Metatranscriptomics and Pyrosequencing Facilitate Discovery of Potential Viral Natural Enemies of the Invasive Caribbean Crazy Ant, *Nylanderia pubens*

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**Abstract**

**Background:** *Nylanderia pubens* (Forel) is an invasive ant species that in recent years has developed into a serious nuisance problem in the Caribbean and United States. A rapidly expanding range, explosive localized population growth, and control difficulties have elevated this ant to pest status. Professional entomologists and the pest control industry in the United States are urgently trying to understand its biology and develop effective control methods. Currently, no known biological-based control agents are available for use in controlling *N. pubens*.

**Methodology and Principal Findings:** Metagenomics and pyrosequencing techniques were employed to examine the transcriptome of field-collected *N. pubens* colonies in an effort to identify virus infections with potential to serve as control agents against this pest ant. Pyrosequencing (454-platform) of a non-normalized *N. pubens* expression library generated 1,306,177 raw sequence reads comprising 450 Mbp. Assembly resulted in generation of 59,017 non-redundant sequences, including 27,348 contigs and 31,669 singlets. BLAST analysis of these non-redundant sequences identified 51 of potential viral origin. Additional analyses winnowed this list of potential viruses to three that appear to replicate in *N. pubens*.

**Conclusions:** Pyrosequencing the transcriptome of field-collected samples of *N. pubens* has identified at least three sequences that are likely of viral origin and, in which, *N. pubens* serves as host. In addition, the *N. pubens* transcriptome provides a genetic resource for the scientific community which is especially important at this early stage of developing a knowledgebase for this new pest.

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**Introduction**

Metagenomic analysis [1] coupled with new generation sequencing technologies have revolutionized the way in which entomologists can search for potential pathogens for use as insect biological control agents. Historically, discovery of pathogens of insects relied upon arduous explorations for unhealthy or dying insects followed by the identification and isolation of the microbe(s) responsible; a process that often took years to complete [2,3]. Metagenomic analysis not only reduces discovery time to days, but it is also capable of identifying numerous pathogenic organisms simultaneously. Indeed, metagenomic analysis may even be utilized to examine insects retrospectively from archived specimens. A number of sequencing projects of environmental samples of insects have demonstrated successful discovery of viruses that show promise as insect control agents [4,5,6,7,8].

*Nylanderia pubens* (Forel), previously *Paratrechina pubens* [9], is an invasive ant species that in recent years has developed into a serious pest problem in the Caribbean and United States [10,11]. A rapidly expanding range, explosive localized population growth, and control difficulties have elevated this ant to pest status. Professional entomologists and the pest control industry in the United States are urgently trying to understand its biology and develop effective control methods [12,13,14]. Efforts have primarily focused on pursuing development of insecticide-based control strategies [15], as well as the effort presented here to identify self-sustaining, biological control agents specific to *N. pubens*. While viruses can be important biological control agents against pest insect populations [16], none are known to infect *N. pubens*. Therefore, the objective of this research was to employ a metagenomics approach coupled with 454-based sequencing technology to examine the *N. pubens* transcriptome for viral infections based on sequence homology/identity with known viral sequences. The ultimate goal is the exploitation of viral discoveries as biologically-based agents for controlling *N. pubens*. As an added benefit, transcriptome sequencing provides a genetic resource for
the scientific community which is especially important at this early stage of developing a knowledgebase for *N. pubens*.

**Methods**

**Ants**

*N. pubens* colonies were obtained from field sites located in Desoto (April, 2011), Hildborough (April, 2011), Alachua (March through May, 2011), and Duval (March, 2011) counties in Florida and subsequently maintained in the laboratory. No specific permits were required to collect these field specimens because they did not occur in locations protected in any way. The field collections did not involve endangered or protected species. Colonies were reared separately in nesting tubes described by Oi and Williams [17] and fed frozen crickets, live housefly larvae, 10% sucrose solution, and water. Identifications of ants were made based on characters listed in Trager [18] and LaPolla et al. [9]. However, there is uncertainty regarding species assignment of these ants currently being reported from Florida, Texas, Louisiana and Mississippi based on morphometric and DNA sequence data. Nonetheless, the current consensus regards these invasive ants as the same species (D. Gotzek, personal communication). Representative voucher specimens were collected and retained in 95% ethanol at the USDA-ARS, Center for Medical, Agricultural and Veterinary Entomology, Gainesville, Florida.

**mRNA extraction, purification, library construction, and sequencing**

Total RNA was extracted from samples of colonies of *N. pubens* by the Trizol (Invitrogen, Carlsbad, CA) method according to the manufacturer’s instructions. Samples were taken from nine colonies. A total of 609 ants of different life stages (workers, alates, queens, larvae, pupae, and eggs) were used to prepare the total RNA. RNA quality of each preparation was assessed by microfluidic analysis on an Agilent 2100 Bioanalyzer (Agilent, Cary, NC) using the RNA 6000 Nano kit according to the manufacturer’s instructions. Microfluidic assays were completed immediately after RNA extraction using a 1 μl volume of purified sample. RNA samples of acceptable quality were pooled and used as source material for mRNA purification. mRNA was isolated from the total RNA sample using the Oligotex mRNA Mini Kit (Qiagen, Valencia, CA) following the manufacturer’s instructions. The isolated mRNA was then utilized to prepare a non-normalized fragment library suitable for 454 platform sequencing using the NEBNext mRNA Sample Pre Reagent Set 2 (New England BioLabs, Ipswich, MA) following the manufacturer’s protocol. The library was used as template for emulsion PCR using the GS Titanium LV emulsion PCR Kit (Lib-L, Roche, Mannheim, Germany) following the manufacturer’s instructions. DNA beads generated from the emulsion PCR reactions were used for Titanium plate 454 sequencing, using the GS Titanium Sequencing Kit XLR70 (Roche). *De novo* assembly was performed for the generated sequencing data using the Newbler software (Roche).

**Bioinformatic analysis**

An initial assembly of the sequences was performed with Newbler Assembler Version 2.3 (454 Life Science, Branford, CT), employing masking and trimming sequencing repeats, primers and/or adaptors used in cDNA library preparation. These hybridized sequences (contigs and leftover singletons) were further assembled with Paracel Transcript Assembler version 3.0.0 (PTA; Paracel Inc., Pasadena, CA).

In PTA, all sequences were masked for universal and species-specific vector sequences, adaptors, and PCR primers used in cDNA library creation. *Escherichia coli* contamination and mitochondrial and ribosomal RNA genes were identified and removed from input sequences using default settings to ascertain the novelty of the sequences. The poly (A/T) tails and intrinsic repeats, such as simple sequence repeats and short interspersed elements (SINE), were annotated prior to clustering and assembly. Low base-call quality data were trimmed from the ends of individual sequences and sequences <75 bp were excluded from consideration during initial pair-wise comparisons. After cleanup, sequences were passed to the PTA clustering module for pair-wise comparison and then to the CAP3-based PTA assembly module for assembly. The PTA assembly was performed based on the sequences of the contigs and the leftover singletons generated from the Newbler assembly.

Large-scale homology database searches of the PTA sequence data set were conducted against the National Center for Biotechnology Information (NCBI) NR and NT databases using BLAST (blastx and blastn) [19] with an in-house computational pipeline. To obtain a more accurate and complete description of potential gene function for each queried sequence, the top 100 BLAST hits were retrieved. Sequences with the best scoring BLAST hit (≤1e⁻⁰⁷) and the corresponding gene ontology (GO) classification were annotated to the queried sequence [20]. GO term assignments were binned according to the categories, biological processes, cellular components, and molecular functions. BLAST results and GO term assignments were completed in BlastQuest, an SQL database developed by the Interdisciplinary Center for Biotechnology Research, University of Florida, that facilitates similarity-based sequence annotation with gene ontology information [21]. In addition, the sequences were characterized with respect to functionally annotated genes by BLAST searching against NCBI specific reference sequences (RefSeq) for *Homo sapiens* (38,556 sequences), *Drosophila* (21,099 sequences) and *Formicidae* (74,540 sequences). Queries were considered to have a clear homolog of the searched organism when e-values were ≤1e⁻⁴, the length of the aligned segment was ≥50 bp, and identity >85%, which essentially eliminated spurious hits while preventing elimination of medium-sized proteins.

**Data Availability**

Raw 454 reads and assembled contigs were deposited in the NCBI database. The *N. pubens* sequence data are publicly available and accessible through the NCBI website accession numbers Ant_454Assem_NCBI.sqn: JP773711 - JP820231.

**Characterization of ESTs with viral identity**

Sequences identified as exhibiting significant viral homology/identity were selected from the *N. pubens* annotated list and further evaluated in an attempt to establish their origin—viral, host, or otherwise. Evaluations were also conducted to ascertain whether identified viral sequences were simply being ingested by the ants or were replicating (i.e., *N. pubens* was serving as host). Figure 1 summarizes the step-by-step decision tree employed to determine the likelihood that a given EST was of viral origin. Note that the process illustrated in figure 1 served only as a general guide to identify sequences of non-viral origin and non-replicating (e.g., ingested) viral sequences. Based on previous studies [5,7], this winnowing method significantly improves the virus discovery process.

**RNA virus confirmation**

Oligonucleotide primers were designed to each EST with significant (e-score ≤1e⁻⁴) identity to the RNA viral sequences
(Table 1), PCR was conducted with RNase-treated DNA extracted from the same *N. pubens* colonies used in expression library creation. If PCR yielded an amplicon, it was concluded that the sequence was likely an *N. pubens* gene and experiments concerned with this sequence were terminated because all of the RNA virus sequences exhibited identity with single-stranded RNA viruses that do not integrate in their host genome (no DNA stage generated). Sequences not generating an amplicon by PCR were next

**Table 1.** Oligonucleotide primers designed to each corresponding *N. pubens* sequence with viral identity.

| Designation | Forward oligonucleotide primer (5′ → 3′) | Reverse oligonucleotide primer (5′ → 3′) |
|-------------|------------------------------------------|------------------------------------------|
| Assem.6302.C1 | p1179 TGGCTTCCAGGCTACATGAGTACCTCA | p1180 AGGATATAAATCAGGGAACACATGACATCCA |
| G492870015H2 | p1255 CCAGGACGAATCTGACCTAGCTACGTCG | p1256 GCTGGCCATACCCGCAACGCG |
| G492870015JQT | p1256 CCCAGCAAAATCTGACCTACGTCGTTG | p1257 ATCCGCAACCTGGCATGATC |
| 492870015PUMA | p1257 GCTGCTTCCGCGGCTACCTACGTCG | p1258 ATGCTGATGAGTCGACTGAG |
| G492870015QXGN | p1260 ACACCTGAACCGGCTACCTACGTCG | p1261 GCTGGCCATACCCGCAACGCG |
| G492870015ORGU | p1261 ATCCGCAACCTGGCATGATC | p1262 GTGGGCTAACCTGCAACGCG |
| G492870015QXJ | p1262 GTGGGCTAACCTGCAACGCG | p1263 ATGCTGATGAGTCGACTGAG |
| Assem.15438.C1 | p1263 GTGGGCTAACCTGCAACGCG | p1264 GCTGGCCATACCCGCAACGCG |
| Assem.10577.C1 | p1264 GCTGGCCATACCCGCAACGCG | p1265 ATGCTGATGAGTCGACTGAG |
| Assem.3776.C1 | p1265 ATGCTGATGAGTCGACTGAG | p1266 GCTGGCCATACCCGCAACGCG |
| Assem.2829.C1 | p1266 GCTGGCCATACCCGCAACGCG | p1267 ATGCTGATGAGTCGACTGAG |
| Assem.13129.C1 | p1267 ATGCTGATGAGTCGACTGAG | p1268 GCTGGCCATACCCGCAACGCG |
| Assem.13541.C1 | p1268 GCTGGCCATACCCGCAACGCG | p1269 ATGCTGATGAGTCGACTGAG |
| Assem.13287.C1 | p1269 ATGCTGATGAGTCGACTGAG | p1270 GCTGGCCATACCCGCAACGCG |
| Assem.8702.C1 | p1270 GCTGGCCATACCCGCAACGCG | p1271 ATGCTGATGAGTCGACTGAG |
| Assem.3776.C1 | p1271 ATGCTGATGAGTCGACTGAG | p1272 GCTGGCCATACCCGCAACGCG |
| Assem.2829.C1 | p1272 GCTGGCCATACCCGCAACGCG | p1273 ATGCTGATGAGTCGACTGAG |
| G492870015QXJ | p1273 ATGCTGATGAGTCGACTGAG | p1274 GCTGGCCATACCCGCAACGCG |
| G492870015QXJ | p1274 GCTGGCCATACCCGCAACGCG | p1275 ATGCTGATGAGTCGACTGAG |
| Assem.4695.C1 | p1275 ATGCTGATGAGTCGACTGAG | p1276 GCTGGCCATACCCGCAACGCG |
| Assem.19410.C1 | p1276 GCTGGCCATACCCGCAACGCG | p1277 ATGCTGATGAGTCGACTGAG |
| Assem.16207.C1 | p1277 ATGCTGATGAGTCGACTGAG | p1278 GCTGGCCATACCCGCAACGCG |
| Assem.13720.C1 | p1278 GCTGGCCATACCCGCAACGCG | p1279 ATGCTGATGAGTCGACTGAG |
evaluated by RT-PCR. Ants from field-collected colonies were evaluated by RT-PCR for the presence of sequence template. If amplification was observed in 100% of the samples, it was assumed that the sequence was of host (or other) origin and further experiments with the sequence were terminated. Viral infections rarely exhibit an incidence of 100% among field-collected arthropods [22]. If amplification was observed in less than 100% of the samples, tagged-RT-PCR was conducted with the appropriate oligonucleotide primers to detect the replicating genomic strand [23]. This method permits discrimination of each genome strand without carryover effects causing false positive detection of either strand [23]. Tagged-RT-PCR employed the use of the appropriate oligonucleotide primer (Table 1) appended at the 5′ end with a TAG sequence (5′GGCCCGTCATGGTGCAATAA) that was used in a cDNA-synthesis reaction (forward primer for positive strand viruses and reverse primer for negative strand viruses). Two-step RT-PCR was employed to amplify a portion of the genome strand. First, 1 μl (50 ng) of total RNA was mixed with 10 mM dNTPs, 1 μM of the appropriate tagged oligonucleotide primer, heated to 65°C for 5 minutes, and then placed on ice for at least 1 minute. First strand buffer and Superscript reverse transcriptase (RT, Invitrogen) were then added and the reaction mixture was incubated at 55°C for 1 hour before inactivating the RT at 70°C for 15 minutes. Unincorporated oligonucleotides were digested with 10 units of Exonuclease I (New England Biolabs, Ipswich, MA) at 37°C for 1 hour. The reaction was terminated by heating to 80°C for 20 minutes. PCR was subsequently conducted in a 25 μl volume containing 2 mM MgCl₂, 200 μM dNTP mix, 0.5 units of Platinum Taq DNA polymerase (Invitrogen), 0.2 μM of each oligonucleotide primer, and 5 μl of the cDNA preparation. PCR products were separated on a 1% agarose gel and visualized by SYBR-safe (Invitrogen) staining. If the replicating strand of the virus was not detected, it was assumed the ant was not serving as host to the putative virus (e.g., may have simply been ingested) and further experiments with the sequence were terminated. If the replicating strand was detected by tagged-PCR, the amplicon was cloned and its sequence verified by Sanger sequencing. In these instances, sequences were considered to be likely from a virus infecting N. pubens and for the purposes of this study further experiments were terminated.

DNA virus confirmation

Field-collected N. pubens colonies (Alachua and Hillsborough counties, Florida) were evaluated by PCR for the presence of sequence template. If amplification was observed in 100% of the samples, it was assumed that the sequence was of host origin and
**Results and Discussion**

454 sequencing data assembly and annotation

The single production GS-FLX Titanium 454 platform sequencing run (two half plates) of the non-normalized *N. pubens* expression library generated 1,306,177 raw sequence reads comprising 450 Mbp (Fig. 2). *De novo* assembly of the raw data with Newbler yielded 22,044 contigs and 232,338 singletons. Subsequent assembly by PTA resulted in generation of 59,017 non-redundant sequences, including 27,948 contigs (average size 794 bp) and 31,669 singlets (average size 295 bp). Among these sequences, 27.9% (16,458) were greater than 500 bp and 72.1% (42,533) were greater than 300 bp. BLASTX analysis of these non-redundant sequences identified 25,898 (43.9%) with significant (e-value ≤ 1e-5) similarity and 33,119 (56.1%) returned no significant similarity. A significant percentage (47%) of the gene sequences (12,174) identified were found to be unique to *N. pubens*.

When categorized taxonomically based on the significant (expectation score ≤ 1e-5) BLAST results, 98.1% (25,405) of the *N. pubens* annotated sequences exhibited identity with genes from Animalia (Fig. 2). The remaining 1.9% of the sequences exhibited identity with known genes from Monera (1.6%), Fungi (0.19%), Plantae (0.07%), Protista (0.02%), and Viruses (0.08%). Further analysis showed that 97.2% of the annotated sequences exhibited identity with genes from organisms within the phylum Arthropoda. Subsequent categorization (Fig. 2) showed that 97.9% of the Arthropoda genes exhibited identity to Hymenoptera (bees, wasps, and ants) and among these, 92.7% to Formicidae (ants). Among the sequences that showed identity to ant genes, the majority (70.3%) exhibited identity with ants within the subfamily, Formicinae (specifically, *Camponotus floridanus*). The remaining sequences with identity to ants included the subfamilies Ponerinae (12.8%) and Myrmicinae (16.7%). Thus, the majority of the annotated *N. pubens* sequences yielded identity with its taxonomically nearest neighbor represented in the Genbank database, specifically, *C. floridanus* are both members of the Formicineae.

Categorization of each *N. pubens* annotated sequence (n = 25,898) returning a known function was summarized by assigning a gene ontology (GO) term when available (biological processes, cellular components, or molecular functions). GO analysis was also completed for reference sequences of *Homo sapiens*, *Drosophila melanogaster*, and Formicidae for comparison. The distribution of *N. pubens* genes among GO categories was essentially similar regardless of taxon (Fig. 4). In each of the three main GO categories, metabolic process, cellular process, cell, cell part, and binding terms were dominant (each category represented >5% of the annotated gene functions). Also well represented were proteins involved in regulation of biological process, biological regulation, catalytic activity, and organelle (each representing 3.5 to 5% of the annotated gene functions). As expected, the most significant departures were observed between *Homo sapiens* and *N. pubens* GO categories. Conversely, the gene representation was more consistent among *Drosophila melanogaster*, Formicidae, and *N. pubens*.

Viral discovery

BLAST analysis of the 59,017 non-redundant sequences yielded from the *N. pubens* library resulted in the identification of 51 sequences of putative viral origin (Table S1 and Table 2). Among them, 31 sequences did not meet the threshold for significance (an expectation score > 1e-4) and were not examined further to establish their source, viral, host, or otherwise (Table S1). However, despite expectation scores greater than 1e-4, some of these sequences could represent viruses that infect *N. pubens*. Indeed, in a similar study, putative viral sequences from an expression library from the red imported fire ant, *Solenopsis invicta*, which were not considered significant by BLAST analysis, were ultimately found to be of viral origin [5,7]. Database underrepresentation or distant relationships may contribute to a non-significant BLAST expectation score. Among the *N. pubens* sequences with BLAST expectation scores greater than 1e-4 (Table S1), 24 indicated identity with DNA viruses and 7 to RNA viruses. Many of the putative DNA (41.9%) and RNA (71.4%) viral genes were from virus families with a host range that does not normally include arthropods.

Twenty sequences from the *N. pubens* expression library yielded significant BLAST expectation scores of putative viral origin (Table 2); nine sequences were similar to DNA and eleven to RNA virus sequences. Among the potential DNA virus sequences, six exhibited identity with bacterial-infecting phage in the Podoviridae, Siphoviridae, and Myoviridae and were excluded from further experimentation (Table 2). Three sequences exhibited strong similarity with protein genes 3, 4, and 5 of the polydnavirus, Hyposoter didymator virus. This finding could potentially represent
the discovery of an ancestral polydnavirus. However, PCR conducted with oligonucleotide primers (Table 1) designed toward each of these sequences (Assem.6302.C1, 49287O01EPUMA, and G49287O01BKDVJ) resulted in amplification (of anticipated size) from all *N. pubens* field colonies examined (*n* = 8). Polydnaviruses are known to infect insect parasitoids in a limited number of subfamilies of the Braconidae and Ichneumonidae [25]. As these viruses integrate into the host genome, portions of the wasp’s genome are thought to incorporate into the virus genome—most likely because they proved useful for successful parasitism [25,26,27]. Thus, although sequence identity suggests polydnaviral origin, it is more likely that these sequences represent related insect (specifically Hymenopteran) genes.

The remaining eleven sequences were related to genes of RNA viruses (Table 2); six negative and five positive sense, single-stranded RNA virus genes were identified. The negative-sense RNA viruses included six viruses from two families (*Bunyaviridae* and *Rhabdoviridae*) and the positive-sense virus genes were all related to unclassified viruses, four of which infect arthropods.

Sequences Assem.2829.C1, G49287O01APTQA, Assem.16207.C1, and Assem.13720.C1 were considered to be of non-viral origin because 100% of the *N. pubens* colonies examined from the field (*n* = 6 to 8) produced amplicons of anticipated size from a DNA template with oligonucleotide primers designed to each respective sequence, suggesting that they were likely to be part of the ant genome (Fig. 1). Conversely, oligonucleotide primers designed to sequences Assem.13541.C1, Assem.13129.C1, Assem.4695.C1, Assem.19410.C1, Assem.13287.C1, Assem.8702.C1, and Assem.3776.C1 failed to yield an amplicon from *N. pubens* DNA template.

*N. pubens* field colonies were examined for the presence of sequences Assem.13541.C1, Assem.13129.C1, Assem.4695.C1, and Assem.19410.C1 by RT-PCR. Oligonucleotide primer p1258 (but not p1257) designed to Assem.13541.C1 successfully produced cDNA and subsequently an amplicon of anticipated size by PCR in 100% of the *N. pubens* field colonies evaluated suggesting that this gene sequence was likely of ant origin (Table 3). Similar results were also observed for sequences Assem.13129.C1, Assem.4695.C1 and Assem.19410.C1. However, amplicons were generated from all field samples evaluated regardless of which oligonucleotide primer (p1241/p1242, p1197/p1198, p1199/p1200, respectively) was used for cDNA synthesis. These results suggest the presence of ambisense RNA template possibly of viral origin. Assem.19410.C1 exhibited identity with a bunyavirus (Table 3) that are known to contain
Table 2. *N. pubens* transcriptome sequences yielding a significant (≤1e-6) expectation score from BLAST analysis and exhibiting viral identity.

| Designation      | Size    | E-value | Gene                     | Virus                      | Genome       | Family         | Host range       |
|------------------|---------|---------|--------------------------|----------------------------|--------------|----------------|-----------------|
| Assem.6302.C1    | 954     | 6.05e-06| Hypothetical protein 4    | Hyposoter didymator virus  | dsDNA        | Polydnaviridae | Invertebrates    |
| 49287001PUMA     | 335     | 1.40e-20| Hypothetical protein 3    | Hyposoter didymator virus  | dsDNA        | Polydnaviridae | Invertebrates    |
| G49287001KDVJ    | 483     | 1.14e-14| Hypothetical protein 5    | Hyposoter didymator virus  | dsDNA        | Polydnaviridae | Invertebrates    |
| G49287001SH2T    | 329     | 1.04e-31| Virion structural protein | Myxococcus phage Mx8       | dsDNA        | Podoviridae    | Bacteria         |
| G49287002FSJQT   | 329     | 1.04e-31| Virion structural protein | Myxococcus phage Mx8       | dsDNA        | Podoviridae    | Bacteria         |
| G49287001A0XGN   | 520     | 2.66e-20| Hypothetical protein      | Listonella phage phhISIC    | dsDNA        | Siphoviridae   | Bacteria         |
| G49287002GATUG   | 335     | 1.23e-06| DNA polymerase            | Pseudomonas phage           | dsDNA        | Myoviridae     | Bacteria         |
| Assem.15438.C1   | 503     | 1.35e-07| Hypothetical protein      | Vibrio phage                | dsDNA        | Myoviridae     | Bacteria         |
| Assem.10577.C1   | 392     | 2.03e-07| Hypothetical protein      | Aeromonas phage             | dsDNA        | Myoviridae     | Bacteria         |
| Assem.2829.C1    | 1628    | 1.25e-17| RNA polymerase            | Uukuniemi virus             | ss(+)RNA     | Bunyaviridae   | Vertebrates, plants |
| Assem.13129.C1   | 490     | 3.99e-19| RNA polymerase            | Uukuniemi virus             | ss(+)RNA     | Bunyaviridae   | Vertebrates, plants |
| Assem.16207.C1   | 711     | 1.32e-06| RNA polymerase            | Gouleako virus              | ss(+)RNA     | Bunyaviridae   | Vertebrates, plants |
| Assem.13541.C1   | 395     | 1.17e-14| L protein                 | Strawberry crinkle virus    | ss(+)RNA     | Rhabdoviridae | Vertebrates, plants |
| G49287001A0PTQA  | 427     | 6.82e-14| Polymerase                | Iranian maize mosaic        | ss(+)RNA     | Rhabdoviridae | Vertebrates, plants |
| Assem.19410.C1   | 413     | 8.79e-07| Polymerase                | West Caucasian bat virus    | ss(+)RNA     | Rhabdoviridae | Vertebrates, plants |
| Assem.3776.C1    | 1328    | 2.62e-25| Nonstructural polyprotein | Solenopsis invicta virus 2  | ss(+)RNA     | Unclassified  | Invertebrates    |
| Assem.13287.C1   | 570     | 2.18e-14| Polyprotein               | Kelp fly virus              | ss(+)RNA     | Unclassified  | Invertebrates    |
| Assem.8702.C1    | 1282    | 3.48e-14| Nonstructural polyprotein | Solenopsis invicta virus 2  | ss(+)RNA     | Unclassified  | Invertebrates    |
| Assem.4695.C1    | 953     | 4.02e-11| RNA polymerase            | Rice grassy stunt virus     | ss(+)RNA     | Unclassified  | Plants           |
| Assem.13720.C1   | 402     | 3.40e-06| Capsid protein            | Nilaparvata lugens          | ss(+)RNA     | Unclassified  | Invertebrates    |

Each sequence designation, its corresponding size, BLAST results (expectation value, gene and virus relatedness), and characteristics (nucleic acid composition, family and host range) are provided. DNA viruses are consolidated in the upper panel and RNA viruses in the lower panel.

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Table 3. Proportion of *N. pubens* field-collected colony samples producing an amplicon with oligonucleotide primers (see Table 1) designed to each sequence indicated.

| Sequence       | PCR (DNA template) [n] | RT-PCR (RNA template) [n] | Tagged RT-PCR (RNA template) [n] |
|----------------|------------------------|---------------------------|----------------------------------|
| Assem.2829.C1  | 100 [8]                | -                         | -                                |
| G49287001A0PTQA| 100 [8]                | -                         | -                                |
| Assem.16207.C1 | 100 [6]                | -                         | -                                |
| Assem.13720.C1 | 100 [6]                | -                         | -                                |
| Assem.13541.C1 | 0 [8]                  | 100 [6]                   | -                                |
| Assem.13129.C1 | 0 [11]                 | 100 [8]*                  | -                                |
| Assem.4695.C1  | 0 [13]                 | 100 [16]*                 | -                                |
| Assem.19410.C1 | 0 [13]                 | 100 [16]*                 | -                                |
| Assem.13287.C1 | 0 [6]                  | 66 [6]                    | 7.7 [26]                         |
| Assem.8702.C1  | 0 [6]                  | 66 [6]                    | 2 [50]                           |
| Assem.3776.C1  | 0 [6]                  | 50 [6]                    | 7.7 [26]                         |

In each column, the percentage of samples and the total number evaluated (n) yielding an amplicon are provided.

*For* positive strand viruses, the forward tagged primer was used to detect the negative (or replicative) viral genome strand. Similarly, for negative strand viruses the reverse tagged primer was used to detect the positive (or replicative) viral genome strand.

Table 3. Proportion of *N. pubens* field-collected colony samples producing an amplicon with oligonucleotide primers (see Table 1) designed to each sequence indicated.

| Sequence       | PCR (DNA template) [n] | RT-PCR (RNA template) [n] | Tagged RT-PCR (RNA template) [n] |
|----------------|------------------------|---------------------------|----------------------------------|
| Assem.2829.C1  | 100 [8]                | -                         | -                                |
| G49287001A0PTQA| 100 [8]                | -                         | -                                |
| Assem.16207.C1 | 100 [6]                | -                         | -                                |
| Assem.13720.C1 | 100 [6]                | -                         | -                                |
| Assem.13541.C1 | 0 [8]                  | 100 [6]                   | -                                |
| Assem.13129.C1 | 0 [11]                 | 100 [8]*                  | -                                |
| Assem.4695.C1  | 0 [13]                 | 100 [16]*                 | -                                |
| Assem.19410.C1 | 0 [13]                 | 100 [16]*                 | -                                |
| Assem.13287.C1 | 0 [6]                  | 66 [6]                    | 7.7 [26]                         |
| Assem.8702.C1  | 0 [6]                  | 66 [6]                    | 2 [50]                           |
| Assem.3776.C1  | 0 [6]                  | 50 [6]                    | 7.7 [26]                         |

In each column, the percentage of samples and the total number evaluated (n) yielding an amplicon are provided.

*For* positive strand viruses, the forward tagged primer was used to detect the negative (or replicative) viral genome strand. Similarly, for negative strand viruses the reverse tagged primer was used to detect the positive (or replicative) viral genome strand.

*All* (100%) of these field samples yielded an amplicon when either the forward or reverse oligonucleotide primer was used in cDNA synthesis suggesting the presence of ambisense RNA.”
periods of active replication [30,31]. Therefore, the presence of both genome strands would not be anticipated among 100% of field-collected colonies.

The most promising virus leads appear to be sequences Assem.13287.C1, Assem.8702.C1, and Assem.3776.C1 (Table 3). These sequences exhibited identity to unclassified positive-strand RNA viruses. Specifically, Assem.13287.C1 exhibited identity with the 3C-like protease region of polyprotein 1 (5’-proximal ORF) of Kelp fly virus [32] and Assem.8702.C1 exhibited identity with the helicase region (ATPase chaperone functionality) of Kelp fly virus [32] and Assem.8702.C1 exhibited identity with the 3C-like protease region of polyprotein 1 (5’-proximal ORF) of Solenopsis invicta virus 3 (SINV-3) [4]. Assem.3776.C1 also exhibited identity with SINV-3 at the interface of the 3C-protease and RNA-dependent RNA polymerase. Oligonucleotide primers designed to these sequences failed to yield an amplicon by PCR (DNA template) among field-collected N. pubens colonies. However, 50 to 66% of field-collected colonies did generate an amplicon of anticipated size by RT-PCR with oligonucleotide primers designed to each sequence. The replicative form of the genome (minus or negative strand) was detected in a small percentage of field-collected N. pubens colonies providing strong evidence that these sequences correspond to positive-strand RNA viruses that infect N. pubens. All of these sequences exhibited identity to viruses that infect arthropods (Diptera and Hymenoptera). Thus, detection of Assem.13287.C1, Assem.8702.C1, and Assem.3776.C1 by targeted-RT-PCR among a small percentage of field-collected N. pubens colonies suggests that these sequences are of viral origin and appear to replicate in N. pubens. Sanger sequencing of these amplicons verified their identity. Although not used as part of our screening method, host gene induction in response to viral infection can also be employed to identify viral infections indirectly. Indeed, a number of virus-induced genes/pathways were detected in the N. pubens transcriptome, including Toll like receptors, STAT transcription factor and Jak kinase. However, Vago, an important participant in the control of viral load in Drosophila, was not detected [33].

Pyrosequencing of entomological samples in an effort to discover viruses [7] and other organisms [34] for use as insect microbial control agents has proven an efficient method. For example, searches for virus infections in S. invicta by traditional means were conducted for decades without a single discovery [35,36,37,38] which ultimately delayed the available use of viruses as natural enemies to control the ant. However, using the metagenomics approach, three RNA viruses were discovered in S. invicta which offer a new approach to control these ants [39]. Pyrosequencing of the N. pubens transcriptome from numerous colony sources collected from different locations throughout the state of Florida has, similarly, identified 51 sequences of putative viral origin; at least three of these sequences appear to be from virus sources that replicate in N. pubens. Efforts to acquire additional genome sequence and characterize the biology and impact of these potential viruses have begun in earnest.

Although the primary objective of the N. pubens pyrosequencing project was discovery of viruses for use in controlling this ant, the transcriptome of this pest ant was also elucidated and is now available publicly. This genetic resource will prove helpful in many future studies to understand and control, N. pubens, especially at this incipient stage of its expanding U.S. infestation. The N. pubens transcriptome represents the 7th ant species (and only the 2nd species in the Formicinae) to undergo a large sequencing project. These data will facilitate comparative genomic studies with the genomes of Camponotus floridanus (Formicinae), Harpagognathus salitator (Ponerinae), Solenopsis invicta (Myrmicinae), Pogonomyrmex barbatus (Myrmicinae), Lasidahema humile (Dolichoderinae), and Atta cephalotes (Myrmicinae) [24,40,41,42,43].

Supporting Information

Table S1 N. pubens transcriptome sequences yielding a non-significant (> 1e−4) expectation score from BLAST analysis with viral identity. Each sequence designation, corresponding BLAST results (gene and virus relatedness), and characteristics (nucleic acid composition, family and host range) are provided. DNA viruses are consolidated in the upper panel and RNA viruses in the lower panel.

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Author Contributions

Conceived and designed the experiments: SMV DHO EAB. Performed the experiments: SMV DHO XXT. Analyzed the data: SMV DHO EAB FY XXT. Wrote the paper: SMV DHO EAB FY XXT.

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