- **1** Insights into the involvement of spliceosomal mutations in myelodysplastic
- 2 disorders from an analysis of SACY-1/DDX41 in *Caenorhabditis elegans*

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Author Summary

33	Mutations affecting spliceosomal proteins are frequently found in hematological malignancies.
34	DDX41/Abstrakt is a metazoan-specific spliceosomal DEAD-box RNA helicase recurrently mutated in
35	inherited and relapsing myelodysplastic syndromes and acute myeloid leukemia. The genetic
36	properties and genomic impacts of disease-causing mutations in spliceosomal proteins have been
37	uncertain. Here we conduct a comprehensive molecular genetic analysis of the C. elegans DDX41
38	ortholog, SACY-1. Our results reveal that multiple sacy-1/DDX41 missense mutations, including the
39	R525H human oncogenic variant, exhibit antimorphic activity that likely compromises the function of
40	the spliceosome. The genomic consequences of SACY-1 depletion include splicing- splicing-

41 independent and splicing-dependent alterations in the transcriptome.

42

ABSTRACT

43	Mutations affecting spliceosomal proteins are frequently found in hematological malignancies,
44	including myelodysplastic syndromes and acute myeloid leukemia. DDX41/Abstrakt is a metazoan-
45	specific spliceosomal DEAD-box RNA helicase found to be recurrently mutated in inherited
46	myelodysplastic syndromes and in relapsing cases of acute myeloid leukemia. The genetic properties
47	and genomic impacts of disease-causing missense mutations in DDX41 and other spliceosomal
48	proteins have been uncertain. Here we conduct a comprehensive molecular genetic analysis of the C.
49	elegans DDX41 ortholog, SACY-1. Our results reveal general essential functions for SACY-1 in both
50	the germline and the soma, as well as specific functions affecting germline sex determination and cell
51	cycle control. Certain sacy-1/DDX41 mutations, including the R525H human oncogenic variant, confer
52	antimorphic activity, suggesting that they compromise the function of the spliceosome. Consistent with
53	these findings, sacy-1 exhibits synthetic lethal interactions with several spliceosomal components, and
54	biochemical analyses suggest that SACY-1 is a component of the C. elegans spliceosome. We used the
55	auxin-inducible degradation system to analyze the impact of SACY-1 on the transcriptome using RNA
56	sequencing. SACY-1 depletion impacts the transcriptome through splicing-independent and splicing-
57	dependent mechanisms. The observed transcriptome changes suggest that disruption of spliceosomal
58	function induces a stress response. Altered 3' splice site usage represents the predominant splicing
59	defect observed upon SACY-1 depletion, consistent with a role for SACY-1 as a second-step splicing
60	factor. Missplicing events appear more prevalent in the soma than the germline, suggesting that
61	surveillance mechanisms protect the germline from aberrant splicing.

62

INTRODUCTION

Mutations affecting components of the spliceosome are frequently found in hematological 63 64 malignancies, including myelodysplastic syndromes (MDS; Yoshida et al. 2011; reviewed by Yoshida 65 and Ogawa 2014; Coltri *et al.* 2019), which comprise a heterogeneous set of myeloid neoplasms 66 characterized by anemia and cytopenia that progress to acute myeloid leukemia (AML) to varying 67 degrees (Tefferi and Vardiman 2009). The genetic properties and genomic impacts of disease-causing 68 missense mutations in DDX41 and other spliceosomal proteins have been uncertain. Nonetheless, 69 mutations affecting spliceosomal components are predictive of poor clinical outcomes in AML patients 70 (Papaemmanuil et al. 2016). Exactly how mutations in spliceosomal components contribute to 71 malignancy is uncertain, but an attractive model is that aberrant splicing may interrupt tumor 72 suppressor activity. Importantly, genome sequencing data in patients is currently being used in the 73 clinic to generate personalized prognoses, with the idea of optimally targeting existing therapies and 74 generating new treatment strategies (Grinfeld et al. 2018). One potential therapeutic approach under 75 development is the discovery of splicing inhibitors (Effenberger et al. 2017; Kim and Abdel-Wahab 76 2017; DeNicola and Tang 2019). Although mutations affecting several spliceosomal proteins appear to 77 be beneficial to tumor cells, excessive splicing abnormalities are likely to be lethal to all cells. Splicing 78 inhibitors have been demonstrated to target tumor cells with splicing mutations by inducing excessive 79 splicing abnormalities, but cells with intact splicing machinery appear to be resistant to these agents 80 (Seiler *et al.* 2018). In fact, several new splicing inhibitors are currently in clinical trials. 81 The spliceosomal components frequently affected in MDS include the biochemically well-82 defined factors SF3B1, SRSF2, and U2AF1 (Yoshida et al. 2011: reviewed by Yoshida and Ogawa 83 2014). More recent studies have implicated DDX41 (Ding et al. 2012; Lewinsohn et al. 2015; 84 Polprasert et al. 2015; Cardoso et al. 2016; Li et al. 2016; Diness et al. 2018; reviewed by Maciejewski 85 et al. 2017), a DEAD-box RNA helicase highly conserved in metazoans, whose precise biochemical

function in the spliceosome is less well understood. DDX41 appears to be specifically recruited to the

catalytically active C complex (Jurica *et al.* 2002; Bessonov *et al.* 2008), which performs the second
step of splicing in which the 5' and 3' exons are ligated and an intronic lariat is released. DDX41 is
one of many spliceosomal proteins specific to metazoans and not found in budding yeast (Bessonov *et al.* 2008).

91 Whole genome sequencing studies suggest that DDX41 mutations are associated with 92 hematological malignancies that are considered to be different clinical entities. For example, 93 examination of clonal evolution of relapsed AML cases identified DDX41 as one of several genes 94 found to be mutated in secondary tumors but not the primary tumors, suggesting it plays a role as a 95 tumor suppressor and might be causative for disease progression (Ding et al. 2012). By contrast, 96 studies of familial acute myeloid leukemia syndromes suggest that preexisting germline DDX41 97 mutations in trans to newly arising somatic mutations cause the development of hematological 98 malignancies (Polprasert et al. 2015; Cardoso et al. 2016; Lewinsohn et al. 2016; Li et al. 2016). 99 Germline biallelic DDX41 missense mutations were recently reported in two siblings that exhibited 100 intellectual disability, psychomotor delays, and facial and skeletal dysmorphologies, with one sibling 101 presenting with childhood leukemia (Diness et al. 2018). Other work suggests that DDX41 might be a 102 multifunctional protein; in addition to its nuclear function in RNA splicing, it has been suggested to 103 function as a cytoplasmic DNA sensor in a signaling pathway in the cytoplasm that detects infecting 104 double stranded DNA and initiates an antiviral interferon response (Zhang et al. 2011; Parvativar et al. 105 2012; Stavrou et al. 2015, 2018; reviewed by Jiang et al. 2017). However, more recent work suggests 106 that cyclic GMP-AMP synthase (cGAS) functions as the major DNA sensor and is several orders of 107 magnitude more effective in inducing interferon beta synthesis than DDX41 (Sun *et al.* 2013). Two 108 studies, one of DDX41 and another of its Drosophila ortholog, Abstrakt, suggested a role in regulating translation of the cyclin-dependent kinase inhibitor p21^{WAF1/CIP1} (Peters *et al.* 2017) and the Inscuteable 109 110 protein (Irion et al. 2004), respectively, though the exact mechanism for these activities has not been 111 elucidated and indirect effects acting at the level of splicing were not addressed in these studies.

112 To better understand the highly conserved functions of *DDX41*, we undertook a comprehensive 113 molecular genetic analysis of its ortholog, sacy-1 in the nematode Caenorhabditis elegans. Our prior 114 studies identified the DEAD-box helicase SACY-1 as a negative regulator of oocyte meiotic 115 maturation functioning in the germline upstream of the TIS11 CCCH zinc-finger RNA-binding 116 proteins OMA-1 and OMA-2 (Kim et al. 2012). Genetic analysis also established roles for SACY-1 in 117 regulating the hermaphrodite sperm-to-oocyte switch and in preventing necrotic cell death of gametes. 118 Genetic experiments further suggested an essential role for sacy-1(+) in early embryos and larvae that 119 appeared to be maternally rescued. At the time of our original study, searchable databases of the 120 scientific literature had not yet annotated DDX41 (or its Drosophila ortholog, Abstrakt) as 121 spliceosomal components identified by proteomics. We therefore did not recognize that SACY-1 was 122 likely involved in splicing.

123 In this study, we undertook a comprehensive molecular genetic analysis of SACY-1's functions 124 in C. elegans. Our results are most consistent with an essential role for SACY-1 in spliceosome 125 function. Further, our genetic results reveal that certain sacy-1 mutations appear to confer a dosage-126 sensitive antimorphic activity, most consistent with the possibility that they compromise the function 127 of the spliceosome by perturbing the function of other spliceosomal proteins. The oncogenic R525H 128 mutation in human DDX41 was introduced into the C. elegans genome using CRISPR-Cas9 genome 129 editing and found to exhibit weak antagonistic activity. Consistent with these findings, sacy-1 exhibits 130 genetic interactions with associated spliceosomal components, and biochemical analyses suggest that 131 SACY-1 is a component of the *C. elegans* spliceosome. Depletion of SACY-1 in the germline or soma 132 was found to have major impacts on the transcriptome through splicing-independent and splicing-133 dependent mechanisms. Alterations in 3' splice site selection represent the most prevalent changes in 134 splicing patterns observed following SACY-1 depletion, consistent with its function as a component of 135 the spliceosomal C complex. Missplicing events are more prevalent upon SACY-1 depletion in the 136 soma than in the germline, leading us to suggest that surveillance mechanisms protect the germline

- 137 from aberrant splicing. The observed gene expression changes observed after SACY-1 depletion
- 138 suggest that perturbations of spliceosomal function might induce a stress response. Our results, taken
- together with a recent study of *sftb-1/SF3B1* (Serrat *et al.* 2019), highlight the potential of the *C*.
- 140 *elegans* system for examining the molecular genetic properties of disease-causing mutations affecting
- 141 highly conserved components of the spliceosome.

142

MATERIALS AND METHODS

143 C. elegans strains and genetic analysis

- 144 The genotypes of strains used in this study are reported in Supporting Information, Table S1. Genes 145 and mutations are described in WormBase (www.wormbase.org; Harris et al. 2013) or in the indicated 146 references. Culture and genetic manipulations were conducted at 20°C unless specified otherwise. The 147 following mutations were used: LGI-fog-1(q253ts), dpy-5(e61), gld-1(tn1478), unc-13(e51), unc-148 13(e1091), lin-41(n2914), lin-41(tn1541[gfp::tev::s-tag::lin-41]), sacy-1(tm5503), sacy-1(tn1385), 149 sacy-1(tn1479), sacy-1(tn1480), sacy-1(tn1481Mog), sacy-1(tn1482), sacy-1(tn1602), sacy-1(tn1603), 150 sacy-1(tn1604), sacy-1(tn1605), sacy-1(tn1606), sacy-1(tn1607), sacy-1(tn1608), sacy-1(tn1609), 151 sacy-1(tn1610), sacy-1(tn1611), sacy-1(tn1612), sacy-1(tn1615), sacy-1(tn1616), sacy-1(tn1617), 152 sacy-1(tn1632[3xFLAG::PreScission protease site::gfp::tev::s-tag::sacy-1]), sacy-153 1(tn1880[aid::gfp::tev::myc::sacy-1]), and sacy-1(tn1887); LGII-tra-2(e2020), ieSi57[eft-154 *3p::TIR1::mRuby::unc-54 3'UTR + Cb unc-119(+)], ieSi64[gld-1p::TIR1::mRuby::gld-1 3'UTR + Cb* 155 unc-119(+)]; LGIII-unc-119(ed3); LGIV-unc-24(e138), fem-3(e1996), and dpy-20(e1282); LGV-acy-156 4(ok1806), her-1(hv1y101), emb-4(sa44), unc-51(e369), and fog-2(oz40). The following 157 rearrangements were used: hT2[bli-4(e937) let-?(q782) qIs48] (I;III), tmC18[dpv-5(tmIs1236) +158 pmyo-2::mCherry] I (Dejima et al. 2018), mIn1[dpy-10(e128) mIs14] II, and tmC12[egl-9(tmIs1194)] 159 + pmvo-2:: Venus V (Dejima et al. 2018). The following transgenes were used: tnEx37[acv-4(+) +160 sur-5::gfp, tnEx159[gfp:sacy-1 + pDPMM0016B(unc-119(+))]. 161 For the analysis of genetic interactions between sacy-1(tn1481) and fem-3(e1996), non-Unc 162 non-Dpy non-GFP animals from sacy-1(tn1481)/hT2[bli-4(e937) let-?(q782) qIs48]; fem-
- 163 *3(e1996)/unc-24(e138) dpy-20(e1282)* were individually cultured and scored for germline phenotypes.
- 164 Following scoring, the *fem-3* genotype of each animal was scored by conducting PCR with primers
- 165 *fem-3* F2 and *fem-3* R2 and sequencing the products.

166	To map the cold-sensitive (15°C) and temperature-sensitive (25°C) phenotypes of sacy-
167	1(tn1480), 34 Unc non-Dpy recombinants were obtained from sacy-1(tn1480)/dpy-5(e61) unc-
168	13(e1091) heterozygotes. The recombinant chromosomes were bred to homozygosity and scored for
169	the presence or absence of the sacy-1(tn1480) mutation by conducting PCR with primers H27M09.1F1
170	and H27M09.1R4 and sequencing purified PCR products with primer H27M09.1F2. We found that 7
171	of the 34 recombinants contained sacy-1(tn1480) and were cold-sensitive and temperature-sensitive.
172	By contrast, 27 recombinants were <i>sacy-1(+)</i> and grew at 15°C and 25°C. These data indicate that
173	sacy-1(tn1480) mutation is inseparable from the cold-sensitive and temperature-sensitive phenotypes
174	(e.g., within ~0.06 map units). In addition, 32 Dpy non-Unc recombinants were selected. Interestingly,
175	all the homozygous recombinants were fertile at both 15°C and 25°C, including the 22 recombinants
176	that contained the sacy-1(tn1480) mutation. Although these dpy-5(e61) sacy-1(tn1480) recombinants
177	grew at 15°C and 25°C, they produced appreciable numbers of dead embryos and grew more slowly
178	than their sacy-1(+) counterparts. This result suggests that the dpy -5(e61) mutation suppresses the
179	cold-sensitive and temperature-sensitive phenotypes of sacy-1(tn1480). Previous work has shown that
180	mutant alleles of collagen genes can suppress temperature-sensitive mutations in other gene products
181	possibly by triggering a stress response (Levy et al. 1993; Maine and Kimble 1993; Nishiwaki and
182	Miwa 1998). That <i>dpy-5(e61)</i> suppresses <i>sacy-1(tn1480)</i> was further shown by constructing <i>dpy-</i>
183	5(e61) sacy-1(tn1480)/sacy-1(tn1480) unc-13(e1091) heterozygotes (n=30), of which 20 exhibited the
184	sacy-1(tn1480) sperm-defective phenotype at 25°C and 10 were fertile. Thus, dpy-5(e61) exhibits
185	semidominance for its effects on body morphology and for suppression of sacy-1(tn1480). To examine
186	the dominant Him phenotype of sacy-1(tn1480) and its interaction with sacy-1(tn1887), we compared
187	the percentage of males produced at 25°C by dpy-5(e61)/sacy-1(tn1480) unc-13(e1091) and dpy-
188	5(e61) sacy-1(tn1887)/sacy-1(tn1480) unc-13(e1091) heterozygotes.
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191 **RNA** interference

192 Genome-wide RNA interference (RNAi) screening employed the Ahringer feeding library (Kamath et 193 al. 2003) using the RNAi culture media described by Govindan et al. (2006) at 22°C. The empty vector 194 L4440 was used as a control. The identity of RNAi clones was verified by DNA sequencing. Gene-195 specific RNAi was performed by placing gravid hermaphrodites on RNAi medium seeded with 196 double-stranded RNA (dsRNA)-expressing E. coli (Timmons and Fire 1998). The gravid 197 hermaphrodites were immediately treated with 20% bleach to release the F1 embryos. Phenotypes 198 were assessed 3-4 days later. For quantification of phenotypes, sterility and gamete degeneration were 199 scored in the F1 generation, and embryonic lethality was scored in the F2 generation produced by the 200 RNAi-treated F1 animals.

201

202 Immunofluorescence, fluorescent labeling, and microscopy

203 Dissected gonads were fixed in 3% paraformaldehyde as described (Rose *et al.* 1997). Fixed gonads

were stained with rabbit anti-RME-2 antibody (Grant and Hirsh 1999; kindly provided by B. Grant,

205 Rutgers University, 1:50), a mixture of two purified mouse monoclonal anti-MSP antibodies (Kosinski

et al. 2005, each at 1:300), rabbit anti-phospho-histone H3 (Ser10) antibody (Millipore, 1:400).

207 Secondary antibodies were Alexa 488-conjugated donkey anti-rabbit antibodies (Jackson

208 ImmunoResearch, 1:500) and Cy3-conjugated goat anti-mouse antibodies (Jackson ImmunoResearch,

209 1:500). 4', 6-diamidino-2-phenylindole (DAPI) was used to detect DNA. DIC and fluorescent images

210 were acquired on a Zeiss motorized Axioplan 2 microscope with either a 40x Plan-Neofluar (numerical

aperture 1.3) or a 63x Plan-Apochromat (numerical aperture 1.4) objective lens using a AxioCam

212 MRm camera and AxioVision software (Zeiss).

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215

216 Genome editing

217	CRISPR-Cas9 genome editing used pRB1017 to express single guide RNA (sgRNA) under control of
218	the C. elegans U6 promoter (Arribere et al. 2014). The sequences of all oligonucleotides used are
219	listed in Table S2. To generate sgRNA clones, annealed oligonucleotides were ligated to BsaI-digested
220	pRB1017 plasmid vector, and the resulting plasmids were verified by Sanger sequencing. pDD162
221	served as the source of Cas9 expressed under control of the <i>eef-1A.1/eft-3</i> promoter (Dickinson <i>et al.</i>
222	2013). Indels were targeted to exon 2 of sacy-1 using sacy-1 sgRNA7 (pCS520). The injection mix
223	contained pCS520 (25 ng/µl), pDD162 (50 ng/µl), and Pmyo-2::tdTomato (4 ng/µl). sacy-1(tn1602-
224	tn1612) were recovered from injections into DG3913 lin-41(tn1541[gfp::tev::s-tag::lin-41]) and sacy-
225	1(tn1615–1617) were recovered from injections into the wild type (strain N2).
226	An N-terminal gfp fusion to endogenous sacy-1, sacy-1(tn1632[3xflag::PreScission protease
227	<i>site::gfp::tev::s-tag::sacy-1])</i> , was constructed using <i>sacy-1</i> sgRNA1 (pCS486) and a repair template
228	generated by conducting the PCR with oligonucleotide primers sacy-1 5HAF and sacy-1 3HAR, using
229	a gfp::sacy-1::tev::s-tag recombineered fosmid (SK212; Kim et al. 2012) as template. Genome editing
230	employed the <i>dpy-10</i> co-conversion method (Arribere <i>et al.</i> 2014). The injection mix contained pJA58
231	(7.5 ng/µl), AF-ZF-827 (500 nM), pCS486 (50 ng/µl), repair template (50 ng/µl), and pDD162 (50
232	$ng/\mu l)$ and was injected into wild-type worms. Correct targeting was verified by conducting PCR with
233	primer pairs GFP_7215 and H27M09.1_R5 and GFP_1094R and H27M09.1_seqF1 followed by DNA
234	sequencing.
235	An N-terminal auxin-inducible degron (aid) fusion to sacy-1, sacy-
236	1(tn1880[aid::gfp::myc::sacy-1]), was constructed using sacy-1 sgRNA1 and a repair template
237	generated by conducting the PCR with oligonucleotide primers sacy-1 AID5F and sacy-1 AID3R using
238	a wee-1.3::aid::gfp::myc clone (pCS575, C. Spike, unpublished results) as template. The injection mix
239	was prepared as described above and was injected into CA1352 worms. sacy-1 (sacy-
240	1(tn1880[aid::gfp::myc::sacy-1]) was identified by screening the progeny of 414 F1 Roller animals for

241	GFP fluorescence.	Correct targeting y	was verified by a	conducting PCI	R with primer	nairs GFP	R1 and
441	OFT HUDIUSUCHUU.		was vernieu uv v			Dans Ori	IXI anu

242 H27M09.1 F5 and GFP F1 and H27M09.1 R5 followed by DNA sequencing.

- The R525H mutation in DDX41 was imported into *C. elegans* (e.g., SACY-1[R534H]) using
- genome editing (Paix et al. 2014) with sacy-1 sgRNA11 and sacy-1 sgRNA12 and a single-stranded
- repair oligonucleotide (*sacy-1* GM1), which introduces the R534H mutation and two synonymous
- changes to alter the protospacer adjacent motif and to facilitate screening using an introduced AvaI
- restriction site. The injection mix contained pJA58 (7.5 ng/µl), AF-ZF-827 (500 nM), sacy-1 sgRNA11
- 248 (25 ng/µl), sacy-1 sgRNA12 (25 ng/µl), sacy-1 GM1 (500 nM), and pDD162 (50 ng/µl) and was

249 injected into wild-type worms. Edited loci were verified by PCR and DNA sequencing using primers

- sacy-1 seq F1 and sacy-1 seq R1.
- 251

252 Antibody production, purification, and western blotting

253 sacy-1 cDNA sequences were cloned into the E. coli expression vector pMal-c2 to create an inducible 254 fusion protein wherein maltose binding protein was fused to amino acids 411–578 of SACY-1 255 (MBP::SACY-1(411–578)). MBP::SACY-1(411–578) was column and gel-purified and used to 256 immunize rabbits. Immunizations and sera collection were performed using standard protocols 257 (Cocalico Biologicals, Inc., Reamstown, PA). Rabbit antibody (R217) was affinity purified and was 258 suitable in western blots with partially purified SACY-1 preparations. Hybridoma cell lines producing 259 anti-GFP monoclonal antibodies 12A6 and 4C9 (Sanchez et al. 2014) were obtained from the 260 Developmental Studies Hybridoma Bank and prepared as described (Tsukamoto et al. 2017). Proteins 261 were separated using NuPAGE 4-12% Bis-Tris gels (Invitrogen) and visualized after western blotting. 262 Blots were blocked with 5% nonfat dried milk. Primary antibodies used to detect proteins were 263 affinity-purified rabbit anti-SACY-1(411–578) R217 antibody (100 ng/ml) and rabbit anti-GFP 264 NB600-308 antibody (Novus Biologicals; 250 ng/ml). The secondary antibody used for western blots 265 was peroxidase-conjugated donkey anti-rabbit antibody (Jackson ImmunoResearch; 1:30,000).

266 Detection was performed using SuperSignal West Femto Maximum Sensitivity Substrate (Thermo267 Scientific).

268

269 SACY-1 tandem affinity purification

270 Tandem affinity purification of SACY-1 was conducted using strains DG4068 and DG4070 using 271 modifications of a previously described protocol (Tsukamoto et al. 2017). Immunopurified proteins 272 were precipitated with 16.7% trichloroacetic acid (TCA), washed with acetone at -20°C, and briefly 273 separated on a 12% NuPAGE Bis-Tris gel, stained with Colloidal Blue Staining Kit (Invitrogen). 274 Lanes were subdivided into eight gel slices and mass spectrometry was performed at the Taplin 275 Biological Mass Spectrometry Facility (Harvard Medical School) using an LTQ Orbitrap Velos Pro 276 ion-trap mass spectrometer (Thermo Fisher Scientific). Protein identification used the Sequest software 277 program (Thermo Fisher Scientific) to match the fragmentation pattern of tryptic peptides to the C. 278 *elegans* proteome. The data were filtered to a 1–2% peptide false discovery rate. File S1 reports the 279 mass spectrometry results and the additional filtering criteria for identifying non-specific interactions.

280

281 RNA sequencing

The auxin-inducible degradation system (Zhang *et al.* 2015) was used to deplete SACY-1 using strain backgrounds in which TIR1 was expressed in the germline (CA1352) or soma (CA1200).

Experimental (DG4700 and DG4703) and control strains (CA1352 and CA1200) were grown on

285 peptone-enriched nematode growth medium with NA22 as a food source. Embryos were isolated by

alkaline hypochlorite treatment (20% bleach and 0.5 N NaOH), washed in M9 buffer and allowed to

hatch overnight in the absence of food. For each of three biological replicates, 60,000 L1-stage larvae

were cultured on two 150- by 15-mm petri dishes containing peptone-enriched medium with OP50.

289 The worms were grown to the young adult stage and harvested by washing off the plates with M9 and

then placed on fresh plates containing peptone-enriched medium and 2 mM auxin seeded with OP50.

Plates were cultured in the dark at 20°C for 24 hours. The worms were then harvested and washed with
M9 repeatedly to reduce the presence of *E. coli*. Total RNA was isolated using TRIzol LS Reagent
(Invitrogen, Carlsbad, CA) and the RNAeasy Micro Kit (QIAGEN, Valencia, CA). Poly(A)+ RNA
was selected from 1 µg of total RNA using the NEBNext Ultra Kit (New England Biolabs, Ipswitch,
MA). Libraries were prepared and sequenced by Genewiz (South Plainfield, NJ). Paired-end reads of
150 base pairs (bp) were obtained on an Illumina HiSeq4000 instrument with an average depth greater
than 31 million reads per sample.

298

299 *Bioinformatics*

300 After trimming adapters with Trim Galore (v0.6.0) and cutadapt (v1.18), reads were assessed for 301 quality with FastQC (v0.11.8), mapped to the WBcel235/ce11 genome with STAR (v2.7.2a) guided by 302 gene annotations defined in Ensembl (release 97) and sorted and indexed with samtools (v1.7). Gene-303 level abundance was estimated for Ensembl defined annotations using the featureCounts function in 304 the Bioconductor package Rsubread (v1.28.1). An average of 28 million high-quality (MAPQ > 55) 305 reads mapped to annotated genes within each sample. Principal component analysis and inspection of 306 5' vs 3' read coverage indicated that one soma control sample (CA1200-2) contained degraded RNA 307 and was excluded from further analysis. Differential gene expression of Ensembl defined genes was 308 determined using DESeq2 (v1.26.0). P values were adjusted for multiple test correction using 309 Benjamini–Hochberg procedure. The fold change, adjusted p values, the mean number of counts across 310 samples and the number of complementary DNA fragments per kilobase of transcript per million 311 mapped reads (FPKM) were used to define differentially expressed genes. Gene ontology (GO) data 312 were obtained from WormBase release WS273 and analyzed taking length bias into account using the 313 Goseq (v1.38.0) package. Novel transcripts in each of the high quality samples and in the previously 314 published GSE57109 (Ortiz et al. 2014) dataset were identified using StringTie (v2.0.4) and merged 315 together with the Ensembl annotations to generate a comprehensive annotation set. These annotations

- 316 were used with RMATS (v4.0.2 turbo) to determine statistically significant differences for splicing
- 317 events between conditions [expressed as false discovery rates (FDRs)]. Coverage data was visualized
- 318 with Gviz (v 1.30.0). A custom R script with details for the analysis and figure generation is available
- 319 at https://github.com/micahgearhart/sacy1.
- 320
- 321 Data availability
- 322 Strains and reagents are available upon request. RNA sequencing data have been deposited in NCBI's
- 323 Gene Expression Omnibus and are accessible through accession number XXXX.

324

RESULTS

325 Synthetic lethal interactions between a sacy-1 reduction-of-function allele and several genes

326 encoding spliceosomal proteins

327 In prior studies, we recovered reduction-of-function (rf) sacy-1 mutant alleles as suppressors of acy-4 328 sterility (Kim *et al.* 2012). ACY-4 is an adenylate cyclase that functions in the gonadal sheath cells to 329 promote oocyte growth and meiotic maturation (Govindan et al. 2009; Nadarajan et al. 2009). acy-4 330 null mutants are sterile, whereas sacy-1(rf); acy-4(0) mutants are fertile, and this suppression was 331 shown to involve reduced sacy-1 function in the germline (Kim et al. 2012). sacy-1(rf) mutants were 332 also found to suppress the self-sterility of *fog-2* null mutants that results from a failure to produce 333 sperm. A sperm-to-oocyte switch governs hermaphrodite germline sex determination. In wild-type 334 hermaphrodites, the first differentiating germ cells produce sperm in the L4 larval stage, whereas, later 335 differentiating germ cells exclusively produce oocytes in the adult stage. The suppression of fog-2 336 sterility suggested that *sacy-1* has a function to promote the oocyte fate. Analysis of the *sacy-1(tm5503)* 337 deletion allele, which is a likely null allele (see below), indicated that sacy-1 also functions to prevent 338 necrotic degeneration of sperm and oocytes and is required for fertility of hermaphrodites and males 339 (Kim et al. 2012).

340 To better understand the function of *sacy-1* in multiple germline processes, we conducted a 341 genome-wide RNAi screen for loci that enhance sacy-1 mutant phenotypes, such as sterility or 342 embryonic lethality. Specifically, we screened for loci which caused severe phenotypes when knocked 343 down by RNAi in the sacy-1(tn1385rf) genetic background but not in the wild type. We screened 344 18.101 RNAi clones from the Ahringer RNAi library and identified five clones that enhanced sacv-1 345 mutant phenotypes when they are knocked down (Table 1). The five RNAi clones target the transcripts 346 of three genes (Table 1): mog-2 (one clone), Y111B2A.25 (one clone), and emb-4 (three clones). To test 347 whether sacy-1 expression and/or localization is affected by RNAi of mog-2, Y111B2A.25, or emb-4, 348 we conducted RNAi of these genes in *sacy-1(tm5503)* mutant animals expressing the rescuing

<i>gfp::sacy-1</i> transgene (<i>tnEx159</i>). In no case did we observe that an RNAi treatment altered the
expression or localization of the GFP::SACY-1 transgene; the expression level and predominant
nuclear localization of GFP::SACY-1 after RNAi was similar to that of the control animals (S. Kim
and D. Greenstein, unpublished results). This result suggests that the RNAi treatments enhance sacy-
1(tn1385) mutant phenotypes through effects independent of SACY-1 expression and localization.
mog-2(RNAi) induces a higher penetrance of sterility, gamete degeneration, and embryonic
lethality in the sacy-1(tn1385) mutant genetic background in comparison to the wild type (Table 1;
Figure S1). mog-2 encodes the U2 snRNP protein A' (Zanetti et al. 2011), which is a constitutive
component of the spliceosome (Jurica et al. 2002; Bessonov et al. 2008, 2010; Herold et al. 2009).
Similarly, Y111B2A.25(RNAi) specifically enhances the penetrance of multiple sacy-1 mutant
phenotypes, including sterility, gamete degeneration, and embryonic lethality (Table 1; Figure S1).
Y111B2A.25 is annotated as a pseudogene (Agarwal et al. 2010; www.wormbase.org). Y111B2A.25 is
part of an operon, and the expressed sequence tag (EST) data show that the Y111B2A.25 locus is
transcribed, but the transcript lacks protein-coding ability. In C. elegans ~40 bp of sequence identity is
sufficient to induce off-target RNAi effects (Rual et al. 2007). Use of the Basic Local Alignment
Search Tool (BLAST) indicates that Y111B2A.25(RNAi) might target the cacn-1 locus, which encodes
a spliceosomal protein and shares ~200 bp of sequence identity with Y111B2A.25. To test whether the
enhanced sterility induced by Y111B2A.25(RNAi) in the sacy-1(tn1385) genetic background might be
explained by an off-target effect to cacn-1, we conducted cacn-1(RNAi) and found that the cacn-
1(RNAi) induces complete sterility in both the sacy-1(tn1385) and wild-type animals (Table 1 and
Figure S1). Interestingly, under the <i>cacn-1(RNAi)</i> condition, the <i>sacy-1(tn1385)</i> animals show
additional phenotypes, such as high penetrance of a protruding vulva (Pvl) phenotype and slow growth
compared to the wild type, suggesting a genetic interaction between <i>cacn-1</i> and <i>sacy-1</i> that might be

374	1(tn1385rf) genetic background but not in the wild type through weaker interference with cacn-1. To
375	test this possibility, we systematically reduced the efficacy of the cacn-1(RNAi) response by serially
376	diluting the <i>cacn-1(RNAi)</i> -inducing bacteria with bacteria containing the empty vector control (L4440).
377	Consistent with the possibility that Y111B2A.25(RNAi) targets cacn-1, limiting the efficacy of the
378	cacn-1(RNAi) response revealed specific enhancement of sterility, gamete degeneration, and
379	embryonic lethality in the sacy-1(tn1385) genetic background (Table S3). Notably, the response to
380	limited <i>cacn-1(RNAi)</i> exhibited by <i>sacy-1(tn1385)</i> animals was remarkably similar to their response to
381	Y111B2A.25(RNAi) (Table S3). The human and Drosophila orthologs of CACN-1 have been identified
382	as components of spliceosomal C complexes (Jurica et al. 2002; Bessonov et al. 2008, 2010; Herold et
383	al. 2009; Fica et al. 2019). Like DDX41/Abstrakt, Cactin is recruited to the C complex of the
384	spliceosome.
385	In addition to mog-2 and Y111B2A.25, we identified three different RNAi clones targeting the
386	emb-4 locus as strong enhancers of the sacy-1(tn1385) sterility and gamete degeneration phenotypes
387	(Table 1; Figure S1). emb-4 encodes a nuclear protein orthologous to human
388	Aquarius/IBP160/KIAA0560/fSAP164, an intron-binding spliceosomal protein with a helicase-like
389	domain (Sam et al. 1998; Jurica et al. 2002; Bessonov et al. 2008; Herold et al. 2009; De et al. 2015;
390	Haselbach et al. 2018). To extend these RNAi results, we examined genetic interactions between sacy-
391	1 and emb-4, employing the emb-4(sa44) reduction-of-function allele. When combined with the sacy-
392	1(tm5503) null allele, we observed enhancement of lethal vulval rupture and protruding vulva (Pvl)
393	phenotypes in sacy-1(tm5503); emb-4(sa44) double mutants (Table 2A). We also observed
394	enhancement of these phenotypes in sacy-1(tn1385); emb-4(sa44) double mutants, which were derived
395	from sacy-1(tn1385)/+; emb-4(sa44) parents (Table 2A). Interestingly, the F1 progeny of fertile sacy-
396	1(tn1385); emb-4(sa44) homozygous adults exclusively produced dead embryos or arrested L1-stage
397	larvae, unlike each of the single mutants, which were highly fertile (Table 2B). Taken together these

genetic interactions between *sacy-1* and three genes encoding spliceosomal proteins suggest that
 multiple *sacy-1* mutant phenotypes might result from compromised functions of the spliceosome.

400

401 SACY-1 is a component of the C. elegans spliceosome

402 To characterize SACY-1-associated proteins, we conducted tandem affinity purifications using strains 403 in which we used CRISPR-Cas9 genome editing to insert 3xFLAG and eGFP affinity tags at the 404 SACY-1 N-terminus, separated by a PreScission protease recognition sequence (Figure S2). The 405 resulting sacy-1(tn1632[3x flag::PreScission::gfp::tev::s-tag::sacy-1]) strain was viable and fertile 406 and exhibited no apparent germline or somatic defects. Although 3xFLAG::GFP::SACY-1 is expressed 407 in all cells, it is particularly abundant in the female germline. Thus we conducted purifications from 408 protein lysates prepared from adult animals in which the germline was feminized (experiment I) and 409 also from adult hermaphrodites (experiment II). In both experiments, we found that 410 3xFLAG::GFP::SACY-1 associated with 55 proteins defined as spliceosomal proteins in other systems 411 (Table 3; Jurica et al. 2002; Bessonov et al. 2008). The spliceosomal proteins identified in our 412 genome-wide RNAi screen for sacy-1 enhancers (MOG-2, EMB-4, and CACN-1), were very well 413 represented in our purifications (~35–56% peptide coverage; Table 3). We also detected 9 additional 414 spliceosomal proteins in the purification from the female but not the hermaphrodite genetic 415 background (Table S4), but this might be a consequence of the fact that more protein extract was used 416 in that experiment. We also detected 28 other proteins in our tandem affinity purifications (Table S5). 417 Homologs of several of these factors have been implicated in the regulation of RNA splicing, including 418 NRDE-2 (Jiao et al. 2019), CIR-1 (Maita et al. 2005; Kasturi et al. 2010), and CDK-12 (Rodrigues et 419 al. 2012). These biochemical studies, taken together with the results from the genome-wide RNAi 420 screen, suggest that both specific and pleiotropic defects conferred by *sacv-1* mutant alleles might 421 result from spliceosomal defects.

422

423 A sacy-1 reduction-of-function mutation enhances the Tumorous phenotype of a gain-of-function 424 glp-1/Notch allele

425 Prior work showed that mutational or RNAi treatments affecting the function of multiple spliceosomal 426 components could enhance weak gain-of-function (gf) mutations in *glp-1/Notch*, resulting in the 427 ectopic proliferation of undifferentiated germ cells in the proximal gonad arm (Mantina et al. 2009; 428 Kerins et al. 2010; Wang et al. 2012), which is referred to as a proximal proliferation or Tumorous 429 phenotype. Thus, we constructed double mutants between the sacy-1(tn1385) mutation and the glp-430 *l(ar202gf)* mutation (Pepper *et al.* 2003). When analyzed in the young adult stage at 15°C (40 hrs 431 post-L4), very few *glp-1(ar202*gf) adult hermaphrodites (~0.8%) were observed to exhibit a proximal 432 proliferation phenotype with undifferentiated germ cells in the proximal gonad arm (Table 4). By 433 contrast, many sacy-1(tn1385rf); glp-1(ar202gf) adults (~51%) exhibited a Tumorous phenotype 434 (Table 4). This phenotype was not observed in *sacy-1(tn1385rf)* single mutants (Table 4). This result is 435 consistent with the idea that the sacy-1(tn1385rf) mutation, though homozygous viable and fertile 436 (brood size ~350; Kim et al. 2012), compromises the function of the spliceosome, as assessed in a 437 sensitized genetic background.

438

Reduction-of-function sacy-1 mutations in C. elegans affect highly conserved residues in the DEAD-box and helicase domains

To better understand the functions and activities of the highly conserved SACY-1/DDX41 protein (Figure 1B), we conducted forward genetic screens for new *sacy-1* mutations, taking advantage of the fact that reductions of *sacy-1* function by mutation or RNAi can suppress the self-sterility of *fog-2* null mutations (Kim *et al.* 2012), which is caused by a failure to produce sperm (Schedl and Kimble 1988). Thus, we conducted a non-complementation screen for new mutations that enable fertility in trans to the *sacy-1(tn1385)* reduction-of-function allele in the *fog-2(oz40)* genetic background (Figure S3). In a screen of 15,577 haploid genomes, we isolated five new *sacy-1* missense alleles (*tn1479–tn1483*;

448 Figure 1; see Table 5 for a list of all *sacy-1* alleles central to this work and their properties). All the 449 sacy-1 missense mutations isolated thus far in C. elegans alter highly conserved amino acids, and 450 several of these mutations are nearby or in subdomains of the DEAD-box affected by DDX41 451 mutations found in human neoplasms (Figure 1B). The sacy-1 missense alleles were modeled onto the 452 crystal structures of DEADc and the HELICc domains of DDX41 (Schütz et al. 2010; Omura et al. 453 2016) and found likely to be surface accessible (Figure 1C), suggesting that the mutant alleles might 454 interfere with the function of other protein components of the spliceosome. Consistent with this idea, 455 the non-complementation screen resulted in the isolation of three novel sacy-1 alleles (tn1479, tn1480, 456 and *tn1481*), which appear to confer antagonistic activities, likely at the level of the spliceosome (see 457 below). 458 Similar to the *sacy-1(tm5503*) deletion allele which removes exons 2 and 3 and a portion of 459 exon 4 (Figure 1A; Kim et al. 2012), sacy-1(tn1479) homozygous hermaphrodites were sterile and 460 displayed the gamete degeneration phenotype, but with reduced penetrance (Table 6; Figure S4). 461 Consistent with the idea that sacy-1(tm5503) defines the null phenotype, an antibody specific to a 462 portion of the DEAD-box domain downstream of the *tm5503* deletion (residues 411–578) fails to 463 detect a protein product in extracts from sacy-1(tm5503) adults (Figure S2). To further define the sacy-464 1 null phenotype, we used CRISPR/Cas9 genome editing to generate indels upstream of the DEAD-465 box-encoding regions by targeting Cas9 double-strand DNA breaks to exon 2 with an efficient sgRNA. 466 We generated *sacy-1* indels in both wild-type as well as *lin-41(tn1541[gfp::lin-41])* hermaphrodites, 467 the latter serving to provide a marker for oocyte development (Spike *et al.* 2014a,b). In these 468 experiments, we generated 14 new sacv-1 alleles (tn1602-tn1612 and tn1615-tn1617). Of these, 13 469 displayed the gamete degeneration phenotype, again consistent with this representing the null 470 phenotype. Not surprisingly, GFP::LIN-41 levels declined and the protein became undetectable as 471 oocytes degenerated (D. Greenstein, unpublished results). One of the new candidate null alleles, sacy-472 *1(tn1615)*, was sequenced and found to result from a 10 bp deletion at the end of exon 2, which is

473	predicted to introduce a stop codon prior to the DEAD-box domain, consistent with a null mutation
474	(Figure 1B). Among the CRISPR-Cas9-induced alleles, <i>sacy-1(tn1617)</i> was exceptional in that it was
т/т	(Figure TD). Among the CRIST R-Casy-induced ancies, sucy-f(m1017) was exceptional in that it was
475	homozygous viable and fertile, though slow growing, despite the fact that the deletion removes the
476	initiation codon of sacy-1 (Figure 1A). This exceptional allele might utilize an alternative start codon
477	just prior to the DEAD-box domain, although this possibility was not explored. Since null mutations in
478	<i>sacy-1</i> result in hermaphrodite sterility, there is the possibility that maternal <i>sacy-1(+)</i> activity
479	contributes to the development of the germline and soma. Indeed, when the gamete degeneration
480	phenotype is delayed through germline feminization, the mating of sacy-1(tm5503) null females to
481	wild-type males produces embryos that arrest prior to morphogenesis (Kim et al. 2012).
482	One notable difference between sacy- $1(tn1479)$ and the sacy- $1(tm5503)$ null allele is that the
483	majority of <i>sacy-1(tn1479</i>) adult hermaphrodites die by vulval rupture at 20°C (Table 6; Figure S4).
484	This phenotype is only observed in a small minority of <i>sacy-1(tm5503)</i> hermaphrodites at 20°C but
485	becomes the predominant phenotype at 15°C (Table 6). This observation suggests that sacy-1(tn1479)
486	is more severe than a null allele, as might happen if the SACY-1[G504E] product is nonfunctional but
487	also antagonizes a protein complex containing SACY-1, likely the spliceosome.
488	Unlike the sacy-1(tm5503) null allele, which is sterile, three of the newly isolated mutations
489	(tn1480, tn1482, and tn1483) are self-fertile as homozygotes at 20°C similar to the sacy-1(tn1385rf)
490	mutant. However, sacy-1(tn1480) and sacy-1(tn1483) homozygotes exhibited additional defects; both
491	appeared to grow more slowly than the wild type at this temperature. In addition, sacy-1(tn1483) adult
492	hermaphrodites had smaller gonad arms, suggesting effects on germline proliferation (T. Tsukamoto
493	and D. Greenstein, unpublished results). Finally, sacy-1(tn1480) confers temperature-dependent gain-
494	of-function pleiotropic defects that will be described below.
495	

- 496
- 497

498 Novel sacy-1 mutant alleles appear to antagonize essential functions of the spliceosome 499 A recessive gain-of-function sacy-1 mutation masculinizes the hermaphrodite germline: In the non-500 complementation screen for sacy-1 mutant alleles, we isolated a novel sacy-1 allele, tn1481, with a 501 masculinization of germline (Mog) phenotype (Table 6; Figure 2). All sacy-1(tn1481) homozygous 502 hermaphrodites produce excess numbers of sperm but no oocytes (n=125). Staining of dissected 503 gonads from adults showed that whereas all wild-type hermaphrodite gonad arms examined (n=64)504 expressed both the major sperm protein (MSP) and the RME-2 oocyte yolk receptor, all sacy-505 1(tn1481) gonad arms (n=178) expressed only MSP but not RME-2 (Figure 2). In our non-506 complementation screen (Figure S3), we also isolated a *gld-1* Mog allele, *tn1478*, as a dominant 507 suppressor of fog-2(oz40). gld-1(tn1478) results from the same G248R amino acid substitution reported for the *gld-1(q93)* Mog allele (Francis *et al.* 1995a,b; Jones and Schedl 1995). Thus it was 508 509 important to determine whether the sacy-1(tn1481[P222L]) mutation was the cause of the Mog 510 phenotype. This was ascertained by crossing a GFP::SACY-1 transgene (*tnEx159*) into the sacy-511 *1(tn1481)* genetic background. We found that all *sacy-1(tn1481)*; *tnEx159* [*gfp::sacy-1+unc-119(+)*] 512 hermaphrodites (n=30) produced oocytes and sperm and were self-fertile. This result established that 513 the P222L mutation in SACY-1 causes the Mog phenotype. 514 The suppression of *fog-2* sterility by reduction-of-function *sacy-1* mutations is consistent with

515 sacy-I(+) possessing a function that promotes the oocyte fate; this function is non-essential, however, 516 because the strongest loss-of-function sacy-1 alleles are able to produce oocytes, which nevertheless 517 undergo necrotic degeneration. Thus, the sacy-l(tn1481) Mog phenotype suggests this mutant allele, 518 although recessive, possesses an activity antagonistic to this oocyte-promoting function in the sperm-519 to-oocvte switch. To genetically characterize *sacv-1(tn1481)* further, we analyzed the phenotype of 520 sacv-1(tn1481)/sacv-1(tm5503 null) heterozygotes. Whereas all sacv-1(tn1481) homozygotes (n=50) 521 displayed a Mog phenotype, all sacy-1(tn1481)/sacy-1(tm5503) heterozygotes (n=48) produced both 522 oocytes and sperm and were self fertile. This result suggests that the *sacy-1(tn1481)* Mog phenotype is

dosage sensitive and that the mutant allele appears to be a recessive gain-of-function mutation that
might antagonize proteins that normally function with SACY-1, likely other components of the
spliceosome as discussed below.

526 In C. elegans, a genetic hierarchy controls germline sex determination (Figure 3). The failure of 527 sacy-1(RNAi) to suppress the sterility of the dominant strongly feminizing tra-2(e2020) mutation, 528 which deletes GLD-1 binding sites within the *tra-2 3'-UTR*, was interpreted in the context of a model 529 in which sacy-1(+) promotes the oocyte fate in opposition to fog-2 and gld-1 at the level of tra-2 (Figure 3; Kim et al. 2012). Because the evaluation of potential interactions between sacy-1 and tra-2 530 531 relied on *sacy-1(RNAi)*, there was the concern that this treatment reduced but did not eliminate the 532 function of sacy-1. Thus, we reevaluated the interaction between tra-2 and sacy-1 genetically. In the 533 first approach, we combined the *sacy-1(tm5503)* null allele with *tra-2(e2020)*. We analyzed the sexual 534 fate of the germline by staining dissected gonads from adult animals with oocyte (RME-2) and sperm 535 (MSP) markers. Whereas all wild-type gonad arms examined (n=30) expressed RME-2 and MSP, all 536 gonad arms of sacy-1(tm5503); tra-2(e2020) animals (n=26) expressed only RME-2 and not MSP 537 (Figure 4). This result is consistent with the model in which *sacv-1* promotes the oocyte fate by 538 promoting the function of *tra-2*. Although the germlines of *sacy-1(tm5503)*; *tra-2(e2020)* adults were 539 feminized, oocytes underwent meiotic maturation constitutively, consistent with the finding that sacy-1 540 is a negative regulator of meiotic maturation (Kim et al. 2012). sacv-1(tm5503); tra-2(e2020) animals 541 did however exhibit a highly penetrant ovulation defect, which caused endomitotic oocytes to 542 accumulate in the gonad arm (Emo phenotype; Figure 4). 543 To extend these observations, we examined germline sexual fates in dissected gonads from 544 sacv-1(tn1481Mog): tra-2(e2020) adults. Whereas all wild-type gonad arms examined (n=21) 545 expressed MSP and contained sperm, none of the sacy-1(tn1481Mog); tra-2(e2020) gonad arms 546 (n=37) expressed MSP or contained sperm. We noted that the Emo phenotype was less penetrant in 547 sacy-1(tn1481Mog); tra-2(e2020) gonad arms (46% penetrance). These results are consistent with the

- possibility that *sacy-1(+)* promotes the oocyte fate by promoting the function of *tra-2* in the germline
- and suggests that *sacy-1(tn1481*Mog) may interfere with this function.
- 550 Interestingly, recessive loss-of-function mutations in six genes, mog-1-6, cause a Mog 551 phenotype and encode spliceosomal components (Graham and Kimble 1993; Graham et al. 1993; Puoti 552 and Kimble 1999, 2000; Belfiore et al. 2004; Zanetti et al. 2011). Mutation and RNAi depletion of 553 many splicing factors have been observed to result in a Mog phenotype, suggesting that the germline 554 sex determination process is particularly sensitive to disruptions in RNA splicing (Mantina et al. 2009; 555 Kerins et al. 2010; Wang et al. 2012; Novak et al. 2015). Prior studies focusing on the C. elegans soma 556 were interpreted in the context of a model in which mog-1-mog-6 might function at the level of fem-3 557 through 3'UTR-dependent translational regulation (Gallegos et al. 1998; Figure 3); however, the 558 experiments in that study did not address the regulation of *fem-3* in the germline. We previously 559 showed that sacy-1(tm5503); fem-3(e1996) adult XX animals had feminized germlines (Kim et al. 560 2012). To examine the genetic relationship between sacy-1 and fem-3 further, we generated sacy-561 *1(tn1481Mog); fem-3(e1996)* double mutants. We observed that 92% (n=23) of *sacy-1(tn1481Mog);* 562 fem-3(e1996) animals were feminized. Mating of sacv-1(tn1481Mog); fem-3(e1996) females (n=29) to 563 wild-type males resulted in the production of embryos that failed to hatch (n=4140, 99.8%) or arrested 564 as larvae (n=7, 0.2%). This result indicates that two copies of sacy-1(tn1481) in the maternal germline, 565 but not one [e.g., sacy-1(tn1481)/sacy-1(tm5503) heterozygotes are fertile] are incompatible with 566 embryonic development. We found that 8% (n=2) of sacy-1(tn1481Mog); fem-3(e1996) animals 567 (n=25) produced oocytes and sperm and a few dead embryos (one of these animals produced sperm in 568 one gonad arm but not the other). This result suggests that the sacv-1(tn1481) mutation can promote 569 sperm development independent of zygotic *fem-3(+)* activity, consistent the genetic epistasis pathway 570 (Figure 3; Zanetti and Puoti 2013). We did observe that a reduction in *fem-3* dosage could suppress the 571 sacy-1(tn1481) Mog phenotype (n=60). Specifically, whereas 65% (n=39) of sacy-1(tn1481); fem-572 3(e1996)/+ animals were Mog, exclusively producing sperm, 33% (n=20) produced sperm and

573	oocytes, and one animal (2%) was feminized. Since SACY-1 genetically and biochemically interacts
574	with components of the spliceosome, we suggest that the sacy- $1(tn1481)$ mutation antagonizes
575	functions of the spliceosome needed for germline sex determination and oogenesis.

576

577 The gain-of-function sacy-1(tn1480) allele confers multiple pleiotropic phenotypes in a temperature-578 *dependent manner:* Interestingly, *sacy-1(tn1480)* displayed both cold-sensitive (15°C) and 579 temperature-sensitive (25°C) defects. At 15°C, sacy-1(tn1480) homozygotes (n=55), which were the 580 offspring of heterozygous parents grown at 15°C, laid dead eggs and produced arrested larvae (91%) or 581 produced very few progeny (9%). At 25°C, sacy-1(tn1480) hermaphrodites were sterile and produced 582 abnormal and fertilization-defective sperm (Figure 5). sacy-1(tn1480) hermaphrodite sterility is 583 rescued by mating with wild-type males at 25°C. In addition to producing oocytes, some sacy-584 1(tn1480) adult hermaphrodites continued to produce sperm, as swollen germ cells specified in the 585 male fate were detected distal to the loop region in 42% of gonad arms examined (Figure 5; n=12). 586 Thus, in addition to the other phenotypes it confers, the sacy-1(tn1480) mutation perturbs the sperm-to-587 oocyte switch at 25°C. We also observed that sacy-1(tn1480) conferred a dominant high-incidence of 588 males (Him) phenotype; *sacy-1(tn1480)/+* heterozygous hermaphrodites produced 1.6% male progeny 589 (n=3759), as compared to 0.1% for the wild-type control (n=6044; p<0.01, Fisher's exact test). Genetic 590 mapping showed that the temperature-dependent pleiotropic defects were inseparable from the *sacv*-591 1(tn1480) mutation (see Materials and Methods). Like sacy-1(tn1481), the defects conferred by sacy-592 1(tn1480) were dosage sensitive. At 25°C, nearly all sacy-1(tn1480)/sacy-1(tm5503) heterozygotes 593 (99%, n=88) were fertile (the average brood size was 132 ± 64 , n=34); however, the majority of sacy-594 1(tm5503) homozygotes (85.3%, n=231) they produced burst as adults (Table 6). This result suggests 595 that maternal sacy-1(tn1480) activity can antagonize the spliceosome in the absence of zygotic sacy-596 1(+) function. By these genetic criteria, sacy-1(tn1480) exhibits recessive and weakly dominant gain-597 of-function properties, depending on the phenotype examined.

598 The oncogenic DDX41 R525H mutation confers weak antagonistic activity in C. elegans

599	The R525H mutation in DDX41 has been reported in myeloid leukemias both as newly arising somatic
600	mutations specific to the neoplastic cells, as well as inherited germline mutations (Polprasert et al.
601	2015; Lewinsohn et al. 2016; Sébert et al. 2019). Thus, it was of interest to examine the impact of the
602	analogous mutation (R534H) on sacy-1 function. Interestingly, a substitution at the adjacent amino
603	acid (G533R) results in the sacy-1(tn1385) reduction-of-function mutation (Figure 1). Thus, we used
604	genome editing to introduce the R534H mutation in the C. elegans genome (see Materials and
605	Methods). By several criteria, sacy-1(tn1887[R534H]) homozygotes were indistinguishable from the
606	wild type. Neither did sacy-1(tn1887) suppress acy-4 sterility (n=140; sacy-1(tn1887); acy-4(ok1806)
607	brood size was 1.5 ± 2.3) nor did it suppress <i>fog-2</i> sterility (n=48). All unmated <i>sacy-1(tn1887); fog-</i>
608	2(oz40) gonad arms examined (n=96) exhibited stacked oocytes, which indicates that the sacy-
609	1(tn1887) mutation does not derepress meiotic maturation in the absence of sperm, like strong
610	reduction-of-function mutations do. Further, the sacy-1(tn1887[R534H]) did not enhance the
611	Tumorous phenotype of a gain-of-function <i>glp-1/Notch</i> allele (Table 4). The brood size of <i>sacy</i> -
612	l(tn1887) (204 ± 37; n=20) at 25°C was indistinguishable from that of the wild type (202 ± 62, n=30;
613	p>0.8, two-sample Z-test). Also, the incidence of males (0.2%, n=4087) was similar to that observed in
614	the wild type (0.1%, n= 6044). When placed in trans to the sacy-1(tm5503) null mutation, sacy-
615	1(tn1887)/sacy-1(tm5503) heterozygotes were found to be fertile at all temperatures examined (15°,
616	20°, and 25°C (n=104)). However, we did observe that sacy-1(tn1887) significantly enhanced the
617	dominant Him phenotype of sacy-1(tn1480) (p<0.05, Fisher's exact test); sacy-1(tn1887)/sacy-
618	1(tn1480) heterozygotes produced 5.7% males at 25°C (n=1483) as compared to 1.6% for the +/sacy-
619	1(tn1480) control (n=3759). The brood size of sacy-1(tn1887)/sacy-1(tn1480) heterozygotes at 25°C
620	$(74 \pm 65, n=20)$ was also significantly lower than that of the +/sacy-1(tn1480) control (188 ± 68, n=20;
621	p<0.001, two-sample Z-test). Because, <i>sacy-1(tn1480)/sacy-1(tm5503)</i> heterozygotes do not exhibit an

enhanced Him phenotype at 25°C (0.7% males, n=4493), the *sacy-1(tn1887*[R534H]) mutation appears
to possess a weak antagonistic activity.

624

625 *Impact of sacy-1 on the transcriptome*

Depletion of SACY-1 using the auxin-inducible degradation system: RNA sequencing studies using human patient samples suggested a role for DDX41 in splice site selection for a small number of

human genes (Polprasert *et al.* 2015). Thus, we sought to address the impact of SACY-1 on the

629 transcriptome by exploiting the power of the *C. elegans* system for transcriptomics under genotypically

and experimentally well-controlled conditions. We chose to use the auxin-inducible degradation

631 system (Zhang *et al.* 2015) to acutely deplete SACY-1 in the adult stage and thus avoid indirect

632 impacts on the transcriptome arising as a developmental consequence of strong loss-of-function

633 phenotypes (e.g., germline degeneration and cell fate changes). Because the genetic analysis

634 established requirements for sacy-1(+) function in both the germline and soma, we used strains bearing

635 germline (CA1352 *ieSi64*) or somatically expressed (CA1200 *ieSi57*) TIR1 F-box proteins to deplete

636 AID::GFP::SACY-1 in each tissue individually (Figure S5 and Figure S6). Depletion of

637 AID::GFP::SACY-1 in the germline starting at approximately the L3 stage phenocopied the gamete

638 degeneration phenotype in a small proportion of the animals (3 of 270; Figure S5). This result is

639 consistent with genetic mosaic analysis showing that the gamete degeneration phenotype is cell

640 autonomous (Kim *et al.* 2012), and it also highlights the difficulty of recapitulating null phenotypes

641 through auxin-inducible degradation. When AID::GFP::SACY-1 is depleted in the germline starting at

642 the L4 stage, many of their F1 progeny arrest as embryos or larvae, consistent with the idea that

643 maternally contributed *sacy-1(+)* activity is essential. Animals that escape the lethality and progress to

644 adulthood often display the germline degeneration phenotype (40%; n=139). To deplete

AID::GFP::SACY-1 in the soma, we placed L4 larvae on media containing 2 mM auxin. The resulting

646 F1 progeny grew very slowly, taking approximately 4–6 days to reach adulthood (instead of 2.5 days)

647 and were sterile (Figure S5). Taken together, depletion of SACY-1 using the auxin-inducible 648 degradation system resulted in a reduction-of-function condition less severe than the null phenotype 649 but more severe than conferred by the reduction-of-function missense alleles (Table 5). 650 For analysis of transcriptomes, we grew adult hermaphrodites to the young adult stage on 651 normal growth media and then transferred them to media containing 2 mM auxin for 24 hours before 652 preparing total RNA for RNA sequencing. Examination of the worms showed that AID::GFP::SACY-653 1 was depleted from the targeted tissues (Figure S6). Total RNA was prepared from each of three 654 biological replicates and their respective controls, which were the parent strains expressing TIR1 in the 655 germline (*ieSi64*) or soma (*ieSi57*) also treated with auxin. Poly(A+) mRNA was sequenced using 150 656 bp paired-end reads and the sequencing reads were aligned to the genome. Principal component 657 analysis (PCA) revealed that the biological replicates clustered together (Figure 6A), which is 658 indicative of experimental reproducibility. However, PCA indicated that the control strains for the 659 germline (CA1352) and soma (CA1200) depleted samples did not cluster together, which indicates that 660 under these conditions the two strain backgrounds exhibit marked differences in their transcriptomes 661 (Figure 6A), a finding that was further confirmed with a more granular assessment of mRNA 662 expression level differences of individual genes (Figure S7A). Thus, in our analysis we compared the 663 germline- and soma-depleted SACY-1 transcriptomes only to their respective controls. 664

665 *Changes in transcript abundance following SACY-1 depletion:* We observed two classes of 666 transcriptome alterations upon depletion of SACY-1 in the germline or soma: changes in transcript 667 abundance and alterations in splicing patterns. In terms of transcript abundance, we observed 242 668 down-regulated genes (two-fold down-regulation, adjusted p<0.05, FPKM in soma control \geq 2.5, mean 669 counts across samples > 25) in the RNA samples depleted for somatic SACY-1 (Figure 6, B and C; 670 File S2). Notably these down-regulated genes included many cuticle collagen genes and genes 671 affecting cuticular morphology and body size (*col-17, col-41, col-46, col-47, col-90, col-128, col-149*,

672	<i>dpy-3</i> , <i>dpy-4</i> , <i>dpy-5</i> , <i>dpy-6</i> , <i>dpy-8</i> , <i>dpy-9</i> , <i>dpy-13</i> , <i>dpy-20</i> , <i>lon-3</i> , <i>mlt-7</i> , <i>qua-1</i> , <i>rol-6</i> , <i>rol-8</i> , <i>sqt-1</i> , and
673	sqt-2). Consistent with this observation, the top enriched gene ontology (GO) term for transcripts
674	reduced in abundance in the SACY-1 soma-depleted samples was "cuticle development involved in
675	collagen and cuticulin-based cuticle molting cycle" (Figure S7B). We also observed 242 up-regulated
676	genes (two-fold up-regulation, adjusted p<0.05, FPKM in SACY-1 soma-deplete \geq 2.5, mean counts
677	across samples > 25) in the SACY-1 soma-depleted samples (Figure 6, B and C; File S2). The top
678	enriched GO term for transcripts with increased abundance in the SACY-1 soma-depleted samples
679	related to cellular responses to heat stress, the unfolded protein response, and innate immune responses
680	(Figure S7), suggesting that the organism might perceive the reduction of $sacy-1(+)$ function as a
681	stressor and might then mount a response that then alters the transcriptome.
682	In the SACY-1 germline-depleted samples, we observed 126 down-regulated genes (two-fold
683	down-regulation, adjusted p<0.05, FPKM in germline control \geq 2.5, mean counts across samples > 25;
684	Figure 6, B and D; File S2). The top enriched GO terms for transcripts with decreased abundance in
685	the SACY-1 germline-depleted samples included the response to heat stress and the unfolded protein
686	response (Figure S7B), suggesting the response to $sacy-1(+)$ depletion differs between the soma and
687	germline. Among the 311 transcripts increased in abundance in the SACY-1 germline-depleted sample
688	was <i>her-1</i> (two-fold up-regulation, adjusted p<0.05, FPKM in SACY-1 germline-deplete \geq 2.5, mean
689	counts across samples > 25; Figure 6, B, D, and E; File S2). This might be due to an increase in X
690	chromosome non-disjunction in embryos located in the uterus following germline depletion of sacy-
691	1(+), but this possibility was not investigated. Because <i>her-1</i> likely encodes an inhibitory ligand for
692	the TRA-2 receptor in the sex-determination pathway (Perry et al. 1993; Figure 3), we tested whether
693	the her-1(hv1y101) null mutation could suppress the Mog phenotype of the recessive gain-of-function
694	sacy-1(tn1481) mutation, but this proved not to be the case (n=53 gonad arms). Since the genetic
695	epistasis results suggest that SACY-1 promotes the expression of TRA-2 (Figure 3), we examined the
696	effect of SACY-1 depletion in the germline on the levels of <i>tra-2</i> mRNA and the fidelity of its

- splicing. We observed no statistically significant change in *tra-2* mRNA levels or its splicing patterns
 (Figure 6F). We did not examine the expression of TRA-2 protein after SACY-1 depletion because a
 recent study suggested that TRA-2 protein expression in the wild-type germline is below the
 immunofluorescence detection limit of immunofluorescence (Hu *et al.* 2019).
- 701

702 Alteration of splicing patterns following SACY-1 depletion: In the SACY-1 soma-depleted samples, 703 we observed significant (FDR<0.05) alterations in splicing patterns for 1606 transcripts (Figure 7A). 704 These splicing alterations fell into several broad classes: the use of alternative 5' splice sites, the use of 705 alternative 3'splice sites, abnormal splicing within an exon, skipped exons, and retained introns. The 706 largest class of splicing changes was the use of alternative 3' splice sites, consistent with the fact that 707 DDX41 was shown to be recruited to the C complex, which mediates the second step in splicing 708 (Bessonov et al. 2008). Multiple splicing defects were sometimes observed within a single gene. For 709 example, in the case of the RNA-binding protein ETR-1, which has multiple isoforms and is expressed 710 in the soma and germline (Boateng et al. 2017), depletion of SACY-1 results in intron retention and 711 multiple alterations in 3'-splice-site usage (Figure 7B). In the SACY-1 germline-depleted samples we 712 observed significant (FDR<0.05) alterations in splicing for 796 transcripts (Figure 7A). Thus, splicing 713 defects appeared less prevalent in the SACY-1 germline-depleted samples than the SACY-1 soma-714 depleted samples. One possibility is that nonsense mediated decay or other surveillance pathways 715 actively clear misspliced mRNAs from the germline. Some alternative splicing events were observed 716 in both the RNA preparations depleted for SACY-1 in the germline and the soma (Figure 7A). For 717 example, we observed retention of an intron in prdx-6 mRNA in both experiments (Figure 7C). 718 Likewise, we observed alternative 3' splice site selection for the heterochronic gene *lin-28* in both 719 SACY-1-depleted samples (Figure 7D).

720

721 Germline sex-specific splicing patterns and the involvement of SACY-1: As we analyzed the changes 722 in splicing patterns following SACY-1, we noticed changes in alternative splicing patterns that 723 appeared to correlate with the sex of the germline. To investigate this observation more fully, we 724 reanalyzed the data of Ortiz et al. (2014), who reported the transcriptomes of genotypically XX 725 animals exclusively undergoing oogenesis or spermatogenesis as a consequence of containing a loss-726 of-function fog-2 mutation or a gain-of-function fem-3 mutation, respectively. This dataset was 727 previously used to identify alternative isoform usage in the germline (Oritz et al. 2014). In analyzing 728 their dataset for alternative splicing events, we observed 1600 genes for which there was a significant 729 (FDR<0.05) germline sex-specific splicing pattern (Figure S8; File S3). We noted that upon SACY-1 730 depletion in either the soma or germline, oocyte-enriched splicing events were favored (Figure 7A and 731 Figure S8). This result suggests that SACY-1 plays a role in the selection of 3' splice sites for many 732 genes and raises the possibility that the appropriate balance of "spliceoforms" may be play a role in 733 cellular differentiation.

734

DISCUSSION

735	In this study, we report the results of a molecular genetic and biochemical analysis of SACY-1/DDX41
736	in C. elegans, conducted to gain potential insights into how DDX41 mutations might contribute to
737	MDS and AML in humans. DDX41 is a component of the spliceosomal C complex (Jurica et al. 2002;
738	Bessonov et al. 2008), which carries out the second step in splicing. Pre-mRNA splicing is an essential
739	process in eukaryotes (reviewed by Wahl et al. 2009). Thus, the finding that mutations in highly
740	conserved genes encoding spliceosomal components are frequently found in hematological
741	malignancies was unexpected (reviewed by Yoshida and Ogawa 2014). Unfortunately, the genetic
742	properties of disease-causing mutations in spliceosomal proteins have been difficult to assess.
743	Nonetheless, the observation that many of these missense mutations map to highly conserved amino
744	acids, within biochemically defined structural and functional domains, might suggest that they reduce
745	but not eliminate gene function. An alternative possibility is that some of the oncogenic missense
746	mutations might confer antagonistic gain-of-function properties. Indeed, oncogenic mutations in
747	several splicing factors (e.g., SF3B1, U2AF1, SRSDF2, and SRSR2) present in the heterozygous
748	condition (Yoshida and Ogaawa 2014); however, it is probable that newly arising mutations in
749	hematopoetic lineages, in combination with the germline mutations, contribute to neoplastic
750	development.

751 In this regard, the *DDX41* mutations inherited and arising in familial cases of AML are 752 particularly informative (Polprasert et al. 2015). Affected individuals often inherit one copy containing 753 a mutation, which is likely to be a null mutation (e.g., a frameshift mutation, pD140fs). In addition to 754 this DDX41 germline mutation, the neoplasms in these patients invariably contain a second copy with a 755 somatic mutation that arose in the hematopoietic lineage. Several specific DDX41 somatic mutations 756 (e.g., R525H) arise independently with high prevalence, suggesting that they markedly contribute to 757 neoplastic development. These somatic *DDX41* mutations are invariably missense mutations affecting 758 conserved amino acids but have never been found to include candidate null alleles (e.g., frameshift

mutations or premature termination codons). Thus, these somatically arising mutations are unlikely to
eliminate DDX41 function. The presumption is that biallelic mutations disrupting DDX41 function
might be lethal. In line with this view, a recent report found that two related patients with biallelic
DDX41 missense mutations exhibited a more severe syndrome characterized by dysmorphic skeletal
and facial features, psychomotor delays, intellectual disability, and early onset leukemia (Diness *et al.*2018).

765 Here we build upon our prior work (Kim et al. 2012) to conduct a comprehensive molecular 766 genetic analysis of SACY-1/DDX41 function in C. elegans. Recent work of others has used the C. 767 elegans system to gain information on oncogenic mutations in the SF3B1 spliceosomal protein (Serrat 768 et al. 2019). Our results reveal that sacy-1 mutations confer a range of phenotypes, from highly 769 pleiotropic defects affecting the germline and soma to very specific defects affecting cell 770 differentiation and cell cycle regulation. Reduction-of-function sacy-1 alleles are homozygous viable 771 and fertile, yet affect germline sex determination and the regulation of oocyte meiotic maturation (Kim 772 et al. 2012; this work). Animals homozygous for sacy-1 null mutations grow to adulthood but exhibit a 773 gamete degeneration phenotype and are sterile. Feminization of the germline delays oocyte 774 degeneration, which enables *sacy-1* null mutant females to be mated and produce embryos. However, 775 these embryos invariably die, revealing an essential maternal requirement for development. Maternally 776 provided sacy-1(+) must in turn be sufficient for homozygous sacy-1 null mutant animals (produced 777 from heterozygous parents) to grow to adulthood.

Several *sacy-1* mutant alleles exhibit genetic properties that suggest they can counteract *sacy-* 1(+) function, potentially by compromising the function of the spliceosome. Most notable among these mutations is *sacy-1(tn1481)*, which confers a masculinization of the germline phenotype resulting from the overproduction of sperm to the exclusion of the oocyte fate. Since multiple reduction-of-function *sacy-1* alleles suppress the feminization of the germline phenotype caused by null mutations in *fog-2* (Kim *et al.* 2012; this work), *sacy-1(+)* must possess a function to promote the oocyte fate. This oocyte-promoting function of sacy-1(+) might not be essential because sacy-1 null mutants produce oocytes, which nonetheless undergo necrotic degeneration. The caveat here is that sacy-1 null mutants only develop to adulthood because of maternally provided sacy-1(+), which might be sufficient to promote oogenesis. In any case, the sacy-1(tn1481) mutation might disrupt an oocyte-promoting function either by interfering with maternally provided SACY-1 activity or proteins with which it associates.

790 Several observations are consistent with the possibility that *sacy-1(tn1481)* interferes with or 791 compromises the function of the spliceosome in germline sex determination in a dosage-sensitive 792 manner. This view is also supported by our tandem affinity purification results showing that SACY-1 793 is a component of the C. elegans spliceosome and genetically interacts with spliceosomal components. 794 Significantly, multiple C. elegans spliceosomal components, functioning at different steps of the 795 splicing reaction, can mutate to a masculinization of germline phenotype (reviewed by Zanetti and 796 Puoti 2013). Interestingly, when placed in trans to a sacy-1 null mutation (e.g., tm5503), the resulting 797 sacy-1(tn1481)/sacy-1(tm5503) heterozygous animals are viable and fertile and can be maintained 798 indefinitely as a heterozygous strain. This highly informative genetic result suggests that a single dose 799 of *sacy-1(tn1481*) can mediate all its essential functions, but that two doses of the mutant protein 800 interferes with the normal mechanisms of germline sex determination (i.e., sacy-1(tn1481) is a 801 recessive gain-of-function antimorphic mutation). Although sacv-1(tn1481)/sacv-1(tm5503) 802 heterozygous adult hermaphrodites are fertile, when sacv-1(tn1481); fem-3(e1996) females are mated 803 to wild-type males, they produce embryos that invariably fail to hatch. Similar results are obtained 804 with the six other mog genes (Graham and Kimble 1993; Graham et al. 1993), which encode 805 spliceosomal proteins (Puoti and Kimble 1999; Puoti and Kimble 2000; Belfiore et al. 2004; Kasturi et 806 al. 2010; Zanetti et al. 2011). Interestingly the P222L amino acid substitution found in sacy-1(tn1481) 807 is adjacent to the O motif, which participates in nucleotide binding and hydrolysis (Schütz *et al.* 2010). 808 It is tempting to speculate that SACY-1/DDX41 might promote remodeling of spliceosomes during the 809 splicing reaction and that two doses of SACY-1 P222L might disrupt these rearrangements. In a 810 similar vein, sacy-1(tn1479[G504E]) exhibits a phenotype more severe than a null allele, suggesting 811 that this mutation in the helicase domain too might possess a dosage-sensitive antimorphic activity. 812 Whether missense alleles in human DDX41 also possess antimorphic activity will require additional 813 work, including biochemical analyses. 814 We imported the oncogenic DDX41[R525H] mutation into C. elegans using CRISPR-Cas9 815 genome editing (R534H in SACY-1). Our analysis revealed that in C. elegans, this mutation possesses 816 very weak antagonistic activity. While it is possible that the genetic properties of this mutation might 817 differ between C. elegans and mammalian systems, it is worthwhile noting that the oncogenic variant 818 must support the high levels of proliferation characteristic of neoplastic cells. While all the sacv-1 819 mutations we isolated in forward genetic screens occur at conserved amino acids, none of them match 820 the oncogenic mutations thus far isolated (Polprasert et al. 2015; Cardoso et al. 2016; Lewinsohn et al. 821 2016; Li et al. 2016; Diness et al. 2018; Sébert et al. 2019). This observation is consistent with the 822 idea that the oncogenic mutations are at most weakly antagonizing or weak reduction-of-function 823 mutations, and thus would not have been isolated in forward genetic screens that require a substantial 824 reduction in function.

825 An attractive idea is that oncogenic mutations affecting the spliceosome contribute to 826 neoplastic development through effects on gene expression occurring through alterations in RNA 827 splicing, as well as effects on the transcriptional machinery or RNA stability (Yoshimi *et al.* 2019). 828 Consistent with this idea, RNAi depletion of the C. elegans spliceosomal protein RSR-2 affects 829 transcript levels without major impacts on splicing (Fontrodona et al. 2013). In this study, we 830 examined the effects of SACY-1 depletion at the adult stage on the transcriptome. In the soma and the 831 germline, we observed splicing-independent impacts on the abundance of many transcripts. The gene 832 expression changes observed suggest that depletion of SACY-1 might elicit a stress response. If 833 analogous processes occur after perturbations of the spliceosome in humans, such stress responses

37

- might contribute to oncogenesis and could represent therapeutic targets. In C. elegans, we observed
- 835 instances of missplicing following SACY-1 depletion, though missplicing events more prevalent when
- 836 SACY-1 was depleted from the soma as compared to the germline. The extent to which gene
- 837 expression changes and splicing alterations contribute to the various *sacy-1* mutant phenotypes will
- require further study but will likely provide insights relevant to spliceosomal perturbations in humans.

839

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1103Table 1 RNAi of sacy-1 enhancer loci increase the penetrance of germline or lethal1104phenotypes in sacy-1(tn1385) reduction-of-function mutants

1105

RNAi ^a	Genotype	Sterile (%) ^b	Gamete degeneration ^b (%)	Embryonic lethal (%) ^c
	Wild type	0	0	1
L4440 (control)		(n=338)	(n=338)	(n=674)
	sacy-1(tn1385)	0	0	1
		(n=256)	(n=256)	(n=502)
	Wild type	4	0	7
$m_{0} = 2(11, 2D16)$		(n=172)	(n=54)	(n=574)
<i>mog-2</i> (II-3D16)	sacy-1(tn1385)	43	47	95
		(n=254)	(n=72)	(n=426)
	Wild type	22	1	84
Y111B2A.25		(n=230)	(n=94)	(n=463)
(III-6G22)	sacy-1(tn1385)	95	16	93
		(n=272)	(n=110)	(n=42)
	Wild type	100	35	ND
agen 1d (11 0E00)		(n=164)	(n=40)	
<i>cacn-1</i> ^d (II-9E09)	sacy-1(tn1385)	100	81	ND
		(n=224)	(n=52)	
	Wild type	2	0	12
emb-4 (V-12E12)		(n=184)	(n=110)	(n=337)
emb-4 (v-12E12)	sacy-1(tn1385)	89	42	N.D.
		(n=176)	(n=82)	
	Wild type	4	0	13
$a_{1} = h + A (11 + 1) = 1 + A (11 + 1) = A (1$		(n=288)	(n=96)	(n=421)
emb-4 (V-12E14)	sacy-1(tn1385)	91	80	ND
		(n=202)	(n=120)	
	Wild type	4	0	9
	- JF -	(n=210)	(n=96)	(n=433)
emb-4 (V-12E16)	sacy-1(tn1385)	91	87	ND
	5409 2(012000)	(n=326)	(n=140)	

^aRNAi clones showing genetic interactions with *sacy-1* are listed with the target gene name in

1107 italics and the location of the clone in the RNAi library in parentheses. The identity of clones was

1108 verified by DNA sequencing.

^bSterility and gamete degeneration were scored by DIC microscopy on the first day of adulthood

1110 24 hours post-L4 at 22°C. Gonad arms were scored as sterile if they did not produce embryos and

1111 exhibited defects in gametogenesis. Number of gonad arms scored is reported.

- 1112 ^cEmbryonic lethality was measured by conducting daily egg lays over the reproductive lifespan
- and determining the number of embryos that failed to hatch by 48 hours after egg laying. The
- 1114 number of embryos scored is reported.
- 1115 *dcacn-1* was not initially identified as an enhancer of *sacy-1* during the genome-wide RNAi screen
- 1116 because *cacn-1(RNAi*) results in complete sterility in both *sacy-1(tn1385*) and wild-type animals.
- 1117 However, we determined that RNAi to the *Y111B2A.25* pseudogene likely targets *cacn-1* (see text
- 1118 for details).
- 1119 ND, not determined.

1120 Table 2 Genetic interactions between *sacy-1* and *emb-4*

1121

1122 A. Enhancement of germline and somatic *sacy-1* mutant defects

Genotype	Vulval rupture (%)	Sterile and Pvl ^a (%)	Sterile ^a (%)	Fertile (%)
<i>sacy-1(tm5503)</i> (n=284)	3	1	96	0
sacy-1(tm5503)/+; emb-4(sa44) ^b (n=205)	0	0	0	100
sacy-1(tm5503); emb-4(sa44) ^b (n=242)	83	16	1	0
<i>sacy-1(tn1385)</i> (n=278)	0	0	0	100
sacy-1(tn1385)/+; emb-4(sa44) ^b (n=201)	0	0	0	100
<i>sacy-1(tn1385); emb-4(sa44)</i> ^{b,c} (n=144)	36	4	3	57 ^d

^aSterile animals exhibit the *sacy-1(lf)* gamete degeneration phenotype.

^bThe *hT2(qIs48)* balancer chromosome, which is dominantly marked with GFP, was used to differentiate between *sacy-1(tm5503)* heterozygotes and homozygotes.

1126 ^cThe progeny of *sacy-1(tn1385)/hT2(qIs48)*; *emb-4(sa44)* hermaphrodites; the balancer

1127 chromosome provides maternal sacy-1(+) function. The fertile F1 progeny of these animals are 1128 maternal-effect lethal, see Table 2B, below.

^dThese adult hermaphrodites produce a majority of embryos that fail to hatch, see Table 2B.

1130

1131 B. Enhancement of embryonic lethality

Genotype ^a	Embryonic lethal (%)	L1 lethal (%)	Viable (%)
<i>sacy-1(tn1385)</i> (n=949)	1	0	99
<i>emb-4(sa44)</i> (n=1348)	4	0	96
<i>sacy-1(tn1385); emb-4(sa44)</i> ^b (n=620)	97	3	0

1132 ^aThe number of embryos examined.

^bThe F1 progeny of fertile *sacy-1(tn1385); emb-4(sa44)* parents derived from the *sacy-*

1134 *1(tn1385)/+; emb-4(sa44)* heterozygotes analyzed in Table 2A above.

1135 **Table 3. Spliceosomal proteins associated with SACY-1 using tandem affinity**

1136 purification

_

		Protein Coverage (%) ^a	
		Experiment I	Experiment II
Protein		Tandem IP	Tandem IP
		female	hermaphrodite
		background ^b	background ^b
SACY-1	DDX41/Abstrakt/recruited to C-	78.9	76.5
	complex		
Spliceosomal proteins	Human protein/Spliceosome		
	subcomplex		
PRP-19	PRP19hPRP19/CDC5L complex ^c	62.0	55.5
MOG-2 ^d	U2A'/17S U2 snRNP ^c	56.1	42.3
SKP-1	SNW1/SKIP/hPRP19/CDC5L-	52.9	40.9
	related complex		
EMB-4, isoform b	Aquarius/KIAA0560/Intron-	50.4	30.6
	binding complex RNA helicase ^{c, e}		
PRP-17	hPRP17/WD40-domain step 2	48.3	33.7
	factor ^{c, f}		
CYN-13	Cyclophilin E/PPIE/Intron-binding	46.8	37.8

complex^e

CDC-5L	CDC5L/hPRP19/CDC5L complex ^c	45.8	26.2
Y69A2AR.21	CCDC12/Recruited to B-complex ^c	45.6	52.1
RBM-22	RBM22/ hPRP19/CDC5L-related complex ^{c, g}	45.1	43.9
EFTU-2 ^d	U5 116K°	45.1	37.2
ISY-1	hIsy1/Intron-binding complex ^e	40.8	34.8
PLRG-1	PLRG1/PRL1/hPRP19/CDC5L complex ^c	39.1	27.3
RBMX-2 ^d	CGI-79/SNU17/IST3/hRes complex ^c	38.7	20.9
CYN-10	PPIL3b/peptidyl-prolyl isomerase recruited to C-complex ^c	37.9	23.6
RBM-25	RBM25/affects alternative splicing/putative U1 snRNP component	37.5	25.8
RNP-6, isoform a	PUF60/U2AF65-related promotes alternative splicing ^h	37.2	49.3
SYF-2	GCIP p29/SYF2/Recruited to C- complex ^c	37.2	29.9

PRP-8	PRPF8/PRP8/U5 220K ⁱ	37.0	29.4
EMB-4, isoform a	Aquarius/KIAA0560/Intron- binding complex RNA helicase ^{c, d}	36.8	28.6
DDX-35	DDX35/DHX35/Recruited to C- complex ^c	36.8	19.8
SNRP-40.1	snRNP40/U5 40K ^c	36.6	27.5
RBM-39 ^d	RBM39/protein recruited to A-complex ^c	35.9	31.0
CACN-1	Cactin/C-complex ^{c, j}	35.9	13.4
ACIN-1	Acinus/Exon junction complex ^c	35.2	13.1
M03F8.3	hSYF3/CRNKL1/hPRP19/CDC5L- related protein	35.0	24.3
ZK1098.1	PRP40/FBP11 ^k	34.5	28.7
C50D2.5	SF3b14/SF3b6/17S U2 snRNP protein	33.3	8.7
SYF-1	hSyf1/Xab2/Intron-binding complex ^e	32.4	28.0
EMB-4, isoform e	Aquarius/KIAA0560/Intron- binding complex RNA helicase ^{c, e}	31.7	18.4
F33D11.10	EIF4A3/ Exon junction complex ^c	31.3	17.0

REPO-1	SF3a66/SF3A2/17S U2 snRNP protein ^{c,1}	30.6	5.9
Y54G2A.12	hPRP17/hCDC40/Step 2 factor	29.9	27.7
PNN-1	Pinin/Exon junction complex ^c	28.7	20.3
BCAS-2	Spf27/BCAS2/hPRP19/CDC5L complex ^c	28.6	20.2
EMB-4, isoform d	Aquarius/KIAA0560/Intron- binding complex RNA helicase ^{c, e}	28.3	27.9
RSP-7	p54 SR protein ^m	27.2	27.4
CWC-15	CCAP2/CWC15/hPRP19/CDC5L complex ^c	27.0	22.6
LET-858	KIAA1604/CWC22/Recruited to B-complex ^c	26.5	16.5
R07E5.1	GPATCH1/ECGP ⁿ	26.0	11.2
RSR-2	SRm300/SRRM2	24.0	21.2
MOG-3	CCDC49/CWC25/recruited to C- complex ^o	21.3	20.0
CYN-12	PPIL1/ peptidyl-prolyl isomerase/hPrp19/CDC5L-related protein	20.1	20.1

F25B4.5	PRPF39/PRP39/U1 snRNP	20.1	10.0
	auxiliary protein ^p		
SNRP-40.2	U5 snRNP 40Kc	19.6	16.0
C16C10.4	SAP18/Exon junction complex ^c	17.5	7.8
DDX-23	U5 snRNP 100K/PRP28¢	17.1	7.5
R08D7.1	BUD13/hRes complex ^c	16.6	3.7
TEG-4	SF3b130/SF3B3/17S U2 snRNP ^c	14.9	8.8
SFTB-1 ^s	SF3b155/SF3B1/17S U2 snRNP ^c	14.7	1.2
F53H1.1, isoform a	SF3b125/17S U2 snRNP ^q	14.5	3.4
RNP-6, isoform b	PUF60/U2AF65-related promotes	13.6	20.3
	alternative splicing ^h		
PRPF-4, isoform a	PRP4 kinase ^r	12.5	7.1
RNP-3	U2B"/17S U2 snRNPc	12.4	12.4
F53H1.1, isoform e	SF3b125/17S U2 snRNP ^q	11.9	5.1
PRP-21	SF3a120/SF3A1/PRP21/17S U2	11.8	7.0
	snRNP protein ^c		
SNRP-200 ^d	U5 200K ^c	11.1	7.7
SFTB-2	SF3b145/SF3B2/17S U2 snRNP ^c	10.1	1.5

	RNP-5	RNPS1/EJC ^c	10.0	18.0
1137	^a Peptide covera	age in a single gel slice assessed by	mass spectrometry in the vario	ous SACY-1
1138	purifications fr	om DG4068	PreScission::gfp::tev::s::sacy-1]]) fog-1(q253)
1139	extract and DG	4070 <i>sacy-1(tn1632)</i> extract. Only	proteins showing at least 10%	coverage in the
1140	tandem immun	opurification (IP) from the high-sp	eed supernatant are shown. The	he complete
1141	data on which '	Table 3 is based, including filtering	criteria, are presented in File S	51.
1142	^b Experiments I	and II utilized 515 mg and 280 mg	total protein, respectively, from	n a high-speed
1143	supernatant for	r tandem IP.		
1144	^c Bessonov <i>et al</i>	. 2008.		
1145	^d MOG-2, EFTU-	2, RBMX-2, RBM-39, SFTB-1, TEG-4	and SNRP-200 are reported ir	1 Table 3 and
1146	not filtered out	despite their low representation (2	14.2%, 4.4%, 3.6%, 6.9%, 2.0%	, 1.1% and 4.5%
1147	coverage, respe	ectively) in the LIN-41 tandem IP fr	om high-speed supernatant (T	sukamoto <i>et al</i> .
1148	2017) because	they were reported as splicing fact	ors. F33D11.10 is reported in T	fable 3 despite
1149	its low represe	ntation (10.3% coverage) in the OC	IC-5 tandem IP from high-spee	d supernatant
1150	(unpublished d	ata) because it was reported as a sj	plicing factor.	
1151	^e De <i>et al.</i> 2015;	Haselbach <i>et al</i> . 2018.		
1152	^f Bertram <i>et al</i> .	2017.		
1153	^g Zhang <i>et al</i> . 20	17.		
1154	^h Page-McCaw e	et al. 1999; Van Buskirk and Schüpb	ach 2002; Hastings <i>et al.</i> 2007.	
1155	ⁱ Grainger and E	leggs 2005.		
1156	Fica <i>et al</i> . 2019			
1157	^k Kao and Silicia	ino 1996.		
1158	¹ Keikhaee <i>et al</i> .	2014.		

^mChaudhary *et al.* 1991; Longman *et al.* 2000.

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- ¹¹⁶⁰ ⁿBessonov *et al.* 2010.
- 1161 °Fabrizio *et al.* 2009.
- 1162 ^pLi *et al.* 2017.
- 1163 qWill *et al.* 2002.
- 1164 ^rSchneider *et al.* 2010.

Gonad arms containing mitotic undifferentiated

	germ cells in the proximal gonad arm ^a
<i>sacy-1(tn1385</i> [G533R])	$0 (n=256)^{b}$
glp-1(ar202)	$0.8 (n=364)^{b}$
sacy-1(tn1385[G533R]); glp-1(ar202)	49.6 (n=415) ^b
sacy-1(tn1887[R534H]); glp-1(ar202)	0 (n=62)

1165 Table 4 *sacy-1(tn1385rf)* enhances the *glp-1(ar202)* Tumorous phenotype

Strain

^aThe percentage of young adult hermaphrodites were examined by DIC microscopy approximately 40

1167 hours post-L4 at 15°C. The number of gonad arms scored is listed.

^bIn addition, dissected and fixed gonad from the same stage were stained for the phosphohistone

1169 H3(Ser10) M-phase marker. Of 25 sacy-1(tn1385); glp-1(ar202) gonads scored, 16 (64%) contained

1170 phosphohistone H3-positive undifferentiated germ cells in the proximal gonad arm. The average

1171 number of proximal phosphohistone H3-positive germ cells in the Tumorous gonads was 22 ± 12 .

1172 None of the sacy-1(tn1385) (n=18) or glp-1(ar202) (n=31) dissected gonads examined contained

1173 phosphohistone H3-positive undifferentiated germ cells in the proximal gonad arm.

1174	Table 5 sacy-1	alleles relevant to	this study
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Allele	Alteration	Phenotypes	Inferred activity
Loss-of-function all	eles		
$sacy-1(tm5503)^{a}$	619 bp deletion	Sterile, gamete	Likely null
		degeneration	
<i>sacy-1(tn1615)</i> ^b	10 bp deletion	Sterile, gamete	Likely null
		degeneration	
sacy-1(tn1385) ^a	G533R	Viable and fertile,	Reduction of function
		suppresses acy-4 sterility,	
		suppresses fog-2 sterility	
sacy-1(tn1391) ^a	G473R	Viable and fertile, supresses	Reduction of function
		acy-4 sterility, suppresses	
		fog-2 sterility	
sacy-1(tn1440) ^a	G331R	Viable and fertile,	Reduction of function
		suppresses acy-4 sterility,	
		suppresses fog-2 sterility	
sacy-1(tn1482) ^b	D506N	Viable and fertile,	Reduction of function
		suppresses fog-2 sterility ^c	
sacy-1(tn1483) ^b	G269E	Viable and fertile,	Reduction of function
		suppresses fog-2 sterility ^c	
Alleles with antagor	nistic activity		
<i>sacy-1(tn1479)</i> ^b	G504E	Sterile, adult lethal	sacy-1(tn1479) ^b
		(rupture) or gamete	
		degeneration	
sacy-1(tn1480) ^b	H527Y	Viable and fertile at 20°C	Loss of function with
		and suppresses fog-2	antimorphic activity,

		sterility. ^c Embryonic lethal	dominant Him
		or larval arrest at 15°C.	
		Sterile and	
		spermatogenesis-defective	
		at 25°C	
<i>sacy-1(tn1481)</i> ^b	P222L	Sterile, masculinization of	Recessive gain-of-
		germline	function
			Loss of function with
			antimorphic activity
<i>sacy-1(tn1887)</i> ^b	R534H	Viable and fertile.	Weak antagonistic
			activity; enhances the
			dominant Him phenotype
			of <i>sacy-1(tn1480)</i>

^aKim *et al.* 2012.

1176 ^bThis work.

^cSuppression of *acy-4* sterility was not tested.

Allele	Class	Gamete	Vulval	Mog ^{a,b}	T ^c
		Degeneration ^a	Rupture ^a		
sacy-1(tm5503)	Strong loss-of-function	5.6	94.4	0	15
(n=143)					
sacy-1(tm5503)	Strong loss-of-function	96.9	3.1	0	20
(n=96)					
sacy-1(tm5503)	Strong loss-of-function	98.0	2.0	0	25
(n=106)	(from <i>tm5503/</i> +)				
<i>sacy-1(tm5503)</i> ^d	Strong loss-of-function	14.7	85.3	0	25
(n=231)	(from <i>tm5503/tn1480</i>)				
sacy-1(tn1615)	Strong loss-of-function	97.8	2.2	0	20
(n=92)					
sacy-1(tn1479)	Strong loss-of-function with	11.2	88.8	0	20
(n=170)	antagonistic activity				
sacy-1(tn1481)	Recessive gain-of-function	0	0	100	20
(n=125)					

1178 Table 6 sacy-1 mutant alleles with antagonistic activity

^aThe percentage of adult hermaphrodites exhibiting the reported phenotype is shown. Adults were

scored 24 hr post-L4 at 20° or 25°C or 48 hr post-L4 at 15°C.

1181 ^bsacy-1(tn1481) adult hermaphrodites produce large numbers of sperm but no oocytes and are sterile.

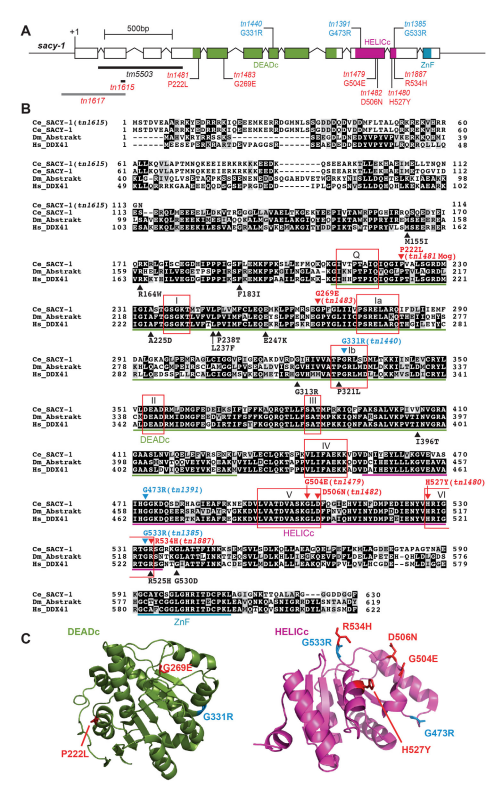
^cGrowth temperature in °C.

^dThe *sacy-1(tm5503)* progeny of *sacy-1(tm5503)/unc-13(e1091) sacy-1(tn1480)* hermaphrodites grown

1184 at 25°C.

1185

LEGENDS TO FIGURES

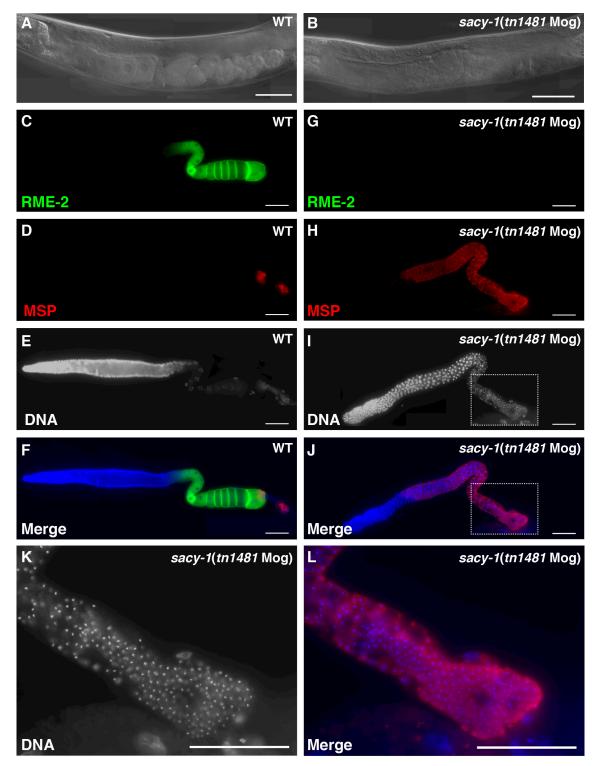


1186

Figure 1 (A) The structure of *sacy-1*. Newly isolated mutations reported in this study are displayed in

1188 red font beneath the exons. The mutations in blue font shown above the exons were reported

- 1189 previously (Kim et al. 2012). The extent of two deletions, tm5503 and tn1615, that result in sacy-1
- 1190 null mutations are shown with black bars. A third deletion, *tn1617*, which is a reduction-of-function
- 1191 mutation, is shown with a gray bar. (B) A protein sequence alignment of SACY-1 (NP_491962.1),
- 1192 Drosophila Abstract (NP_524220.1), and human DDX41 (NP_057306.2). Mutations isolated in *C*.
- 1193 *elegans* are shown above that sequence, whereas the human mutations associated with myelodysplastic
- syndromes are shown beneath the human sequence. Conserved domains [DEAD-box domain
- 1195 (DEADc), helicase domain (HELICc), and zinc finger domain (ZnF)] and motifs (Q, I Ia, Ib, III, IV, V,
- and VI) are indicated as described by Henn *et al.* (2012). (C) The locations of SACY-1 missense
- 1197 mutations are shown on structures of the DDX41 DEADc (Omura et al. 2016) and HELICc (Schütz et
- 1198 *al.* 2010) domains. The side chains of the amino acids in the human structure are labeled with amino
- acid numbering that corresponds to the SACY-1 missense mutations in this study.

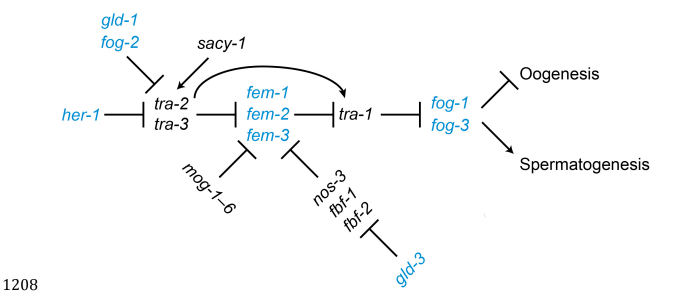


1200

1201 Figure 2 sacy-1(tn1481) adult hermaphrodites exhibit a masculinization of germline (Mog) sterile

- 1202 phenotype. DIC images of wild-type (A) and *sacy-1(tn1481)* (B) adult hermaphrodites. The wild-type
- animal contains oocytes and sperm and produces embryos but the *sacy-1(tn1481*) animal only
- 1204 produces sperm (arrow). (C–L) Dissected gonads stained for the RME-2 yolk receptor (C, G), the

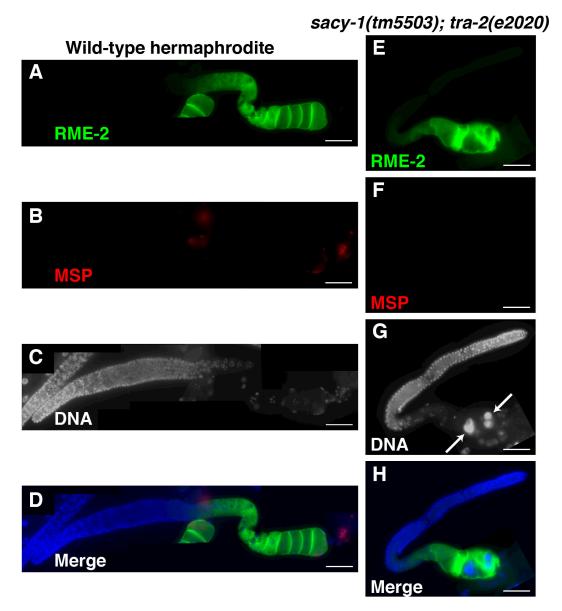
- 1205 major sperm protein (D, H), or DNA (E, I, and K). Merged images are also shown (F, J, and L). The
- 1206 sacy-1(tn1481) mutant overproduces sperm to the exclusion of oocytes and is sterile. This phenotype is
- 1207 completely penetrant. Bars, 50 μm.



1209 Figure 3 The *C. elegans* germline sex determination pathway. Genes promoting the male and female

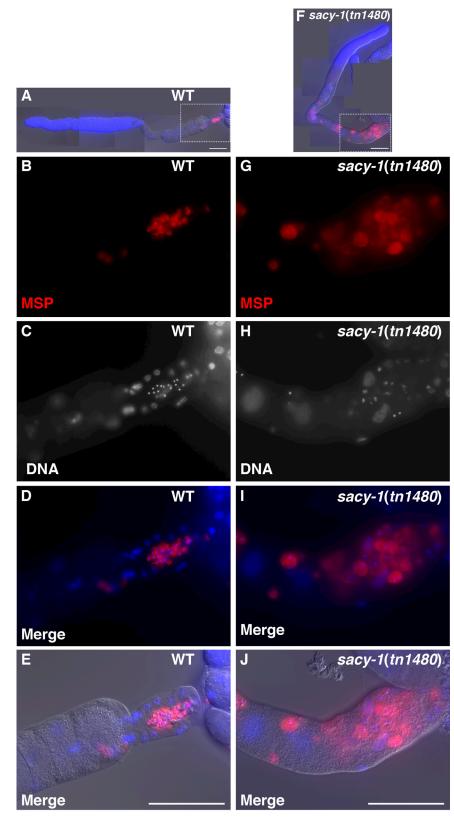
1210 fate are shown in blue and black, respectively. *sacy-1* promotes the oocyte fate antagonistically to *fog-*

1211 2, which promotes spermatogenesis.

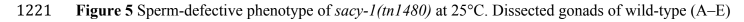


¹²¹²

- 1213 Figure 4 Genetic interactions between *sacy-1* and *tra-2* in germline sex determination were analyzed
- 1214 by combining the strong germline-feminizing dominant *tra-2(e2020)* mutation with the *sacy*-
- 1215 *I(tm5503)* null mutation. Dissected gonads of wild-type hermaphrodites (A–D) and *sacy-1(tm5503)*;
- 1216 *tra-2(e2020)* females were analyzed by staining for the oocyte RME-2 yolk receptor (A and E) and the
- 1217 major sperm protein (B and F). DNA was detected with DAPI (C and G). Merged images are also
- 1218 shown (D and H). Note gonads from *sacy-1(tm5503); tra-2(e2020)* females do not express MSP and
- 1219 frequently contain endomitotic oocytes (arrows). Bars, 50 μm.







1222 and sacy-1(tn1480) (F–J) stained with anti-MSP antibodies (B, G) and DAPI to detect DNA (C, H).

- 1223 Merged images are also shown (A, D, E, F, I, and J). At 25°C, sacy-1(tn1480) hermaphrodites produce
- swollen and abnormal sperm, which are incapable for fertilization. Note that defective sperm are also
- 1225 found near the bend region in *sacy-1(tn1480)* adults (F) indicating that there is a defect in the sperm-
- 1226 to-oocyte switch. Bars, 50 μm.

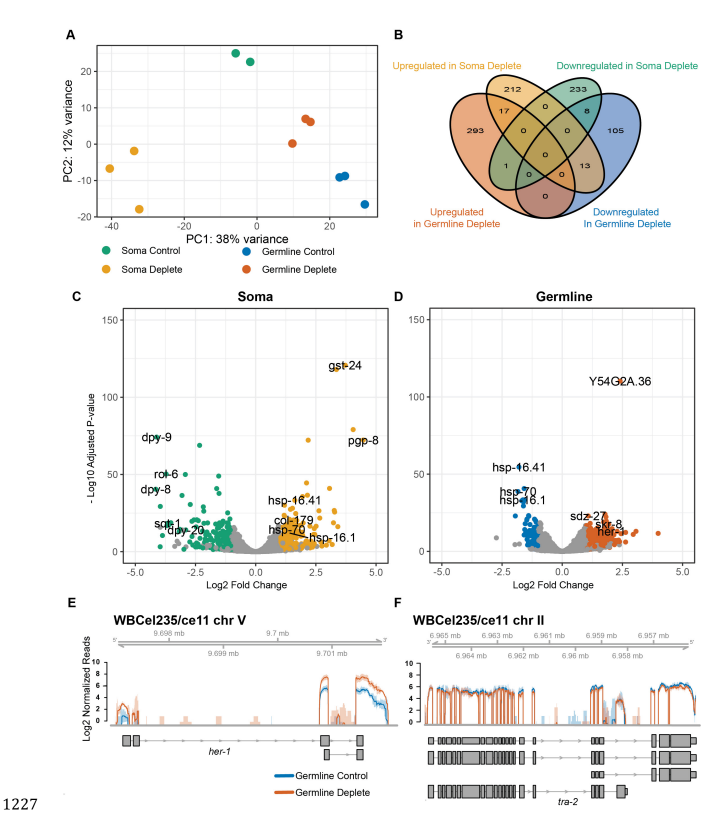


Figure 6 Transcriptome changes upon SACY-1 depletion. (A) PCA comparison of RNA-seq data of
controls strains and the experimental samples in which SACY-1 was depleted in the germline or soma,
as indicated. Three biological replicates were analyzed for each sample; however, one of the control

- samples for the soma depletion exhibited evidence of RNA degradation and was excluded from the
- 1232 analysis. (B) A Venn diagram showing the limited overlap of upregulated genes (2-fold; adjusted
- 1233 p < 0.05, FPKM deplete > 2.5 and mean counts > 25) and downregulated (2-fold; adjusted p < 0.05,
- 1234 FPKM control > 2.5 and mean counts > 25) genes in the RNA-seq datasets. (C, D) Volcano plots
- showing the log2 fold change in expression versus the -log10 of the adjusted p value of genes
- 1236 following SACY-1 depletion in the soma (C) or germline (D). (E, F) The normalized coverage of
- 1237 sequencing reads across *her-1* (E) and *tra-2* (F) following depletion of SACY-1 in the germline. The
- solid lines represent the mean of the biological replicates and shaded regions represent the
- 1239 corresponding confidence interval. Note, the pattern of *tra-2* splicing is not affected.

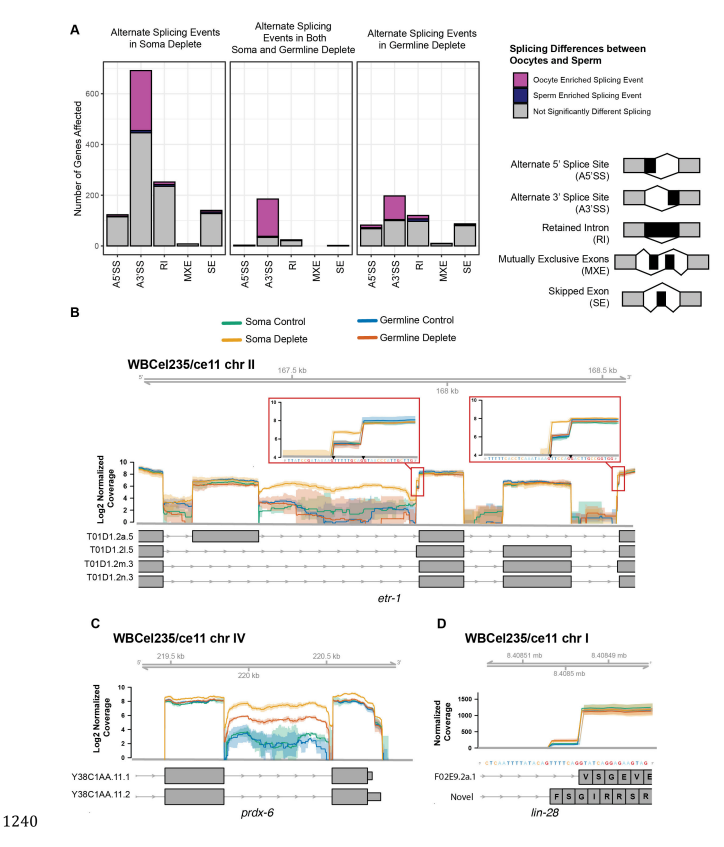


Figure 7 Quantification of altered splicing patterns upon SACY-1 depletion. (A) Bar graphs showing
the number of genes with statistically significant (FDR<0.05) changes in splicing patterns. The legend

- 1243 at the right depicts the nature of the observed splicing changes: A5'SS= Alternate 5' Splice Site,
- 1244 A3'SS = Alternate 3' Splice Site, RI= Retained Intron, MXE=Mutually Exclusive Exons, SE=Skipped
- 1245 Exon (B–D) Examples of alterations in splicing patterns following SACY-1 depletion in the germline
- 1246 or soma as indicated. The *etr-1* gene shows pronounced intron retention and two alternatively spliced
- 1247 3' sites in the SACY-1 soma depleted (gold) sample (B). A subset of *etr-1* transcript annotations are
- 1248 shown. The *prdx-6* gene has a retained intron in the SACY-1 soma depleted (gold) and germline
- 1249 depleted (red) samples (C). The soma and germline depleted samples have an increase in the usage of
- 1250 alternate splice acceptor in the *lin-28* gene that results in an altered reading frame (D). The solid lines
- 1251 represent the mean of the biological replicates and shaded regions represent the corresponding
- 1252 confidence interval.