

1 **Sequencing RNA from old, dried specimens reveals past viromes and properties of long-**
2 **surviving RNA**

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4 *Running title: RNA survives well in biological specimens without freezing or fixation*

5
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15 **ABSTRACT**

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17 Recovery of virus sequences from old samples provides an opportunity to study virus evolution
18 and reconstruct historic virus-host interactions. Studies of old virus sequences have mainly
19 relied on DNA or on RNA from fixed or frozen samples. The millions of specimens in natural
20 history museums represent a potential treasure trove of old virus sequences, but it is not clear
21 how well RNA survives in old samples. We experimentally assessed the stability of RNA in
22 insects stored dry at room temperature over 72 weeks. Although RNA molecules grew
23 fragmented, RNA yields remained surprisingly constant. RT-qPCR of host and virus RNA
24 showed minimal differences between dried and frozen specimens. To assess RNA survival in
25 much older samples we acquired *Drosophila* specimens from North American entomological
26 collections. We recovered sequences from known and novel viruses including several coding
27 complete virus genomes from a fly collected in 1908. We found that the virome of *D.*
28 *melanogaster* has changed little over the past century. Galbut virus, the most prevalent virus
29 infection in contemporary *D. melanogaster*, was also the most common in historic samples.
30 Finally, we investigated the genomic and physical features of surviving RNA. RNA that survived
31 was fragmented, chemically damaged, and preferentially double stranded or contained in
32 ribonucleoprotein complexes. This showed that RNA - especially certain types of RNA – can
33 survive in biological specimens over extended periods in the absence of fixation or freezing and
34 confirms the utility of dried specimens to provide a clearer understanding of virus evolution.

35 **IMPORTANCE**

36 RNA from old specimens has been instrumental for understanding the origin and evolution of
37 RNA viruses. Most studies have relied on relatively rare fixed or frozen samples, likely because
38 researchers assume that RNA doesn't survive well. Using experimentally dried insects and dried
39 specimens from museums, we show that RNA can in fact persist for decades or longer without
40 freezing or fixation. We found that the virome of the model fruit fly *Drosophila melanogaster* has
41 remained largely stable over the last century. But we also discovered completely new viruses
42 from past infections. Double-stranded RNA preferentially survived, but double strandedness did
43 not account for all surviving RNA, and RNA incorporated into proteinaceous complexes also
44 survived better. This work confirms the value of the millions of dried specimens stored in natural
45 history and archaeological collections for understanding the evolutionary history of RNA viruses.

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47

48 **INTRODUCTION**

49

50 The study of ancient DNA has been a useful way to better understand the evolutionary
51 history of cellular organisms and their pathogens¹⁻⁴. RNA viruses cause many important
52 diseases, but the study of old RNA has mainly relied on relatively rare fixed or frozen
53 specimens^{5,6}. Nevertheless, such specimens have been useful for reconstructing the early
54 history of pandemic pathogens^{7,8}. Other studies have taken advantage of endogenized RNA
55 virus sequences, which capture the sequences of viruses that existed long ago⁹⁻¹¹. The
56 understanding of virus evolution would benefit from the recovery of additional old RNA virus
57 sequences. But how well RNA survives in the absence of freezing or fixation is unclear.

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59 There are reasons to believe that RNA may not survive well over extended periods. For
60 one thing, RNA is short-lived during normal cellular function. The half-life of messenger and
61 ribosomal RNAs are measured in minutes or hours in a living cell¹². RNA molecules are shorter
62 than genomic DNA and more prone to hydrolysis at physiological conditions¹³. Marketing of
63 RNA-stabilizing reagents further reinforces the idea that RNA is unstable. Studies of historic
64 RNA may therefore be uncommon simply because researchers assume that RNA does not
65 survive well in old specimens.

66

67 However, there are an increasing number of studies that indicate that RNA may in fact
68 survive well over long periods¹³. Examples include recovery of a single stranded (ss) RNA virus
69 genome from 750-year-old barley, sequencing of a double stranded (ds) RNA virus genome
70 from 1000-year-old corn, and profiling of RNA from a 130-year-old extinct Tasmanian tiger^{14–16}.
71 These studies indicate that RNA can survive for long periods and hint at the potential broad
72 utility of old specimens to study RNA viruses from hundreds or thousands of years ago.

73

74 There are nearly 300 million archived arthropod specimens in North American museums
75 alone¹⁷. Dried beetles, butterflies, ants and fruit flies have yielded useful DNA but to our
76 knowledge RNA has not been sampled from such specimens^{18–24}. Amongst the diverse
77 organisms contained in entomological collections are flies in the *Drosophilidae* family²⁴.
78 *Drosophila melanogaster* has been instrumental to our understanding of genetics, evolution,
79 and disease, and the virome of contemporary *D. melanogaster* has been described in detail^{25–}
80 ²⁹.

81

82 In this study we used two approaches to investigate RNA survival. First, we
83 experimentally dried and froze flies and mosquitoes and measured RNA yield, length, and
84 detectability over 72 and 52 weeks. Next, we obtained *Drosophila* specimens from museum
85 collections. Using next generation sequencing we characterized surviving viral and host RNA.
86 We recovered viral genome sequences from over 100 years ago and identified properties of
87 RNA associated with survival. We followed strict lab protocols to minimize contamination by
88 contemporary nucleic acid and present many lines of evidence to support the idea that our
89 conclusions are based on RNA from the actual samples. Our findings challenge the assumption
90 that RNA does not survive well and confirm the usefulness of dried biological specimens as a
91 source of RNA to study past viral infection.

92

93 **RESULTS**

94

95 **Viral and host RNA persisted in dried biological specimens**

96

97 Entomological collections contain millions of dried specimens, but it's not clear to what
98 extent RNA survives in dried insects, or more generally in dried biological specimens. To assess
99 RNA survival in such samples, we recovered RNA from flies and mosquitoes that had been
100 pinned and stored at room temperature or that had been frozen. We used adult *D. melanogaster*

101 and *Aedes aegypti* from outbred colonies maintained in our insectary^{30,31}. Both populations
102 contained individuals persistently infected by one or more viruses. We sampled *D.*
103 *melanogaster* over 72 weeks and *Ae. aegypti* over 52 weeks. We quantified the concentration
104 and size distribution of recovered RNA and used RT-qPCR to quantify levels of specific host
105 and viral RNAs.

106
107 RNA concentrations recovered from individual flies were variable but we were able to
108 recover RNA from all dried samples (**Fig. 1A**). We used a multiple linear regression model
109 (MLR) to assess the relationship between concentration, time and storage condition. There was
110 a small but statistically significant decrease in RNA concentration over time ($F = 4.9$, $p = 0.03$)
111 but surprisingly RNA yields from dried and frozen samples did not significantly differ ($F = 0.84$, p
112 $= 0.36$). We assessed the size distribution of recovered RNA and found that RNA grew
113 increasingly fragmented in both dried and frozen flies (**Fig. 1B, Supp. Fig. 1**). In a multiple
114 linear regression model with the same predictors as above, there was a statistically significant
115 decrease in RNA length over time ($F = 19.8$, $p = 4.3 \times 10^{-5}$) but no significant difference in length
116 between dried and frozen samples ($F = 0.4$, $p = 0.53$).

117
118 Host messenger RNA (mRNA) remained reliably detectable by RT-qPCR in both dried
119 and frozen samples. In *D. melanogaster*, we detected ribosomal protein L32 (RpL32) mRNA in
120 72 of 72 frozen flies and 68 of 72 dried flies (**Fig. 1C**). When we used shorter-range primers that
121 amplified a 56 bp region of RpL32 instead of the 110 bp region amplified by our standard
122 primers, 71 of 72 dried flies were positive (**Fig. 1C**).

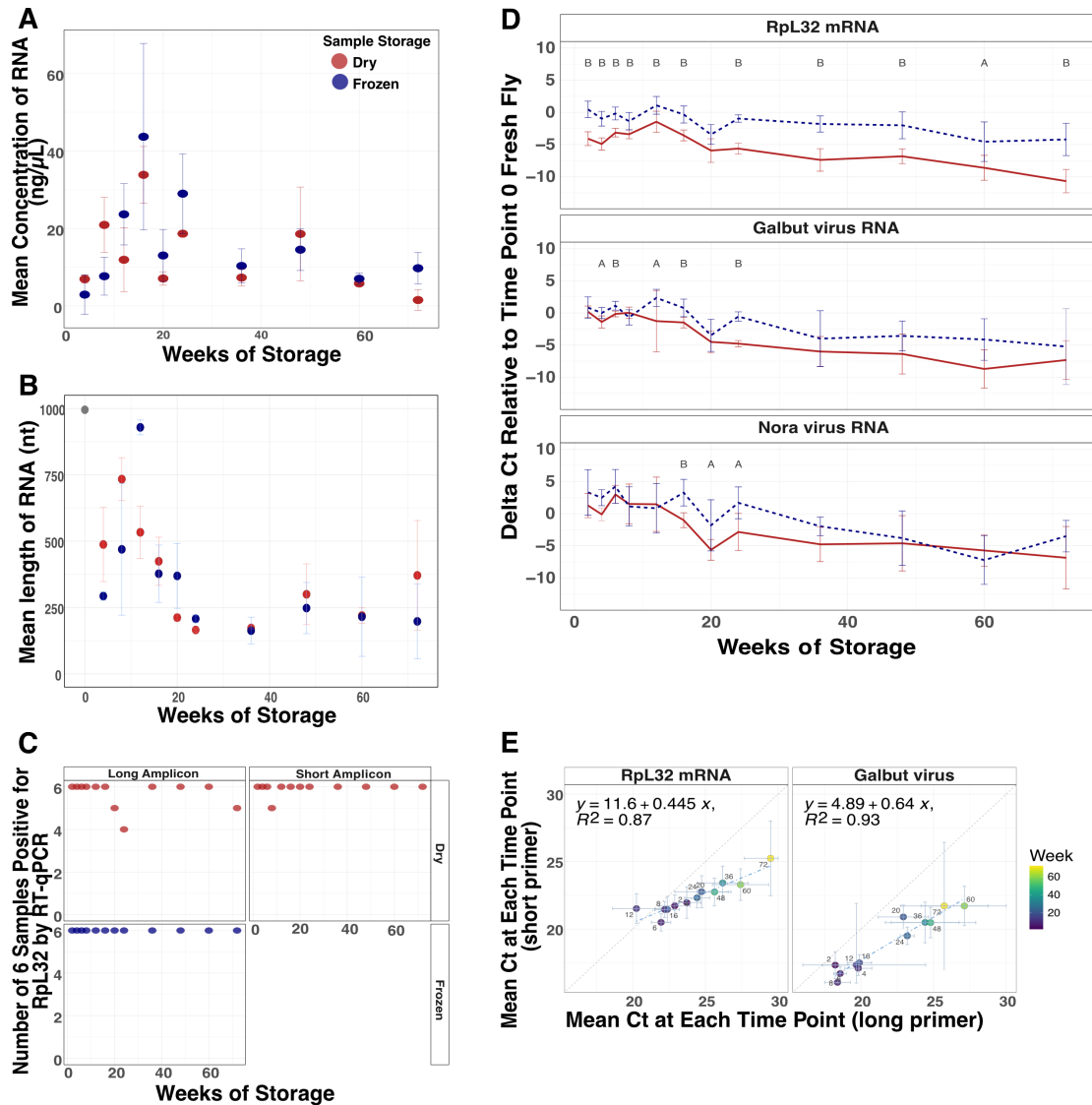
123
124 Although there was no statistically significant difference in RNA concentration and length
125 between dried and frozen flies, there was variability in detected levels of specific host and viral
126 RNA targets (**Fig. 1D; Supp. Fig. 2**). Levels of host and viral RNA measured by RT-qPCR
127 decreased as storage time increased. Levels of RpL32 mRNA were higher in frozen flies than
128 they were in the dried flies (**Fig. 1D**).

129
130 Each qPCR run included one positive control: RNA from known infected flies, and three
131 negative controls: an extraction blank (a no sample extraction), a no-sample RT control, and a
132 no template qPCR control. The use of negative controls corresponding to each step of the
133 extraction and detection process would have allowed us to determine when cross-contamination

134 had been introduced if it had been. All controls behaved as expected, confirming the absence of
135 cross-contamination during extraction and RT-qPCR (**Supp. Fig. 3A**).

136

137 The FoCo-17 *D. melanogaster* population we sampled contained individual flies variably
138 infected by galbut virus, La Jolla virus, Nora virus, and Thika virus^{32,33}. The prevalence of each
139 virus varied in this outbred population as did viral RNA levels in infected individuals. As with
140 RpL32 mRNA, average levels of detected viral RNAs decreased over time in both dried and
141 frozen flies (**Fig. 1D; Supp. Fig. 2A**). Levels of galbut virus dropped by 69x in frozen samples
142 and 181x in dried samples between 0 and 72 weeks. After 24 weeks, there was no significant
143 difference between galbut virus RNA levels in dried and frozen flies. The overall pattern was
144 similar for the three other viruses (**Fig. 1D; Supp Fig. 2**). Shorter range primers detected galbut
145 virus RNA in fewer qPCR cycles, unsurprisingly confirming that short PCR products work better
146 for fragmented RNA (**Fig. 1E**).



147

148 **Figure 1: RNA survived in dried flies over 72 weeks.** (A) Average RNA concentration and
 149 (B) length of RNA from dried and frozen flies over 72 weeks. N = 3 female *D. melanogaster*
 150 from each storage condition at each timepoint. (C) Number of samples positive of 3 male and 3
 151 female flies at each timepoint for RpL32 mRNA via RT-qPCR using primers that target long (110
 152 nt) and short (56 nt, dried only) regions of RpL32 mRNA. (D) Average difference of RpL32
 153 mRNA, galbut virus and Nora virus RNA levels relative to fresh *D. melanogaster* collected at the
 154 initial timepoint. A two-tailed t-test was used to determine the statistical significance between
 155 dried and frozen means at every time point (A < 0.05, B < 0.001). (E) Comparison of mean Ct
 156 value at each time point of dried *D. melanogaster* using primers that target a long (x-axis) or

157 short (y-axis) region of RpL32 mRNA and galbut virus. Error bars represent mean plus and
158 minus the standard deviation. N = 3 male and 3 female at each time point.

159

160 RNA also survived well in dried mosquitoes (**Supp. Fig 4**). An MLR was used to assess
161 whether time and/or storage condition impacted mosquito RNA concentration. Over time (F =
162 2.09, p = 0.15) and between dried and frozen mosquitoes (F = 2.37, p = 0.13) there was no
163 statistically significant difference in mosquito RNA concentrations (**Supp. Fig. 4A**). A caveat is
164 that we neglected to save fresh mosquitoes at time 0 as we did with flies, so RNA levels were
165 normalized to levels in the 4-week frozen samples, which had likely already experienced
166 degradation. There was no statistically significant difference in RNA length over time after 4
167 weeks (F = 0.75, P = 0.39) but there was by storage condition (F = 79.3, p = 2.5×10^{-13} ; **Supp.**
168 **Fig. 4B**).

169

170 We used RT-qPCR to assess levels and detectability of specific host and viral RNAs in
171 stored mosquitoes. We detected actin mRNA in all 78 dried and all 78 frozen mosquitoes
172 (**Supp. Fig. 4C**). Actin mRNA levels remained consistent over time with no difference between 4
173 weeks and 52 weeks in the frozen samples and a 6x decrease in the dried samples. This
174 mosquito population harbored verdadero virus and Guadeloupe mosquito virus^{30,34}. Viral RNAs
175 decreased over time in the dried and frozen specimens. In the dried mosquitoes, virus levels
176 dropped between 13x for verdadero virus and 18x for Guadeloupe mosquito virus between 4
177 and 52 weeks. In frozen mosquitoes there was no difference in verdadero virus levels between
178 4 weeks and 52 weeks but there was a 5x decrease for Guadeloupe mosquito virus (**Supp. Fig.**
179 **4D-E**).

180

181 The mosquito positive and negative controls behaved as expected except for the cDNA
182 positive control sample at week 28, which was negative for Guadeloupe mosquito virus. This
183 may reflect a true negative infection status, given that our cDNA positive control samples were
184 individual mosquitoes from a variably infected outbred population confirmed to have a
185 verdadero virus infection (**Supp. Fig. 3B**).

186

187 ***Drosophila* obtained for RNA metagenomic sequencing were 10-125 years old and from**
188 **entomological collections across the United States and Canada**

189

190 Having determined that RNA survives well in experimentally dried insects, we turned our
191 attention to much older samples. We obtained 46 specimens from entomological collections
192 across the United States and Canada (**Table 1**). Thirty-two had been assigned as *D.*
193 *melanogaster* based on morphology, nine as *D. simulans*, and five as unknown drosophilids.
194 Collection dates ranged from 1896 to 2011 (**Fig. 2A; Table 1; Supp. Table 1**). We extracted
195 RNA from 29 specimens following a standard destructive protocol, which yielded RNA
196 concentrations from 0 to 152 ng/μL (**Fig. 2B; Table 1; Supp. Table 1**). Some samples were
197 loaned on the condition that they not be destroyed, so we followed a non-destructive RNA
198 isolation protocol for the remaining samples³⁵. This used an overnight incubation with proteinase
199 to liberate internal contents of the specimen while leaving morphology intact. The non-
200 destructive extractions yielded more RNA than the standard protocol ($p = 0.03$), with
201 concentrations ranging from 4 to 120 ng/μL (**Fig. 2B; Table 1; Supp. Table 1**). However, when
202 samples from Hawai'i, which had unusually low yields, were removed there was no statistically
203 significant difference between RNA concentration obtained from the two methods ($p = 0.59$;
204 **Supp. Fig. 5**). Four samples that had been stored in 70% ethanol required rehydration prior to
205 extraction (**Supp. Fig 5**).

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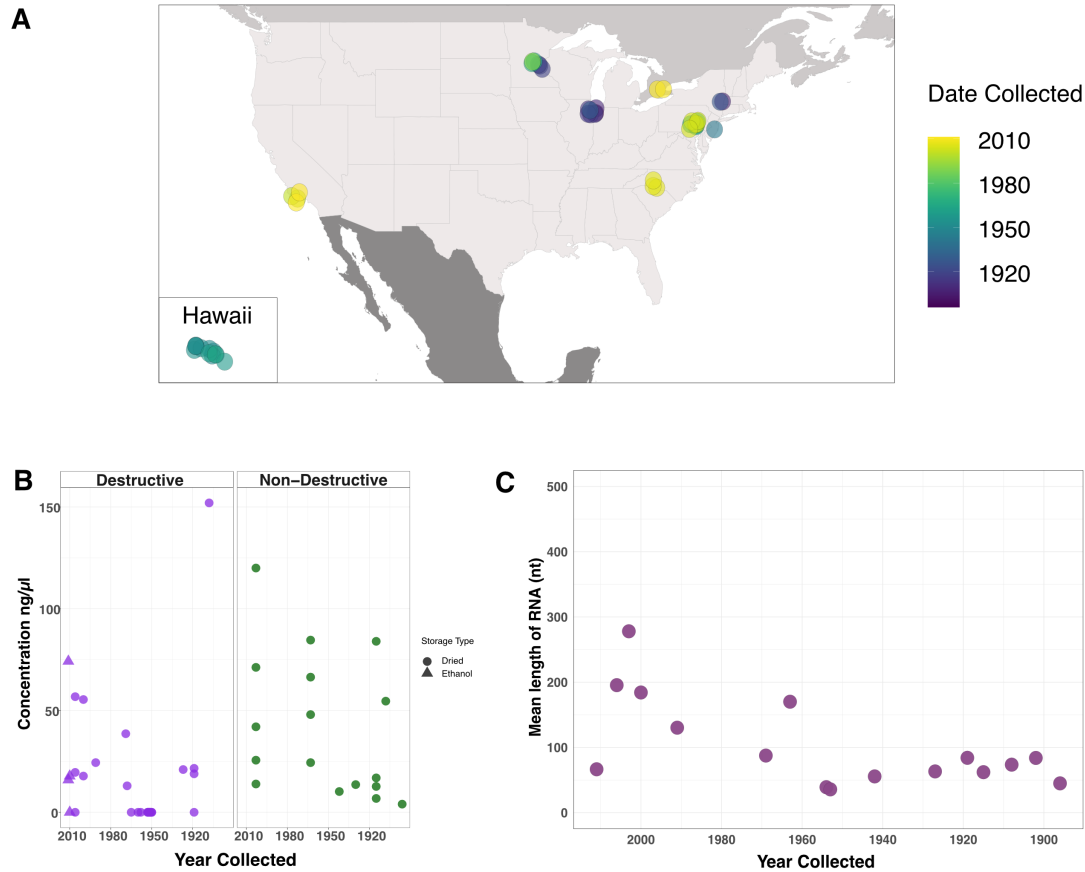
224 **Table 1: Museum specimen metadata.**

Sequencing Based Species ID	Date Collected	Location (1)	Donating Institution	BioSample (2)	Sample ID (3)	Museum Accession	Storage Type	Extraction Method	Concentration (ng/μl) (4)
<i>D. melanogaster</i>	1896	Urbana, IL	Illinois Natural History Survey	SAMN38071661	Illinois_1896	1004284	Dried	Non-Destructive	4
<i>D. melanogaster</i>	1908	Albany, NY	New York State Museum	SAMN38071662	NewYork_1908	-	Dried	Destructive	152
<i>D. melanogaster</i>	1908	Urbana, IL	Illinois Natural History Survey	SAMN38071654	Illinois_1908	1004277	Dried	Non-Destructive	54.6
<i>D. melanogaster</i>	1915	Urbana, IL	Illinois Natural History Survey	SAMN38071656	Illinois_1915_1	1004279	Dried	Non-Destructive	84
<i>D. putrida</i>	1915	Urbana, IL	Illinois Natural History Survey	SAMN38071657	Illinois_1915_2	1004280	Dried	Non-Destructive	6.8
<i>D. melanogaster</i>	1915	Urbana, IL	Illinois Natural History Survey	SAMN38071658	Illinois_1915_3	1004281	Dried	Non-Destructive	12.7
<i>D. melanogaster</i>	1915	Urbana, IL	Illinois Natural History Survey	SAMN38071659	Illinois_1915_4	1004282	Dried	Non-Destructive	16.9
<i>D. melanogaster</i>	1919	St. Paul, MN	University of Minnesota, Insect Collection	SAMN38090828	Minnesota_1919_1	-	Dried	Destructive	0
<i>D. melanogaster</i>	1919	St. Paul, MN	University of Minnesota, Insect Collection	SAMN38071687	Minnesota_1919_2	-	Dried	Destructive	21.6
<i>D. melanogaster</i>	1919	St. Paul, MN	University of Minnesota, Insect Collection	SAMN38071688	Minnesota_1919_3	-	Dried	Destructive	18.9
<i>D. melanogaster</i>	1927	Albany, NY	New York State Museum	SAMN38071663	NewYork_1927	-	Dried	Destructive	21
<i>D. melanogaster</i>	1930	Urbana, IL	Illinois Natural History Survey	SAMN38071660	Illinois_1930	1004283	Dried	Non-Destructive	13.6
<i>D. melanogaster</i>	1942	Glassboro, NJ	Illinois Natural History Survey	SAMN38071655	NewJersey_1942	1004278	Dried	Non-Destructive	10.2
<i>D. melanogaster</i>	1950	Oahu, HI	University of Hawaii, Insect Museum	-	Hawaii_1950_1	-	Dried	Destructive	0
<i>D. melanogaster</i>	1950	Oahu, HI	University of Hawaii, Insect Museum	-	Hawaii_1950_2	-	Dried	Destructive	0
<i>D. melanogaster</i>	1951	Honolulu, HI	University of Hawaii, Insect Museum	-	Hawaii_1951_1	-	Dried	Destructive	0
<i>D. melanogaster</i>	1951	Oahu, HI	University of Hawaii, Insect Museum	-	Hawaii_1951_2	-	Dried	Destructive	0
<i>D. melanogaster</i>	1952	Kauai, HI	University of Hawaii, Insect Museum	-	Hawaii_1952_1	-	Dried	Destructive	0
<i>D. melanogaster</i>	1953	Kauai, HI	University of Hawaii, Insect Museum	SAMN38071677	Hawaii_1953_1	-	Dried	Destructive	0
<i>D. melanogaster</i>	1963	Lebanon Co, PA	Frost Entomological Museum	SAMN38071668	Pennsylvania_1963_1	-	Dried	Non-Destructive	66.4
<i>D. melanogaster</i>	1963	Lebanon Co, PA	Frost Entomological Museum	SAMN38071669	Pennsylvania_1963_2	-	Dried	Non-Destructive	24.4
<i>D. melanogaster</i>	1968	St. Cloud, MN	University of Minnesota, Insect Collection	SAMN38071684	Minnesota_1968	-	Dried	Destructive	13
<i>D. melanogaster</i>	1969	St. Cloud, MN	University of Minnesota, Insect Collection	SAMN38071685	Minnesota_1969	-	Dried	Destructive	38.6
<i>D. melanogaster</i>	1991	St. Cloud, MN	University of Minnesota, Insect Collection	SAMN38071686	Minnesota_1991	-	Dried	Destructive	24.4
<i>D. putrida</i>	2003	Lebanon Co, PA	Frost Entomological Museum	SAMN38071673	Pennsylvania_2003_1	4908	Dried	Non-Destructive	71.2
<i>D. putrida</i>	2003	Lebanon Co, PA	Frost Entomological Museum	SAMN38071672	Pennsylvania_2003_2	4887	Dried	Non-Destructive	13.9
<i>S. pallida</i>	2003	Lebanon Co, PA	Frost Entomological Museum	SAMN38071675	Pennsylvania_2003_3	5867	Dried	Non-Destructive	42
<i>S. pallida</i>	2003	Lebanon Co, PA	Frost Entomological Museum	SAMN38071676	Pennsylvania_2003_4	5873	Dried	Non-Destructive	120
<i>S. pallida</i>	2003	Lebanon Co, PA	Frost Entomological Museum	SAMN38071674	Pennsylvania_2003_5	5863	Dried	Non-Destructive	25.6
<i>D. melanogaster</i>	2006	Davidson, NC	The Davidson College Entomology Collection	SAMN38071666	NorthCarolina_2006_1	-	Dried	Destructive	19.6
<i>D. melanogaster</i>	2006	Davidson, NC	The Davidson College Entomology Collection	SAMN38071667	NorthCarolina_2006_2	-	Dried	Destructive	56.8
<i>D. melanogaster</i>	2006	Davidson, NC	The Davidson College Entomology Collection	-	NorthCarolina_2006_3	-	Dried	Destructive	0
<i>D. melanogaster</i>	2010	Puslinch Township, Ontario, Canada	University of Guelph, Centre for Biodiversity Genomics	SAMN38071680	Canada_2010_1	PHDIP1072-11	Ethanol	Destructive	0
<i>D. simulans</i>	1952	Kauai, HI	University of Hawaii, Insect Museum	-	Hawaii_1952_2	-	Dried	Destructive	0
<i>D. simulans</i>	1952	Kauai, HI	University of Hawaii, Insect Museum	-	Hawaii_1952_3	-	Dried	Destructive	0
<i>D. simulans</i>	1953	Klauea, HI	University of Hawaii, Insect Museum	SAMN38071678	Hawaii_1953_2	-	Dried	Destructive	0
<i>D. simulans</i>	1958	Maui, HI	University of Hawaii, Insect Museum	-	Hawaii_1958	-	Dried	Destructive	0
<i>D. simulans</i>	1960	Oahu, HI	University of Hawaii, Insect Museum	-	Hawaii_1960	-	Dried	Destructive	0
<i>D. simulans</i>	1963	Lebanon Co, PA	Frost Entomological Museum	SAMN38071670	Pennsylvania_1963_3	-	Dried	Non-Destructive	84.6
<i>D. simulans</i>	1903	Lebanon Co, PA	Frost Entomological Museum	SAMN09071671	Pennsylvania_1903_4	-	Dried	Non-Destructive	49
<i>D. simulans</i>	1965	Oahu, HI	University of Hawaii, Insect Museum	SAMN38071679	Hawaii_1965	-	Dried	Destructive	0
<i>D. simulans</i>	2000	Santa Barbara, CA	Santa Barbara Museum of Natural History	SAMN38071683	California_2000_1	-	Dried	Destructive	17.8
<i>D. simulans</i>	2000	Santa Barbara, CA	Santa Barbara Museum of Natural History	SAMN38071682	California_2000_2	-	Dried	Destructive	55.4
<i>D. simulans</i>	2010	Puslinch Township, Ontario, Canada	University of Guelph, Centre for Biodiversity Genomics	SAMN38071681	Canada_2010_2	PHDIP946-11	Ethanol	Destructive	17.8
<i>D. simulans</i>	2011	Ventura Co, CA	University of Guelph, Centre for Biodiversity Genomics	SAMN38071664	California_2011_1	BBDIV1549-12	Ethanol	Destructive	74.2
<i>D. simulans</i>	2011	Ventura Co, CA	University of Guelph, Centre for Biodiversity Genomics	SAMN38071665	California_2011_2	BBDIV1550-12	Ethanol	Destructive	15.9

(1) All specimens from USA unless otherwise noted
(2) Specimens that did not yield sufficient sequencing library were not sequenced and thus have no BioSample ID
(3) Our study sample ID
(4) RNA concentrations below the limit of detection on the Qubit were assigned a concentration of 0

225
226 RNA from samples collected after or before 1960 had average lengths of 159 and 60 nt,
227 respectively (**Fig. 2C**). Although there was variability in RNA quantity and quality, the RNA
228 obtained from these specimens was of suitable length and concentration for downstream
229 analyses.

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231



232

233 **Figure 2: Museum sample collection locations and characteristics of extracted RNA.** (A)

234 Map of the United States and Canada indicating sample collection locations. (B) RNA

235 concentration from individual museum flies using either the destructive (purple) or non-

236 destructive (green) extraction method. (C) Mean length of RNA molecules from a subset of

237 museum specimens encompassing the range of years specimens obtained for this study had

238 been collected.

239

240 We screened museum samples for galbut virus RNA and RpL32 mRNA using our short-

241 range primers. Sixteen of 46 samples (35%) were positive for galbut virus and 21 (46%) were

242 positive for RpL32 mRNA (**Supp. Fig. 6A, Supp. Table 1**). Positive status was confirmed using

243 melting temperature and agarose gel electrophoresis. The small size of PCR products made

244 them difficult to distinguish from primer dimers on agarose gels, so it is possible that there were

245 false negative calls. All negative control samples were negative as expected (**Supp. Fig. 6B**).

246

247 **Metagenomic sequencing of museum samples and molecular species assignment**

248

249 We prepared shotgun sequencing libraries from museum sample RNA. Thirty-six
250 samples yielded libraries suitable for sequencing. Sequencing yielded an average of 48 million
251 reads per sample (range: 0.4-238 million). Several samples were sequenced multiple times to
252 increase recovery of complete virus sequences.

253

254 The records for some museum samples did not include morphological taxonomic
255 assignments below the family level, so we used competitive mapping to generate molecular
256 taxonomic assignments. We mapped quality and adapter trimmed reads to a set of 545
257 representative cytochrome oxidase subunit-1 (CO1) sequences from across the Drosophilidae
258 family to generate a molecular species identification for each sample. This molecular species
259 assignment corroborated the museum morphology-based assignment for most samples (**Supp.**
260 **Table 2**). Four samples labeled as *D. melanogaster* in museum records were revealed to be *D.*
261 *simulans* by CO1-mapping. *D. melanogaster* and *D. simulans* are common, sympatric, and
262 difficult to distinguish morphologically, so such misassignment is expected³⁶. Five samples did
263 not have species-level taxonomic assignments from the museums. Of these, CO1-mapping
264 indicated that 2 were *Drosophila putrida* and the remaining three were *Scaptomyza pallida*. One
265 fly collected in Illinois in 1915 was also assigned as *D. putrida* (**Supp. Table 1-2**). These
266 assignments are consistent with the known range of *D. putrida* and *S. pallida*³⁷. All fresh frozen
267 and experimental dried samples were assigned as *D. melanogaster* as expected and there were
268 no CO1-mapping reads in the three water negative control datasets.

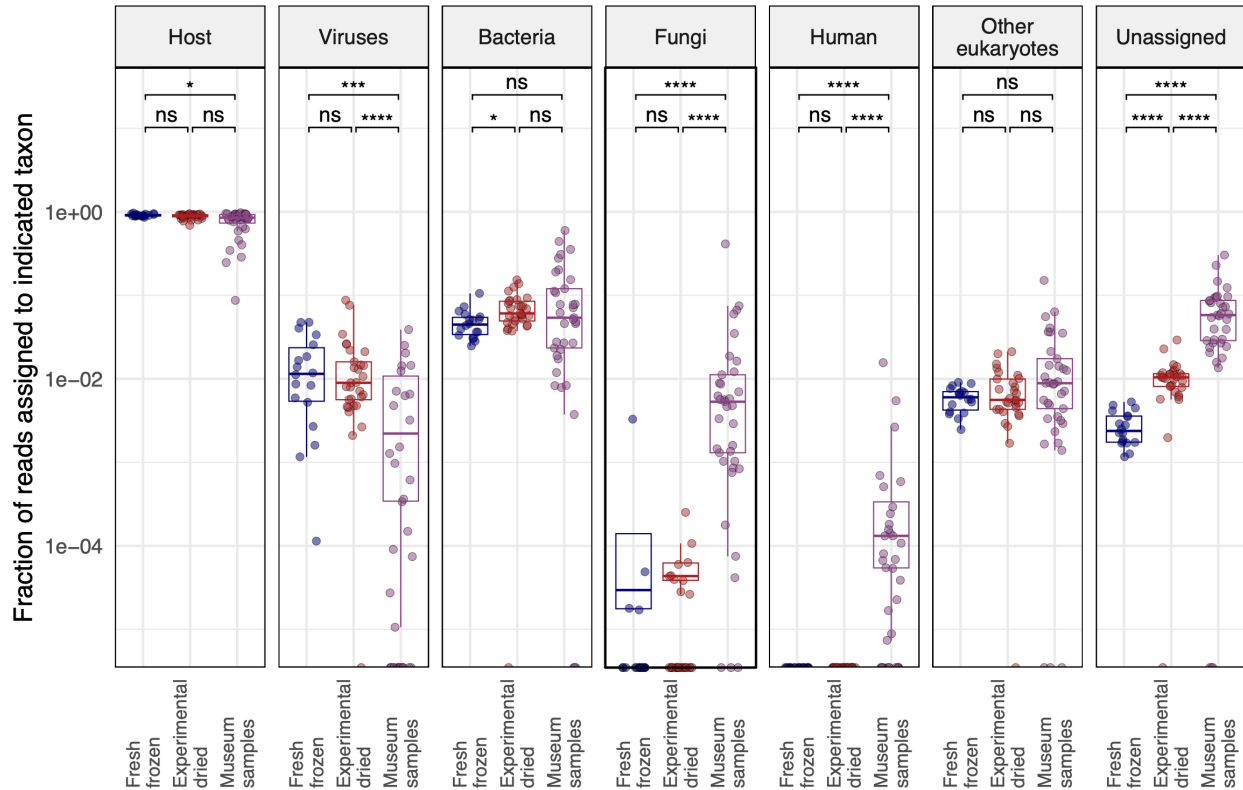
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270 **The overall taxonomic composition of new and old samples was similar**

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272 To broadly assess the source of sequences in our datasets, we taxonomically binned
273 sequences using our lab's metagenomic classification pipeline³⁸. This first identified host-derived
274 reads by mapping to an index composed of the *D. melanogaster*, *D. putrida*, and *S. pallida*
275 genomes^{39,40}. The pipeline then assembled remaining non-host reads and taxonomically
276 assigned the resulting contigs and unassembled reads using a BLASTN search of the NCBI
277 nucleotide database. We tabulated the fraction of reads in each dataset that were assigned to
278 various taxa or that remained unassigned (**Fig. 3**).

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281

282 **Figure 3: The overall taxonomic composition of old and new samples was similar.** The

283 fraction of reads assigned to the indicated taxa are plotted. Each point represents a dataset

284 from an individual fly. Adjusted p -values significance levels from Wilcoxon test are

285 indicated. Host-mapping reads were identified by mapping to combined *D. melanogaster*, *D.*

286 *putrida*, and *S. pallida* genomes. Fractions of reads mapping to other taxa were determined by

287 metagenomic classification of non-host-mapping reads. The “other eukaryotes” category

288 accounts for all eukaryotic taxa apart from fungi and human. Unassigned read fractions account

289 for reads that were not assigned by host-mapping or metagenomic classification.

290

291 Most reads mapped to the fly genomes in all datasets, although the median fraction of

292 host-mapping reads fell from 90.5% in fresh samples to 85% in museum samples (**Fig. 3**). This

293 decrease was accompanied by a corresponding increase in unassigned reads. Only 0.2% of

294 reads were unassigned in fresh datasets, compared to 5.5% in museum datasets ($p=2.3 \times 10^{-6}$).

295 This increase in unassignable reads is likely attributed to the fact that short reads from

296 fragmented RNA contain less information for host mapping or taxonomic assignment. There

297 were on average fewer virus-mapping reads in museum samples, which could reflect the fact

298 that our FoCo-17 population was persistently infected by multiple viruses or could reflect

299 decreased relative survival of viral RNA in old samples. There were increases in the proportion
300 of fungi- and human-mapping reads in museum datasets. Fungal reads in old samples may
301 have derived from saprophytic fungi. Human-mapping reads in museum datasets may have
302 originated from handling of specimens by curators, accumulation of dust on samples, or an
303 increased fraction of contamination-derived sequences in old datasets. The median fraction of
304 human-mapping reads in old datasets was 5.4×10^{-5} and the maximum fraction was 1.6% of
305 reads, in a dataset from Hawai'i. We concluded that some of the reads in our libraries, including
306 human-assigned reads, may have derived from contamination. However, these accounted for a
307 small fraction of datasets and the overall taxonomic composition of new and old datasets was
308 similar (**Fig. 3**).

309

310 **Entomological specimens harbored diverse viral sequences including previously** 311 **unknown viruses**

312

313 We recovered at least one virus sequence from 21 of the 36 sequenced museum
314 samples and two or more virus sequences from 5 samples (**Fig. 4; Table 2**). All but three of the
315 virus sequences corresponded to known *Drosophila*-infecting viruses^{25,26}. Galbut virus was the
316 most common virus, with six samples producing coding-complete genomes and six samples
317 producing partial galbut virus genomes (**Fig. 4; Table 2**). The next most common viruses were
318 *D. melanogaster* sigma virus and vera virus, which were detected in three samples each (**Fig. 4;**
319 **Table 2**). The sample with the most viral sequences was collected in 1908 in Illinois, USA: this
320 fly yielded three coding-complete and one partial virus sequence.

321

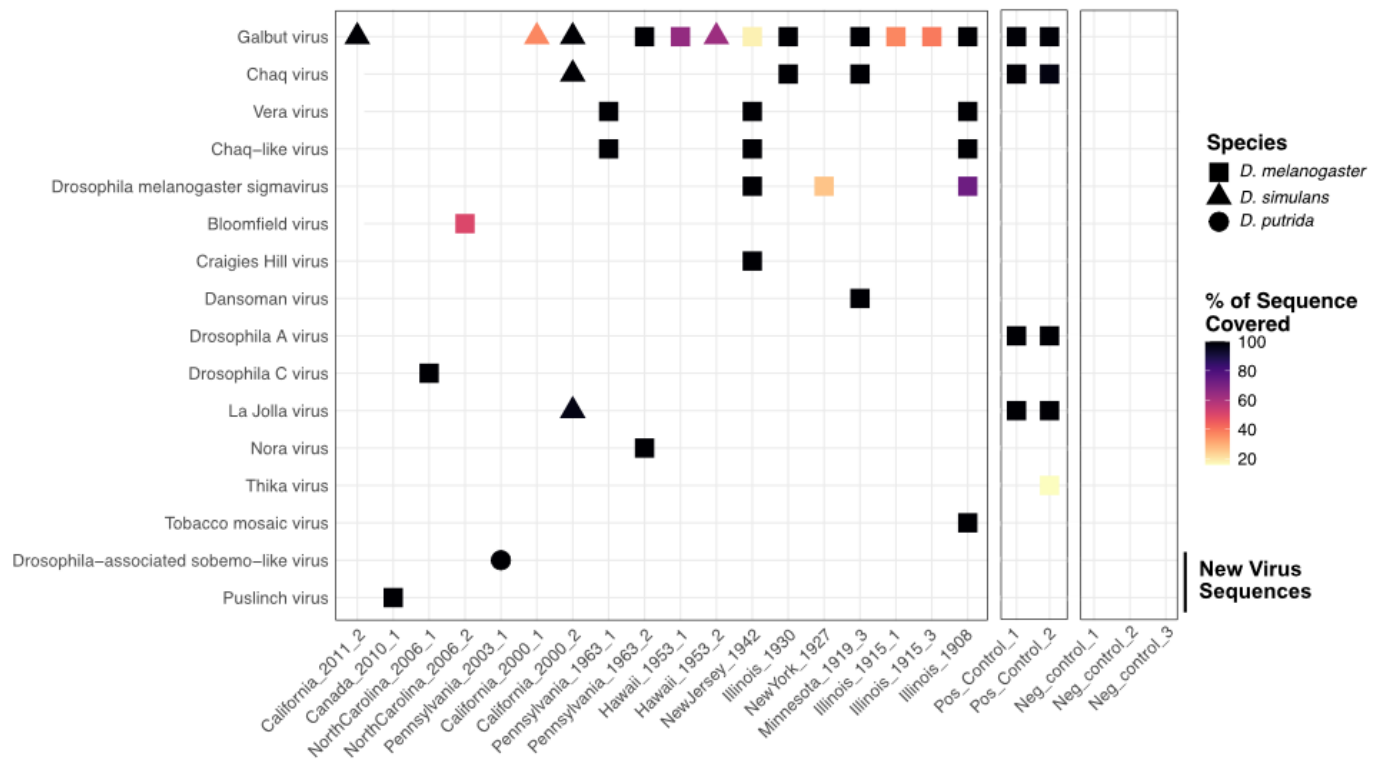
322 We identified two coding-complete genomes from previously undescribed viruses. The
323 first was a bunyavirus from a *D. melanogaster* collected in Ontario, Canada in 2010, which we
324 named Puslinch virus. The Puslinch virus L protein shares only ~35% pairwise amino acid
325 identity with its nearest known relatives, which were viruses in the genus Herbevirus (**Fig. 4;**
326 **Fig. 5; Table 2**). The other new virus sequence was a sobemo-like virus from a *D. putrida*
327 collected in Pennsylvania, USA in 2003 (**Fig. 4; Fig 5; Table 2**).

328

329 Additionally, we recovered a complete tobacco mosaic virus (TMV) genome with high
330 coverage depth (850x) from the 1908 *D. melanogaster* specimen. We speculated that this
331 sequence may have originated from contamination of the specimen, perhaps during handling by
332 a smoker⁴¹.

333

334 Positive control libraries, constructed from RNA from pooled FoCo-17 flies, contained the
 335 expected virus sequences (**Fig. 4**). To avoid potential read contamination via index hopping, we
 336 sequenced positive control libraries on a separate sequencing run from the museum
 337 specimens⁴². Negative control datasets were included in each run and contained fewer than 15k
 338 reads after trimming of adapter and low-quality bases. Negative control datasets contained zero
 339 reads mapping to any of the virus sequences present in any of the other datasets (**Fig. 4**)³⁹.



340

341

342 **Figure 4. Entomological museum specimens harbored diverse viral sequences.** Heatmap
 343 showing virus sequences recovered from individual museum samples. Shape corresponds to
 344 host species and color corresponds to the percent of reference sequence covered (**Table 2**).
 345 For segmented viruses, the average coverage across all segments is shown. Viruses detected
 346 in positive control datasets, made from pooled FoCo17 flies, are shown. No reads mapping to
 347 any of these viruses were present in any of the three negative control datasets.

348

349 **Table 2: Virus sequences identified in *Drosophila* museum collection specimens.**

Date Collected	Location	Sample ID	Virus	GenBank Sequence Accession	Nearest GenBank ⁷	% Query Coverage ^{2,3}	% Identity ^{4,5}	Completeness	Average Coverage ⁶	Estimated Evolutionary Rate ⁸
Known										
1908	Illinois	Illinois_1908	<i>Drosophila melanogaster</i> sigma virus	-	MH384277	73	99.4	Partial sequence	2	-
1908	Illinois	Illinois_1908	Tobacco mosaic virus	OR820564	KY810785	100	99.7	Coding complete sequence	850	7.13E-05
1908	Illinois	Illinois_1908	Vera virus	OR820566, OR820565	MT742171, MT742172	100	99.9 - 100	Coding complete sequence	9-34	1.09E-05 - 0.00E-06
1908	Illinois	Illinois_1908	Galbut virus	OR820562, OR820561, OR820563	OR729093, MT742165, OR729068	100	98.9 - 99.3	Coding complete sequence	7-14	6.43E-05 - 1.00E-04
1908	Illinois	Illinois_1908	Cha-q-like virus	OR820560	MT742173	100	99.9	Coding complete sequence	18	6.36E-06
1915	Illinois	Illinois_1915_1	Galbut virus	-	MT742164, MT742161, MT742162	37	96.1	Partial sequence	4	-
1915	Illinois	Illinois_1915_3	Galbut virus	-	MT742164, MT742161, MT742162	39	96.5	Partial sequence	1	-
1919	Minnesota	Minnesota_1919_3	Dansoman virus	OR820572, OR820573	MH384295, MH384270	100	97.4 - 98.3	Coding complete sequence	40-46	2.93E-04 - 3.93E-04
1919	Minnesota	Minnesota_1919_3	Cha-q virus	OR820571	MH384367	100	96.9	Coding complete sequence	317	4.11E-04
1919	Minnesota	Minnesota_1919_3	Galbut virus	OR820575, OR820574, OR820576	MT742164, MT742165, OR729068	100	99.4 - 99.6	Coding complete sequence	247-523	4.55E-05 - 9.31E-05
1927	New York	NewYork_1927	<i>Drosophila melanogaster</i> sigma virus	-	NC_038281	26	99.2	Partial sequence	2	-
1930	Illinois	Illinois_1930	Cha-q virus	OR820567	MH384311	100	97.1	Coding complete sequence	32	4.88E-04
1930	Illinois	Illinois_1930	Galbut virus	OR820569, OR820568, OR820570	OR729094, MT742165, OR729070	100	89.3 - 99.7	Coding complete sequence	29-50	3.64E-05 - 1.19E-03
1942	New Jersey	NewJersey_1942	Craigies Hill virus	OR820579, OR820578	MH384377, MH384349	100	90.4 - 98.1	Coding complete sequence	22-40	2.95E-04 - 1.46E-03
1942	New Jersey	NewJersey_1942	Galbut virus	-	MT742164, MT742161, MT742162	18	97.1	Partial sequence	17	-
1942	New Jersey	NewJersey_1942	Cha-q-like virus	OR820577	MT742173	100	99.9	Coding complete sequence	490	9.21E-06
1942	New Jersey	NewJersey_1942	Vera virus	OR820582, OR820581	MT742171, MT742172	100	99.9 - 100	Coding complete sequence	463-686	1.58E-05 - 0.00E-06
1942	New Jersey	NewJersey_1942	<i>Drosophila melanogaster</i> sigma virus	OR820580	MH384306	100	99.2	Coding complete sequence	46	1.88E-04
1953	Hawai'i	Hawaii_1953_1	Galbut virus	-	MT742164, MT742161, MT742162	65	92.4	Partial sequence	3	-
1953	Hawai'i	Hawaii_1953_2	Galbut virus	-	MT742164, MT742161, MT742162	63	99.7	Partial sequence	3	-
1963	Pennsylvania	Pennsylvania_1963_1	Vera virus	OR820593, OR820592	MT742168, MT742172	100	99.6 - 100	Coding complete sequence	2512-2727	6.55E-05 - 0.00E-06
1963	Pennsylvania	Pennsylvania_1963_1	Cha-q-like virus	OR820591	MT742173	100	99.8	Coding complete sequence	5323	3.64E-05
1963	Pennsylvania	Pennsylvania_1963_2	Galbut virus	OR820588, OR820587, OR820589	MH384303, MH384304, MH384276	100	96.4 - 99.2	Coding complete sequence	8-81	1.73E-04 - 7.98E-04
1963	Pennsylvania	Pennsylvania_1963_2	Nora virus	OR820590	JX220408	100	97.1	Coding complete sequence	16	5.98E-04
2000	California	California_2000_2	La Jolla virus	OR820598	MH384285	97	95.35	Partial sequence	11	5.81E-03
2000	California	California_2000_2	Cha-q virus	OR820594	MT742163	100	82.8	Coding complete sequence	62	1.01E-02
2000	California	California_2000_2	Galbut virus	OR820596, OR820595, OR820597	MH384283, MH384336, MH384366	100	83.5 - 92.4	Coding complete sequence	150-271	6.00E-04 - 2.07E-02
2000	California	California_2000_1	Galbut virus	-	MT742164, MT742161, MT742162	37	95	Partial sequence	2	-
2006	North Carolina	NorthCarolina_2006_1	<i>Drosophila</i> C virus	OR820583	OK188767	100	98.3	Coding complete sequence	264	1.86E-03
2006	North Carolina	NorthCarolina_2006_2	Bloomfield virus	-	MF416371, KP714091, KP714093, KP714094	50	99.6 - 100	Partial sequence	0.4-2	-
2011	California	California_2011_2	Galbut virus	OR820600, OR820599, OR820601	MH384283, MH384336, MH384366	100	83.8 - 99.4	Coding complete sequence	16-18	2.00E-03 - 2.61E-02
Novel										
2003	Pennsylvania	Pennsylvania_2003_1	<i>Drosophila</i> -associated sobemo-like virus	OR820603, OR820602	UYL94340.1, QHA33877.1	100	38.1 - 70.6	Coding complete sequence	84-136	-
2010	Ontario, CAN	Canada_2010_1	Puslinch virus	OR820586, OR820585, OR820584	YP_009362026.1, YP_010840683.1, YP_009362024.1	100	32.8 - 39.1	Coding complete sequence	81-188	-

¹ Nearest GenBank sequence is provided for RdRp of *Drosophila*-associated sobemo-like virus and the L, M and S segment of Puslinch virus.

² % Query coverage from BLASTN alignment

³ % Query coverage from BLASTN alignment for novel virus sequences was determined based on nearest GenBank sequence.

⁴ nt, percentage nucleotide identity to closest GenBank sequence identified via BLASTn.

⁵ For novel virus sequences, % nt identity to the reference sequence is shown.

⁶ Average mapped read coverage across contigs as reported in Geneious.

⁷ Calculated using the nearest GenBank sequence with date collected data.

351 **Relatedness of old virus sequences to contemporary ones**

352

353 Sequences from museum samples of previously described viruses ranged from 83.5% to
354 100% identical to existing sequences with 38% of the sequences sharing $\geq 99\%$ pairwise
355 identity to existing sequences (**Table 2**). Notably, all vera virus RNA 1 sequences, including one
356 from 1908, were 100% identical to their closest available existing sequence, though the vera
357 virus RNA 2 and chaq-like sequences from the same samples were not identical to existing
358 sequences.

359

360 We initially considered a Bayesian approach to generate evolutionary rate estimates but
361 found our data had insufficient temporal signal⁴³. We instead generated initial estimates of
362 evolutionary rates using differences in sequence identity and sampling times for the most
363 closely related contemporary sequences (**Supp. Fig. 7; Table 2**,). This may underestimate
364 rates due to saturation and might overestimate rates because the old sequences are likely not
365 the direct ancestors of the contemporary sequences^{44–46}. Sequences without a suitably similar
366 contemporary sequence (e.g., USA 2000 and 2011 galbut virus *D. simulans* RNA 2 and 3) were
367 not analyzed in this way. Estimated rates were generally higher in sequences from more recent
368 samples (2000-2011) and lower in sequences from older samples, with sequences from 1908
369 showing the lowest rates (**Supp. Fig. 7**). This observation is consistent with the time-dependent
370 rate phenomenon, in which evolutionary rate estimates decrease as the timescale of
371 measurement increases^{44,47}.

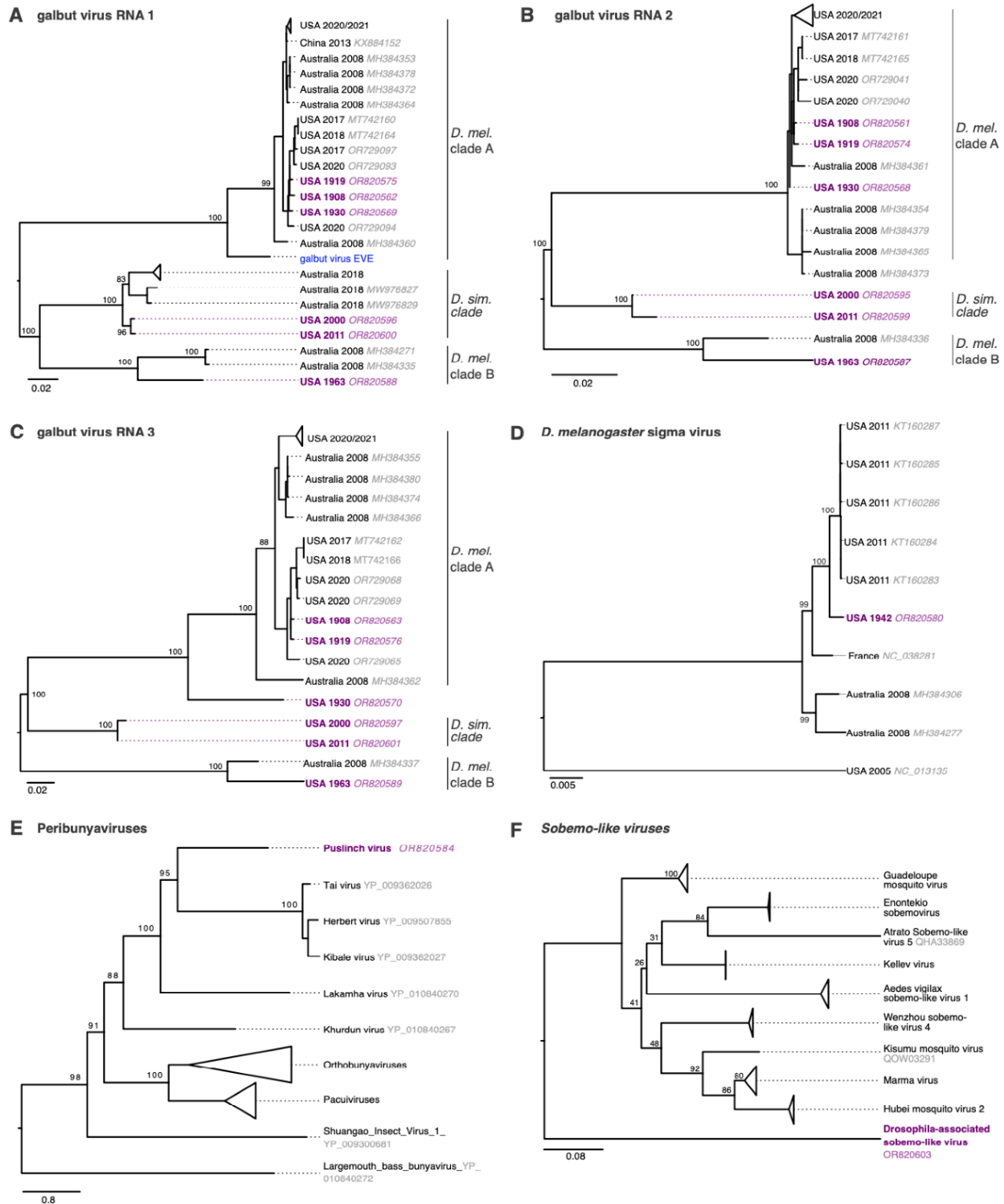
372

373 The most common contemporary viral infection of *D. melanogaster* is galbut virus, the
374 success of which is likely attributable to efficient biparental vertical transmission and minimal
375 apparent fitness costs^{26,30,48}. Detection of galbut virus in museum samples by sequencing
376 matched detection by RT-qPCR (**Table 1**). In maximum likelihood trees, three clades of galbut
377 virus were evident: two clades (A and B) consisted of sequences from *D. melanogaster* while
378 the third clade consisted of sequences from *D. simulans* (**Fig. 5A-C**). An endogenized galbut
379 virus RNA 1 sequence was on its own branch most closely related to clade A (**Fig. 5A**)⁴⁹. All the
380 historic galbut virus sequences fell within existing galbut virus diversity and clustered with other
381 sequences from the same host i.e., the two *D. simulans* sequences clustered with previously
382 described *D. simulans* sequences and the *D. melanogaster* sequences with known *D.*
383 *melanogaster* RNA1 sequences (**Fig. 5A**). Interestingly, the RNA1 and RNA2 sequence from
384 Illinois 1930 clustered with clade A but this virus's RNA 3 was situated on a separate branch.

385 The 1930 Illinois RNAs 1 and 2 were 99% identical to contemporary sequences in GenBank
386 while RNA 3 was only 89% identical. This phylogenetic discordance is consistent with galbut
387 virus reassortment⁵⁰.

388

389 The sigma virus sequence from 1942 clustered within the previously described diversity
390 of *D. melanogaster* sigma viruses (**Fig. 5D**). Puslinch virus and *Drosophila*-associated sobemo-
391 like virus sequences were situated on their own branches on trees of related sequences (**Fig.**
392 **5E-F**). These new virus sequences are highly divergent and may establish new genera or higher
393 order taxa.



394

395

396

397

398

Figure 5: Trees showing museum sample virus sequences in the context of related contemporary sequences. (A) Maximum likelihood trees for galbut virus RNA 1 (RdRp) sequences, (B) galbut virus RNA 2 sequences, (C) galbut virus RNA3 sequences, (D) *D. melanogaster* sigma virus sequences, (E) L protein sequences for viruses in the family

399 *Peribunyaviridae*, and (F) putative RNA-dependent RNA polymerase protein sequences related
400 to the new *Drosophila*-associated sobemo-like virus. Trees in (A)-(D) are based on nucleotide
401 alignments and trees in (E) and (F) are based on protein sequence alignments. Purple color
402 indicates museum sample-derived sequence generated in this study. Accession numbers are
403 noted except when groups of closely related sequences are collapsed. All trees are midpoint
404 rooted. Blue tip indicates the sequence of an endogenized galbut virus RNA 1 sequence.

405

406 **Certain types of RNA survived better in old samples**

407

408 We performed a variety of analyses to characterize the molecular and genomic
409 properties of surviving RNA. An initial indication that not all types of RNA survived equally came
410 from the strandedness of virus-mapping reads. We constructed libraries using a protocol that
411 preserved information about the RNA strand from which reads derived⁵¹. Galbut virus is a
412 partitivirus, which have dsRNA genomes⁵². But in infected flies, nearly all galbut virus RNA was
413 positive (+) sense (98.2% in fresh frozen flies; **Fig. 6; Supp. Fig. 8**). This is presumably
414 because most galbut virus RNA in infected cells is messenger RNA (mRNA). The fraction of
415 +strand galbut virus RNA decreased in older samples. In experimentally dried samples an
416 average of 89.1% of reads were from +strand RNA, and this value fell to 74.1% in museum
417 specimens ($p = 1.4 \times 10^{-4}$; **Fig. 6**). The decreasing fraction of +strand galbut virus RNA could be
418 explained by the preferential survival of double-stranded galbut virus RNA in older samples.

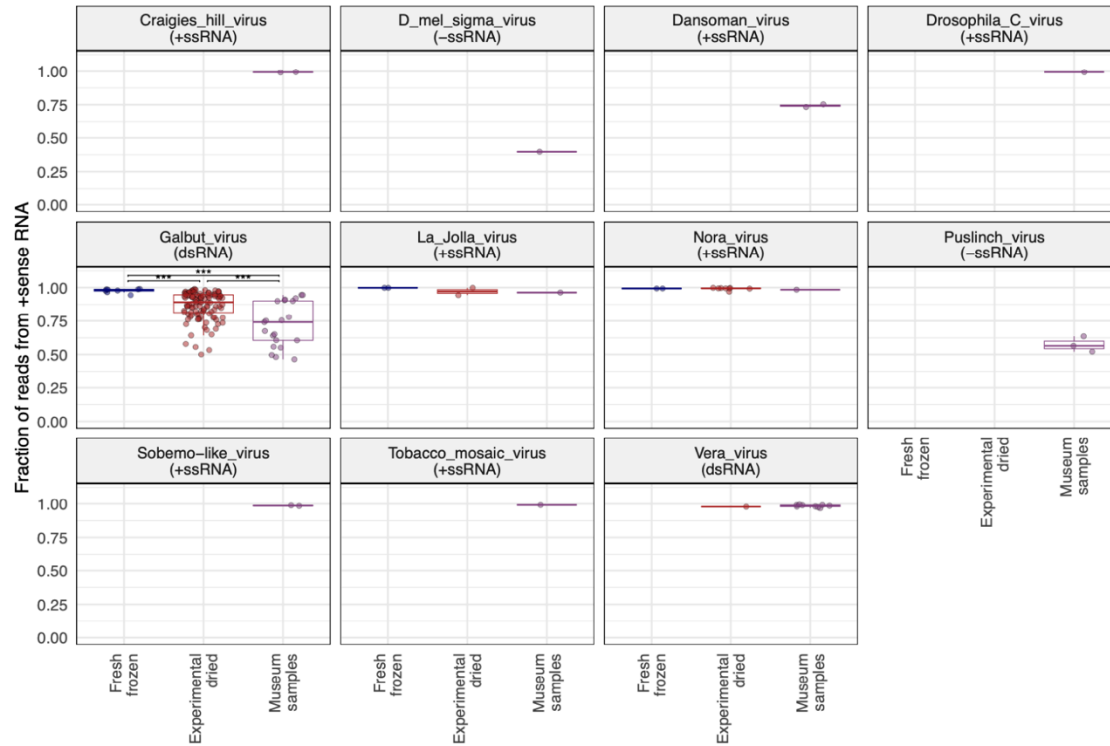
419

420 Coverage of galbut virus in museum datasets derived from all segments and was not
421 concentrated in regions of RNA 1 corresponding to the PCR primers we use to detect galbut
422 virus in our lab (**Supp. Fig. 8**). The lack of higher coverage in PCR product regions supports the
423 conclusion that galbut virus sequences did not result from contamination by contemporary PCR
424 products.

425

426 Other viruses had different fractions of +strand RNA that were consistent with the
427 genome type of each virus (**Fig. 6**). For instance, a majority of reads from *D. melanogaster*
428 sigma virus, a negative (-) strand RNA virus, were from -strand RNA²⁹. RNA from Nora virus (a
429 +strand virus) remained almost totally +strand in all samples³³. The fraction of +strand vera virus
430 RNA, another partitivirus, remained high in old samples (**Fig. 6**). This indicated that it was not
431 simply dsRNA that survived in older samples.

432



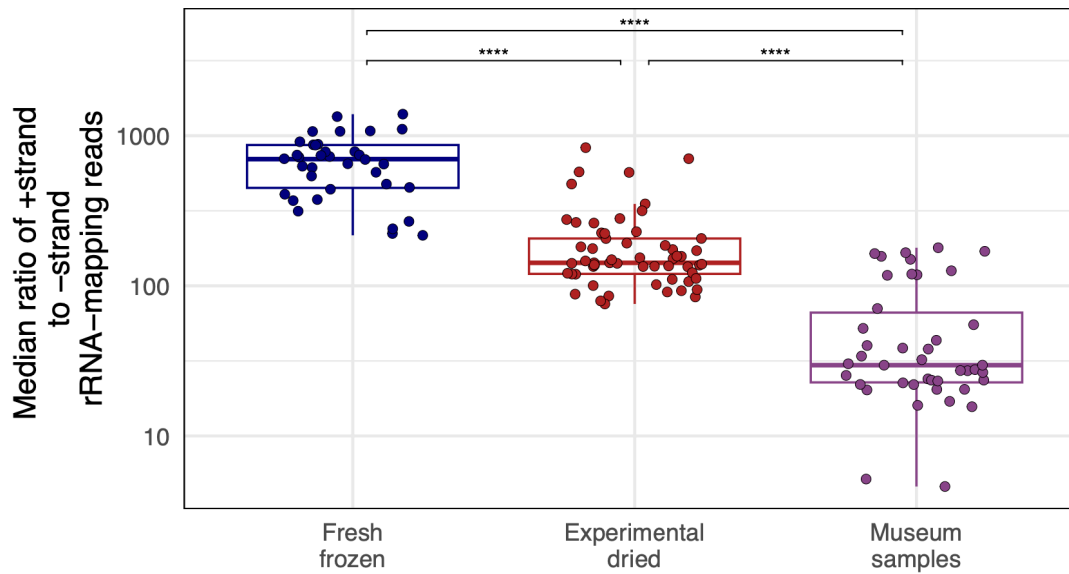
433

434 **Figure 6: The strandedness of galbut virus RNA in old samples is consistent with**
 435 **preferential survival of dsRNA.** The fraction of virus-mapping reads that originated from
 436 +strand RNA is plotted. Each point represents a dataset from an individual *D. melanogaster* fly.
 437 Significance levels of Wilcoxon test adjusted *p*-values are indicated: ns: $p > 0.05$; *: $p \leq 0.05$;
 438 **: $p \leq 0.01$; ***: $p \leq 0.001$; ****: $p \leq 0.0001$. Vera virus and other viruses were not present in
 439 sufficient numbers in different datasets to statistically evaluate how their strand ratios changed
 440 over time.

441

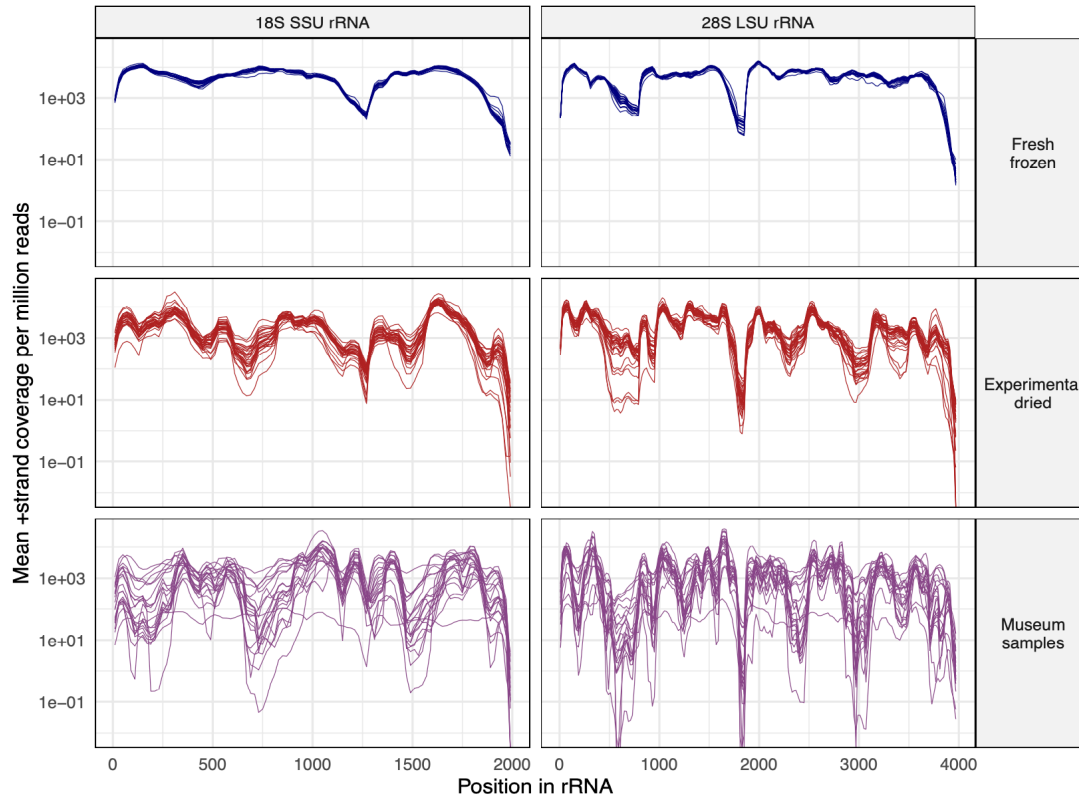
442 Nevertheless, there was additional evidence from host-mapping reads that double-
 443 strandedness contributed to preferential RNA survival. Most host-mapping RNA derived from
 444 ribosomal RNA (rRNA), and most rRNA-mapping reads derived from the +strand (**Fig. 7**). In
 445 fresh samples, there was on average 698x more +strand rRNA than -strand rRNA. The small
 446 amount of -sense rRNA could have derived from anti-sense transcription of rRNA genes or
 447 pseudogenes⁵³. In experimentally dried and museum samples, the +strand:-strand rRNA ratio
 448 dropped to 142:1 and 30:1, respectively ($p = 8.2 \times 10^{-13}$ and 2.8×10^{-22} ; **Fig. 7**). This drop was
 449 driven by a relative decrease in +strand rRNA and a relative increase in -strand rRNA in older
 450 samples (**Supp. Fig. 9**). This pattern could be explained by the preferential survival of
 451 sense:antisense rRNA duplexes. However, even in old samples, there was 30x more +strand

452 rRNA than –strand (**Fig. 7**), so sense:antisense duplexes could not account for all surviving
453 rRNA.
454



455
456 **Figure 7: Strand ratios of surviving ribosomal RNA were consistent with preferential**
457 **survival of dsRNA.** The median ratio of +strand to –strand coverage of rRNA-mapping reads.
458 Points represent individual-fly datasets. Adjusted *p*-values significance levels from Wilcoxon test
459 are indicated.

460
461 Coverage levels across rRNAs varied more in old samples than they did in new
462 samples, where coverage was relatively even (**Fig. 8**). For instance, coverage surrounding
463 position 1500 of the 18S rRNA and position 3000 of the 28S rRNA had consistently lower
464 coverage in old samples (**Fig. 8**). In contrast, in fresh samples coverage over these same
465 regions was similar to average coverage.



466

467

468 **Figure 8: Certain regions of ribosomal RNA were underrepresented in old samples.** The
469 mean +strand rRNA coverage per million reads in 20 bp windows is plotted for *D. melanogaster*
470 datasets. Each line represents a dataset from an individual fly. SSU: small subunit (18S) rRNA;
471 LSU: large subunit (28S) rRNA.

472

473 To investigate why some rRNA regions might not survive as well as others, we took
474 advantage of a high resolution cryo-electron-microscopy structure of the *D. melanogaster* 80S
475 ribosome⁵⁴. This structure includes ribosomal proteins and RNA (**Fig. 9A**). It is possible to
476 resolve individual rRNA bases in the structure and to determine whether they are interacting
477 with other bases, and whether they are present in the structure (**Fig. 9B**). We used this structure
478 and RNApdbe software to individually assign the 5965 rRNA bases to one of four categories:
479 paired with other rRNA bases (62.9%), unpaired (29.2%), present in higher-order secondary
480 structures like pseudoknots (4.5%), or missing from the 3D structure altogether (3.4%; **Fig. 9B**)
481 ⁵⁴⁻⁵⁶.

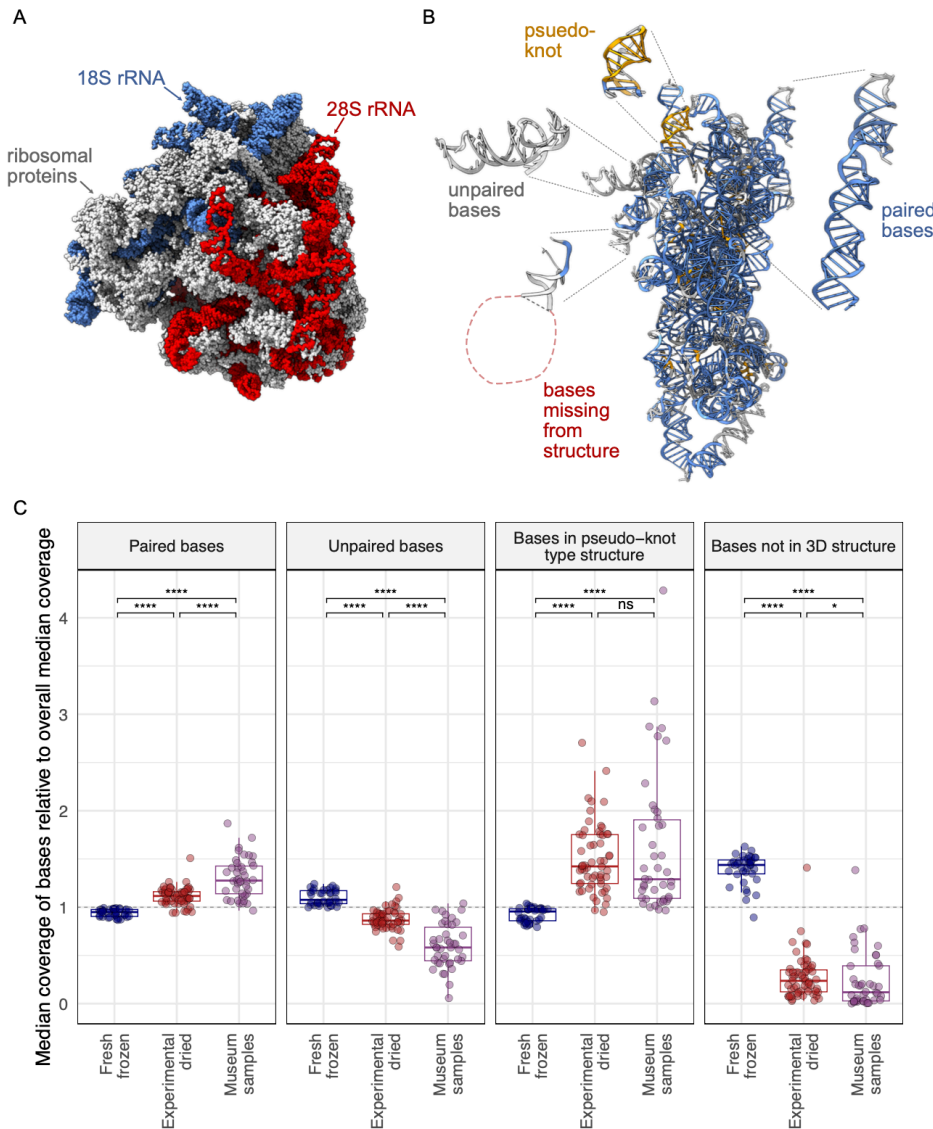
482

483 We calculated coverage of bases in each of these four categories relative to total
484 median coverage (**Fig. 9C**). Paired bases had higher average coverage in older samples ($p =$

485 1.9×10^{-19} in museum samples vs. fresh ones). Similarly, bases in higher-order secondary
486 structures like pseudoknots had higher coverage in old samples ($p = 3.3 \times 10^{-17}$ relative to fresh).
487 In contrast, unpaired bases had lower average coverage in old samples ($p = 3.8 \times 10^{-20}$ vs. fresh
488 samples). Bases that were not captured in the 3D structure were the least well represented in
489 older samples, with coverage levels 8x lower than overall average coverage in museum
490 samples ($p = 1.0 \times 10^{-19}$ relative to fresh frozen). Experimentally dried sample coverage levels
491 were generally intermediate between fresh and museum samples, consistent with their
492 intermediate age (**Fig. 9C**).

493

494 These patterns suggested that the molecular environment surrounding rRNA bases
495 influenced the likelihood that they would survive. Being base-paired or in a higher-order RNA
496 secondary structure protected bases, enabling them to survive over longer periods. In contrast,
497 unpaired bases or bases not present in the 3D structure - presumably because they were
498 outside of the protective environment of the ribosome - were less likely to survive.



499

500 **Figure 9: Different types of ribosomal RNA exhibited differential survival in old samples.**

501 (A) The structure of the *D. melanogaster* 80S ribosome from Anger at al, showing ribosomal

502 proteins and the 18S and 28S ribosomal RNAs. (B) The 18S rRNA with individual bases color

503 coded to indicate whether they are paired with other rRNA bases (blue), unpaired (grey), or

504 involved in higher order pseudo-knot type structures (gold). Some bases were not captured in

505 the structure (red). Bases were binned into categories using RNApdbee software. The 18S

506 rRNA is rotated relative to panel A. (C) The median coverage level of bases in the indicated

507 categories relative to total median coverage is plotted. Each point represents a dataset from an

508 individual *D. melanogaster* fly. Significance levels of Wilcoxon test adjusted *p*-values are

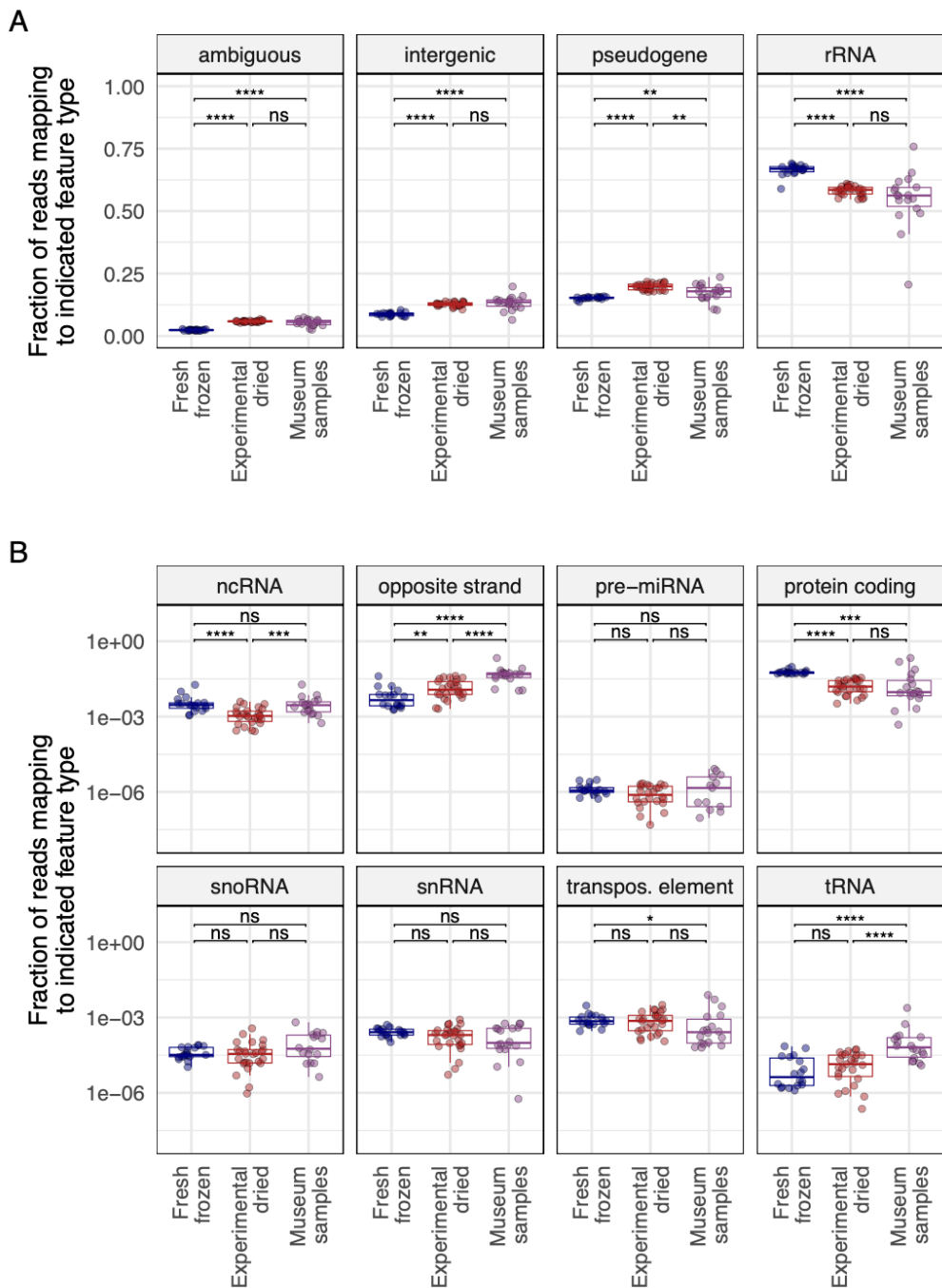
509 indicated.

510

511 **Different types of non-ribosomal RNA survived differentially in old samples**

512

513 We also quantified the extent to which different types of non-ribosomal host RNA
514 survived in old samples. We mapped reads from *D. melanogaster* samples to the *D.*
515 *melanogaster* genome and used the ALFA software to quantify the different types of host RNA
516 present in each dataset⁵⁷. ALFA combines mapping information with genome annotation to
517 assign mapped reads to one of a dozen RNA types (**Fig. 10**). As expected, most reads were
518 categorized as rRNA, though there was relatively less rRNA in older samples (**Fig. 10A**; $p =$
519 5.1×10^{-6} for museum vs. fresh samples). There was also relatively less protein-coding mRNA in
520 older samples (**Fig. 10B**; $p = 5.7 \times 10^{-4}$). Opposite strand RNA levels – that is, reads derived from
521 the RNA strand opposite an annotated feature - were elevated in older samples, consistent with
522 the preferential survival of sense:antisense transcript duplexes ($p = 1.8 \times 10^{-7}$). Highly structured
523 transfer RNA (tRNA) levels were also elevated in older samples ($p = 5.9 \times 10^{-5}$)^{58,59}.



524
 525 **Figure 10: Certain types of RNA survived better in old samples.** The fraction of reads in *D.*
 526 *melanogaster* datasets mapping to the indicated RNA types is plotted. Each point represents a
 527 dataset from an individual fly. Adjusted *p*-values significance levels from Wilcoxon test are
 528 indicated. (A) Types of RNA present at >5% median abundance. (B) Types of RNA present at
 529 <5% abundance. Abbreviations for different RNA types: ncRNA: non-coding RNA; miRNA:
 530 micro RNA; snoRNA: small nucleolar RNAs; snRNA: small nuclear RNA; tRNA: transfer RNA.

531 **Old RNA was chemically damaged**

532

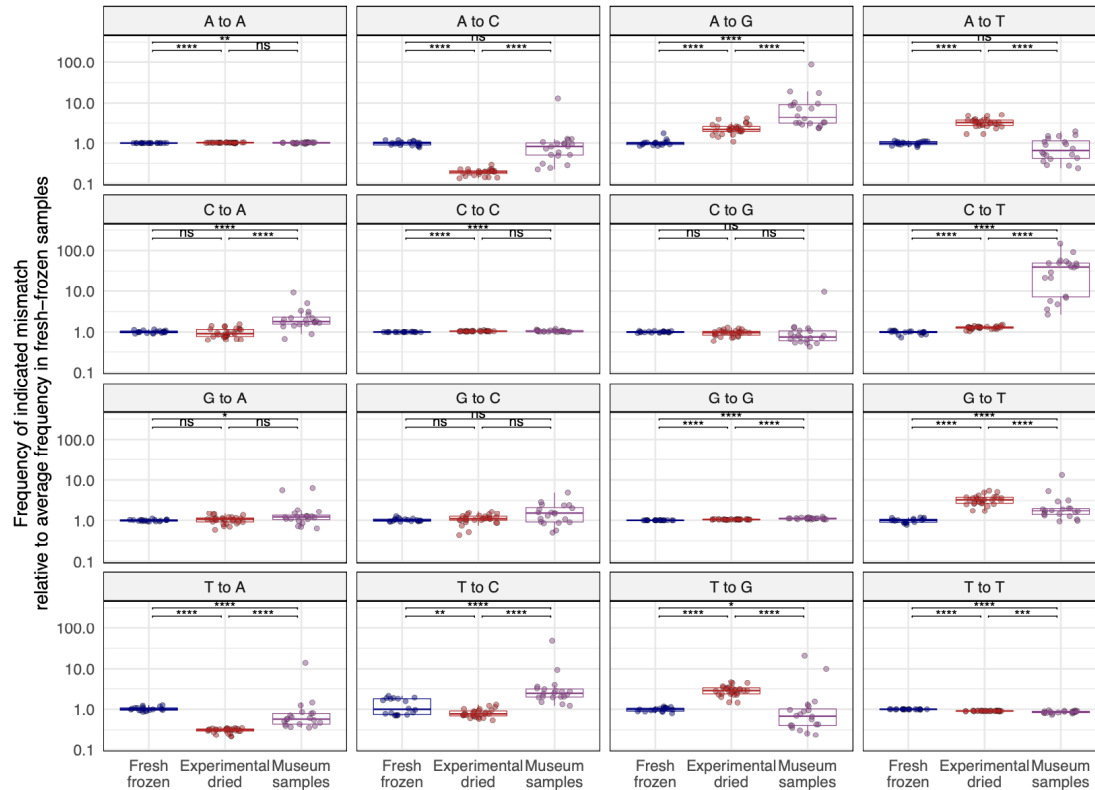
533 Old DNA molecules are fragmented and chemically damaged⁶⁰. Polymerases can
534 misincorporate bases when copying damaged templates, causing damage to manifest as
535 substitutions in library molecules⁶¹⁻⁶³. For example, deaminated cytosines, common in ancient
536 DNA, result in C-to-T substitutions in library molecules. In fact, signatures of chemical damage
537 in sequence reads are taken as evidence that reads derive from old DNA and not from
538 contaminating contemporary DNA.

539

540 We quantified mismatch patterns in mapped reads to look for signatures of chemical
541 damage in old RNA (**Fig. 11**). Mismatch frequencies varied in different sample types. Some
542 differences in mismatch frequencies could be attributed to samples being sequenced on
543 different sequencing runs (we sequenced museum samples separately from other samples to
544 avoid the possibility of read misassignment from index-hopping)⁶⁴. The largest difference
545 between old and new samples was an elevated frequency of C-to-T substitutions in reads from
546 museum samples. The median level of C-to-T mismatches in datasets from museum samples
547 was 40x higher than in fresh-frozen datasets (2.0×10^{-3} frequency of C-to-T mismatches vs.
548 5.0×10^{-5} (**Supp. Fig. 10**; $p=1.0 \times 10^{-10}$). This increased frequency of C-to-T substitutions is
549 consistent with deamination of cytosines in old RNA.

550

551 A-to-G substitutions were the next most elevated type of mismatch in old samples (**Fig.**
552 **11**). A-to-G substitutions occurred at a rate 4.4-fold higher in museum samples than in fresh
553 datasets (5.7×10^{-4} vs. 1.3×10^{-4} ; $p=1.6 \times 10^{-10}$). Such substitutions can result from spontaneous
554 deamination of adenine to hypoxanthine^{60,65}.



555

556 **Figure 11. Old RNA was chemically damaged and exhibits mismatch patterns consistent**
 557 **with cytosine and adenine deamination. Mismatches in rRNA-mapping reads from *D.***
 558 *melanogaster* datasets were quantified and the frequency of each mismatch type relative to the
 559 median frequency in fresh-frozen datasets is plotted. Each point represents a dataset from an
 560 individual fly. Adjusted *p*-values significance levels from Wilcoxon test are indicated.

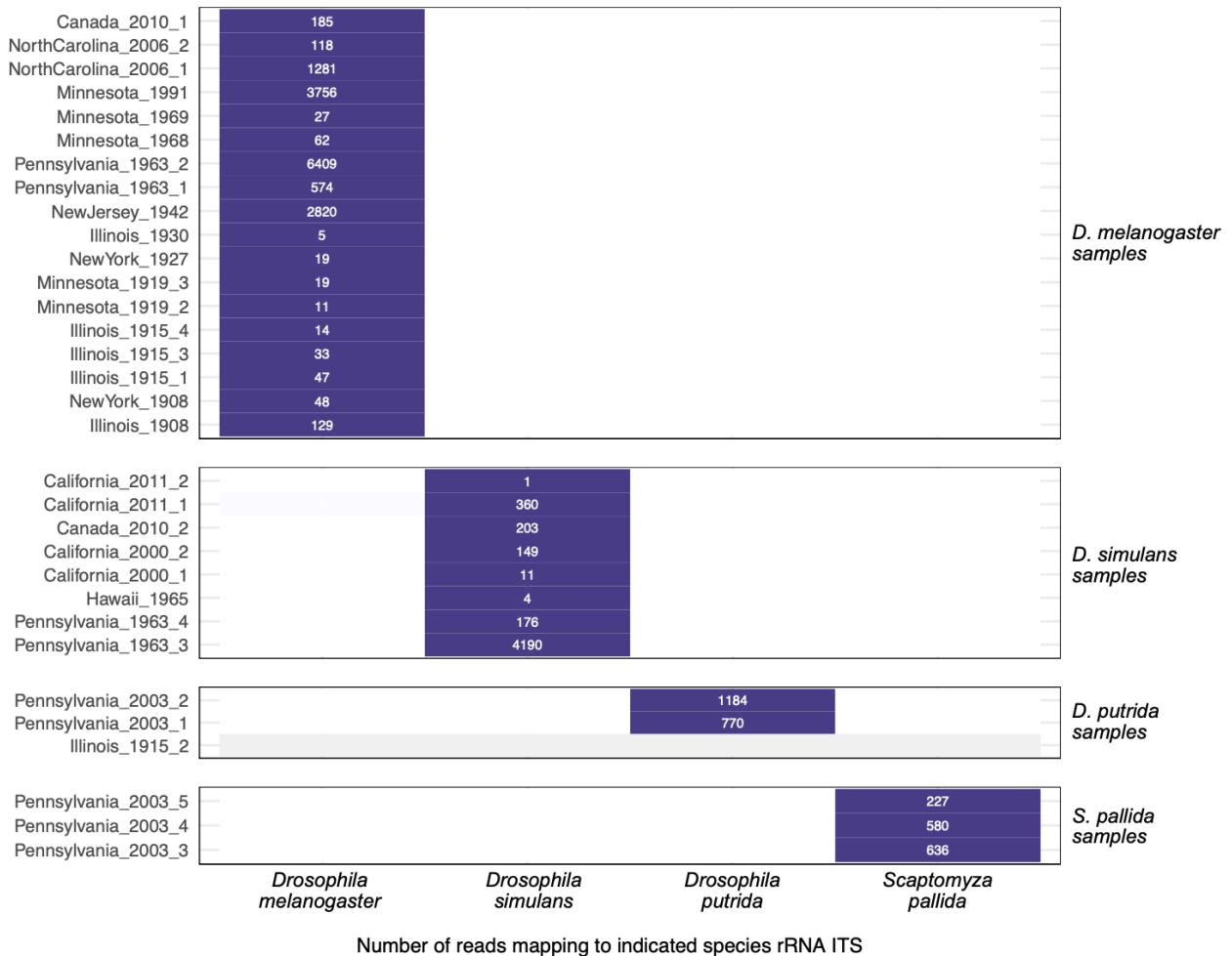
561

562 **Host-mapping reads exhibited species specificity**

563

564 We took advantage of the fact that our museum samples derived from 4 different
 565 species to investigate the possibility that host-mapping reads – specifically rRNA-mapping reads
 566 – might result from contemporary contamination. The likely source of fly-mapping contamination
 567 would be the *D. melanogaster* that we rear and study in our lab⁴⁸. The 18S and 28S ribosomal
 568 RNA sequences of *D. melanogaster* and *D. simulans* share over 99% pairwise identity which
 569 makes them difficult to distinguish by competitive read mapping. We therefore collected and
 570 mapped reads to the relatively variable region of the ribosomal RNA between the 18S and 5.8S
 571 genes (the internal transcribed spacer 1, ITS) of the 4 fly species that we sampled (**Table S1**;
 572 **Fig. 12**; **Supp. Fig. 11**). All ITS-mapping reads in museum datasets mapped to the expected
 573 species ITS sequence and no reads mapped to an unexpected species (**Fig. 12**). Positive and

574 negative control samples behaved as expected, with water negative control datasets containing
 575 no reads mapping to any of these 4 ITS sequences (**Supp. Fig. 11**). If fly rRNA-mapping reads
 576 were from contamination, we would have expected there to be *D. melanogaster*-mapping reads
 577 in the non-*melanogaster* and negative control datasets; instead, there were none. The perfect
 578 concordance of ITS mapping and sample species supported the conclusion that the host-
 579 mapping reads in our datasets were indeed from museum samples.
 580



581
 582
 583 **Figure 12: Host-mapping reads exhibited species specificity.** The number of reads mapping
 584 to the indicated species rRNA internal transcribed spacer 1 (ITS) sequence for each museum
 585 sample dataset is shown. Samples are grouped according to their molecular species
 586 assignment (**Table S1**). Datasets with no ITS-mapping reads are represented with grey.
 587

588 **DISCUSSION**

589

590 In this study we took a two-pronged approach to investigate RNA stability in old
591 biological samples. In flies and mosquitoes stored dry at room temperature for weeks or
592 months, RNA grew increasingly fragmented, but RNA yields did not significantly decrease, and
593 cellular and viral RNA remained detectable. Perhaps the most unexpected result from this
594 experiment was the limited difference in RNA quality and quantity between dried and frozen
595 samples (**Fig. 1; Supp. Figs. 2, 4**). We recovered RNA from dried flies that were from as far
596 back as the 1890s and used this RNA to make sequencing libraries. We recovered coding
597 complete virus sequences from 11 old specimens and identified two novel virus sequences. We
598 also showed that certain types of RNA tended to survive in old samples, including dsRNA and
599 RNA in secondary structures or ribonucleoprotein complexes.

600

601 There were limitations to the experimental arm of our study. First, RNA recovery from
602 individual flies and mosquitoes - even fresh ones - was variable. Second, we used outbred
603 populations of flies and mosquitoes with different levels of virus infection between individuals. In
604 future studies, optimization of our RNA isolation protocol to decrease yield variability between
605 individual flies and mosquitoes would be useful. It would also be better to use pinned specimens
606 from inbred populations with less variable infection phenotypes.

607

608 We found that not all RNA survived equally well and described molecular features that
609 influenced how well different types of RNA survived. dsRNA survived better than ssRNA.
610 Surviving RNA duplexes derived from intermolecular base pairing, such as sense-antisense
611 duplexes (**Figs. 6, 7A, 10B**) and from intramolecular base pairing, such as paired rRNA bases
612 (**Fig. 9C**). The preferential survival of dsRNA likely reflects decreased susceptibility to
613 hydrolysis⁶⁶. However, double strandedness could not explain all RNA survival, as RNA from
614 certain viruses remained largely +strand (**Fig. 6**) and unpaired rRNA survived (albeit less than
615 paired rRNA; **Figs. 7, 9**). It may be that ribonucleoprotein complexes like virus particles and
616 ribosomes provide a protective environment in which RNA can survive longer.

617

618 Like old DNA, old RNA was fragmented and chemically damaged (**Figs. 2C, 11**). And,
619 like DNA, deamination was the most evident type of damage (**Fig. 11**). Other types of damage
620 may be common in old RNA but undetectable by sequencing. The extent to which
621 ribonucleases, deaminases, and other enzymes involved in normal RNA processing continue to

622 function in dead cells is unclear, but it is likely that most RNA fragmentation and damage in old
623 samples results from spontaneous chemical rather than enzymatic degradation.

624

625 Environmental factors like temperature and humidity likely play a major role in the
626 survival of RNA. It was notable that none of the 12 specimens from tropical Hawai'i yielded
627 enough RNA to be detected fluorometrically and only two generated sequenceable libraries
628 (**Fig. 2b, Table 1 & Supp. Fig. 5**). Humidity might be more important than temperature, as RNA
629 was recovered from 1000-year-old corn in Arizona, where temperatures fluctuate to extremes,
630 but humidity is very low¹⁵. If this were true, RNA might survive better in drier environments and
631 in quickly desiccating samples like the small insects that are the focus of this work. Even for
632 much larger samples like woolly mammoths, dehydration has been proposed to promote
633 preservation of nuclear architecture and biological macromolecules⁶⁷.

634

635 The virus sequences we recovered revealed that the *D. melanogaster* virome has not
636 changed much over the last 120 years. All the viruses that we detected in *D. melanogaster*,
637 except for Puslinch virus, had already been described in contemporary *D. melanogaster*^{25,26}. For
638 the most part the old sequences were closely related to contemporary sequences (**Table 2**). For
639 example, the viruses infecting the fly collected in 1908 were >98% identical to known *D.*
640 *melanogaster* viruses (**Fig. 4; Table 2**).

641

642 The discrepant phylogenetic positions of the 1930 galbut virus segments is consistent
643 with reassortment⁶⁸. The 1930 galbut virus RNA 3 sequence that is positioned on its own branch
644 may correspond to an extinct RNA 3 lineage, or simply one that has not been sampled before
645 (**Fig. 5C**). Similarly, the endogenous galbut virus RNA 1 sequence in these trees may represent
646 an extinct lineage or an ancestor of contemporary galbut virus RNA 1 sequences (**Fig. 5A**)⁴⁹. A
647 potential benefit of large-scale sequencing of old samples is that it could provide estimates of
648 how frequently virus lineages go extinct from a particular host.

649

650 Contamination from contemporary nucleic acid has been a long-standing concern for
651 those sequencing ancient DNA⁶⁹. There was evidence of low-level contamination in our
652 datasets (**Fig. 3**), and it is difficult to completely avoid contamination in NGS datasets, even
653 when sequencing new samples^{70,71}. But, there were numerous lines of evidence to support the
654 idea that our conclusions were based on sequences from actual old RNA and not from
655 contemporary contamination. Old RNA was short and chemically damaged; properties that are

656 shared with old DNA⁶⁰. Characteristics of RNA from experimentally dried samples were
657 generally intermediate between those of fresh RNA and museum RNA, consistent with their
658 intermediate age (**Figs. 6-11**). We recovered different virus sequences from different old
659 samples, and reads in individual datasets supported the same distinct virus sequences (**Table**
660 **2**). The virus sequences we recovered were for the most part known *Drosophila*-infecting
661 viruses. Although many virus sequences from old samples were similar to existing
662 contemporary sequences, most were not identical (**Table 2**). The exceptions were three vera
663 virus RNA 1 sequences that were identical to existing sequences. But the RNA 2 and chaq-like
664 sequences from these samples were not identical to any contemporary sequences. The two
665 new virus sequences we recovered were divergent, but their closest relatives were from other
666 insects (**Table 2; Fig. 5**). The tobacco mosaic virus sequence from the 1908 dataset was an
667 exception, which we can't explain except to suggest that this specimen was contaminated
668 during handling by a smoker, as individual cigarettes were found to contain as many as 10⁹
669 TMV RNA copies⁴¹. There was no peak in galbut virus coverage corresponding to our diagnostic
670 RT-qPCR amplicons, which would be a prime candidate source of contaminating galbut virus
671 sequences (**Supp. Fig 6**)^{30,48}. All museum samples that were positive for galbut virus by RT-
672 qPCR and generated a sequenceable library were also positive by sequencing (**Supp. Table 1**).
673 Negative control datasets were uniformly free of virus and rRNA-mapping reads (**Fig. 4; Supp**
674 **Fig. 11**). Galbut virus sequences from *D. simulans* museum samples clustered with
675 contemporary *D. simulans* sequences, and the same for *D. melanogaster* sequences (**Fig. 5**).
676

677 Indeed, the species-specificity of host and virus sequences (**Figs. 5 & 12**) provides
678 some of the strongest support for the idea that our analyses are based on nucleic acid from the
679 actual old samples. If our conclusions were based on contaminating contemporary sequences, it
680 would be necessary to invoke a complicated mechanism where contaminating host and virus
681 nucleic acids sorted themselves by species into each sample (**Figs 5 & 12**). The overall
682 apparent lack of contaminating virus and host sequences was consistent with the fact that all
683 extractions were done in biosafety cabinets following a pre-PCR to post-PCR workflow,
684 including DNase treatment before RT-qPCR and library preparation.
685

686 RNA can persist in biological samples over decades or centuries without freezing or
687 fixation. The half-life of RNA in dead cells is clearly much longer than in living cells. The millions
688 of dried specimens in museums represent a valuable source of RNA for reconstructing historic
689 virus-host interactions^{14,15}. It is tempting to speculate that the long-term survival of certain RNA

690 might approach that of DNA⁷². It is time to move beyond the incorrect idea that old, dried
691 samples are not a good source of useful RNA.

692

693 **MATERIALS AND METHODS**

694

695 **Sample Collection**

696 *Pinned Specimens:* FoCo-17 *D. melanogaster* were collected and pinned using standard
697 entomological collection techniques with storage at room temperature or stored frozen at -80°C
698 ⁷³. Poza Rica *Aedes aegypti* mosquitoes were reared using standard lab techniques until
699 adulthood⁷⁴. Prior to blood feeding mosquitoes were collected and pinned or frozen.

700

701 *Museum Specimens:* Samples were obtained from museums and other institutions containing
702 entomological specimens located in the United States of America and Canada. Altogether, 46
703 drosophilid samples, mainly *D. melanogaster* or *D. simulans*, were selected for the study.

704 Museum species assignments were checked using molecular data as described below. One
705 sample from California, California_2000_2, was initially identified as *D. melanogaster*, but our
706 sequence data indicated that it was *D. simulans* (these species can be difficult to distinguish
707 morphologically). Location, date of sample collection, sample storage type, extraction method,
708 RNA quality and RNA extraction concentration are summarized in **Figure 2, Table**

709 **1 and Supplemental Table 1**

710

711 **Sample Workflow**

712 Deliberate protocols were used to minimize contamination during sample processing and library
713 construction. Sample processing and library construction moved from dedicated pre-PCR rooms
714 to a post-PCR room. RNA extractions were performed in a dedicated pre-PCR sample
715 extraction room in a Class II type B2 biosafety cabinet which protects both personnel and
716 samples. Initial library preparation steps, including reverse transcription, ligation, and setup of
717 library amplification reactions were completed in a separate PCR setup room in an AirClean
718 PCR workstation equipped with a HEPA filter (AirClean Systems, AC600 Series). Finally, library
719 amplification and subsequent cleanup and pooling steps were performed in a third post-PCR lab
720 space.

721

722 **RNA Extraction**

723 *Non-Destructive Sampling:* For samples for which destructive sampling was prohibited, we
724 adapted the protocol described by Santos et. al., 2018³⁵. Briefly, individual samples were placed
725 in a PCR tube containing 200µL of solution containing 200mM Tris HCl, 250mM NaCl, 25mM
726 EDTA, 0.5% SDS and 400 µg/mL proteinase K. Samples were then incubated at 56°C for 16
727 hours. After incubation the supernatant was moved to a new tube for RNA isolation and the
728 specimen was transferred to a cryovial containing 80% ethanol. RNA was purified using the
729 Zymo RNA Clean & Concentrator Kit as described below.

730

731 *Ethanol Stored Sample Rehydration:* Four samples arrived in 70% ethanol. To prepare these
732 samples for extraction, each specimen was rehydrated using a gradient of decreasing
733 concentrations of ethanol. The specimens were suspended in 200µL 70%, 50%, 30% and 10%
734 ethanol for 15 minutes on ice with agitation every 2-3 minutes prior to a final incubation in water
735 for 15 minutes before RNA extraction following the destructive sampling method described
736 below.

737

738 *Destructive Sampling:* A phenol/chloroform approach with mechanical disruption was used for
739 RNA isolation of the experimentally dried and frozen samples as well as the museum
740 specimens that could be destructively sampled. Briefly, each specimen was placed in a 2.0mL
741 tube with a small BB (McMaster Carr, 1598K22) and 500µL of Trizol was added and incubated
742 for 5 minutes. Tubes were then shaken in a TissueLyser (Qiagen, 9003240.) at 30 Hz for 3
743 minutes. 100µL of chloroform was added, thoroughly mixed and incubated for two minutes prior
744 to being spun at 12,000xg for 10 minutes. The aqueous phase was retained for RNA purification
745 using the Zymo RNA Clean & Concentrator-5 kit following manufacturer directions for the
746 capture of small RNA fragments with slight modification (Zymo Research, R1013). Briefly,
747 225µL of sample was mixed with 225µL 100% ethanol and 225µL RNA binding buffer. Samples
748 were transferred to spin columns with collection tubes and spun. To capture small RNAs, the
749 flow through from the first spin was retained and mixed with an equal amount of 100% ethanol.
750 This was transferred to the spin column and centrifuged. A wash using RNA wash buffer was
751 done prior to a DNase treatment following manufacturer recommendations (Zymo Research,
752 E1010). After DNase treatment, two more washes were complete prior to elution in 30µL of
753 nuclease free water. All spins were performed at 9,000xg for 1 minute unless otherwise noted.
754 For all extraction types, FoCo-17 male and female flies were used as a positive control and an
755 empty tube was used as a negative control.

756

757 **RNA Quality and Quantification**

758 RNA quality was assessed using a nanodrop spectrophotometer (Thermo Scientific
759 ND2000USCAN) and RNA concentration was determined using the Qubit high-sensitivity RNA
760 reagent (ThermoFisher, Q32852). RNA length was measured using the Agilent TapeStation
761 High Sensitivity RNA screening tapes and reagents (Agilent, 5067-5579 & 5067-5580).

762
763 **RT-qPCR**

764 *Pinned Samples:* Several targets were assessed for presence/absence and evidence of
765 degradation using RT-qPCR (primer sequences in **Supplemental Table 3**). cDNA was
766 generated using 5.5µL RNA, 10µM random hexamer primers (IDT), 10µM dNTPs and water to
767 13µL and incubated at 65°C for 5 minutes. Next, 4µL 5X FS buffer, 1µL 0.1M DTT and 1µL
768 reverse transcriptase was added and incubated at 50°C for 60 minutes followed by a heat
769 inactivation step at 80°C for 10 minutes after which each sample was diluted in 90µL nuclease-
770 free water. qPCR was conducted using NEB Luna Universal qPCR Mastermix following
771 manufacturer recommendations and cycling conditions (NEB, M3003).

772
773 **PCR controls**

774 RNA from a pool of FoCo-17 flies was reverse transcribed to create a cDNA positive control for
775 fly samples. For mosquito RT-qPCR, positive control cDNA was reverse transcribed from
776 individual colony mosquitoes. We included these positive controls and three negative controls in
777 all qPCR runs. Negative controls consisted of an extraction blank (a no sample extraction), a
778 no-sample RT control, and a no template qPCR control. The use of negative controls at each
779 step would have allowed us to determine when cross-contamination had been introduced if it
780 had been. Positive/negative status was determined using the melt curve (within 1 degree of the
781 positive control) for each sample and target. Agarose gel electrophoresis of qPCR products was
782 used to determine positive/negative status when Ct and melt curve were ambiguous.

783
784 *Museum Collections:* We used RT-qPCR to determine if any samples had detectable levels of
785 galbut virus RNA and RpL32 mRNA. To increase our ability to detect highly fragmented RNA,
786 we designed a second set of primer that target a small region of the galbut virus RNA 1 and *D.*
787 *melanogaster* mRNA RpL32. A 2% ethidium bromide agarose gel was used to confirm
788 positive/negative status. Galbut virus positive female and male FoCo-17 flies were used as
789 positive controls and water was used as a negative control.

790 **Supplemental Table 3: Primer sequences used in this study.**

Primer pair name (#s in lab collection)	Forward	Reverse	Reference
Galbut virus (long) (#1600/1601)	CCGTGAAGCAAGGAATCAAT	TGCCGATTTTCTGCTCTTTT	³⁰
Galbut virus- short (1948/1949)	AGAAGATGTGCTGTAGTGACAC	CGTGGAATTCCGAACGGCTA	This study
RpL32- long (1012/1013)	TGCTAAGCTGTCGCACAAATGG	TGCGCTTGTTGATCCGTAAC	⁷⁵
RpL32- short (1950/1951)	GCGCTTGTTGATCCGTAAC	GCCGCTTCAAGGGACAGTAT	This study
Thika virus (1604/1605)	CAGCAGGTCCCTTGCTAAAG	TGGTCAGCATATGACCGAAA	³⁰
Nora virus (1592/1593)	GCACCTGGTCGATTGAATCC	CGTTCAGGGCATAGTCAAGC	³⁰
La Jolla virus (1692/1693)	ACCGTATGGCGTCGTACTTC	AAAGTATCAGCAGCGCGAAT	³⁰
Actin (#1826/1827)	CGTTCGTGACATCAAGGAAA	GAACGATGGCTGGAAGAGAG	⁷⁶
Verdadero virus (#1662/1663)	ATATGGGTTCGTGTCGAAAGC	CACCCCGAAATTTTCTTCAA	³⁰
Guadeloupe mosquito virus (#1688/1689)	TTGTAAATCCCCCTGTGCTC	CGCAAAAATTGGGTTTGAGT	This study

791

792 **NGS Library Preparation**

793 Museum collection samples were prepared for sequencing using the Kapa Biosystems Kapa
794 RNA Hyper Prep kit following manufacturer recommendations with modification of the
795 fragmentation/priming step. Since the RNA was already highly fragmented, samples were
796 minimally incubated at 65°C for 1 minute during the fragmentation/priming step (Roche,
797 08098093702). Due to low starting concentrations, 12 cycles were used for library amplification.
798 FoCo-17 fly RNA was used as a positive control and water was used as a negative control for

799 library preparation and sequencing. Controls were sequenced on a separate run to avoid index
800 hopping from the positive control sample⁷⁷. Sample pooling was determined using High
801 Sensitivity DNA Qubit reagents and final library quality control was done using an Agilent D1000
802 HS Tapestation and KapaQuant reagents following manufacturers recommendations (Roche,
803 07960140001). Museum samples were sequenced on three NextSeq 500 runs; one high output
804 75 cycle run and two mid output 150 cycle runs. All museum specimens were run on the first
805 NextSeq run, runs two and three consisted of samples that we wanted to increase coverage of
806 near complete virus sequences. Experimentally dried samples were sequenced on a high output
807 75 cycle run. The two positive control samples were sequenced on a 300 cycle MiSeq run.
808 Additional fresh frozen samples were sequenced on a NextSeq mid output 150 cycle kit.

809

810 **Data Analysis**

811

812 *Experimental Collections:* All statistical analysis and data visualization, unless otherwise noted,
813 was done using R/RStudio. Data cleaning and data visualization were done using the tidyverse
814 package⁷⁸. All scripts are available in the GitHub repository linked below. A p-value of $\alpha = 0.05$
815 or below was used as our significance threshold. RNA Concentration and Length: Four
816 individual multiple linear regression models (MLRs) were constructed to test for statistical
817 significance in RNA concentration and length between dried and frozen flies and mosquitoes
818 over 72- and 52-weeks. The predictor time was treated as a continuous variable and the
819 predictor storage type was converted to factor. A balanced one-way analysis of variance
820 (ANOVA) was used to test for significance using the car package⁷⁹. The performance package
821 was used to test model assumptions⁸⁰. RT-qPCR: Raw qPCR data files were cleaned and
822 compiled using the SangerTools R package and a custom R script⁸¹. Two-tailed Students T-
823 tests from the rstatix package were used to determine statistical significance of dried and frozen
824 Ct values⁸². P-values were adjusted for multiple hypothesis testing.

825

826 *Identification of virus sequences in old samples:* We used our lab's previously described
827 metagenomic classification pipeline (https://github.com/stenglein-lab/taxonomy_pipeline) to
828 identify and validate virus sequences in NGS datasets³⁰. In brief, adapters and low quality reads
829 were trimmed using cutadapt 3.5⁸³. Fastqc was used to assess post-collapsed read
830 quality 0.11.9⁸⁴. Host reads were removed using bowtie2 2.4.5 and the remaining reads were
831 assembled using spades 3.15.4^{85,86}. Virus sequences were identified using BLASTn 2.12.0 and
832 a BLASTx search using diamond 2.0.14 was used to identify novel sequences^{87,88}. Draft

833 sequences were validated by remapping of trimmed reads using bwa mem aligner version
834 0.7.17⁸⁹. Final sequences were submitted to the NCBI nucleotide database and raw NGS data
835 to the NCBI sequence read archive repository.

836

837 This metagenomic classification pipeline was also used to taxonomically assign non-host
838 contigs and unassembled reads. Counts assigned to individual taxa were normalized to the
839 number of quality- and adapter-trimmed reads.

840

841 *Molecular species identification using cytochrome oxidase subunit 1 competitive mapping.* The
842 sequence of this mitochondrial-encoded gene is commonly used to distinguish related
843 species⁹⁰. We used the *D. melanogaster* CO1 coding sequence from NC_024511 as a BLASTN
844 query to the NCBI nucleotide database to identify other drosophilid CO1 sequences⁹¹. We
845 restricted results to sequences from Drosophilidae and to alignments with >90% query coverage
846 and E-value <1x10⁻⁵⁰. We used cd-hit-est to collapse sequences that shared >99% pairwise
847 nucleotide identity to create a set of 545 representative CO1 sequences⁹². We mapped adapter
848 and quality trimmed reads to these sequences using bowtie2 v.2.4.4 in end-to-end mode and
849 retained alignments with a mapping quality ≥ 20.

850

851 *Maximum Likelihood Trees:* All available galbut virus and sigma virus sequences from individual
852 flies (and not pools) with collection date metadata were downloaded from GenBank. Sequences
853 in the Puslinch virus phylogeny include all available L protein sequences in the NCBI RefSeq
854 protein database from the *Peribunyaviridae* family. Sequences in the sobemo-like virus tree
855 include RdRp protein sequences from viral genomes identified by a BLASTN search of the
856 NCBI nucleotide database. Alignments were generated using MAFFT v7.490 with default
857 parameters. IqTree v1.0 (<http://iqtree.cibiv.univie.ac.at>) was used to generate maximum
858 likelihood trees⁹³. The following parameters were used on the web interface version of IqTree:
859 substitution model- auto, Free-rate Heterogeneity- yes, bootstrap analysis- ultrafast with all
860 other options set to default. FigTree was used for tree visualization
861 (<http://tree.bio.ed.ac.uk/software/figtree/>). Trees were midpoint rooted.

862

863 *Analysis of strandedness of virus-mapping reads.* To tabulate the numbers of positive and
864 negative strand reads mapping to virus sequences, we mapped quality- and adapter-trimmed
865 reads to assembled virus sequences using the bwa mem aligner version 0.7.17⁸⁹. The

866 orientation of each mapped read was determined from the 0x10 bit flag of the resulting sam
867 format output.

868

869 For all analyses of virus- and host -mapping reads, values were summarized and visualized
870 using the tidyverse R packages⁷⁸. Wilcoxon tests were used to statistically evaluate differences
871 between groups using the rstatix R package. In all cases values were determined to be non-
872 normally distributed by Shapiro-Wilk test⁸². *P*-values were adjusted for multiple testing using the
873 Holm–Bonferroni method and were indicated on plots using the ggpubr R package using the
874 following significance levels: ns: $p > 0.05$; *: $p \leq 0.05$; **: $p \leq 0.01$; ***: $p \leq 0.001$; ****: $p \leq$
875 0.0001 ⁹⁴. Analyses were implemented as nextflow workflows available
876 at: https://github.com/LKeene/Old_Collections_Figures/⁹⁵.

877

878 *Analysis of ribosomal rRNA-mapping reads.* Adapter- and quality-trimmed reads from *D.*
879 *melanogaster* datasets were mapped to the 18S and 28S rRNA sequences from Anger et al
880 (chains B5 and A2 from⁵⁴) using bwa mem aligner version 0.7.17⁸⁹. Per-base, per-strand
881 coverage was calculated using the BamToCov tool v2.7.0⁹⁶. The *D. melanogaster* 80S ribosome
882 structure was visualized using ChimeraX v.1.7.1⁹⁷. Bases were binned into one of four
883 categories (paired, non-paired, higher-order structure, absent from structure) by feeding this
884 structure (protein data bank accession 4V6W) into RNApdbee 2.0⁵⁵.

885

886 *Analysis of non-ribosomal RNA types.* Adapter- and quality-trimmed reads from *D.*
887 *melanogaster* datasets were mapped to the *D. melanogaster* genome version BDGP6.32⁹⁸
888 using bwa mem aligner version 0.7.17⁸⁹. Mapping output and genome annotation were input
889 into ALFA v1.1.1 to tabulate the fractions of reads mapping to each of the annotated genomic
890 biotypes⁵⁷.

891

892 *Analysis of mismatch signatures of chemical damage.* Trimmed reads from *D. melanogaster*
893 datasets were mapped to rRNA as above. Mismatches of bases with a basecall quality score
894 >30 were tabulated.

895

896 *Competitive mapping to ribosomal internal transcribed spacer sequences.* We collected
897 sequences between the 18S and 5.8S rRNA genes for the four museum species. Sequences
898 correspond to Genbank accessions: *D. melanogaster*: NR_133558.1:2861-722; *D. simulans*:
899 NGVV02000020.1:5755-5071; *D. putrida*: AF184042.1:153-611; *S. pallida*:

900 JAECXP010000218.1:7393-7904. We mapped adapter and quality trimmed reads to these
901 sequences using bowtie2 v.2.4.4 in end-to-end mode and retained alignments with a mapping
902 quality ≥ 30 .

903

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918

919 **DATA AVAILABILITY**

920 All metadata, RT-qPCR raw data, plots, and analysis and visualization code can be found at:
921 https://github.com/LKeene/Old_Collections_Figures. Raw sequence data has been uploaded to
922 the NCBI SRA database; assembled viral sequences have been deposited in GenBank; both
923 can be found under NCBI BioProject accession PRJNA1034757.

924

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