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1 Data Note for: *GigaScience*

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11 ***De novo* transcriptome assemblies of four xylem sap-feeding insects**

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35 **Abstract**

36 **Background:** Spittle bugs and sharpshooters are well-known xylem sap-feeding insects and
37 vectors of the phytopathogenic bacterium *Xylella fastidiosa* (Wells), a causal agent of Pierce’s
38 disease of grapevines and other crop diseases. Specialized feeding on nutrient-deficient xylem
39 sap is relatively rare among insect herbivores, and only limited genomic and transcriptomic
40 information has been generated for xylem-sap feeders. To develop a more comprehensive
41 understanding of biochemical adaptations and symbiotic relationships that support survival on a
42 nutritionally austere dietary source, transcriptome assemblies for three sharpshooter species and
43 one spittlebug species were produced.

44 **Findings:** Trinity-based *de novo* transcriptome assemblies were generated for all four xylem-sap
45 feeders using raw sequencing data originating from whole-insect preps. Total transcripts for each
46 species ranged from 91,384 for *Cuernia arida* to 106,998 for *Homalodisca liturata* with transcript
47 totals for *Graphocephala atropunctata* and the spittlebug *Clastoptera arizonana* falling in
48 between. The percentage of transcripts comprising complete open reading frames ranged from
49 60% for *H. liturata* to 82% for *C. arizonana*. BUSCO analyses for each dataset indicated quality
50 assemblies and a high degree of completeness for all four species.

51 **Conclusions:** These four transcriptomes represent a significant expansion of data for insect
52 herbivores that feed exclusively on xylem sap, a nutritionally deficient dietary source relative to
53 other plant tissues and fluids. Comparison of transcriptome data with insect herbivores that
54 utilize other dietary sources may illuminate fundamental differences in the biochemistry of
55 dietary specialization.

56 **Keywords:** Transcriptome, RNA-seq, Trinity, Insect herbivory, Endosymbiont

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57 Data description

58 Background

59 Resource partitioning among herbivorous insects spans a continuum between specialists that feed
60 on one or a few plant species to generalists that are able to utilize hundreds of species belonging
61 to multiple plant families. A further element of plant partitioning involves the particular location
62 on a plant or tissue type from which an insect feeds [1]. The diversity of plant feeding strategies
63 has evolved along with specialized anatomical features such as mouthparts and digestive
64 systems, unique enzyme complements for processing plant compounds, and partnerships with
65 symbiotic microbiota that contribute to nutritional gain of the host insect. The transcriptome
66 assemblies presented here include four species that feed exclusively on sap from xylem vessels, a
67 relatively rare form of plant feeding from a source that among plant tissues is the most deficient
68 in nitrogen and carbon content [2]. Three of the transcriptomes represent sharpshooter species
69 (Cicadellinae) and the fourth belongs to a spittlebug (Clastopterae), all members of the
70 hemipteran suborder Auchenorrhyncha. Their piercing-sucking mouthparts tap into xylem
71 vessels from which sap is consumed in copious quantities to compensate for its low nutritional
72 value. Sharpshooters are recognized for their efficient assimilation of limited nutrients in xylem
73 sap [3], but putative biochemical mechanisms that enable specialization on xylem sap are
74 unknown. Also unclear is whether the respective roles in host nutrition played by the dual
75 primary endosymbionts are consistent among xylem feeders [4]. Comparison of transcriptomes
76 of four xylem-feeding insects will provide additional knowledge and insight into the survival of
77 ecological specialists on a nutritionally impoverished dietary source.

78 Samples

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4 79 The spittlebug *Clastoptera arizonana* Doering was collected in 2014 from a wild population
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7 80 infesting grapevines in Yavapai County, Arizona and established as a glasshouse colony for eight
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9 81 months prior to sample collection in Maricopa, AZ. Samples of the sharpshooter *Cuerna arida*
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11 82 Oman and Beamer (tribe Proconiini) were collected in 2015 by sweep net from a wild population
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14 83 in mixed vegetation in Cochise County, Arizona. The smoke-tree sharpshooter *Homalodisca*
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16 84 *liturata* Ball (Proconiini) was collected in 2015 from *Euphorbia tirucalli* L. plants in Phoenix,
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19 85 AZ. The blue-green sharpshooter *Graphocephala atropunctata* (Signoret) (Cicadellini) was
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21 86 collected in 2013 from a wild population in Orange County, California and maintained as a
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24 87 glasshouse colony on basil (*Osimium basilicum* L.) until samples were collected in 2015. Adult,
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26 88 whole-body samples from all four species were homogenized separately in RNeasy®
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28 89 (Ambion/Life Technologies, Carlsbad, CA) and stored at -20°C. Total RNA extraction, library
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31 90 generation (TruSeq RNA Sample Preparation Kit v2; Illumina Inc., San Diego, USA) and
32
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34 91 sequencing (Illumina HiSeq2000 or HiSeq2500) were performed at the University of Arizona
35
36 92 Genomics Center in Tucson, AZ (<http://uagc.arl.arizona.edu>).

93 94 **Data filtering**

95 The total number of reads, data quantity, and short read archive (SRA) numbers for each of the
96 four xylem-feeding insects are shown in Table 1. For each data set, raw quality was assessed and
97 filtered using both FastQC and Trimmomatic (v 0.32) using the parameters
98 ILLUMINACLIP:TruSeq3-PE.fa:2:30:10 LEADING:10 TRAILING:20
99 SLIDINGWINDOW:4:25 MINLEN:36 to remove adaptor sequence and filter by quality score.

100 **Transcriptome assembly**

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101 All raw data for each insect transcriptome were run through the following pipeline. Prior to
102 assembly, the three replicate samples were concatenated and read abundance was normalized to
103 50X coverage using the *in silico* normalization tool in Trinity v. 2.0.6 [5] to improve assembly
104 time. Each of the datasets were assembled in Trinity using the default parameters, with the
105 addition of the ‘-jaccard clip’ flag to reduce the generation of transcript fusions from non-strand
106 specific data. Open reading frames (ORFs) were predicted using Transdecoder [5]. The
107 transcriptomes were filtered, sorted, and prepared for NCBI transcriptome shotgun assembly
108 (TSA) submission as previously described [6].

109 **Annotation**

110 Functional annotation for each of the transcriptomes was performed at the peptide level using a
111 custom pipeline [6] that defines protein products and assigns transcript names. Predicted proteins
112 and peptides were analyzed using InterProScan 5 [7], using the ‘-iprlookup’ and ‘-goterms’
113 flags, to search all available databases, including Gene Ontology (GO). Each transcriptome was
114 annotated using BLASTp against the UniProt Swiss Prot database (downloaded 11 February
115 2015). Annie [8], a program that cross-references SwissProt BLAST and InterProScan5 results to
116 extract qualified gene names and products, was used to generate the transcript annotation file.
117 The resulting .gff3 and .tbl files were further annotated with functional descriptors in
118 Transvestigator [9].

119
120 **Transcriptome Quality and Comparisons**

121 Each of the assembled transcriptomes for the four xylem-feeding insects was used in a reciprocal
122 tBLASTx search to identify similarities between the four species and their transcriptome

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4 123 assemblies. The final, filtered transcriptomes were made into nucleotide BLAST databases using
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6 124 NCBI Blast+ (v 2.2.30) *makeblastdb* tool and all tBLASTx searches were performed using an e-
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9 125 value cutoff of $1e^{-3}$. The tBLASTx results (Table 3) indicate similarities between the four xylem-
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12 126 feeder transcriptomes, with the lowest (38%) occurring between species (e.g. the spittlebug and
13
14 127 all sharpshooters) and the highest (84%) between the sharpshooters *H. liturata* and *C. arida* [12].
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17 128 Additional transcriptome metrics between assemblies show a high percentage of reads mapping
18
19 129 back to each transcriptome (Table 2) indicating successful assemblies. The TransRate [10] scores
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22 130 range from 0.16 to 0.42 and BUSCO v. 1.1.b1 (benchmarking universal single-copy orthologs)
23
24 131 results using the arthropod gene set (downloaded December 19, 2015) [11] indicate the four
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27 132 transcriptomes have a moderate to high level of completeness. It should be noted that both the
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30 133 TransRate value (0.16) and BUSCO results for *H. litruata* suggest this transcriptome may
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32 134 contain more partial transcripts than the other three assemblies.
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35 135 **Availability of supporting data**

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38 136 The filtered and annotated transcriptomes have been deposited in GenBank as a TSA under the
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41 137 accessions and BioProject numbers found in Table 1. Datasets further supporting the results of
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43 138 this article are available in the *GigaScience* repository, GigaDB [12].
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46 139 **Abbreviations**

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49 140 ALARC: Arid Land Agricultural Research Center; BUSCO: Bench-marking Universal Single-
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52 141 Copy Orthologs; GO: Gene Ontology; ORF: Open Reading Frame; SRA: Short Read Archive;
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54 142 TSA: Transcriptome Shotgun Assembly; USDA-ARS: United States Department of Agriculture-
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57 143 Agricultural Research Service
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145 **Competing Interests**

146 The authors declare that they have no competing interests.

147 **Authors Contribution**

148 SJC and CCC conceived and performed the experiments; EET analyzed the data and evaluated
149 the conclusions; EET, SJC, and CCC wrote the manuscript. All authors approved the final
150 manuscript.

151 **Acknowledgements**

152 The authors thank Tom Perring and Darcy Reed at UC Riverside for providing samples of *G.*
153 *atropunctata* for transcriptome evaluations. Bioinformatic analysis was performed on computing
154 resources available at ALARC. Mention of trade names or commercial products in this article is
155 solely for the purpose of providing specific information and does not imply recommendation or
156 endorsement by the US Department of Agriculture. USDA is an equal opportunity provider and
157 employer.

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166 three species of leafhoppers in response to the dilute nutrient content of xylem fluid. *J.*
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218 **Table 1.** Accession numbers for sequence reads and assembled transcripts for four species of
 219 xylem-feeding insects.

Sample	Reads	Size (Gb)	Short Read Archive	BioSample	BioProject
<i>Homalodisca liturata</i>	18,936,520	18.9	SRX1451710	SAMN04293489	PRJNA303151
			SRX1451711	SAMN04293490	“
			SRX1451712	SAMN04293491	“
<i>Clastoptera arizonana</i>	19,038,998	17.8	SRX1451715	SAMN04293493	PRJNA303152
			SRX1451717	SAMN04293494	“
			SRX1451718	SAMN04293495	“
<i>Cuerna arida</i>	14,667,040	18.3	SRX1451216	SAMN04292971	PRJNA303150
			SRX1451218	SAMN04292972	“
			SRX1451467	SAMN04292973	“
<i>Graphocephala atropunctata</i>	16,868,134	8.2	SRX1411425	SAMN04208332	PRJNA299492
			SRX1411426	SAMN04208333	“
			SRX1411427	SAMN04208334	“

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 223 **Table 2.** Comparison of transcriptome assembly statistics and BUSCO analysis results for four
 224 xylem-feeding insects.

	<i>H. liturata</i>	<i>C. arizonana</i>	<i>C. arida</i>	<i>G. atropunctata</i>
Assembly				
Normalized reads	9,468,260	9,519,499	10,714,375	32,429,458
Total no. transcripts	106,998	93,845	91,384	97,830
Average transcript length and range	954 (224 – 30,062)	1,232 (224 – 29,936)	901 (224 – 20,095)	962 (224 – 17,082)
Total assembled bases	102,317,189	115,686,868	79,785,471	94,141,447
N50	1,650	2,510	1,560	1,692
% GC	37	31	37	39
% mapping	84	91	88	95
TransRate Score	0.16	0.28	0.25	0.42
BUSCO				
Complete (%)	60	82	68	66
Duplicated (%)	23	42	26	24
Fragmented (%)	23	9.2	17	19
Missing (%)	15	8	14	13

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Table 3. Total percent matches from tBLASTx reciprocal searches. Transcriptome used as query on the left, and nucleotide database tBLASTx against which the query was performed is shown at the top.

	Nucleotide Database (% similarity)			
	<i>H. liturata</i>	<i>C. arizonana</i>	<i>C. arida</i>	<i>G. atropunctata</i>
<i>Homalodisca liturata</i>	--	42.94	76.31	56.2
<i>Clastoptera arizonana</i>	40.9	--	38.35	38.58
<i>Cuerna arida</i>	83.9	43.4	--	58.36
<i>Graphocephala atropunctata</i>	56.86	40.3	56.1	--

e-value $\leq 1^{E-3}$

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