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Suchita Rastogi  
Washington University School of Medicine in St. Louis

Ben Borgo  
Washington University School of Medicine in St. Louis

Nanette Pazdernik  
Washington University School of Medicine in St. Louis

Paul Fox  
Washington University School of Medicine in St. Louis

Elaine R. Mardis  
Washington University School of Medicine in St. Louis

See next page for additional authors

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Authors
Suchita Rastogi, Ben Borgo, Nanette Pazdernik, Paul Fox, Elaine R. Mardis, Yuji Kohara, Jim Havranek, and Tim Schedl
Caenorhabditis elegans glp-4 Encodes a Valyl Aminoacyl tRNA Synthetase

Suchita Rastogi,* Ben Borgo,* Nanette Pazdernik,* Paul Fox,* Elaine R. Mardis,* Yuji Kohara,†
Jim Havranek,* and Tim Schedl*†,1
*Department of Genetics, Washington University School of Medicine, St Louis, Missouri 63110, and †National Institute of Genetics, Mishima 411-8540, Japan

ABSTRACT Germline stem cell proliferation is necessary to populate the germline with sufficient numbers of cells for gametogenesis and for signaling the soma to control organismal properties such as aging. The Caenorhabditis elegans gene glp-4 was identified by the temperature-sensitive allele bn2 where mutants raised at the restrictive temperature produce adults that are essentially germ cell deficient, containing only a small number of stem cells arrested in the mitotic cycle but otherwise have a morphologically normal soma. We determined that glp-4 encodes a valyl aminoacyl transfer RNA synthetase (VARS-2) and that the probable null phenotype is early larval lethality. Phenotypic analysis indicates glp-4(bn2ts) is partial loss of function in the soma. Structural modeling suggests that Gly296Asp results in partial loss of function by a novel mechanism: aspartate 296 in the editing pocket induces inappropriate deacylation of correctly charged Val-tRNAval. Intragenic suppressor mutations are predicted to displace aspartate 296 so that it is less able to catalyze inappropriate deacylation. Thus glp-4(bn2ts) likely causes reduced protein translation due to decreased levels of Val-tRNAval. The germline, as a reproductive preservation mechanism during unfavorable conditions, signals the soma for organismal aging, stress and pathogen resistance. glp-4(bn2ts) mutants are widely used to generate germline deficient mutants for organismal studies, under the assumption that the soma is unaffected. As reduced translation has also been demonstrated to alter organismal properties, it is unclear whether changes in aging, stress resistance, etc. observed in glp-4(bn2ts) mutants are the result of germline deficiency or reduced translation.

Germline stem cells produce daughter cells that differentiate into gametes. The generation of a sufficiently large number of gametes for reproductive success requires significant germline stem cell activity. The Caenorhabditis elegans hermaphrodite exhibits this extensive stem cell activity. At hatching, each hermaphrodite contains two germ cells that proliferate to more than 800 germ cells per gonad arm in the young adult and that continue to proliferate during the subsequent progeny production period (Hubbard and Greenstein 2005; Pazdernik and Schedl 2013). To identify genes that specify the germline stem cell fate as well as genes that are necessary for stem cell proliferation and cell-cycle progression, genetic screens have been conducted in C. elegans for mutations that result in an abnormal Germ Line Proliferation (Glp) phenotype. Two genes identified in these screens are of particular interest, glp-1 and glp-4. Mutations in the glp-1 gene were used to identify the Notch pathway as the key signaling system between the somatic niche and germ cells for the specification of the germline stem cell fate (Austin and Kimble 1987, 1989; Yochem and Greenwald 1989). In glp-1 loss of function mutants, germline stem cells prematurely enter meiosis to form gametes. Temperature-sensitive (ts) glp-1 alleles have been identified and used to further understand the role of GLP-1 signaling in the proliferation vs. meiosis decision at the cellular level and to identify other genes involved in the stem cell fate decision through suppressor and enhancer screens (e.g., Austin and Kimble 1987; Fox and Schedl 2015; Maine and Kimble 1989). The glp-4 gene was identified by the ts allele bn2; when homozygous mutant embryos or L1 larvae are grown at the restrictive temperature (25°C) adults contain only approximately 12 germ cells, which appear to be arrested in prophase of...
the mitotic cell cycle (Beanan and Strome 1992). When glp-4(bn2ts)
mutant adults are shifted back to the permissive temperature, cell-cycle
arrest is reversed and extensive proliferation occurs, restoring the nor-
mal pattern of meiotic development and oogenesis. The regeneration of
the germline indicates that the arrested cells correspond to the germline
stem cell population. glp-4 was suggested to interact with GLP-1 sig-
naling as mutations reported to be alleles of glp-4 were isolated in a
screen for enhancers of the premature meiotic entry phenotype of glp-1
partial loss of function (Qiao et al. 1995). glp-1(ts) and glp-4(bn2ts)
mutant adults grown throughout larval development at the restrictive
temperature have 1% or less the normal number of germ cells but have
a soma that is morphologically indistinguishable from wild type at the
level of light microscopy (Austin and Kimble 1987; Kodoyianni et al.
1992; Beanan and Strome 1992), suggesting that only the germline is
affected in the mutants. These properties have led glp-4(bn2ts) and glp-
1(ts) mutants to be widely used to generate populations of germline
deficient adult hermaphrodites for molecular biology experiments to
identify genes whose RNAs and proteins are enriched in the germline
(e.g., Rosenquist and Kimble 1988; reviewed in Reinke 2006).

Germline proliferation in C. elegans has been linked to rapid organismal
aging. Elimination of all germ cells by laser ablation results in lifespan
extension (Hsin and Kenyon 1999), which appears to be unrelated to
the production of oocytes or sperm, whereas mutants that result in germ
cell overproliferation cause lifespan shortening (Arantes-Oliveira et al.,
2002). Stimulated by the finding that loss of germ cells leads to lifespan
extension, researchers have investigated the role of germ cells in signaling
for regulation of other organismal characteristics. However, removal of
the germline by laser ablation is technically challenging and very low through-
put. Therefore, researchers have turned to the use of glp-1(ts) and glp-
4(bn2ts) and temperature shifts to generate populations of adult
hermaphrodites that have very few germ cells. For example, glp-4(bn2ts)
mutants, as well as glp-1(ts) mutants, have been used to examine the role of
the germline in aging, stress resistance, pathogen resistance and fat metab-
olism (e.g., Wang et al. 2008; TeKippe and Aballay 2010; Greer et al.
2010; Labbadia and Morimoto 2015). Nevertheless, interpretation of findings
after use of glp-4 and glp-1 mutants to assess the role of germ cell prolif-
eration on organismal properties relies on the assumption that the mu-
tant conditions do not also affect somatic tissues in a way that could
influence aging, stress resistance, pathogen resistance and metabolism. This
assumption has not been fully investigated.

The molecular identity and null phenotype of glp-4 are unknown. This
limits our understanding of its role in germline cell proliferation,
potential interaction with GLP-1 signaling, and interpreting results when
glp-4(bn2ts) is used to generate germ cell—deficient adults for
studies of organismal properties such as aging and stress resistance. We
used whole-genome sequencing to demonstrate that glp-4 encodes the
valyl aminocyl transfer RNA (tRNA) synthetase VARS-2 and deter-
mined that the null phenotype is early larval lethality, consistent with a
major disruption of protein synthesis. Based on phenotypic analysis we
find that glp-4(bn2ts) is partially deficient for valine-tRNA synthetase
function in the soma at the restrictive temperature, thus raising the
question whether changes in organismal properties in glp-4(bn2ts) mu-
tants are due to germ cell deficiency or reduced protein synthesis.

MATERIAL AND METHODS

C. elegans strains

All strains were cultivated on nematode growth medium plates seeded
with E. coli strain OP50 (Brenner 1974) and were maintained at 20°C
unles indicated. The strains used in these studies are as follows: N2
(Bristol) as wild type, glp-4(bn2ts) (grown at 15°C), glp-4(bn2bn39), and
glp-4(bn2bn40) were obtained from Susan Strome. Enhancer of glp-1
(bn18), om14, was obtained from Eleanor Main. Y87G2A.5 (bn3947)
was obtained from the Japanese National Biosource Project and bal-
covered over hlin1[unc-54], aggf-1(ok1736)hlin1[unc-101(e241)] and
F22G12.5(ok2367) were obtained from the Caenorhabditis Genetics
Center. unc-75(e950) glp-4(bn2ts) and unc-73(c950) unc-101(m1) were
constructed by standard methods. Phenotypes were scored using a
dissecting microscope or Nomarski DIC microscopy with a 40 or
63x objective.

Molecular methods

Genomic DNA was isolated by a minor modification of the CTAB
(bn2ts) was obtained by paired-end reads (average 30x coverage) using
an Illumina Genome Analyzer through The McDonnell Genome In-
stitute, Washington University, St Louis MO. Following read pair align-
ment to the C. elegans reference genome with BWA (Li and Durbin
2009), five coding sequence changes in a previously mapped genetic
interval were identified in glp-4(bn2ts) that differed from the N2 wild-
type reference sequence (WormBase.org, release WS248): C54C8.5
(glct-5), nt position 12454271, G->A, Trp->Stop; F22G12.5, nt position
13168458, G->C, His->Asp; Y87G2A.5 (formerly called vurs-2, see below),
nt position 13556729, C>T, Gly->Asp; Y6BS3.1a (aggf-1), nt position
13616715, C>T, Asp->Asn; and Y71A12B.17 (gadr-5), nt position
13985048, A>G, Lys->Glu. To identify the molecular lesion in the glp-4(bn2ts)
intragenic revertants, we isolated total RNA from each mutant strain. Reverse-transcription polymerase chain re-
action with three sets of overlapping primers for Y87G2A.5 was then
used to generate the corresponding coding region cDNA, which was
Sanger sequenced. Feeding RNA interference (RNAi) was performed
(Lee et al. 2007) for each of the five candidate genes with the use of
clones obtained from the Ahringer library (Rual et al. 2004), with the
rrf-1(pk1417) background to largely limit knockdown to the germline
(Kumsta and Hansen 2012). RNA in situ hybridization was per-
formed as described (Motohashi et al. 2006).

Genetics methods

Complementation testing: We tested deletion alleles for three of the five
candidate genes by placing them in trans to unc-75(e950) glp-4(bn2ts),
verifying the genotype at 15°C, and then shifting to 25°C to examine the
phenotype. Only in the case of Y87G2A.5 (bn3947) unc-75(e950) glp-4
(bn2ts) were abnormal phenotypes observed.

Isolation of suppressors of glp-4(bn2ts): We isolated recessive suppres-
sors of glp-4(bn2ts) with an ethyl methanesulfonate mutagenesis
dscreen similar to the method described by Beanan and Strome (1992),
except that the screen for fertility was at 24°C. We assessed linkage of the
suppressor mutation to glp-4(bn2ts) by constructing unc-75(e950) glp-4
(bn2ts); suppressor+ strains and screening for unc-75 (e950) glp-4(bn2ts)
animals that were fertile at 24°C in the following generation. We recovered 26 suppressors from ~1.1 x 108 F2 generation
worms, which included suppressors that were both linked and unlinked
to glp-4(bn2ts). Linked suppressor bn2oz283 was examined further.

Analysis of glp-1 enhancer om14: A screen for extragenic enhancers of
glp-1(bn18) was performed previously in an effort to identify new
genes involved in GLP-1 Notch signaling (Qiao et al. 1995); among the
mutations identified was om14, which was reported to be an allele of
glp-4. We used complementation and linkage analysis to test whether
**om14** was an allele of **glp-4**. First, **om14**/*unc-75(e950) glp-4(bn2ts)* animals were generated at 15° and shifted to 25°. We found that at 25° heterozygotes were fertile and segregated ~1/4 of progeny that displayed the oogenesis defective phenotype of **om14** (Qiao et al. 1995), and ~1/2 of progeny that were nonUnc fertile hermaphrodites. Second, **om14** was placed in **trans** to **unc-75(e950) unc-101(m1)**, markers that are tightly linked to **glp-4**. In the next generation we found that ~1/4 of the **Unc** animals ([unc-75(e950) unc-101(m1)] displayed the oogenesis defective phenotype of **om14**. Based on finding that **om14** and **glp-4 (bn2ts)** complement and that **om14** is unlinked to the right arm of chromosome I, **om14** appears not to be an allele of **glp-4**.

### Comparative structural modeling of **glp-4** valine-tRNA synthetase

**Construction of the model:** The 2.9 angstrom crystal structure of the valine tRNA-synthetase (valRS) from *Thermus thermophilus* [PDB id: 1IVS (Fukai et al. 2003)] was used as a template for constructing a structural model of the valRS from *C. elegans* (**glp-4** VARS-2, see below). Protein sequences for **valRS** and **glp-4** VARS-2 were aligned using ClustalW (Larkin et al. 2007). Three general issues were addressed to generate a homology model: 1) substitution of amino acids at aligned positions; 2) deleted regions; and 3) inserted regions.

To address the first issue, we used RosettaDesign (Leaver-Fay et al. 2011; Kaufmann et al. 2010) to computationally substitute all nonidentical, aligned residues on the valRS structural template to their identity in **glp-4** VARS-2. The repulsive term in the Rosetta full-atom scoring function was drastically reduced, and the amino acids at altered positions were mutated *in silico* to adopt the **glp-4** VARS-2 sequence. Next, several rounds of rotational isomer (rotamer) sampling (Bower et al. 1997) were used to resolve steric clashes whereas the repulsive component of the Lennard-Jones term was gradually increased to ~60% of its standard value (Kaufmann et al. 2010). Finally, once all substitutions had been made and clashes resolved, gradient minimization (Payne et al. 1992) was performed on each domain in several iterations.

With each iteration, the repulsive term was slowly increased to its standard value to alleviate remaining clashes. Although some significant clashes remained, they were outside the regions of interest for this study, and no further effort was made to remove them.

Second, for elements of **valRS** that are not present in **glp-4** VARS-2, we manually deleted residues from the PDB file, leaving "gaps" in the structure. One side of the gap was manually chosen as the stationary partner (or target to which the other end of the gap would be aligned) whereas the other side was considered the mobile partner. Selection of the stationary and mobile partners was done depending on which end appeared less likely to be hindered by the surrounding structure. A single amino acid was added onto the mobile partner that overlapped with the targeted stationary partner. Harmonic constraints that forced the overlap residue (on the mobile partner) and the template residue (on the stationary partner) to be superimposed were then enforced, and the structure minimized in several steps with gradually increasing repulsive energy. This allowed efficient closure of the deletion gaps.

Third, for inserted stretches of amino acids that were present in **glp-4** VARS-2 but not in **valRS**, we added them within Rosetta by a well-developed process called fragment assembly (Misura et al. 2006). Briefly, fragment assembly employs a user-generated library of 3 or 9 amino acid peptide structures extracted from the protein database. These fragments are selected to be of similar sequence as the insertion region. A Monte Carlo sampling algorithm selects the combination of short fragments that both optimize the Rosetta fullatom scoring function and satisfy the starting and ending points for the insertion in the structural model. The fragment insertion that results in the best overall score is retained and manually inspected for the best fit, which in this case was the lowest root-mean square deviation (RMSD) compared with the template. RMSD is a cumulative measure of the mean distance between equivalent atoms in the model and template, which is routinely used to compare the similarity of two regions of homologous protein structure.

Next, we positioned the valyl-adenylate substrate in the editing site by structurally aligning a valyl-adenylate from PDB 1IVS onto to the 5'-O-(N-(L-threonyl)-sulfamoyl) adenosine substrate in PDB 1WK9.

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**Figure 1** Relative positions of genetic loci and mutant alleles within the genetic and physical map regions around **glp-4**. (A) Displays the genetic map on the right arm of Chromosome I with morphologic loci in the region used to localize **glp-4** indicated (top) and the physical map (bottom) of the region with genes that contain missense mutations from whole genome sequencing indicated. (B) Shows the intron-exon structure of Y87G2A.5, which encodes the cytoplasmic valyl aminoaeryl transfer RNA synthetase VARS-2. (C) Shows an expansion of exon 3 with the relative position of **glp-4** genetic lesions characterized in this paper indicated. Arrows in (B) and (C) are included to indicate the direction of transcription, with 3’ to the left. Diagrams are drawn to scale based on nucleotide position on Chromosome I. The tm3947 deletion spans nucleotide positions 13,555,971 to 13,556,472. The bn2, bn39, bn40 and o2283 point mutations occur at positions 13,556,729, 13,556,285, 13,556,288 and 13,556,310, respectively.
Figure 2 Alignment of C. elegans glp-4 VARS-2_C.elegans with human and Thermus thermophilus orthologs. Alignment performed with ClustalW (Larkin et al. 2007). Protein domains are color coded—the split Rossman fold in blue, CP1 domain in magenta, editing domain in green, alpha peptide in purple, tRNA recognition domain 1 in turquoise, helical bundle in pink, and tRNA recognition domain in yellow; this color code is used again in Figures 4A to indicate the relevant domains in the 3D structures. Point mutations (bn2, oz283, bn40, and bn39) are marked in the alignment in gray, whereas the deletion allele tm3947 is indicated by the dashed line above the alignment. The editing active
three proteins are indicated. Note that full structural reinsertions/deletions integrated, and the valyl-adenylate substrate added, site aspartates D379 and D382 are underlined. Below the alignment, residues that are identical (editing domain only) (Fukunaga and Yokoyama 2005). PDB 1WK9 was then aligned to the model and the coordinates for the valyl-adenylate substrate were transferred to the model.

Finally, once the model amino acids were fully substituted, insertions/deletions integrated, and the valyl-adenylate substrate added, full structural refinement (Misura et al. 2006) using constraints on the template yielded the completed model (see Figure 4). All structural manipulations and calculations were performed using standard and custom Rosetta programs (Leaver-Fay et al. 2011; Kaufmann et al. 2010) and all visualization performed using PyMol (DeLano 2002; The PyMol Molecular Graphics System, version 1.3. http://pymol.org/).

**Modeling mutations and paralogous substitutions:** Amino acid changes were modeled using RosettaDesign. Briefly, we performed Monte Carlo sampling of discrete rotamer side-chain conformations on a fixed backbone. This is referred to as “fixed-backbone” design and involves computationally cycling through predefined, low-energy conformations of a particular amino acid, calculating the whole structure energy for each conformation and finding the one with the lowest energy (Leaver-Fay et al. 2011; Kaufmann et al. 2010). After fixed backbone mutation/substitution, a number of small perturbations in the backbone and side-chain torsion angles are applied in combination with gradient minimization. This allows the backbone to be repositioned along an energy gradient based on new requirements from changing an amino acid identity. In each step, both the tRNA and the bound valyl-adenylate substrate were constrained to their original position (such that they could move, but such motion was heavily penalized by the scoring function). We performed flexible backbone optimization iteratively until no further improvement in energy due to the sequence change was observed (Misura et al. 2006).

**RESULTS**

**Molecular Identity of the glp-4 locus**

The glp-4 locus was previously mapped genetically to an approximately two-megabase interval between markers unc-75 and lev-10 on Chromosome I (Beanan and Strome 1992). We performed whole-genome sequencing (Hobert 2010) of the glp-4(bn2ts) reference allele-containing strain and identified five candidate genes in the mapped region that contained coding sequence changes compared to the Bristol N2 wild-type reference genome (see the section Materials and Methods; Figure 1). To determine which of the five genes encodes glp-4, we performed RNAi knockdown of each in the rrf-1 mutant background, which is proficient for RNAi in the germline but largely defective for RNAi in the soma (Kumsta and Hansen 2012). We then assessed the treated animals at 20°C for a sterility phenotype similar to that observed for glp-4(bn2ts) at 25°C. Only knockdown of Y87G2A.5 displayed a sterile phenotype, showing a strong reduction in germ cell number and little or no gametogenesis. We next performed complementation tests for the three candidate genes in the mapped region where deletion alleles were available from the knockout consortium (C. elegans Deletion Mutant Consortium 2012): Y87G2A.5 (tm3947), agef-1(ok1736) [Y6B3A.1], and F22G12.5 (ok2367). Only Y87G2A.5 (tm3947) failed to complement glp-4(bn2ts). The trans-heterozygotes, although weakly fertile at 15°C, are fully sterile at 25°C with a limited number of germ cells (also see below). Three tightly linked, possibly intragenic, suppressors of the glp-4(bn2ts) temperature sensitive sterility have been identified, bn2bn39, bn2bn40 and bn2oz283 (Beanan and Strome 1992; see the section Materials and Methods). The coding sequence of Y87G2A.5 in each of the suppressors was sequenced; in addition to the original bn2 missense change in exon 3, we found that the suppressors contained a second missense mutation in exon 3 (Figure 1). These findings together provide strong evidence that glp-4, defined by allele bn2, encodes a valyl aminoacyl tRNA synthetase (VARS-2, Y87G2A.5). Based on the large body of literature on glp-4, the gene name will remain glp-4, while the protein will be identified as VARS-2 to indicate its molecular identity.

glp-4 VARS-2 is a cytoplasmic class I valyl aminoacyl tRNA synthetase that catalyzes the transfer of valine to its cognate tRNA for protein synthesis. glp-4 VARS-2 has two tRNA recognition domains (1 and 2); a split class 1 Rossmann-fold, which functions in catalyzing the synthesis of aminoacyl-adenylate and aminoacyl-tRNA; an editing domain; and the connective polypeptide (CP1) domain, which also functions in post-transfer editing (Figure 2, Fukai et al. 2000). tm3947 deletes 502 base pairs and inserts four base pairs, removing part of the editing domain, all of the CP1 and alpha peptide domains and part of the Rossmann-fold. Thus tm3947 is likely a null allele. bn2 is a missense mutation, G296D, in the CP1 domain. Intragenic revertants bn2bn39-T444M, bn2bn40-P443L, and bn2oz283-E436K, are missense mutations in the adjacent editing domain (Figure 2; also see Structural analysis of glp-4 VARS-2).
opposite side of the pocket (red) and distances between aspartate carboxyl or glycine alpha-carbon and the scissile bond (dashed black line) indicated. (C) Close-up view of the editing pocket in indicated. (D) Close-up view of the editing pocket in opposite side of the pocket (red) and distances between aspartate carboxyl or glycine alpha-carbon and the scissile bond (dashed black line) indicated.

**glp-4 VARS-2 has somatic functions**

The soma of *glp-4(bn2ts)* homozygotes grown at the restrictive temperature throughout development does not have any obvious morphological abnormalities by light microscopy (Beanan and Strome 1992). Therefore, it was of interest whether *glp-4 VARS-2* has somatic functions, *tm3947*-null homozygotes arrest during early larval development. Although there is some variability in animal size, based on somatic gonad and germ cell counts (from 4–12 cells) the arrest is in the L1 stage. By contrast, *tm3947* heterozygotes are morphologically normal. These results are consistent with slow growth and larval lethality observed in genome-wide RNAi screens with Y87G2A.5 (Simmer et al. 2003; Nehme et al. 2010; Dunbar et al. 2012). *In situ* hybridization reveals that *glp-4* mRNA is expressed in the intestine and somatic gonad, in addition to strong expression in the germline (Figure 3). Thus *glp-4 VARS-2* is expressed and functions in both the germline and soma.

Based on the germline phenotype, *bn2* is a partial loss of function mutation; *bn2* failure to complement the *tm3947* deletion allele for the reduced germ cell number phenotype at 25° and the reduced germ cell number can be phenocopied by RNAi of *glp-4 VARS-2*. In addition to defects in germline development, *glp-4(bn2ts)* is also partially defective in somatic function. *bn2/tm3947* trans-heterozygotes at 25° are slow growing, reaching adulthood more than a day later than the *tm3947/+* control, and have a protruding vulva that often results in bursting in adults. Thus, notwithstanding the apparently normal somatic morphology of *bn2* homozygotes, *glp-4 VARS-2* activity is partially compromised in the soma of *glp-4(bn2ts)* animals at 25°.

**Structural analysis of *glp-4 VARS-2***

**Structural homology model**: We have used structural analysis to provide insight into (a) the effect of *bn2* on *glp-4 VARS-2* activity; (b) the mechanism by which the intragenic revertants suppress the effect of *bn2*; and (c) the unique function of *glp-4 VARS-2* relative to the valyl-tRNA synthetase paralog VARS-1. The crystal structure of the *T. thermophilus* valRS has been determined (Fukai et al. 2000; 2003). Each of the structural domains identified in valRS is readily discerned in the *glp-4 VARS-2* sequence (Figure 2) and secondary structure predicted with DSSP (Kabsch and Sander 1983). Although the overall primary sequence identity is modest (37%), the aligned predicted secondary structure of the two proteins in placement of helical and sheet segments is in 92% agreement. These findings, along with previous work that successfully generated a homology model with proteins of similar modest sequence identity (Misura et al. 2006), indicate that a reasonably accurate structural homology model for *glp-4 VARS-2* can be generated using valRS as a template. An initial model was generated as described in the Materials and Methods section and assessed by visual examination of the model to look for obvious structural inaccuracies and by computation of the model RMSD, which is a quantitative scalar measure of the generated model backbone’s fit relative to the template backbone. Our structural model for *glp-4 VARS-2* (Figure 4A) has a backbone RMSD of 0.8 angstroms (determined by pyMol alignment) from the starting template, indicating that the crystal structure can be used for homology modeling of the *C. elegans* protein with only modest backbone rearrangement.

**Effect of *bn2* on GLP-4 VARS-2 activity**: During the process of charging tRNAs with amino acids for subsequent translation, an amino acid residue can be misacylated to result in a noncognate tRNA–amino acid pair. Class I tRNA synthetases can correct such errors via post-transfer editing, i.e., the removal of an amino acid that is attached to a noncognate tRNA. This is an essential part of the "double-sieve" mechanism critical for maintaining translational fidelity. Isosteric threonine charged onto tRNAval is the major misacylation that is post-transfer edited by valyl-tRNA synthetases. Although a number of residues have been shown to be critical for coordination and recognition of the amino-acylated tRNA during post-transfer editing, hydrolytic cleavage of the misacylated Thr-tRNAVal substrate is thought to take place in a hydrophilic pocket of the tRNA’s editing domain. This domain contains a pair of conserved aspartates (D379 and D382, in green in Figure 2 and Figure 4) that are about 6.5 and 4.4 angstroms from the scissile bond (Fukai et al. 2000; Fujunaga and Yokoyama 2005). One of these aspartate residues deprotonates an adjacent water molecule to create an activated hydroxyl ion, which nucleophilically hydrolyzes the Thr-adenylate bond of the noncognate amino acid tRNAVal substrate.
The molecular mechanisms appear to be very different. The prior failure to recover new loss of function alleles in trans with bn2 in a noncomplementation screen (Beanan and Strome 1992) was probably because of severely reduced fertility, as we observed for tm3947/bn2 at the permissive temperature, which would be exacerbated by a general reduction in progeny production that typically occurs following chemical mutagenesis.

**Suppression of bn2 G296D by the intragenic revertants:** The glp-4 (bn2bn39) T444M and glp-4 (bn2bn40) P443L suppressor mutations lie in the editing domain, approximately 23 and 21 angstroms from the bn2 G296D mutation, respectively. Each sit in what is likely a 'hinge' region that joins the editing and CP1 domains to the rest of the protein. Modeling of these mutations into the G296D model introduces local steric clashes around residues 443 and 444 that, when resolved, alter the orientation of the beta-sheet linker in which they are contained. In the case of glp-4 (bn2bn39) T444M, minimizing the clashes increases the distance between the bn2 aspartate 296 mutation and the scissile bond from 7.4 to 8.5 angstroms, as the beta-sheet linker is "pushed" away from the editing domain. This is likely due to the physical change when replacing a relatively polar amino acid (threonine) with a larger, hydrophobic residue (methionine). Specifically, the surrounding region may "collapse" to minimize the solvent exposure of methionine, thus shifting the "upstream" beta-sheet. For glp-4 (bn2bn40), the P443L mutation increases the distance between the bn2 aspartate 296 mutation and the scissile bond from 7.4 to 11 angstroms as well as rotating the carboxyl side-chain away from the valyl-adenylate substrate (Figure 4D). Replacement of constraining proline 443 with the larger side chain in leucine results in a major disruption in the beta-sheet linker in which it resides. For the third intragenic suppressor glp-4 (bn2oa2283), E436K, the side chain of lysine 436 faces the external solvent and does not reveal an obvious disruption of the structure and positioning of aspartate 296. Thus the mechanism of suppression by E436K must be more indirect, with the structural model providing little insight. In sum, the intragenic suppressor mutations T444M and P443L result in displacement of the mutant aspartate in the editing pocket through steric effects that are propagated to bn2 G296D (Figure 4D), such that its ability to catalyze inappropriate deacylation of the cognate Val-tRNA is reduced.

Intragenic suppressors of bn2 were isolated at 100- to 1000-fold lower frequency following EMS mutagenesis than typical loss of function frequency, which was interpreted by Beanan and Strome (1992) as the bn2 mutant not being reverted through simple loss of gene activity and that the gene product is likely to be essential. Both inferences were correct based on our findings: (a) glp-4 VARS-2 is an essential gene; and (b) only a small subset of intragenic mutations are expected to suppress bn2, those that displace G296D in the editing pocket and thus decrease...
its inappropriate deacylation activity without otherwise damaging tRNA synthetase function. bn2bn40 was observed to be a stronger intragenic suppressor (weakly dominant, larger brood size) than bn2bn39 (fully recessive) (Benan and Strome 1992), which is consistent with the structural model where bn2bn40 P443L results in a greater displacement of the bn2 aspartate 296 from the valyl-adenylate substrate, compared with bn2bn39 T444M, as well as rotating the carboxyl side-chain away from the scissile bond (Figure 4D).

**Unique function of glp-4 VARS-2**: *C. elegans* contains a second valyl-tRNA synthetase paralog, *vars-1*. To determine why *vars-1* is unable to compensate for the reduced ability of glp-4 *bn2* and *tm3947* to produce Val-tRNAval, we aligned sequences in glp-4 VARS-2 that are in direct contact with the anticodon with the same region in VARS-1, and then mapped the VARS-1 anti-codon recognition residues onto our model (Figure 4A and Figure 5). The model reveals several bulky substitutions in the immediate vicinity of the anticodon in VARS-1. Tyrosine 816 of glp-4 VARS-2 lies in direct contact with the wobble position in the anticodon; VARS-1 has a bulkier tryptophan substitution for residue 816. Two nearby residues in glp-4 VARS-2, proline 813 and threonine 940, also have substitutions in VARS-1 to bulkier amino acids, lysine and glutamine, respectively. Thus, although the anticodon recognition pocket in glp-4 VARS-2 appears to easily accommodate binding of either purines or pyrimidines in the wobble position, the three bulky substitutions in VARS-1 decrease the pocket volume, sterically preventing binding anticonds with larger purines in the wobble position. Therefore VARS-1 is not able to efficiently charge tRNAval that recognizes valine codons GUC and GUU, which provides an explanation for the unique and essential function of glp-4 VARS-2 in viability, germ cell proliferation, and protein synthesis.

**DISCUSSION**

glp-4 encodes the VARS-2 valyl aminoacyl tRNA synthetase [Y87G2A.5] based on (a) genetic mapping of the canonical allele *bn2* to a 2 megabase region surrounding Y87G2A.5, (b) finding the G296D missense change in VARS-2, (c) finding that of all the genes within the genetically mapped region containing missense mutations in the glp-4 (*bn2ts*) strain, only RNAi of Y87G2A.5 phenocopies the glp-4 (*bn2ts*) mutant, (d) finding that the *tm3947* deletion in Y87G2A.5 fails to complement glp-4 (*bn2ts*), and (e) finding that tightly linked glp-4 (*bn2ts*) suppressor mutations are missence changes within VARS-2. Genetic analysis indicates that glp-4 (*bn2ts*) is a partial reduction-of-function mutation while structural modeling suggests that *bn2* G296D in the CP1 domain results in substitution of an aspartate in the editing pocket opposite to the catalytic aspartates, in a position where it can catalyze inappropriate deacylation of the correctly charged Val-tRNAval and thus reduce levels of Val-tRNAval. Structural modeling of the *bn2bn39* T444M and *bn2bn40* P443L intragenic suppressor mutations suggests that the bulkier missense changes result in displacement of the *bn2* aspartate 296 in the editing pocket so that it is not able to efficiently catalyze inappropriate deacylation of Val-tRNAval. glp-4 VARS-2 is essential for somatic and germine development as the likely null phenotype is early larval lethality. The widely used allele *bn2*, in addition to its germine defects at the restrictive temperature, is also defective in somatic development and function as glp-4 (*bn2ts*)/glp-4 null displays slow growth, protruding vulva, and bursting at the vulva. These findings reinforce the caution that a loss-of-function mutant that displays a strong morphologic phenotype in one cell type but no morphologic phenotype in other cell types does not preclude that the mutant is also partially defective in other cell types, although not manifested at a morphologic level. A partial loss-of-function allele of the arginyl-tRNA synthetase *vars-1* was identified in a forward genetic screen for mutations that result in resistance to hypoxia (Anderson et al. 2009). Although *vars-1* mutants, like glp-4 (*bn2ts*) mutants, are morphologically normal, they nevertheless have reduced protein translation rate. Thus by analogy, glp-4 (*bn2ts*) mutants at the restrictive temperature are likely to be partially defective in protein synthesis, which lead to the observed mutant phenotypes.

glp-4 (*bn2ts*) and glp-1 (*ts*) mutants have been used widely to generate germline deficient adults to study the role of signaling between germ cells and somatic tissues for organismal aging, stress resistance, pathogen resistance, and metabolism. Interpretations of results from such studies rely on the assumption that the glp-4 (*bn2ts*) and glp-1 (*ts*) germ cell–deficient animals have a soma that is unperturbed. However, our results indicate that glp-4 (*bn2ts*) at the restrictive temperature is partial loss of function in the soma. In addition, a large body of literature in *C. elegans* has implicated partial inhibition of translation in lifespan extension, stress resistance and pathogen resistance (e.g., Hansen et al. 2007; Pan et al. 2007; Anderson et al. 2009; Wang et al. 2010; Dunbar et al. 2012). glp-4 (*bn2ts*) mutants, although germine deficient, very likely have partial inhibition of translation due to reduced levels of charged Val-tRNAval. Thus it is unclear whether organismal lifespan extension, stress resistance, pathogen resistance, and increased fat accumulation observed in glp-4 (*bn2ts*) mutants is a result of germine deficiency, reduced protein synthesis or a combination of both phenotypes. It should be noted that glp-1 also has functions in the larval/adult soma (Mango et al., 1991; Berry et al., 1997; Singh et al., 2011), which raises concern about interpretation of studies of organismal properties of germine deficient glp-1 (*ts*) mutants. We suggest that attribution of organismal phenotypes to signaling from the germline is best demonstrated through laser ablation of the germine precursors, Z2 and Z3 (Kimble and White 1981).

glp-4 (*bn2ts*) mutants display two intriguing temperature shift phenotypes that can now be correlated with protein translation. Partial disruption of protein translation (at 25°C) through decreased levels of charged Val-tRNAval leads to mitotic cell cycle arrest in the stem cell population, whereas restoration of protein translation in the adult (following shift back to 15°C) results in release of mitotic cell-cycle arrest, extensive stem cell proliferation, and the resulting reformation of the normal germine distal-proximal polarity of meiotic prophase progression and gametogenesis. These phenotypes are remarkably similar to adult reproductive diapause (Angelo and Van Gilst 2009; Seidel and Kimble 2011) where complete food deprivation at the late L4 stage leads to cell cycle arrest among the stem cells, autophagy/apoptosis mediated resorption of the remaining germine and extended lifespan; upon refeeding there is extensive stem cell proliferation and reformation of the polarized meiotic and gametogenic germine organization. Studies of glp-4 (*bn2ts*) may thus contribute to understanding of how physiological perturbations (changes in levels of charged Val-tRNAval and/or protein translation) affect entrance and exit from adult reproductive diapause and used to model reformation of the polarized germine organization in adults that contain only a small stem cell population.

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