role for GHMP kinase family members as mediators of sexual differentiation and dosage compensation and, possibly, other aspects of differentiation and development.

[Keywords: *C. elegans*; XOL-1; sexual differentiation; GHMP kinase; crystal structure]

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Sex determination is the critical and universal developmental pathway underlying sexual reproduction. Its manifestations are pervasive and often conspicuous. Whereas the presence or absence of the Y chromosome dictates male or female development in mammals, sexual fate in the fruit fly *Drosophila melanogaster* and the free-living nematode *Caenorhabditis elegans* is determined genetically by the number of X chromosomes relative to the number of sets of autosomes. In mammals, the primary sex determining gene is *SRY*, which is present only on the Y chromosome and encodes an HMG domain-containing transcription factor. In the fruit fly, the primary sex determination gene *Sex-lethal* (*Sxl*; Maine et al. 1985) is a female-specific *trans*-acting gene

regulator that binds *tra* transcripts and directs alternative splicing (Inoue et al. 1990). The *SRY* (Werner et al. 1995) and *SXL* (Handa et al. 1999) interactions with polynucleotides have been characterized structurally. In *C. elegans*, sexual differentiation is regulated by the expression levels of the developmental switch gene *xol-1*. High and low levels of *xol-1* result in male (XO) and hermaphrodite (XX) development (Fig. 1), respectively. XOL-1 activity is absolutely required for proper sexual differentiation and male viability (Rhind et al. 1995), but its mechanism of action is unknown.

The cooperative activity of at least four X-linked genes, termed X-signal elements, represses expression of *xol-1* (for review, see Meyer 2000a). By doubling the number of X-signal elements, an XX embryo reduces *xol-1* expression by ~10-fold (Rhind et al. 1995), facilitating hermaphrodite development. Two *C. elegans* X-signal elements have been characterized molecularly as follows: *FOX-1* (Hodgkin et al. 1994; Nicoll et al. 1997; Skipper et al. 1999), an RNA-binding protein that may regulate alternate splicing of *xol-1* RNA, and *SEX-1* (Carmi et al. 1998), a nuclear receptor and likely a tran-

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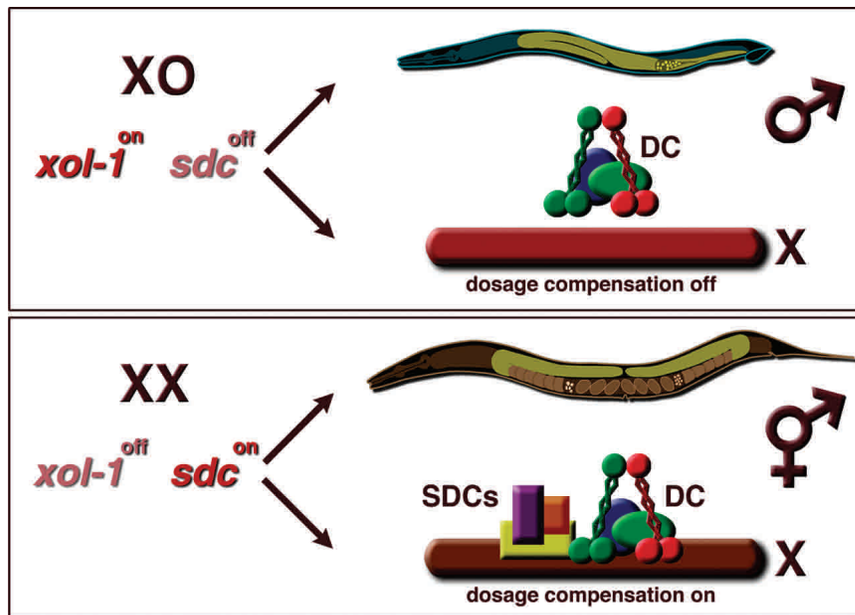
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**Figure 1.** Genetic control of sex determination and dosage compensation in *C. elegans*. *xol-1* is the primary sex-determination switch gene and the direct molecular target of the X-chromosome counting mechanism. (Top) Male (XO). High *xol-1* activity specifies male fate by repressing the activity of *sdc-1*, *sdc-2*, and *sdc-3* genes. Overt male sexual characteristics include a one-armed gonad (yellow) and a highly specialized tail used for mating. The dosage compensation machinery (colored ovals) does not bind to the X chromosome. Thus, X chromosome gene expression is unregulated (red chromosome). DC, dosage compensation complex (i.e., DPY-26, DPY-27, DPY-28, and MIX-1). (Bottom) Hermaphrodite (XX). Low *xol-1* activity specifies hermaphrodite fate by allowing the activation of *sdc-1*, *sdc-2*, and *sdc-3* genes. Overt hermaphrodite sexual characteristics include a two-armed gonad (yellow, with brown oocytes and embryos) and a tapered tail. The *sdc-1*, *sdc-2*, and *sdc-3* gene products (colored rectangles) recruit a large complex of dosage compensation proteins (DC, colored ovals) to both X chromosomes, repressing X gene expression by half (brown chromosome).

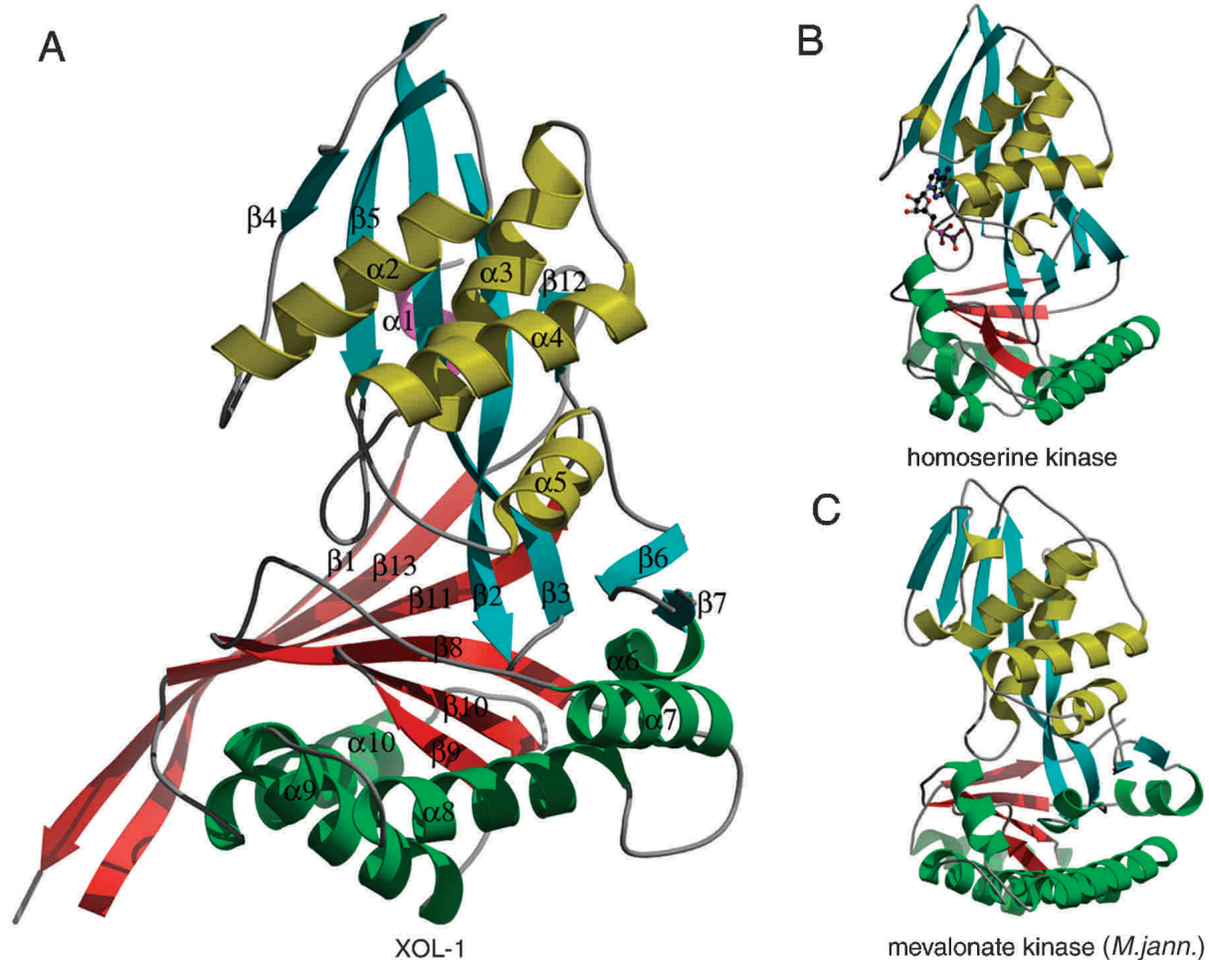


scription factor. Although hermaphrodites and males differ in X chromosome number, the expression of most X-linked genes must be equal to ensure viability. This is accomplished through dosage compensation, which reduces expression of X-linked genes in hermaphrodites to male levels (for reviews, see Wood et al. 1997; Hansen and Pilgrim 1999; Meyer 2000b; Boag et al. 2001). High levels of XOL-1 in males correlate with low SDC-2 expression, preventing dosage compensation (Miller et al. 1988; Rhind et al. 1995). Conversely, low levels of XOL-1 in hermaphrodites correlate with high SDC-2 expression and the assembly on the X chromosome of the dosage compensation complex, which is composed of *sdc*, *dpy*, and *mix-1* gene products (Nonet and Meyer 1991; Chuang et al. 1996; Lieb et al. 1996, 1998; Davis and Meyer 1997; Dawes et al. 1999; Chu et al. 2002). Other genes downstream of *xol-1*, such as *her*, *tra*, and *fem*, whose activities are inversely related in hermaphrodites and males, coordinate sexual differentiation (Goodwin and Ellis 2002). Null mutants of *xol-1* are XO-lethal, inappropriately activating dosage compensation where only one X chromosome is present, whereas XOL-1 overexpression is XX-lethal, deactivating the dosage compensation pathway and elevating the expression of X chromosome genes to lethal levels in hermaphrodites (Rhind et al. 1995).

XOL-1 is an acidic 51-kD nuclear protein (pKa 4.6), whose transcript is expressed at high levels only in pre-comma stage XO embryos (Rhind et al. 1995). XOL-1 transcripts are present in low levels throughout other larval stages in XO animals, but are nearly undetectable in XX larvae and adults of both sexes (Rhind et al. 1995).

Currently, XOL-1 is annotated as a subtilisin-like protease on the basis of primary sequence (<http://www.wormbase.org>). BLAST searches of Genbank failed to identify any homologs that may have provided additional clues as to the function of XOL-1. Thus, we used x-ray crystallography to investigate the function of XOL-1, hypothesizing that the three-dimensional structure of the protein would yield insights into its nature, largely uncharacterized biochemically. The resulting crystal structure of XOL-1 (Fig. 2A) unambiguously and unexpectedly defines XOL-1 as a member of the GHMP kinase family, a family of proteins known to be involved in small molecule metabolism, but not known to participate directly in sexual differentiation or dosage compensation.

Galactokinase, homoserine kinase, mevalonate kinase, and phosphomevalonate kinase were originally identified as prototypic members of the GHMP kinase family and observed to contain a conserved Pro-Xaa<sub>3</sub>-Gly-Leu-Gly-Ser-Ser-Ala-Ala motif (Fig. 3A) that was hypothesized (Tsay and Robinson 1991; Bork et al. 1993), and later proved (Zhou et al. 2000; Krishna et al. 2001; Fu et al. 2002) to be involved in ATP binding. The nucleotide fold of GHMP kinases is distinct from those of other kinases, that is, P-loop, protein, and Hsp70-like kinases (Zhou et al. 2000; Bonanno et al. 2001; Fu et al. 2002; Romanowski et al. 2002; Yang et al. 2002), and binds ATP in both *syn* and *anti*-conformations (Zhou et al. 2000; Fu et al. 2002). GHMP kinases are found in bacteria, archaea, and eukaryotes and contain an unusual left-handed  $\beta$ - $\alpha$ - $\beta$  fold similar to that observed in domain IV of elongation factor G (Zhou et al. 2000). In humans, deficiency of galactokinase, which participates in the



**Figure 2.** Comparison of XOL-1 and GHMP kinase structures. (A) The structure of XOL-1. Ribbon diagram of XOL-1 (PDB ID: 1MG7). Domain 1 consists of  $\beta$ -strands 2–7 and 12 (cyan) and  $\alpha$ -helices 1–5 (yellow). Domain 2 consists of  $\beta$ -strands 8–11 (red) and  $\alpha$ -helices 6–10 (green). Additional strands  $\beta 1$  and  $\beta 13$  and helix  $\alpha 1$  form novel elements not found in known GHMP kinase structures. (B) The structure of homoserine kinase bound to ADP (PDB ID: 1FWK). Domain 1  $\beta$ -strands and  $\alpha$ -helices are cyan and yellow, respectively. Domain 2  $\beta$ -strands and  $\alpha$ -helices are red and green, respectively. ADP is represented by a ball and stick model. (C) The structure of mevalonate kinase (*M. jannaschii*; PDB ID: 1KKH). Colors are as in B.

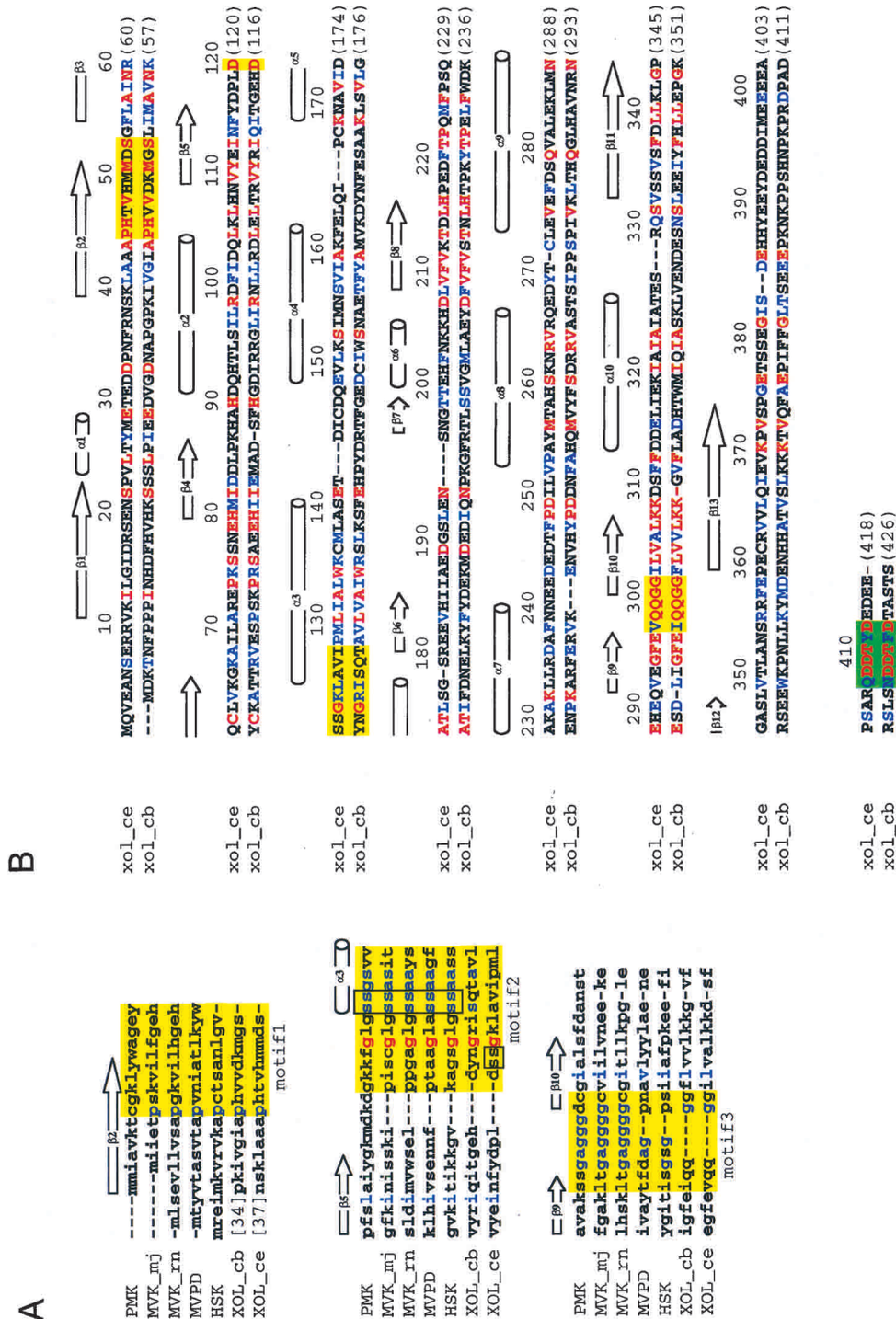
conversion of galactose to glucose, contributes to cataract formation (Monteleone et al. 1971; Beutler 1972; Harley et al. 1972; Levy et al. 1972). Mutations in mevalonate kinase, an enzyme involved in the synthesis of sterols from acetate, are associated with mevalonic aciduria (Hoffmann et al. 1986; Schafer et al. 1992; Houten et al. 1999b) and hyperimmunoglobulinemia D/periodic fever syndrome (Drenth et al. 1999; Houten et al. 1999a, 2001; Cuisset et al. 2001; Rios et al. 2001; Simon et al. 2001). To date, no GHMP kinases have been shown to function in developmental pathways unrelated to metabolism.

## Results

### *XOL-1 is a GHMP kinase family member*

Full-length XOL-1 (417 residues), as well as various N- and C-terminal truncation mutants, were overexpressed

and purified from *Escherichia coli*. N- and C-terminal truncations were designed on the basis of biological data (Rhind et al. 1995) and secondary structure predictions. A C-terminal truncation mutant (residues 1–374), which retained full rescue activity of *xol-1* null males (Rhind et al. 1995), crystallized in the orthorhombic space group  $P2_12_12$  with unit cell dimensions  $a = 116.1 \text{ \AA}$ ,  $b = 86.0 \text{ \AA}$ , and  $c = 80.2 \text{ \AA}$ , with two molecules per asymmetric unit ( $V_m = 2.4 \text{ \AA}^3/\text{Da}$ ). Initial phases and the positions of selenium atoms of a selenomethionine-derivatized XOL-1 (see Materials and Methods) were determined by MAD at  $2.5 \text{ \AA}$ . The structure was further refined by phase extension using native data to  $1.55 \text{ \AA}$ . The final R values for the refined model are  $R_{\text{cryst}} = 20.1$  and  $R_{\text{free}} = 21.2$  (Table 1). The final model contains two XOL-1 molecules per asymmetric unit consisting of 703 amino acid residues and 628 water molecules. Superimposition of monomer 2 (chain B) onto monomer 1 (chain A) yielded an r.m.s. deviation of  $0.17 \text{ \AA}$  for all equivalent backbone atoms,



**Figure 3.** Sequence alignment of XOL-1. (A) Structure-based sequence alignment of XOL-1 within GHMP kinase conserved motif regions. Sequence of XOL-1 (bottom) is aligned with structurally equivalent residues from other GHMP kinases. The three GHMP conserved motifs are highlighted in yellow. In XOL-1, these residues are as follows: motif1, residues P45-S53; motif2, residues D120-P128; and motif3, V298-G302. Tandem series of motif2 are highlighted by boxes. The sequence of XOL-1 is highly divergent, even within conserved motifs, from other GHMP kinases with only one structurally equivalent residue found to be identical (Gly 123 in motif2) throughout the entire sequence. Residues conserved identically in all structures are in red. Highly conserved residues are in blue. Secondary structure is assigned above. PMK, phosphomevalonate kinase; MVK\_mj, mevalonate kinase (*M. jannaschii*); MVK\_rn, mevalonate kinase (*R. norvegicus*); MVPD, mevalonate-5-phosphate decarboxylase; HSK, homoserine kinase; XOL\_c, XOL-1 (*C. briggsae*); XOL\_ce, XOL-1 (*C. elegans*). (B) Alignment of *C. elegans* and *C. briggsae* XOL-1 sequences. The *C. elegans* and *C. briggsae* XOL-1 sequences were aligned by the clustal method (GONNET series). Gaps and insertions incompatible with the structure were edited manually. Residues corresponding to GHMP conserved motifs are highlighted in yellow. A conserved acidic C-terminal motif is highlighted in green. Identically conserved residues are highlighted in red and conservatively substituted residues are in blue. Secondary structure is assigned above.

**Table 1.** Crystallographic data and refinement

	Data collection				
	Native	Se_peak	Se_inflection	Se_remotel	Se_remote2
Space group	P2 <sub>1</sub> 2 <sub>1</sub> 2				
Wavelength (Å)	0.98	0.9796	0.9794	0.9611	0.8947
Resolution (Å)	50.00–1.55	50.00–2.30	50.00–2.00	50.00–2.40	50.00–1.81
Completeness (%)	97.3 (96.9) <sup>a</sup>	99.2 (99.8)	99.4 (99.6)	83.5 (86.4)	97.3 (99.1)
Unique reflections	113,135 (12,340)	67,634 (6,783)	103,251 (10,355)	51,374 (5,279)	151,074 (15,262)
R <sub>merge</sub> (%) <sup>b</sup>	5.4 (54.0)	8.9 (41.1)	7.6 (40.6)	7.2 (35.1)	8.1 (56.6)
I/σ	25.2 (2.4)	14.4 (2.4)	18.8 (3.1)	14.3 (2.4)	21.7 (2.0)
Overall figure of Merit	56.0				

Refinement	
Resolution (Å)	50–1.55
R <sub>cryst</sub> (%) <sup>c</sup>	20.3
R <sub>free</sub> (%) <sup>d</sup>	21.3
Rmsd bonds (Å)	0.006
Rmsd angles (°)	1.40
Average B (Å <sup>2</sup> )	
protein	20.4
water	30.2
Ramachandran plot (%)	
most favored	92.3
additionally favored	7.7
generously allowed	0.0
disallowed	0.0

<sup>a</sup>Numbers in parenthesis refer to the highest resolution shell.

<sup>b</sup>R<sub>merge</sub> =  $[\sum_h \sum_i |I_i(h) - \langle I(h) \rangle| / \sum_h \sum_i I_i(h)] \times 100$ , where  $\langle I(h) \rangle$  is the mean of the I(h) observation of reflection h.

<sup>c</sup>R<sub>cryst</sub> =  $\sum_{hkl} |F_o - F_c| / \sum_{hkl} |F_o|$

<sup>d</sup>R<sub>free</sub> was calculated as for R<sub>cryst</sub>, but on 5% of the data, excluded from the refinement.

indicating no significant differences between the two molecules in the asymmetric unit. The following regions were omitted from the final model for lack of interpretable density: residues 1–7, 29–38, and 190–195 in monomer 1 and residues 1–7, 29–38, and 190–194 in monomer 2.

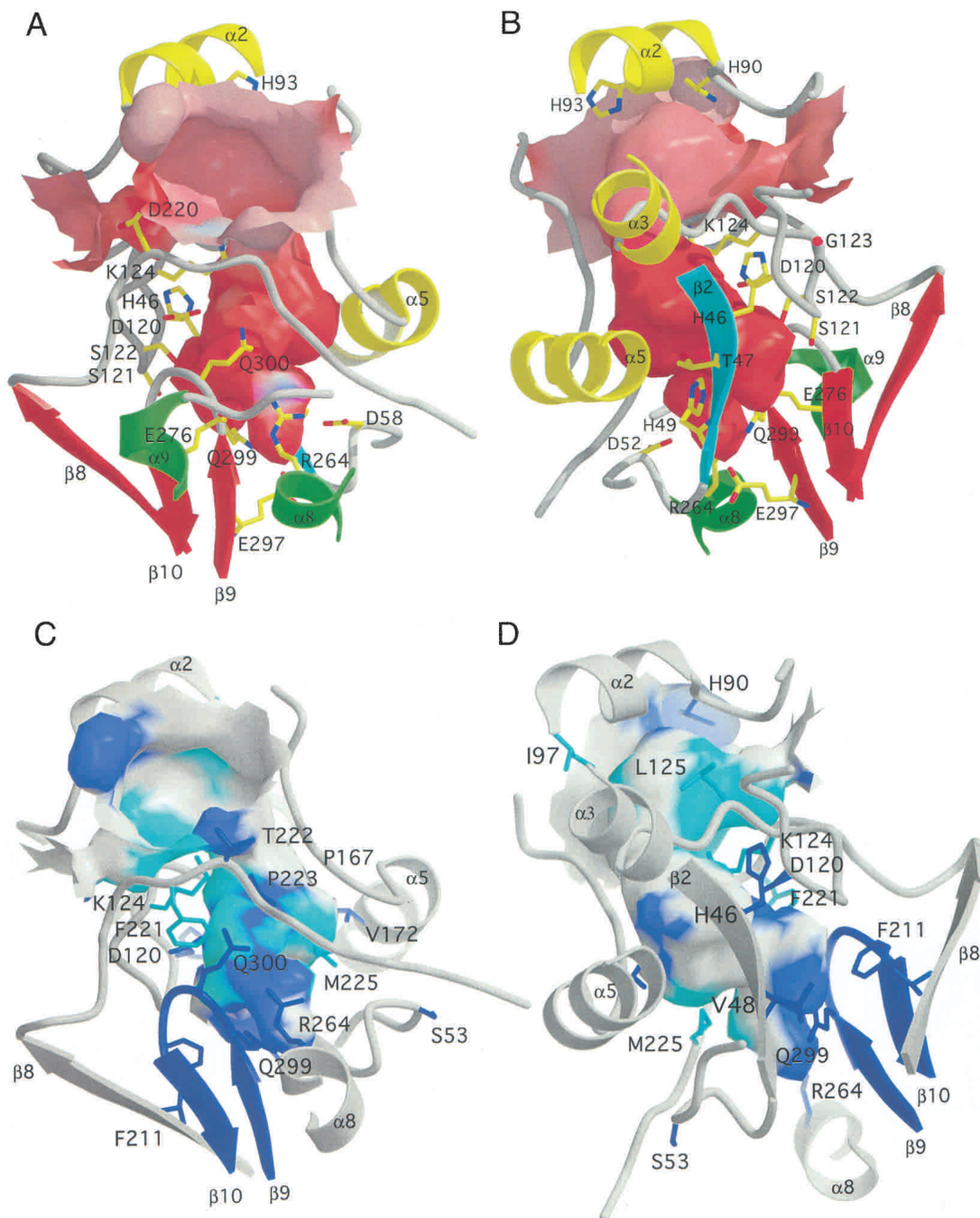
Submission of the XOL-1 atomic coordinates to the Dali server (Holm and Sander 1993; <http://www.ebi.ac.uk/dali>) returned surprisingly high Z-scores of 18.5, 17.1, 15.8, 15.2, and 15.1 for five GHMP kinases; mevalonate kinase (*Rattus norvegicus*; PDB ID:1KVK; Fu et al. 2002), mevalonate kinase (*Methanococcus jannaschii*; PDB ID:1KKH; Yang et al. 2002; Fig. 2B), phosphomevalonate kinase (*Streptococcus pneumoniae*; PDB ID:1K47; Romanowski et al. 2002), homoserine kinase (PDB ID:1FWK; Zhou et al. 2000; Fig. 2C), and mevalonate-5-phosphate decarboxylase (PDB ID:1FI4; Bonanno et al. 2001) with r.m.s. deviations for superimposed C<sub>α</sub> atoms

of 3.8, 3.0, 3.1, 3.5, and 4.0 Å, respectively (Table 2). Despite indisputable structural similarity, XOL-1 displays remarkably little sequence similarity to other members of the family, with overall sequence identities returned from the Dali server of 9%–10%. Even within the GHMP kinase conserved motif1, motif2, and motif3 (Fig. 3A), the sequence is highly divergent. In fact, throughout the entire sequence, only one structurally equivalent residue (Gly 123 in motif2) is absolutely conserved [Figs. 3A (in red), 4B] among XOL-1 and the five GHMP kinase structures deposited in the PDB. Although, across subfamilies of GHMP kinases, sequence identity can be surprisingly low (Burley and Bonanno 2002), sequence identity is sufficiently high within subfamilies to convincingly delineate related enzymes. Thus, XOL-1 is the first GHMP kinase family member to be identified solely from structural information. In addition, the role of XOL-1 in sexual differentiation was

**Table 2.** Statistics from Dali Server

GHMP Kinase	PDB ID	Z score	R.M.S.D.	% identity	No. equivalent residues
Mevalonate kinase ( <i>R. norvegicus</i> )	1KVK	18.5	3.8	9	269
Mevalonate kinase ( <i>M. jannaschii</i> )	1KKH	17.1	3.0	9	242
Phosphomevalonate kinase	1K47	15.8	3.1	9	243
Homoserine kinase	1FWK	15.2	3.5	10	251
Mevalonate 5-phosphate decarboxylase	1FI4	15.1	4.0	9	257

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**Figure 4.** The putative active-site of XOL-1. (A) The cavity that represents the putative active-site of XOL-1 and, in other GHMP kinases, binds ATP is represented as an electrostatic surface nestled between domains 1 and 2. Red is electronegative and blue is electropositive (calculated in GRASP; Nicholls et al. 1991;  $-15$  to  $+15$  kT/e). Secondary structure elements are colored as in Figure 2A. Side chains of charged and polar residues lining the cavity are represented by ball and stick. (B) Same as A, but with view rotated by  $\sim 180^\circ$ . The location of the C $\alpha$  carbon of Gly 123 is represented by a red dot. (C) The cavity that represents the putative active site of XOL-1 is represented as a surface colored by conservation. Side chains of residues identical between *C. elegans* and *C. briggsae* XOL-1 are in dark blue, whereas those conservatively substituted are in cyan with the corresponding colors mapped onto the surface. The strongly conserved region encompassing motif3 ( $\beta 9$ – $\beta 10$ ) is colored dark blue. (D) Same as C, but with view rotated by  $\sim 180^\circ$ .

completely incongruous with known GHMP kinase functions.

The overall secondary structure of XOL-1 is mixed,

containing 10  $\alpha$ -helices and 13  $\beta$ -strands, which fold into an elongated dumbbell-shape (Fig. 2A). As in other GHMP kinases (Zhou et al. 2000; Bonanno et al. 2001; Fu

et al. 2002; Yang et al. 2002; Fig. 2B,C), the GHMP kinase core of XOL-1, which excludes  $\beta$ -strands 1 and 13 formed by the extended N and C termini (Fig. 2A), consists of two domains, each composed of four associated  $\alpha$ -helices, associated  $\beta$ -strands, and a single, short, isolated  $\alpha$ -helical segment. Domain 1 contains a bundle of four  $\alpha$ -helices ( $\alpha 2$ ,  $\alpha 3$ ,  $\alpha 4$ ,  $\alpha 5$ ; Fig. 2A, in yellow), a  $\beta$ -sheet ( $\beta 12$ ,  $\beta 2$ ,  $\beta 3$ ,  $\beta 5$ ,  $\beta 4$ ) with two associated strands ( $\beta 6$ ,  $\beta 7$ ; Fig. 2A, in cyan), and a short, isolated  $\alpha$ -helix ( $\alpha 1$ ; Fig. 2A, in magenta). Domain 2 contains four  $\alpha$ -helices packed side-by-side ( $\alpha 7$ ,  $\alpha 8$ ,  $\alpha 9$ ,  $\alpha 10$ ; Fig. 2A, in green) on one side of the  $\beta$ -sheet ( $\beta 9$ ,  $\beta 10$ ,  $\beta 8$ ,  $\beta 11$ ; Fig. 2A, in red), and a short  $\alpha$ -helix ( $\alpha 6$ ; Fig. 2A, in green), which connects the  $\beta$ -sheets from domains 1 and 2.

#### Identification of a XOL-1 ortholog in *C. briggsae*

*C. briggsae* diverged from *C. elegans* ~50–100 million years ago. The recent, but not yet complete, sequencing of the *C. briggsae* genome (The Sanger Institute, <http://www.sanger.ac.uk>; The Genome Sequencing Center, <http://genome.wustl.edu>) allowed us to predict an apparent ortholog of *xol-1*. The existence of the expressed gene was confirmed by isolating and sequencing corresponding cDNA fragments from *C. briggsae*. Although the overall amino acid sequence identity is low (22% identical, 41% conservatively substituted; Fig. 3B), significant homology is found in key locations. Within strands  $\beta 9$  and  $\beta 10$  and the intervening loop that defines motif3, the sequences are 68% identical and the unusual QQGG sequence is completely conserved (Figs. 3B, 4C,D). The critical glycine (123) in motif2 is also conserved [Figs. 3B (in red), 4B]. Synteny reinforces the view that the *xol-1* genes in *C. briggsae* and *C. elegans* are orthologs. In *C. elegans*, the *xol-1* gene is 10 kb upstream of *dim-1*. Similarly, in *C. briggsae*, *xol-1* and *dim-1* are in close proximity (32 kb) on the same chromosome (see Materials and Methods). Although the biological role of *C. briggsae* XOL-1 is unknown, the identification of a *xol-1* ortholog indicates that the *C. elegans* XOL-1 protein is not simply an isolated case, and that the GHMP kinase fold has likely been adapted by a much more divergent array of protein sequences than had been assumed previously. Studies to analyze the function of *C. briggsae xol-1* are underway.

#### The putative catalytic site of XOL-1 and implications for activity

The putative active site of XOL-1 is a deep cavity nestled in the interface of domains 1 and 2 (Fig. 4), a position analogous to that occupied by the active sites of other GHMP kinases (Fig. 2B). The surface of the cavity is largely electronegative with interspersed electropositive regions defined by Lys 124 and Arg 264 (Fig. 4A,B). The negatively charged residues lining the cavity are Asp 52, Asp 120, Asp 220, Glu 276, and Glu 297 (Fig. 4A,B), only one of which (Asp 120) is apparently conserved in the *C. briggsae* ortholog (Asp 116; Fig. 4C,D). The phosphate-

binding loop (residues 120–129), which connects  $\beta 5$  and  $\alpha 3$ , straddles the back of the cavity (Fig. 4B). Although the XOL-1 motif2 contains two tandem serines (121, 122; Fig. 3A, in boxes), as in other GHMP kinases, the positions of these residues are not analogous. Whereas in other GHMP kinases the tandem serines are incorporated into the first turn of the ensuing  $\alpha$ -helix, these positions are occupied by the hydrophobic residues Ala 132 and Val 133 in XOL-1, and the tandem serines are located within the loop several residues before the start of the corresponding  $\alpha$ -helix,  $\alpha 3$  (Fig. 4A,B). The only structurally equivalent phosphate-binding loop residue conserved between XOL-1 and other GHMP kinases appears to be Gly 123 [Figs. 3A (red within motif2), 4B]. A functional requirement for a strong positive charge at position Lys 124 is suggested by conservative substitution by Arg 120 in the *C. briggsae* ortholog (Fig. 3B, motif2). A constellation of residues (Gly 295 to Lys 309) at the bottom of the cavity formed by the putative active site of XOL-1 is also conserved in the *C. briggsae* ortholog (Figs. 3B, 4C,D;  $\beta 9$  and  $\beta 10$ , in dark blue), a characteristic that may correlate with functional requirements.

In homoserine kinase and mevalonate kinase (*R. norvegicus*), Glu 130 and Glu 193 carboxylates, respectively, coordinate the nucleotide phosphate liganded  $Mg^{2+}$ . Glu 149 in mevalonate kinase (*M. jannaschii*) and Ser 153 carbonyl oxygen and  $O_{\gamma}$  in mevalonate-5-phosphate decarboxylase occupy the analogous position. In phosphomevalonate kinase, nearby Asp 144 and Asp 150 provide possible  $Mg^{2+}$ -chelating moieties. XOL-1 presents a marked structural incongruity with regard to this important  $Mg^{2+}$ -binding region. This location in XOL-1 is entirely hydrophobic, bounded by Val 127, Ile 128, Ile 159, Leu 164, Ile 166, and Ile 190 side chains and devoid of any charged or polar groups, even from the peptide backbone, to coordinate  $Mg^{2+}$ .

The surprising structure of XOL-1 implies that, in this case, the GHMP kinase fold acts as a developmental gene regulator or that sexual differentiation in *C. elegans* is regulated in a manner only remotely considered previously, that is, by a small secondary effector molecule. Thus, the structure suggests new avenues of experimentation that otherwise would likely have been overlooked.

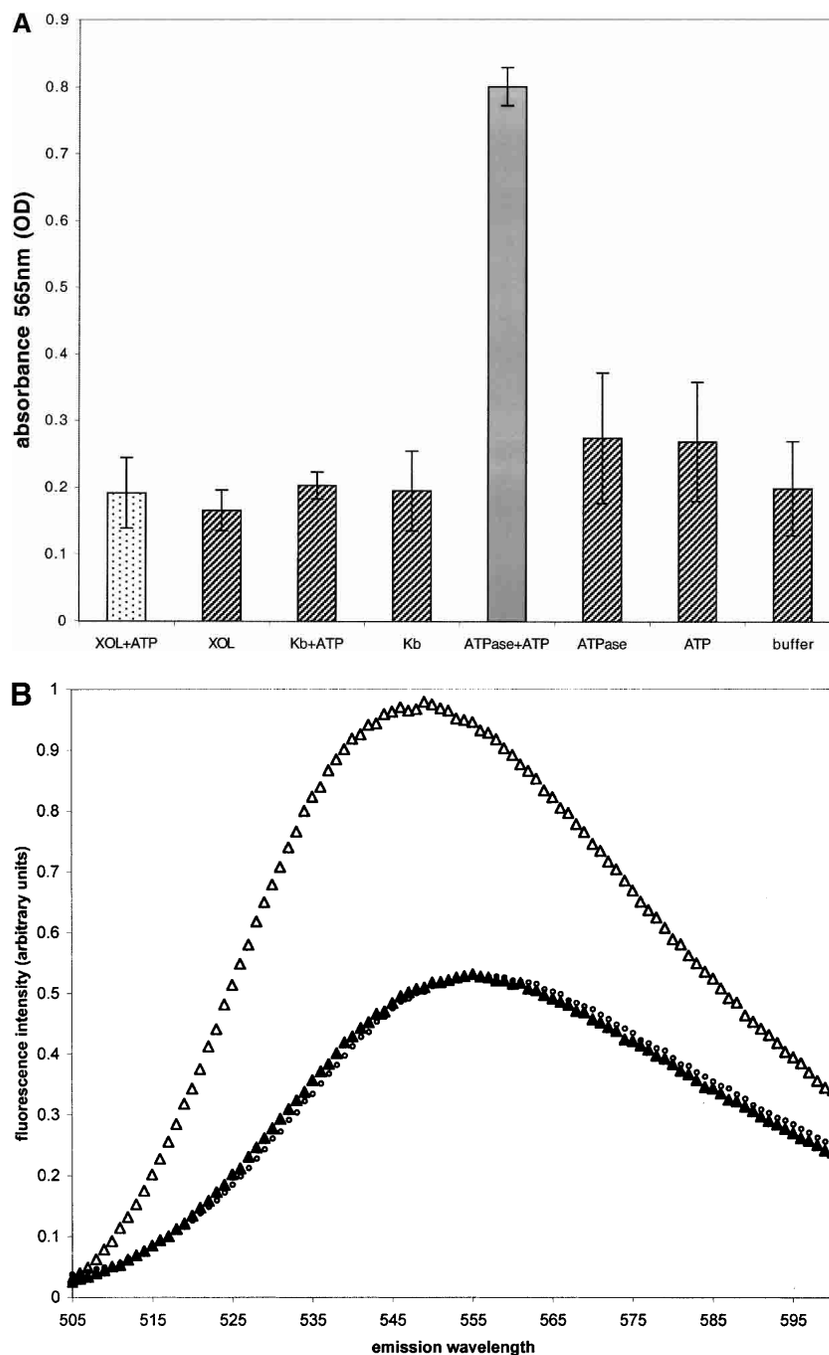
The XOL-1 used in the structure determination (residues 1–374) lacks the highly acidic feminizing C terminus (residues 375–417), yet retains complete ability to rescue *xol-1* null males (Rhind et al. 1995). Therefore, the structure does not provide information as to the function of this short C-terminal domain, which, thus far, has not been observed in other GHMP kinases. However, it can be deduced from the structure that the GHMP kinase core of XOL-1 does not incorporate the acidic C terminus. A conserved acidic motif at the extreme C termini is apparent in the aligned *C. elegans* and *C. briggsae* XOL-1 orthologs (residues 408–413 in *C. elegans* XOL-1; QDDTYD, residues 416–421 in *C. briggsae* XOL-1; NDDTFD), the functional significance of which is unknown (Fig. 3B, highlighted in green).

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*XOL-1 binding to ATP is not detectable*

The characterization of XOL-1 as a member of the GHMP kinase family of proteins led us to test whether XOL-1 could bind ATP. UV cross-linking failed to detect binding of XOL-1 to isotopically labeled ATP (data not shown). Likewise, residual XOL-1-catalyzed ATP hydrolysis was not detectable in an enzyme-coupled colorimetric assay (Fig. 5A). These results were confirmed by fluorescence spectroscopy. The fluorescent probe TNP-ATP, which had been used in studies of human mevalonate kinase (Cho et al. 2001), did not display any signifi-

cant enhancement of fluorescence in the presence of XOL-1 (Fig. 5B). In addition, crystals were both soaked and grown in the presence of a 10-fold molar excess of either ATP or the nonhydrolyzable analog AMP-PNP. However, no additional density was observed in the putative catalytic site in electron density maps calculated from x-ray diffraction data collected from these crystals (data not shown). Therefore, XOL-1 may not be a small molecule kinase, having evolved and adapted a GHMP kinase scaffold to execute a completely novel function. The lack of sequence similarity present between XOL-1 and other GHMP kinases, even within previously iden-



**Figure 5.** XOL-1 ATP hydrolysis and binding. (A) In the presence of XOL-1 (10  $\mu$ g; stippled bar), absorbance at 565 nm (Y-axis) is not significantly increased over the absorbance observed in any negative controls (hatched bars) in an enzyme-coupled colorimetric ATPase assay (see Materials and Methods). Negative controls (hatched bars) XOL-1, H-2K<sup>b</sup> + 1 mM ATP, H-2K<sup>b</sup>, Na<sup>+</sup>/K<sup>+</sup>-ATPase (5 milliunits), 1 mM ATP, buffer. As a positive control (gray bar), Na<sup>+</sup>/K<sup>+</sup>-ATPase (5 milliunits) was incubated with 1 mM ATP. (B) The fluorescence emission of 20 mM ATP-TNP (○) is not significantly increased in the presence of 20  $\mu$ M XOL-1 (▲). As a positive control, 20  $\mu$ M hexokinase was incubated with 20  $\mu$ M TNP-ATP (Δ). Fluorescence emission intensity (Y-axis) is represented as function of wavelength (505–600 nm, X-axis). Excitation wavelength = 410 nm.

tified conserved motifs (Fig. 3A), supports this conclusion. Of note, the left-handed  $\beta$ - $\alpha$ - $\beta$  fold in the N-terminal domain of XOL-1 and other GHMP kinases, termed the ribosomal protein S5 domain 2-like fold, is found in numerous RNA/DNA-binding proteins, raising the intriguing possibility that XOL-1 may interact with polynucleotide substrates, although, for this to be the case, significant conformational change in the active site would seem to be required. Structural similarity has also been noted between an RNA-binding protein domain, for example, domain V of elongation factor G, and the C-terminal domain of homoserine kinase (Zhou et al. 2000). Interestingly, RNA-binding domain 2 of the Sex-lethal protein has a fold similar to that of elongation factor G. The possibility remains that XOL-1-ATP binding may require an additional uncharacterized interaction.

#### Implications of crystal packing for oligomeric structure

Although the biological oligomerization state of XOL-1 is unknown, crystal packing suggests that XOL-1 might be a dimer or even a tetramer. One possible mode of dimerization evident from the crystal packing of XOL-1 involves the reciprocal interactions of domains 1 and 2 from neighboring monomers (Fig. 6A) with a buried surface area of 2146 Å<sup>2</sup> per dimer, slightly less than that calculated for *R. norvegicus* mevalonate kinase (2450 Å<sup>2</sup>; Fu et al. 2002). The additional structural feature formed by the unique N- and C-terminal  $\beta$ -strands ( $\beta$ 1 and  $\beta$ 13) of XOL-1 (Fig. 3A) presents a second possible interface for dimerization. XOL-1 molecules pack in the crystal such that  $\beta$ 1 strands link  $\beta$ -sheets from adjacent monomers to form an extended 12-stranded antiparallel  $\beta$ -sheet comprised of 6 strands from each monomer ( $\beta$ 9,  $\beta$ 10,  $\beta$ 8,  $\beta$ 11,  $\beta$ 13,  $\beta$ 1; Fig. 6B). As a caveat, extended  $\beta$ -sheets that do not correlate with functional dimerization are sometimes observed in crystal structures (Schiffer et al. 1985). In addition, XOL-1 elutes as a monomer during size exclusion chromatography (data not shown).

Crystal structures of homoserine kinase (Zhou et al. 2000) and mevalonate kinase (*R. norvegicus*; Fu et al. 2002) reveal a possible mode of dimerization mediated through interactions between C-terminal domains. However, the crystal structure of XOL-1 does not reflect a similar mode of dimerization, as the molecules are packed in a head-to-toe array with corresponding C-terminal dimerization interfaces distal to each other, an arrangement more similar to that seen in the mevalonate decarboxylase dimer (Bonanno et al. 2001). The two potential modes of XOL-1 dimerization combine to form a tetrameric entity with a large central cavity having a diameter of ~45 Å at its core (Fig. 6C).

#### Discussion

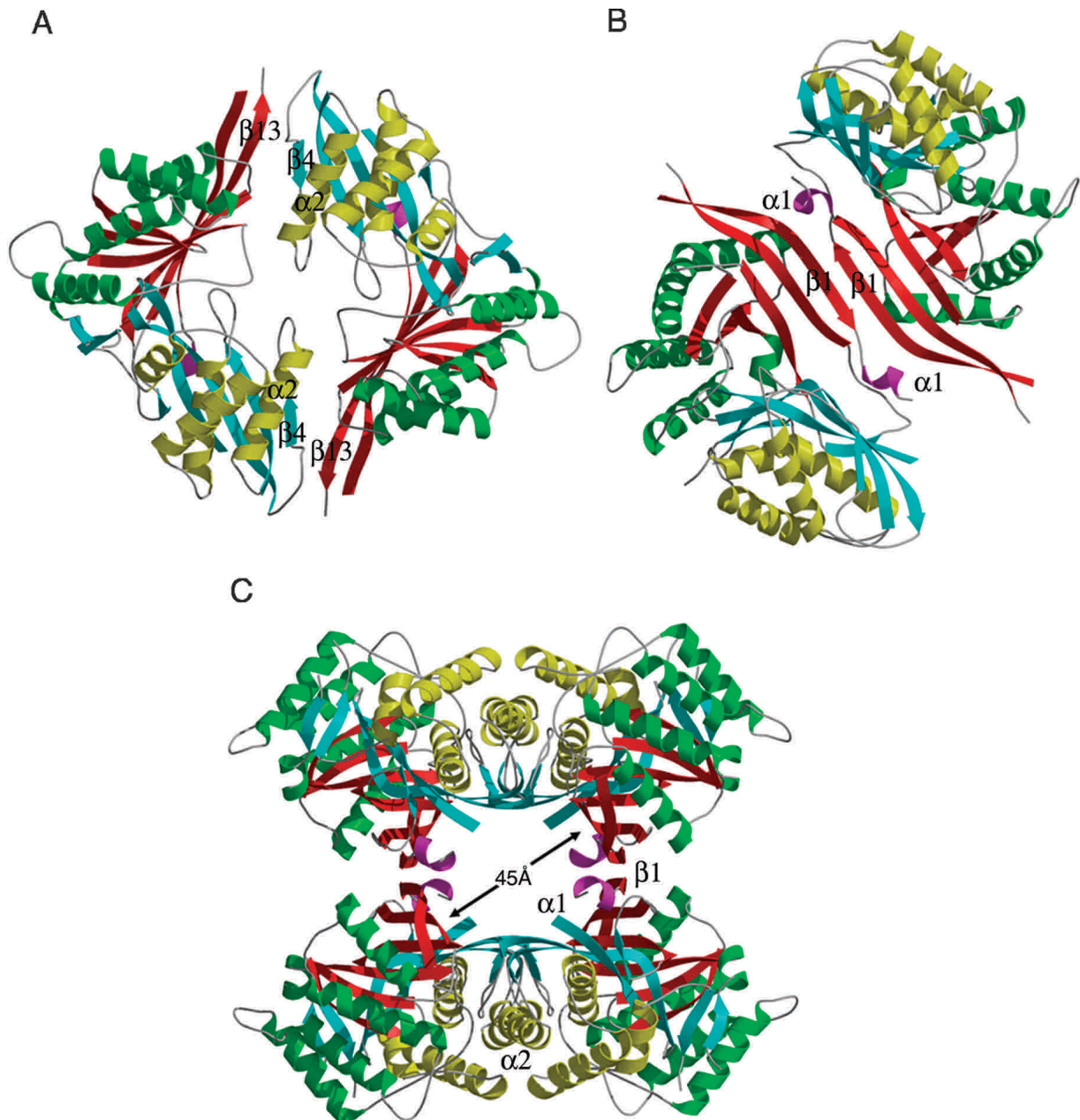
The identification of XOL-1 as a GHMP kinase family member through the elucidation of its crystal structure is difficult to reconcile with its role as a developmental regulator. However, gene regulation by GHMP kinase

family members has precedent in yeast. *Saccharomyces cerevisiae* Gal3p, whose primary structure is highly similar to that of galactokinase, is a galactose- and ATP-dependent transcriptional inducer of *GAL* genes (Platt and Reece 1998; Platt et al. 2000). Moreover, *GAL1* of *Kluyveromyces lactis* is a bifunctional protein that acts as an inducer of *GAL* genes and a galactokinase. (Meyer et al. 1991). As our understanding of how *S. cerevisiae* Gal3p and *K. lactis* *GAL1* induce gene expression is incomplete, their relationship to XOL-1 does not enable us to predict a detailed mechanism; however, circumstantial evidence suggests a mechanism of gene regulation that is conserved across phyla.

The principles of sexual dimorphism and dosage compensation are conserved in multicellular eukaryotes, but the mechanisms are not. For instance, in female mammals, a single X chromosome is inactivated randomly to reduce the overall X-linked expression by half, thereby reducing X chromosome gene expression to male levels. In the fruit fly, the single male X chromosome is hypertranscribed, such that X chromosome gene expression is doubled to female levels. In *C. elegans*, transcription from both hermaphrodite X chromosomes is reduced by half. With only one XOL-1 ortholog identified, the extent to which XOL-like GHMP kinases might regulate sexual fate and dosage compensation in other eukaryotes cannot be assessed. Because dosage compensation in different eukaryotic lineages is thought to have evolved independently, regulation of sexual fate and dosage compensation by XOL-like GHMP kinases may be limited to *C. elegans* and related nematodes. However, XOL-like GHMP kinases may also participate considerably more extensively, not only in the regulation of sexual fate and dosage compensation, but in other modes of development and differentiation. The accelerated rate at which genes involved in sexual differentiation and dosage compensation have evolved may obscure related gene sequences. Presumably, XOL-1 is a case of rapidly divergent evolution as such a remarkable degree of structural similarity would less likely be manifested by a case of convergence. Ambiguities will likely only be resolved when both sequence and structure space are more densely populated. To this end, structural genomic (Bonanno et al. 2001; Burley and Bonanno 2002) and whole-genome sequencing projects will undoubtedly provide indispensable insights.

Nematodes, which may be free-living or parasitic, are among the most numerous animals on earth and populate terrestrial, aquatic, and marine environments in abundance. *C. elegans*, whose entire genome has been sequenced (The *C. elegans* Sequencing Consortium 1998), is the most thoroughly studied nematode species and a useful organism for the study of parasitic nematodes (Burglin et al. 1998; Favre et al. 1998; Ashton et al. 1999; Blaxter and Ivens 1999). Efforts to characterize the genomes of parasitic nematodes (The Filial Genome Project: <http://nema.cap.ed.ac.uk/fgp.html>) have already resulted in the identification of the apparent homolog of *C. elegans* male sex determination gene *her-1* in *Brugia malayi* (Streit et al. 1999). A *her-1* homolog has also been

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**Figure 6.** Crystal packing interactions of XOL-1 monomers. (A) Interface 1 consists of reciprocal interactions of regions 1 and 2 from neighboring monomers with a buried surface area of 2146 Å<sup>2</sup> per dimer. (B) Interface 2 consists of anti-parallel hydrogen bonding between  $\beta 1$ -strands of neighboring monomers, creating a 12-stranded anti-parallel  $\beta$ -sheet (red) comprised of 6 strands from each monomer ( $\beta 9$ ,  $\beta 10$ ,  $\beta 8$ ,  $\beta 11$ ,  $\beta 13$ ,  $\beta 1$ ). (C) The central cavity of the tetramer in the crystal has a 45 Å diameter at its core. Secondary structure elements are colored as in Figure 3.

identified in *C. briggsae* (Streit et al. 1999), and homologs of *C. elegans* hermaphrodite sex determination gene *tra-2* have been identified in *C. briggsae* (Kuwabara and Shah 1994) and *Caenorhabditis ramaneni* (Haag and Kimble 2000).

These findings imply that, despite differing strategies of sexual reproduction between nematode species, some elements of the process are conserved. We infer from these observations that XOL-like GHMP kinases may

play an important role in the development and differentiation of various nematodes. Furthermore, XOL-like GHMP kinases may regulate differentiation in related organisms such as other helminths, for example, schistosomes and, perhaps even more broadly, in other metazoans. It is likely that the precise mechanism of XOL-1 activity will eventually emerge from studies based on the discoveries described herein. In summary, the structure of XOL-1 demonstrates that the GHMP kinase

family of proteins is much more diverse in sequence and function than had been presumed, and that critical pathways in development and differentiation may be controlled by a subfamily of enzymes preliminarily termed XOL-like GHMP kinases.

## Materials and methods

### Protein expression and purification

Full-length XOL-1 and a series of C- and N-terminal truncation mutants were generated by PCR and expressed with N-terminal polyhistidine tags in pRSET-A (Invitrogen) and PET28a (Novagen) vectors. Protein expression in BL21 DE3 strain *E. coli* was induced with 1 mM IPTG at a cell density of ~0.6 OD at 600 nm. Induction took place overnight at room temperature or 16°C. Protein was purified by Ni<sup>2+</sup>-affinity and anion exchange chromatography. N-terminal polyhistidine tags were removed by thrombin or enterokinase cleavage followed by an additional application of anion exchange chromatography. Selenomethionine-labeled protein was generated in the methionine-auxotrophic strain B834 DE3 (Novagen) in a similar manner, but bacteria were grown in methionine-deficient minimal medium supplemented with 0.3 mM L-selenomethionine (Sigma). Incorporation of selenomethionine into the recombinant protein was confirmed by MALDI mass spectrometry.

### Crystallization and data collection

The C-terminal truncation mutant consisting of residues 1–374 (80 mg/mL) crystallized at 4°C from 0.1 M HEPES (pH 7.8), 0.2 M MgCl<sub>2</sub>, 17.5% PEG400. MPD (>5%) was used as an additive to improve crystal size and quality. Crystals were harvested from drops with nylon cryoloops (Hampton Research) and directly cryocooled to –180°C without additional cryoprotectant. Native and far remote data were collected to 1.55 Å resolution at SSRL beamline 11-1 and MAD data (peak, inflection, and remote wavelengths) were collected at ALS beamline 5.0.2 (Table 1). All data were processed and scaled in HKL2000 (Otwinowski and Minor 1997). The space group is orthorhombic P2<sub>1</sub>2<sub>1</sub>2 with unit cell dimensions a = 116.1 Å, b = 86.0 Å, and c = 80.2 Å, and two protein molecules per asymmetric unit (V<sub>m</sub> = 2.4 Å<sup>3</sup>/Da).

### Structure determination and refinement

Positions of Se atoms (20 of 22 possible) were determined and refined in SOLVE and RESOLVE (Terwilliger and Berendzen 1996). Phase extension and automated chain tracing using the native data were performed in ARP/WARP (Lamzin and Wilson 1993). Further refinement was performed in CNS (Brünger et al. 1998) with intervening rounds of manual model building using the program O (Jones et al. 1991). Water molecules were added automatically using CNS (Brünger et al. 1998) and verified by manual inspection in O (Jones et al. 1991). Additional analysis of the final model was performed with PROCHECK (Laskowski et al. 1993). The final model consists of 703 amino acid residues and 628 water molecules (Table 1).

### Domain analysis

The XOL-1 structure has a complex topology. Automated domain analysis programs TOPS (Westhead et al. 1999) and DOMID [Guogusng, Lund University (<http://bioinfo1.mbfys.lu.se/Domid>)] classified it as a single domain protein. However, visual

inspection of the structure hinted at the presence of two domains. Therefore, we used a more time-consuming algorithm (Godzik et al. 1993) on the basis of the analysis of a protein distance map to analyze the domain organization of the structure. A protein domain, by definition, is a compact substructure that would form a stable structure when excised from the whole protein. Upon distance mapping, such domains can be identified as boxes along the diagonal. Visual inspection of the XOL-1 structure distance map identified two domains, located approximately between residues 45 and 195 (domain 1) and 202 and 360 (domain 2). Structures of both domains were submitted separately to DALI (Holm and Sander 1993; <http://www.ebi.ac.uk/dali>) and CE (Shindyalov and Bourne 1998; <http://www.cl.sdsc.edu/ce.html>) structure comparison servers. In both cases, strong structural similarities to the domains of GHMP kinases were detected.

### Cloning of *C. briggsae* xol-1

The *C. elegans* XOL-1 amino acid sequence was compared with the unfinished genomic sequence of *C. briggsae* (The Sanger Institute, <http://www.sanger.ac.uk>; The Genome Sequencing Center, Washington University in St Louis, <http://genome.wustl.edu>) by TBLASTN with standard parameters. A short stretch of potential similarity was identified in *C. briggsae* cosmid CB030H15. Using the coding sequence predicted from the *C. briggsae* genome, gene-specific primers to the most highly conserved regions were used in conjunction with RACE and conventional PCR to isolate cDNA fragments corresponding to the transcribed *xol-1* gene. When aligned with the *C. elegans* XOL-1 amino acid sequence by the clustal method (GONNET series), the ORF from the *C. briggsae* ortholog displays low sequence identity (22%). The *C. briggsae* *dim-1* gene is found on cosmid CB027N19. Cosmids CB030H15 and CB027N19 are both found on contig cb25.fpc4044. In *C. briggsae*, *xol-1* is ~32 kb upstream of *dim-1*. In *C. elegans*, *xol-1* is ~10 kb upstream of *dim-1*.

### ATPase assay

ATPase activity was measured using a maltose phosphorylase-glucose oxidase-horseradish peroxidase enzyme-coupled colorimetric assay (P<sub>i</sub>per Phosphatase Assay, Molecular Probes) according to the manufacturer's protocol. Briefly, XOL-1 (10 µg), MHC Class I H-2K<sup>b</sup> (10 µg; negative control) and Na<sup>+</sup>/K<sup>+</sup>-ATPase (5 milliunits; positive control) were incubated in the presence and absence of 1 mM ATP for various times up to 2 h at 37°C in 1× reaction buffer containing 100 µM Amplex red, 0.6 unit maltose phosphorylase, 0.3 unit glucose oxidase, and 0.6 unit horseradish peroxidase in a volume of 150 µL. Assays were performed and repeated in triplicate. Absorbance at 565 nm was measured in 96-well plates using a Molecular Devices SpectraMax 250 plate reader. Additional negative controls were ATP (1 mM) alone and buffer only.

### Steady-state fluorescence spectroscopy

Data were acquired using a SLM-AMINCO 8100 spectrofluorimeter. The concentration of protein and ligand (TNP-ATP, Molecular Probes) were 20 µM in 20 mM Tris-HCl (pH 8.0), 5 mM MgCl<sub>2</sub> at 25°C (volume = 150 µL). Emission was collected at 90° to the direction of excitation. To correct for fluctuation in the output from the 450 W xenon lamp, the experimental signal was divided by that of a quantum counter (3 g/L rhodamine G in ethylene glycol). To eliminate polarization artifacts, the excitation polarizer was set to the vertical position, and the emission

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polarizer was oriented 54.7° to the vertical (magic angle conditions). An excitation wavelength of 410 nm (8 nm = bandpass) and an emission maximum of 544 nm (8 nm = bandpass) for the bound ligand were determined experimentally. The sensitivity of the emission channel was set to 30% of maximum, resulting in a PMT voltage of 1050 V and a signal gain of 10 (voltage = 560 V and signal gain = 1 for reference channel). Emission spectra were collected from 490 to 600 nm with an integration time of 1 sec. Experiments were performed in triplicate, and spectra were corrected by subtracting the background fluorescence of the protein in the absence of ligand.

**Coordinates** Structure factors and atomic coordinates (accession no. 1MG7) have been deposited in the Protein Data Bank (<http://www.rcsb.org/pdb>).

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