De Novo Foliar Transcriptome of *Chenopodium amaranticolor* and Analysis of Its Gene Expression During Virus-Induced Hypersensitive Response

Yongqiang Zhang¹, Xinwu Pei¹, Chao Zhang¹, Zifeng Lu², Zhixing Wang¹, Shirong Jia¹, Weimin Li¹*

¹ Biotechnology Research Institute, Chinese Academy of Agricultural Sciences, Beijing, People’s Republic of China, ² State Key Laboratory of Grassland Agro-ecosystems, College of Pastoral Agriculture Science and Technology, Lanzhou University, Lanzhou, People’s Republic of China

**Abstract**

**Background:** The hypersensitive response (HR) system of *Chenopodium* spp. confers broad-spectrum virus resistance. However, little knowledge exists at the genomic level for *Chenopodium*, thus impeding the advanced molecular research of this attractive feature. Hence, we took advantage of RNA-seq to survey the foliar transcriptome of *C. amaranticolor*, a *Chenopodium* species widely used as laboratory indicator for pathogenic viruses, in order to facilitate the characterization of the HR-type of virus resistance.

**Methodology and Principal Findings:** Using Illumina HiSeq™ 2000 platform, we obtained 39,868,984 reads with 3,588,208,560 bp, which were assembled into 112,452 unigenes (3,847 clusters and 108,605 singletons). BlastX search against the NCBI NR database identified 61,698 sequences with a cut-off E-value above 10⁻³. Assembled sequences were annotated with gene descriptions, GO, COG and KEGG terms, respectively. A total number of 738 resistance gene analogs (RGAs) and homology sequences of 6 key signaling proteins within the R proteins-directed signaling pathway were identified. Based on this transcriptome data, we investigated the gene expression profiles over the stage of HR induced by *Tobacco mosaic virus* and *Cucumber mosaic virus* by using digital gene expression analysis. Numerous candidate genes specifically or commonly regulated by these two distinct viruses at early and late stages of the HR were identified, and the dynamic changes of the differently expressed genes enriched in the pathway of plant-pathogen interaction were particularly emphasized.

**Conclusions:** To our knowledge, this study is the first description of the genetic makeup of *C. amaranticolor*, providing deep insight into the comprehensive gene expression information at transcriptional level in this species. The 738 RGAs as well as the differentially regulated genes, particularly the common genes regulated by both TMV and CMV, are suitable candidates which merit further functional characterization to dissect the molecular mechanisms and regulatory pathways of the HR-type of virus resistance in *Chenopodium*.

**Citation:** Zhang Y, Pei X, Zhang C, Lu Z, Wang Z, et al. (2012) De Novo Foliar Transcriptome of *Chenopodium amaranticolor* and Analysis of Its Gene Expression During Virus-Induced Hypersensitive Response. PLoS ONE 7(9): e45953. doi:10.1371/journal.pone.0045953

**Editor:** Zhanjiang Liu, Auburn University, United States of America

**Received** June 16, 2012; **Accepted** August 23, 2012; **Published** September 28, 2012

**Copyright:** © 2012 Zhang et al. This is an open-access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

**Funding:** This work was supported by the National Major Special Project of China on New Varieties Cultivation for Transgenic Organisms (grant nos. 2011ZX08001-002 and 2011ZX08004-004), and the Special Fund for Agro-scientific Research in the Public Interest (grant no. 201103026-2). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Competing Interests:** The authors have declared that no competing interests exist.

* E-mail: liwm518@hotmail.com

**Introduction**

During the entire lifetime, plants are attacked by a broad range of pathogens, being the targets of bacteria, fungi, viruses, protozoa as well as nematodes. To combat the ever-changing landscape of biotic interactions, plants have evolved sophisticated and effective defense mechanisms. The hypersensitive response (HR) represents one of the most active resistance responses. This type resistance often occurs following the incompatible interactions between plants and pathogens, and is characterized by rapid necrosis of infected and neighboring host cells [1,2]. Regarding to the well known ‘Gene-for-Gene’ hypothesis [3], the HR depends on a genetic interaction, either directly or indirectly, between the product of a dominant or semidominant resistance (*R*) gene and a corresponding pathogen avirulence (*Avr*) gene product, thereby preventing pathogen growth and spread in plants [1,4,5].

The *R* gene-mediated HR usually is associated with rapid initiation of signal transduction pathways leading to expression of disease resistance responses, such as the oxidative burst, alteration of membrane potentials, increases in lipoxygenase activity, production of antimicrobial compounds, and the expression of defense-related genes [6]. Plants undergoing HR also induce a state of pathogen-nonspecific resistance throughout the plant, a phenomenon termed systemic acquired resistance [7]. This local response is therefore attractive in plant breeding programs, and characterization of host genetic elements involved in HR is assumed to have great potential for developing new strategies to reduce losses from plant disease. Over the past decades, the detailed analysis of many hypersensitive disease resistance systems
has substantially advanced the basic research on HR-type of resistance mechanisms. Progress has been made in characterizing R genes as well as the signal-transduction events that coordinate the R protein-directed HR-type plant defense [4,5,6].

The HR system in Chenopodium spp. is one of the most notable disease resistance mechanisms. Many plant viruses in over a dozen of genera, including bromo-, como-, cucum-, clostero-, potex-, poty- and tobamovirus have been documented to elicit local lesion HR on Chenopodium [9,10]. The broad-spectrum virus resistance suggests Chenopodium are rich in genes that act in restricting viral spread as well as that confer multi-virus resistance to a plant. However, understanding of the HR-type of resistance in Chenopodium is surprisingly incomplete. Due to lack of developed genetic tools such as tissue culture and transformation, only few studies were performed to define the genetic mechanisms of HR in Chenopodium [10–13]. Genomic information on Chenopodium spp. is rather limited, only 1,249 expressed sequence tags (EST) and 358 proteins have been deposited in Genbank as of May 2012.

The next-generation sequencing, referred to as RNA sequencing (RNA-Seq), provides a unique opportunity for genomic exploration in non-model plant species that do not have a reference genome sequence data [14,15]. So far, RNA-Seq has been successfully used for annotation, transcript profiling and single nucleotide polymorphism (SNP) discovery in a number of non-model plant species, such as alfalfa [16], Artemisia annua [17], buckwheat [18], California poppy [19], Eucalyptus grandis [20], grape [21], magnoliid avocado [19], orchid [22] and Pachycladon engyi [23].

In order to facilitate the investigation of the HR-type of virus resistance in Chenopodium, herein we took advantage of RNA-seq to survey the foliar transcriptome of C. amaranticolor, one Chenopodium species being most widely used as laboratory indicator for pathogenic viruses (Figure 1A). Over 3.5 gigabase pairs of high-quality DNA sequence were generated with Illumina technology. After de novo assembly and annotation, a sufficiently large transcriptome database containing 112,452 distinct sequences was built. By using Illumina’s digital gene expression (DGE) platform, this database was immediately applied to analyze the gene expression profiles over the stage of HR induced by platform, this database was immediately applied to analyze the

Table 1. General features of the foliar transcriptome of C. amaranticolor.

| Feature                        | Value                  |
|-------------------------------|------------------------|
| Total number of reads          | 39,868,984             |
| Total base pairs (bp)          | 3,588,208,560          |
| Average read length (bp)       | 90                     |
| Total number of contigs        | 326,027                |
| Mean length of contigs (bp)    | 163                    |
| Total number of scaffolds      | 172,748                |
| Mean length of scaffolds (bp)  | 253                    |
| Distinct clusters              | 3,848                  |
| Distinct singletons            | 108,605                |
| Total distinct sequences       | 112,452                |
| Sequences with E-value<10^−5   | 62,482                 |

*Adaptors and low-quality reads were excluded. doi:10.1371/journal.pone.0045953.t001

Figure 1. Virus induced local lesions on leaves of C. amaranticolor. (A) The 6-week old plant of C. amaranticolor. (B) Leaves inoculated with Tobacco mosaic virus at 40 hours p.i.. (C) Leaves inoculated with Cucumber mosaic virus at 40 hours p.i.. Scale bar = 1.0 cm. doi:10.1371/journal.pone.0045953.g001

These short 21-mers were then assembled into 326,027 contigs (Table 1) with average lengths of 163 bp, and their size distribution is shown in Figure S1A. Using paired-end joining and gap-filling, the contigs were assembled into 172,748 scaffolds with a mean size of 253 bp including 6,034 scaffolds larger than 800 bp (Table 1). These scaffolds were further analyzed by using TGICL software [26], resulting in 112,452 unigenes with 3,847 clusters (mean size: 575 bp) and 108,605 singletons (mean size: 307 bp) (Table 1). In this study, the singleton means a scaffold that failed to match other scaffolds and the unigenes include all the clusters and singletons generated from scaffolds. The size distribution of these distinct sequences is shown in Figure S1B. All files of assembled contigs and unigenes are available by request.
To verify the quality of sequencing data, 12 unigenes were randomly selected and pairs of primers were designed accordingly for RT-PCR amplification. Consequently, each of primer pairs generated a band with the expected size and the identity of all 12 PCR products were confirmed by Sanger sequencing (data not shown).

Annotation of predicted proteins

For annotation, distinct gene sequences were first searched against the NCBI NR database using a cut-off E-value of $10^{-5}$. Although lack of genomic information in *Chenopodium* species, totally 61,698 genes (54.9% of all distinct sequences) including 3,272 clusters and 38,426 singletons showed significant matches (Table S1). As shown in Figure 2, the fraction of singletons that had BLAST matches was lower than for the clusters given their shorter length (mean size of 307 bp). For instance, a 100% of singletons longer than 2,000 bp achieved significant BLAST scores, and the proportion decreased sharply to 50.4% for sequences ranging from 100 to 500 bp (Figure 2A). However, clusters longer than only 1,000 bp totally achieved significant BLAST scores, and the proportion remained about 79.6% for sequences ranged in size from 100 bp to 500 bp (Figure 2B). Notably, a total of 49,970 genes (45.1%) were found no homologues in the current NCBI NR database, similar with the observations which have been previously made for several other transcriptomic studies [27–30]. These orphan sequences may represent novel genes in *C. amaranticolor* or are possibly derived from chimerical sequences (assemblage errors), the cDNA of untranslated regions or non-conserved regions of proteins.

The E-value distribution of the top hits in the NR database showed that 15.3% of the mapped sequences have strong homology (<$1.0E^{-50}$), whereas 94.7% of the homolog sequences ranged between $1.0E^{-3}$ to $1.0E^{-50}$ (Figure 3A). The similarity distribution showed that 20.6% of the query sequences have a similarity higher than 80%, while 79.4% of the hits have a similarity ranging from 18% to 80% (Figure 3B). For species distribution, 27.9% of the distinct sequences have top matches (first hit) with sequences from the *Arabidopsis thaliana*, the genome of which has been sequenced in 2000 [31]. The next closest species were the *Oryza sativa* Japonica Group (11.6%), *A. lyrata* subsp. *Lyrata* (10.6%), *Populus trichocarpa* (7.1%), *Vitis vinifera* (6.6%) and *Zea mays* (3.2%) (Figure 3C). In addition, there were 772 and 152 distinct sequences with the highest homology to genes from *Spinacia oleracea* and *Chenopodium* spp., respectively. A total of 9,119 distinct sequences (14.8%) had the strongest similarity to genes of *O. sativa* Japonica Group and *Zea mays*, two types of monocot species, suggesting the interesting phylogenetic status of *C. amaranticolor*.

**GO, COG and KEGG classification**

Gene Ontology (GO) assignments were used to classify the functions of the predicted *C. amaranticolor* genes. Based on sequence homology, 35,162 sequences can be categorized into 43 functional groups (Figure 4). In each of the three main categories (biological process, cellular component and molecular function) of the GO classification, 'Metabolic process', 'Cell' and 'Catalytic activity' terms are dominant respectively. In agreement with GO assignments of other higher plant, such as *Taxus mairei* [32], *Camellia sinensis* [33], *Brassica juncea* [34] and *Siraitia grosvenorii* [35], we noticed a high-percentage of genes from categories of 'Cellular process', 'Cell part', 'Organelle' and 'Binding' (Figure 4), indicating that these GO terms are significant among all plant species and the mechanism of 'Cellular process' in different plants may be similar.

To further evaluate the completeness of our transcriptome library and the effectiveness of our annotation process, we searched the annotated sequences for the genes involved in the clusters of orthologous group (COG) classifications. In total, out of 61,698 NR hits, 24,147 sequences have a COG classification (Figure 5). Among the 24 COG categories, the cluster for 'General function prediction' represents the largest group (3,714, 15.4%) followed by 'Transcription' (1,968, 8.2%) and 'Replication, recombination and repair' (1,855, 7.7%), while the category of 'Nuclear structure' (5, 0.02%) represents the smallest group (Figure 5). It is noteworthy that 453 unigenes were assigned to the term of 'Defense mechanism'.

To identify the biological pathways that are active in *C. amaranticolor*, we mapped the 61,698 annotated sequences to the reference canonical pathways in Kyoto Encyclopedia of Genes and Genomes (KEGG) [36]. In total, 27,042 sequences were assigned to 119 KEGG pathways. The pathways with most representation by the unique sequences were 'Metabolic pathways' (6,477, 23.95%), 'Biosynthesis of secondary metabolites' (3,795, 14.03%), and 'Plant-pathogen interaction' (2,346, 8.68%),

**Figure 2. Effect of query length on the percentage of significant matches.** (A) Singleton sequences. (B) Cluster sequences. The proportion of sequences with matches (with a cut-off E-value of 1.0E–5) in NR databases is greater among the longer assembled sequences.

doi:10.1371/journal.pone.0045953.g002
providing a valuable genetic resource for the investigation of specific biological processes, functions and pathways in *C. amaranticolor*.

Detection of resistance gene analogs and disease resistance signaling genes

To date, the majority of cloned and functional *R* genes described within the plant kingdom contain a conserved nucleo-
tide binding site (NBS) and a C-terminal leucine-rich repeat (LRR) domain, known as NBS-LRRs [37,38]. NBS-LRR-encoding genes represent one of the largest and most variable gene families in plants [39], with 149 members in genome of A. thaliana [40], 92 in Brassica rapa [41], 400–500 in Medicago [42], 464 and 483 in two genomes of O. sativa [43], 438 in potato [44], 416, 535 and 54 in three woody species poplar, grapevine and papaya, respectively [45,46] (Yang et al., 2008; Porter et al., 2009). Herein, we compared the current transcriptome data with sequences from NCBI nucleotides and EST database to extract the unigenes representing candidate NBS-LRR genes, and in total 738 unigenes were identified as resistance gene analogs (RGAs) (Table S2). In addition, a number of disease resistance signaling proteins, including NDR1, Rar1, Sgt1, EDS1, PAD4 and NPR1, which play key roles in the R proteins-directed signaling pathways [47,48], were also identified from C. amaranticolor for the first time (Table S3). These overall findings establish a solid genetic basis for characterization of the R genes as well as the disease resistance signaling proteins in C. amaranticolor, and will certainly facilitate the future research of Chenopodium in R gene-mediated disease resistance.

**DGE library sequencing**

An immediate application of our transcriptome sequence data includes analysis of gene expression variation over the stage of virus-induced HR in C. amaranticolor leaves. Five cDNA libraries derived from healthy leaves, TMV-6h, TMV-28h, CMV-6h, and CMV-20h were constructed and sequenced by using Illumina HiSeq™ 2000, respectively. After filtering the low quality reads, 12.5, 11.7, 12.2, 12.5, and 12.4 million clean reads were generated from the five samples, respectively (Table 2). To reveal the molecular events behind DGE profiles, these reads were mapped to the C. amaranticolor foliar transcriptome containing 112,453 unigenes. In total of 6,189,719, 5,826,515, 6,121,996, 6,339,408 and 5,948,757 reads derived from healthy leaves, TMV-6h, TMV-28h, CMV-6h and CMV-20h uniquely match with the unigenes in the reference database, respectively (Table 2). Reads mapped to a unique sequence are the most critical subset of the DGE libraries as they can explicitly identify a transcript. In this study, at least 93.93% of sequences in our transcriptome database could be unequivocally identified by the unique match reads (Table 2). To confirm if the number of detected genes increases proportionally to sequencing amount (total tag number), a saturation analysis was performed. As summarized in Figure S2, the results revealed a trend of saturation, in which the number of detected genes almost ceases to increase when the number of reads reaches ~12 million. Next, the locations of DGE reads on reference genes were also evaluated [14], and the evenly distribution showed in Figure S3 indicated good randomness of these reads.

**Genes responding to virus inoculation in C. amaranticolor**

To identify genes showing a significant expression change in C. amaranticolor upon virus infection, the differentially expressed tags between virus-inoculated leaves (TMV-6h, CMV-6h, TMV-28h or CMV-20h) and healthy leaves were identified by an algorithm [49] based on the criteria of significance [False Discovery Rate (FDR)≤0.001 and |log2Ratio|≥1]. Accordingly, a total of 1,022,284 significantly changed tag entities were detected between healthy leaves and TMV-6h. Those tags were mapped to 3,297 genes with 1,636 genes up-regulated and 1,661 genes down-regulated, respectively, termed early-TMV-inducible genes (Figure 6 and Table S4). Between healthy leaves and TMV-28h, a total of 23,247 differentially expressed genes, termed late-TMV-inducible genes (Figure 6 and Table S5). Similarly, between healthy leaves and CMV-6h, 1,383 genes up-regulated and 1,961 genes down-regulated were defined as early-CMV-inducible genes (Figure 6 and Table S6), while between healthy and CMV-20h, a total of 17,458 differentially expressed genes, termed late-CMV-
inducible genes, were identified with roughly the same amount of up-regulated (9,862) and down-regulated genes (7,596) (Figure 6 and Table S7). Collectively, these four datasets showed that numbers of both up-regulated (>2-fold) and down-regulated (<2-fold) genes increased following either TMV- or CMV-inoculation.

Based on the four datasets, multiple comparisons were made to capture genes commonly regulated in *C. amaranticolor* upon different virus infection. As shown in the Venn diagrams (Figure 7A and 7B), a total of 1,238 unigenes including 319 up-regulated unigenes and 919 down-regulated unigenes were common between the early- and late-TMV-inducible genes (Table S8), while 286 up-regulated unigenes and 652 down-regulated unigenes overlapped between the early- and late-CMV-inducible genes (Table S9). Moreover, a total of 927 up-regulated unigenes and 1,038 down-regulated genes were common between the early-TMV- and early-CMV-inducible genes (Table S10), while between the late-TMV- and late-CMV-inducible genes, 6,181 up-regulated unigenes and 5,430 down-regulated genes, a total of 11,611 unigenes overlapped (Table S11). Further comparative analysis of the four datasets defined a core set of 461 unigenes comprising 93 up-regulated unigenes and 368 down-regulated unigenes which were commonly regulated by both TMV and CMV at all aforementioned post-inoculation times (Figure 7A and 7B and Table S12).

Efforts were also made to collect the alternatively regulated genes during the stage of HR induced by TMV or CMV. To this end, we identified that 275 unigenes were down-regulated in CMV-6h but up-regulated in CMV-20h, showing ‘down-up’ expression profiles, whereas 405 unigenes were regulated in the opposite direction (Table S13). For TMV infection, a total of 330 unigenes displayed ‘down-up’ expression profiles in TMV-6h/TMV-28h, while 194 unigenes showed ‘up-down’ expression profiles (Table S14). Further comparisons of these alternatively regulated genes identified 102 commonly ‘down-up’ regulated unigenes and 89 commonly ‘up-down’ regulated unigenes during both TMV and CMV infection (Table S15).

These overall datasets presented a significant number of genes commonly regulated in both TMV- and CMV-infected *C. amaranticolor*, suggesting the host plant might make use of some common pathways for defense against distinct viruses.

**Functional categorization of the identified genes**

Of the identified differentially expressed genes, efforts have been first put forward to functional categorization of the 93 commonly up-regulated unigenes and 368 down-regulated unigenes among the TMV-6h, TMV-28h, CMV-6h and CMV-20h. Apart from the unclassified ones, a total of 49 up-regulated unigenes and 111 down-regulated unigenes were categorized into various levels of GO biological processes, predominately in the ‘metabolism’ GO category including disaccharide, lipid, amino acids, protein, carbohydrate and primary metabolism, while the remaining ones were assigned to the GO processes of stress, defense, signal as well as transport (Table S12). Similarly, functional categorization of the 102 unigenes with ‘down-up’ expression profiles and 89 unigenes with ‘up-down’ profiles showed that, except the unclassified ones, the unigenes were also predominately categorized into the ‘metabolism’ GO category including protein and primary metabolism (Table S15).

To comprehensively assess the biological functions of the differentially expressed genes, all four sets of differentially expressed genes were mapped to KEGG database terms and compared with the whole transcriptome data, with a view to identify significantly enriched metabolic or signal transduction pathways. Among all the genes with KEGG pathway annotation, a total of 1,119, 1,174, 6,064 and 7,571 differentially expressed genes from the datasets of the early-TMV-inducible genes (TMV-6h), the late-TMV-inducible genes (TMV-28h), the early-CMV-inducible genes (CMV-6h) and the late-CMV-inducible genes (CMV-20h), were assigned to 97, 118, 102 and 118 KEGG pathways.
pathways, respectively, suggesting the virus infection induces global alteration of the genes transcription profiles in *C. amaranticolor*.

Specific observations were made for the pathway of plant-pathogen interaction (PPI), in which, a total of 155, 141, 954 and 685 differentially expressed genes derived from TMV-6h, CMV-6h, TMV-28h and CMV-20h, respectively, were enriched. Interestingly, in TMV-6h and CMV-6h which represent the early stage of HR, over 80% of the differentially expressed genes in PPI pathway were significantly down-regulated, including RIN4, RPS2 and RPS5, the key proteins mediate effector-triggered HR cell death [5,50]. Similarly, the host factors of EFR [51,52] and WRKY29 [53,54] in Mitogen-activated protein kinase (MAPK) signaling cascade, the well-known defense marker PR1 [55] and the essential gene COI1 for jasmonate-regulated defense [56] were all down-regulated at the early stage. In contrast, at the late stage of HR (CMV-20h and TMV-28h), the majority (>80%) of the differentially expressed genes enriched in PPI were highly up-regulated, such as RIN4, RPS5 and the MAPK signaling cascade-associated genes MKK4/5 [57] and WRKY25/33 [58–60]. It is noteworthy to mention the 'down-up' expression profiles of the *RIN*4 and *RPS5*, which suggest essential roles of these two genes during virus-induced HR and merit further investigation. In addition, a particular interest was given to NHO1, a non-host resistance gene [61,62], which is the unique gene consistently up-regulated in all four samples, indicating its yet undetermined role in virus resistance. Collectively, these observations defined that number of unigenes in PPI pathway underwent complicated regulation mechanisms during virus-induced HR, leading to a first dynamic picture of the changes in the overall pattern of gene expression in PPI pathway of *C. amaranticolor*.

Conclusions

This study represents the first application of Illumina sequenc- ing technology for genomic studies in *C. amaranticolor*, a host with the broad-spectrum virus resistance. A single run produced 112,453 unigenes with 62,482 sequences having an above cut-off BLAST result. From this foliar transcriptome database, 738 RGAs and a number of sequences represent disease resistance signaling proteins were identified. By using Illumina sequencing-based DGE system, we further analyzed the gene expression profiles during virus-induced HR in *C. amaranticolor* leaves, and identified numbers of candidate genes specifically and commonly regulated by TMV and CMV at early and late stages of the HR. This genome-scale transcriptional information provides a substantial contribution to the sequence resources for *C. amaranticolor* and will particularly aid in understanding the genetic mechanism of the broad-spectrum virus resistance conferred by this host.

Materials and Methods

Plant materials preparation

Leaves of 6~7 week-old *C. amaranticolor* grown in the greenhouse were inoculated with CMV SD strain [63] and TMV U1 [64], respectively. Briefly, infectious sap was prepared with one gram of fresh TMV- or CMV-infected *Nicotiana tabacum* leaves that were macerated in 10 mL of inoculation buffer (50 mM KH2PO4, pH 7.0, 1% Celite), and was mechanically inoculated onto *C. amaranticolor* leaves dusted with carborundum powder (600 mesh). The virus-inoculated leaf tissues were harvested using half leaf method as follows. Half of each virus-inoculated leaf was first sampled at 6 hpi, while the remaining halves were collected at 20 hpi for CMV-inoculated leaves and at 28 hpi for TMV-inoculated leaves, at which time point local lesions became just visible. These samples were accordingly named as TMV-6h, CMV-6h, TMV-28h and CMV-20h. Notably, the leaf tissues of CMV-6h or TMV-6h were selected only when the remaining halves showed significant number of local lesions at 20 hpi (CMV) or 28 hpi (TMV), in order to ensure the tissues of CMV-6h and TMV-6h being fully infected. Leaves of the healthy plants were also harvested as control. All samples were immediately frozen in liquid nitrogen and were stored at −80°C until use.

cDNA library preparation for transcriptome analysis

Total RNA was extracted using TRIzol® reagent (Invitrogen) according to the manufacturer’s protocol. RNA integrity was confirmed with RNA6000 Nano Assay using the 2100 Bioanalyzer (Agilent Technologies, Palo Alto, CA). The samples for transcriptome analysis were prepared using Illumina’s kit following manufacturer’s instruction. Briefly, using oligo (dT) magnetic beads (Invitrogen), poly(A) mRNA was first isolated from 6 μg of total RNA, which was extracted from the mixed materials containing equal amount of healthy leaves, TMV-8h, CMV-8h, TMV-28h, and CMV-20h. The mRNA was then fragmented into

![Figure 7. The generalized Venn diagrams with four datasets of TMV-6h, TMV-28h, CMV-6h and CMV-28h and their intersections. (A) Up-regulated genes and (B) down-regulated genes for each time point post-inoculation with TMV or CMV. All comparisons were made between inoculated and uninoculated (healthy) leaves at each time point. doi:10.1371/journal.pone.0045953.g007](7596)
smaller pieces at 70°C for 5 min in the fragmentation buffer (Ambion). Taking these short mRNA fragments as templates, reverse transcriptase and random hexamer-primer were used to synthesize the first-strand cDNA. This was followed by second strand cDNA synthesis using DNA polymerase I (Invitrogen) and RNaseH (Invitrogen). The resulted short cDNA fragments were purified with QiaQuick PCR extraction kit (Qiagen) followed by end repair, adding poly(A), and ligation of sequencing adaptors with Illumina’s adaptor oligo mix. The fragments were purified for the section of approximate 200 bp long using Qiaquick Gel Extraction Kit (Qiagen) and were further enriched with PCR to create the final sequencing cDNA library.

Analysis of Illumina sequencing results

The cDNA library was sequenced from both of the 5’ and 3’ ends on the Illumina HiSeq™ TM 2000 platform. The fluorescent images deconvolution and quality value calculation were performed using the Illumina data processing pipeline (version 1.6), in which 90 bp paired-end reads were obtained. Before assembly, the raw reads were filtered to obtain high-quality clean reads by removing low quality reads as well as reads with adaptor sequences or containing unknown nucleotides larger than 5%.

De novo assembly of the clean reads was performed using SOAPdenovo software [65]. Briefly, SOAPdenovo first combines the reads with certain length of overlap to form contigs, and the contigs from the same transcript as well as the distances between these contigs were further detected by using paired-end reads. Next, SOAPdenovo connects the contigs using N to represent unknown sequences between each two contigs, resulting in Scaffolds. Finally, paired-end reads are used again for gap filling of scaffolds to generate Unigenes. To determine the sequence direction, the unigenes were aligned by blastx (E-value < 10^{-5}) against the COG database [69] and the KEGG database [36], respectively.

Digital gene expression library preparation and sequencing

Tag library preparation for the distinct *C. amaranticolor* sample (healthy leaves, TMV-6h, TMV-20h, CMV-6h and CMV-20h) was performed in parallel using Illumina gene expression sample preparation kit as described in cDNA library preparation. The generated cDNA library was sequenced using Illumina HiSeq™ 2000, resulting in the raw image data, which, subsequently, was transformed by base calling into sequence data. Reads were mapped to the reference sequences using SOAPaligner/soap2 [63] with a maximum of no more than 2 nucleotides mismatches. The reads mapped to reference sequences from multiple genes were filtered, while the remaining reads were designed as unambiguous tags. For gene expression analysis, the number of expressed reads was counted and normalized using RPKM (reads per kb per million reads) [66].

A statistical analysis of the frequency of each read in the different cDNA libraries was further performed to screen the differentially expressed genes in *C. amaranticolor* upon virus infection. Statistical comparison was performed with a strict algorithm referring to the method described previously [49]. FDR was used to determine the threshold of P value in multiple test and analysis. The threshold as “FDR≤0.001” was introduced to judge the significance of gene expression difference. The identified genes with differentially expression levels were then mapped to terms in GO and KEGG database, looking for the significantly enriched KEGG terms comparing to the genome background.

Data deposition

The nucleotide sequences of raw reads from this study were submitted to NCBI Gene Expression Omnibus under the accession number GSE30451.

Supporting Information

**Figure S1** Summary of the *C. amaranticolor* transcriptomic sequences. (A) Size distribution of Illumina sequencing contigs. (B) Size distribution of distinct sequences after paired-end and gap filling. (TIF)

**Figure S2** Relationship between the number of identified genes and the sequencing amount (total read number). The figures represent DGE reads of healthy leaves (A), TMV-6h (B), TMV-20h (C), CMV-6h (D) and CMV-20h (E), respectively, showing a trend of saturation. When the sequencing amount reaches 12 millions, the number of identified genes almost ceases to increase. (TIF)

**Figure S3** Distribution of DGE reads derived from healthy leaves (A), TMV-6h (B), TMV-20h (C), CMV-6h (D) and CMV-20h (E) on reference genes in the *C. amaranticolor* transcriptome. (TIF)

**Table S1** Top BLAST hits from NCBI nr database. BLAST results against the NCBI nr database for all the clusters and singletons with a cut-off E value above 10^{-5} are shown. (RAR)

**Table S2** Putative resistance gene analogs (RGAs) identified from the foliar transcriptome of *C. amaranticolor*. (XLS)

**Table S3** Putative disease resistance signaling proteins identified from the foliar transcriptome of *C. amaranticolor*. (XLS)

**Table S4** Differentially expressed Genes between TMV-6h and healthy leaves (early-TMV-inducible genes). RPKM: reads per kb per million reads. Raw intensity: the total number of reads sequenced for each gene. FDR: false discovery
rate. The significance of gene expression difference between samples was identified by using FDR < 0.001 and the absolute value of log2Ratio ≤ 1 as the threshold. To calculate the log2Ratio and FDR, we set RPKM value of 0.001 instead of 0 for genes that do not express in one sample.

Table S5 Differentially expressed genes between TMV-28h and healthy leaves (late-TMV-inducible genes).

Table S6 Differentially expressed genes between CMV-6h and healthy leaves (early-CMV-inducible genes).

Table S7 Differentially expressed genes between CMV-20h and healthy leaves (late-CMV-inducible genes).

Table S8 Common genes between the early-TMV-inducible genes and the late-TMV-inducible genes.

Table S9 Common genes between the early-CMV-inducible genes and the late-CMV-inducible genes.

Table S10 Common genes between the early-TMV-inducible genes and the late-CMV-inducible genes.

Table S11 Common genes between the late-TMV-inducible genes and the late-CMV-inducible genes.

Table S12 Common genes among the early-TMV-inducible genes, the early-CMV-inducible genes, the late-TMV-inducible genes and the late-CMV-inducible genes.

Table S13 Genes showed ‘down-up’ or ‘up-down’ expression profiles during the TMV-induced HR.

Table S14 Genes showed ‘down-up’ or ‘up-down’ expression profiles during the CMV-induced HR.

Table S15 Genes showed ‘down-up’ or ‘up-down’ expression profiles during both TMV- and CMV-induced HR.

Acknowledgments

We wish to thank the technical support for Illumina sequencing and initial data analysis from Beijing Genome Institute at Shenzhen, P. R. China. We also thank two anonymous reviewers for their constructive and helpful comments.

Author Contributions

Conceived and designed the experiments: YZ XP ZW SJ WL. Performed the experiments: YZ CZ ZL WL. Analyzed the data: XP ZW SJ WL. Contributed reagents/materials/analysis tools: YZ XP WL. Wrote the paper: YZ WL.

References

1. Heath MC (2000) Hypersensitive responses-related death. Plant Mol Biol 44: 321–334.
2. Shirasu K, Schulze-Lefert P (2000) Regulators of cell death in disease resistance. Plant Mol Biol 44: 371–385.
3. Flor HH (1971) Current status of gene-for-gene concept. Annu Rev Phytopathol 9: 275–296.
4. Dangl JL, Jones JD (2001) Plant pathogens and integrated defence responses to infection. Nature 411: 386–393.
5. Jones JD, Dangl JL (2006) The plant immune system. Nature 444: 323–329.
6. Baker B, Zambrzycki P, Staszkiewicz B, Dinesh-Kumar SP (1997) Signaling in plant-microbe interactions. Science 276: 726–733.
7. Durrant WE, Dong X (2004) Systemic acquired resistance. Annu Rev Phytopathol 42: 153–209.
8. Spelo SH, Dong X (2012) How do plants achieve immunity? Defence without specialized immune cells. Nat Rev Immunol 12: 89–100.
9. CMI/AAB (1984) Descriptions of Plant Viruses, July Commonwealth Mycological Institute and the Association of Applied Biologists. Unwin Brothers Ltd. Old Woking Surrey.
10. Cooper B (2001) Collateral gene expression changes induced by distinct plant viruses during the hypersensitive resistance reaction in Cheno podium amaranticolor. Plant J 26: 339–349.
11. Vicedo G, Fernandez-Piqueras J, Garcia JA (1990) Isozyme profiles associated with the hypersensitive response of Chenopodium foetidum to plum pox virus infection. Physiologia Plantarum 78: 288–291.
12. Schmitz I, Rao AI (1996) Molecular studies on bromovirus capsid protein I. Characterization of cell-to-cell movement defective RNAs variants of Bromus mollis virus. Virology 226: 281–293.
13. Canto T, Palukaitis P (1999) The hypersensitive response to cucumber mosaic virus in Chenopodium amaranticolor requires virus movement outside the initially infected cell. Virology 265: 74–82.
14. Wang Z, Gerstein M, Snyder M (2009) RNA-Seq: a revolutionary tool for transcriptomics. Nat Rev Genet 10: 57–63.
15. Ward JL, Posnala L, Weber CA (2012) Strategies for transcriptome analysis in nonmodel plants. Am J Bot 99: 267–276.
16. Yang SS, Tu ZJ, Cheung P, Xu WW, Lamb RJ, et al. (2011) Using RNA-Seq for gene identification, polymorphism detection and transcript profiling in two alfalfa genotypes with divergent cell wall composition in stems. BMC Genomics 12: 199.
17. Wang W, Wang Y, Zhang Q, Qi Y, Guo D (2009) Global characterization of Artemisia annua glandular trichome transcriptome using 454 pyrosequencing. BMC Genomics 10: 465.
18. Logacheva MD, Kasianov AS, Vinogradov DV, Namgilin TH, Geldman MS, et al. (2011) Dicer sequencing and characterization of floral transcriptome in two species of balsamroot (Papaver). BMC Genomics 12: 30.
19. Wall PK, Leebens-Mack J, Chandler BA, Barakat A, Wolcott E, et al. (2009) Comparison of next generation sequencing technologies for transcriptome characterization. BMC Genomics 10: 347.
20. Novaeas E, Drost DR, Farmerie WG, Pappas GJ Jr, Grattapaglia D, et al. (2008) High-throughput gene and SNP discovery in Eucalyptus grandis, an uncharacterized genus. BMC Genomics 9: 312.
21. Bellin D, Ferrari A, Chimento A, Kaiser O, Levenkova N, et al. (2009) Combining next-generation pyrosequencing with microarray for large scale expression analysis in non-model species. BMC Genomics 10: 355.
22. Su CL, Chao YT, Alex Chang YC, Chen WC, Chen CY, et al. (2011) De novo assembly of expressed transcripts and global analysis of the Phalaenopsis aphrodite transcriptome. Plant Cell Physiol 52: 1301–1314.

23. Collina LJ, Bigg PJ, Voelckel C, Joly S (2008) An approach to transcriptome analysis of non-model organisms using short-read sequences. Genome Informatics 21: 3–14.

24. Scholthof KB, Adkins S, Czyczek H, Palukaitis P, Jacob E, et al. (2011) Top 10 plant viruses in molecular plant pathology. Mol Plant Pathol 2011: 938–954.

25. Li R, Zhu H, Ruan J, Qian W, Fang X, et al. (2010) De novo assembly of human genomes with massively parallel short read sequencing. Genome Res 20: 265–272.

26. Pereda G, Huang X, Liang F, Antonescu V, Sultana R, et al. (2003) TIGR Gene Indices clustering tool (TGICL): a software system for fast clustering of large EST datasets. Bioinformatics 19: 651–652.

27. Mittapalli O, Bai X, Mamidala P, Rajaparap S, Bonello P, et al. (2010) Tissue specific transcriptomics of the exotic invasive insect pest emerald ash borer. PLoS ONE 5: e13760.

28. Liang H, Carlsson JE, Leebens-Mack JH, Wall PK, Mueller LA, et al. (2008) An ultrafast tool for short read alignment. Bioinformatics 25: 1966–1967.

29. Bai X, Mamidala P, Rajarapu SP, Jones SC, Mittapalli O (2011) Transcriptomics of the Bed Bug (Cimex lectularius). PLoS ONE 6: e16336.

30. Zhang WX, Luai JG, Li JG, Yao YH, Zhang CX, et al. (2010) De novo characterization of a whitefly transcriptome and analysis of its gene expression during development. BMC Genomics 11: 400.

31. The Arabidopsis Genome Initiative (2000) Analysis of the genome sequence of the flowering plant Arabidopsis thaliana. Nature 408: 796–815.

32. Hao DC, Ge G, Xiao P, Zhang Y, Yang L (2011) The first insight into the tissue specific tissue transcriptome via Illumina second generation sequencing. PLoS ONE 6: e21220.

33. Shi CY, Yang H, Wei CL, Yu O, Zhang ZZ, et al. (2011) Deep sequencing of the Camellia sinensis transcriptome revealed candidate genes for major metabolic pathways of tea-specific compounds. BMC Genomics 12:131.

34. Sun Q, Zhou GF, Cai YF, Fan YH, Zhu XY, et al. (2012) Transcriptome analysis of stem development in the tumorous stem mustard Brassica juncea var. tumida Tsen et Lee by RNA sequencing. BMC Plant Biol 12: 53.

35. Hou WN, Duan CG, Fang RX, Zhou XY, Guo HS (2011) Satellite RNA analysis of non-model organisms using short-read sequences. Genome Informatics 216: 343.

36. Kanchisa M, Arakii M, Goto S, Hitomi M, Hiraoka M, et al. (2008) KEGG for functional and comparative genomics in Liriodendron. Tree Genet Genom 4: 419–433.

37. Jupe F, Pritchard I, Ehrenberg GJ, MacKenzie K, Cock PJF, et al. (2012) Identification and localisation of the NB-LRR gene family within the potato genome. BMC Genomics 13: 75.

38. Yang S, Zhang X, Yue JX, Tian D, Chen JQ (2008) Recent duplications in Arabidopsis thaliana reveal a small NBS resistance gene family. Mol Genet Genomics 280: 187–198.