Genetic diversity and genome recombination in *Yam mild mosaic virus* isolates

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**Abstract**

*Yam mild mosaic virus* (YMMV) is prevalent in yams (*Dioscorea* spp.) worldwide. To gain an insight into the genetic diversity and molecular evolution of YMMV, 89 isolates from West Africa, Asia, South Pacific, and America were analyzed phylogenetically by using sequence between the 3′-terminal of the coding region of the coat protein and the 5′-terminal of non-coding region at the 3′ end of the YMMV genome. The results revealed that there was a significant genetic diversity among isolates and a clear correlation between the coat protein gene sequence and the geographical origin of YMMV isolates. Of particular, YMMV isolates from North China and South China fell into two different groups. Furthermore, full genome comparison identified four chimeric genome patterns and six putative recombination signals representing four recombination events were detected among 12 genomes of independent isolates from China and Brazil, suggesting a high frequency of genome recombination event.

**Keywords:** *Yam mild mosaic virus* (YMMV), Genetic diversity, Genome recombination

**Background**

*Yam mild mosaic virus* (YMMV), a distinct member of the genus *Potyvirus*, is a major viral agent in yams in Africa, Asia, Caribbean and South America (Mumford and Seal 1997; Fuji et al. 1999; Odu et al. 1999; Bousalem and Dallot 2000; Dallot et al. 2001; Eni et al. 2008; Zou et al. 2011). This virus has flexuous, filamentous particles of approximately 750 nm in length and is transmitted by aphids or mechanical inoculation (Odu et al. 1999). The virus causes mild symptoms of mottle and mosaic on water yam (*Dioscorea alata*), Guinea yams (*D. cayenensis* – *D. rotundata* complex) and Indian yam (*D. trifida*), but no symptoms on white yam (*D. rotundata*) (Mumford and Seal 1997). Its natural host range is restricted to several *Dioscorea* spp., but the virus is also transmitted easily to cowpea (*Vigna unguiculata*) (Odu et al. 1999). It was reported that YMMV isolates from Caribbean island Martinique and French Guyana were divergent in their coat protein (CP) gene sequences (Bousalem et al. 2003). However, little was known about the diversity of the YMMV at whole genome level, simply because only three whole genome sequences of YMMV were available at the time (Simon-Loriere and Holmes 2011; Filho et al. 2013).

The Qinling Mountains-Huaihe River Line, with its west end at E104°15′/N32°18′ and east end at E120°21′/N34°05′ separates China as South (south to the line) and North (north to the line) mainly by its climatic impact. Yams are mainly grown in the provinces of Guangxi and Jiangxi in South China, and Henan, Shandong and Jiangsu in North China. YMMV infection of yam plants in China was first reported in 2010 (Zou et al. 2011). To gain an insight into the composition of genetic population of YMMV, we conducted a nation-wide survey of YMMV in China. Here, we report the identification,
genome sequencing, phylogenetic analysis, and genome recombination analysis of YMMV isolates from China and those reported elsewhere. Our results showed that YMMV genome sequences were diversified among isolates with geographical characteristics, and extensive genome recombination events had taken place in the population of the virus.

Results

YMMV infection was prevalent in yams in the major yam-producing regions in China

During the survey, leaves and tubers of yam plants showing viral disease-like symptoms, i.e., mosaic, chlorosis, vein banding, flecking, leaf puckering, stunting and distortion were collected, with leaves stored at −80 °C for viral identification and tubers grown in the glasshouse for keeping the materials. IC-RT-PCR and sequencing results revealed that 112 out of 365 samples collected from five yam producing regions (Guangxi, Jiangxi, Henan, Jiangsu and Shandong) in China were YMMV-positive, among which 75 were from 132 D. alata, 11 from 96 D. japonica and 26 from 137 D. opposit. The incidence ranged from 11% to 40% in general, varying among yam species with the highest rate for D. alata and lowest for D. japonica (Additional file 1: Table S1). Most YMMV-positive samples (90.5%) were co-infected with other viruses (our unpublished results) and those (9.5%) solely infected with YMMV showed constantly a mild mosaic or mottle or symptomless on leaves (Fig. 1), regardless of species/cultivars and years of sampling. The rest of YMMV-negative samples were infected either with other known viruses, uncharacterized viruses, or unknown agents.

YMMV isolates showed a clear geographical character

To establish evolutionary relationship among the YMMV isolates, sequences from the region (nucleotides 259–262) in the 3′-terminal region of CP to the 5′-terminal region of 3′-nocoding region (NCR) of 26 YMMV isolates obtained in this study and 37 isolates previously reported, and 26 unpublished YMMV isolates (Table 1) were used to construct an unrooted phylogenetic tree using Maximum Likelihood algorithm. Among 14 groups clustered in the phylogenetic tree (Fig. 2), YMMV isolates from China fall into two groups: Group VII that contains isolates from North China, and Group X that contains isolates from South China, both distinguishing themselves from isolates from West Africa (Group II, Group III, Group XI, Group XII, Group XIII, and Group XIV), America (Group IV and Group VI), and other parts of Asia (Group V and Group VIII).

YMMV population was diversified at whole genome level

The genome sequences of isolates representing Group X and Group VII were assembled from the high throughput sequencing (HTS) data of the infected yam leaf samples and validated by RT-PCR and RACE. The whole genomes of the 12 YMMV isolates were from 9521 to 9538 nucleotides (nts) in length excluding the poly (A) tail (GenBank Accession No. KC407674, KC473517, KJ125472–125479, JX470965, KX156847). Eleven of the 12 isolates were the same as the reported Brazilian isolate in terms of processed protein sizes, but with slightly different length in untranslated regions (UTRs) and isolate CN20 differed from the rest of the isolates in that its P1 protein was composed of 321 aa, instead of 320 aa (Additional file 2: Table S2). While motifs of FRNK in the HC-Pro, GDD in the NiLb, and DAG in the CP were conserved in all isolates, residues flanking the motif of the HC-Pro for WX1, WX3, and XZ1 (Group VII) differed from other isolates, and more diversified patterns flanking the CP were seen (Fig. 3).

Phylogenetic tree based on the whole genome sequences of YMMV isolates further showed that isolates from South China and North China grouped separately, with the North China group closer to the Brazilian and the Korean isolates (Fig. 4). By taking advantage of whole genome sequences of multiple isolates in Group...
| Isolate | Year of collection | Geographic location | Host | GenBank accession no. | Reference |
|---------|------------------|---------------------|------|-----------------------|-----------|
| FX1     | 2012             | Fengxian, Jiangsu   | D. opposita | KJ125474             | This work |
| FX2     | 2012             | Fengxian, Jiangsu   | D. opposita | KJ789113             | This work |
| Luzhai  | 2010             | Luzhai, Guangxi     | D. opposita | JF357963             | This work |
| LZ1     | 2010             | Luzhai, Guangxi     | D. alata   | KJ789114             | This work |
| Nanning | 2010             | Nanning, Guangxi    | D. opposita | JF357962             | This work |
| NC1     | 2011             | Nanxian, Jiangxi    | D. alata   | KJ125476             | This work |
| NC2     | 2011             | Nanxian, Jiangxi    | D. alata   | KJ125477             | This work |
| NC3     | 2011             | Nanxian, Jiangxi    | D. alata   | KJ125473             | This work |
| NN1     | 2011             | Nanning, Guangxi    | D. alata   | KJ125472             | This work |
| NN2     | 2010             | Nanning, Guangxi    | D. japonica| KJ789115             | This work |
| NN3     | 2011             | Nanning, Guangxi    | D. alata   | KJ789116             | This work |
| NN4     | 2011             | Nanning, Guangxi    | D. alata   | KJ789117             | This work |
| NN5     | 2011             | Nanning, Guangxi    | D. alata   | KJ789118             | This work |
| SG1     | 2011             | Shouguang, Shandong  | D. opposita| KJ789119             | This work |
| SG2     | 2011             | Shouguang, Shandong  | D. opposita| KJ789120             | This work |
| WM1     | 2010             | Wuming, Guangxi     | D. japonica| KJ789121             | This work |
| WX1     | 2011             | Wenxian, Henan      | D. opposita| KJ125475             | This work |
| WX2     | 2011             | Wenxian, Henan      | D. alata   | KJ789122             | This work |
| WX3     | 2011             | Wenxian, Henan      | D. alata   | KJ125478             | This work |
| WX4     | 2011             | Wenxian, Henan      | D. opposita| KJ789123             | This work |
| WX5     | 2011             | Wenxian, Henan      | D. opposita| KJ789124             | This work |
| WX6     | 2011             | Wenxian, Henan      | D. opposita| KJ789125             | This work |
| WX7     | 2011             | Wenxian, Henan      | D. opposita| KJ789126             | This work |
| WX8     | 2011             | Wenxian, Henan      | D. opposita| KJ789127             | This work |
| XZ1     | 2012             | Xuzhou, Jiangsu     | D. alata   | KJ125478             | This work |
| XZ2     | 2012             | Xuzhou, Jiangsu     | D. alata   | KJ789128             | This work |
| Andong  | 2014             | South Korea         | D. polystachya| KK156847             | Unpublished\a |
| Brazil  | 2009             | Brazil              | D. alata   | JX470965             | (Filho et al. 2013) |
| Br2     | 2000             | Brazil              | D. alata   | AF548498             | (Bousalem et al. 2003) |
| CN1     | 2011             | Yunnan, China       | D. opposita| KC407674             | (Wang et al. 2015) |
| CN20    | 2010             | Yunnan, China       | D. polystachya| KC473517             | (Wang et al. 2015) |
| Col 1   | 1999             | Colombia            | D. alata   | AF548491             | (Bousalem et al. 2003) |
| Col 2   | 1999             | Colombia            | D. alata   | AF548492             | (Bousalem et al. 2003) |
| CR 1    | 1999             | Costa Rica          | D. trifida | AF548499             | (Bousalem et al. 2003) |
| DaBen 1 | –                | Benin               | D. alata   | AJ305471             | Unpublished\a |
| DaCDI 19| –                | Ivory Coast         | D. alata   | AJ305470             | Unpublished\a |
| DaCDI 37| –                | Ivory Coast         | D. alata   | AJ305469             | Unpublished\a |
| DaCmn 42| –                | Cameroon            | D. alata   | AJ305468             | Unpublished\a |
| DaGui 1 | –                | Guinea              | D. alata   | AJ305467             | Unpublished\a |
| DaNig 2 | –                | Nigeria             | D. alata   | AJ305466             | Unpublished\a |
| DaNig 316| –                | Nigeria             | D. alata   | AJ305465             | Unpublished\a |
| DaSLK 1 | –                | Sri-Lanka           | D. alata   | AJ305464             | Unpublished\a |
| DaSLK 2 | –                | Sri-Lanka           | D. alata   | AJ305463             | Unpublished\a |
| DaTog 2 | –                | Togo                | D. alata   | AJ305462             | Unpublished\a |
| Isolate  | Year of collection | Geographic location | Host | GenBank accession no. | Reference |
|---------|-------------------|---------------------|------|-----------------------|-----------|
| DaTog 3 | –                 | Togo                | D. alata | AJ305461           | unpublished	a |
| DaUga 1 | –                 | Ghana               | D. alata | AJ305460           | unpublished	a |
| DaUga 3 | –                 | Ghana               | D. alata | AJ305459           | unpublished	a |
| DaVan 1 | –                 | Vanuatu             | D. alata | AJ305458           | unpublished	a |
| DeSLK 1 | –                 | Sri-Lanka           | D. esculenta | AJ305457      | unpublished	a |
| DrCmn 11| –                 | Cameron             | D. cayenensis-rotundata | AJ305456     | unpublished	a |
| DrGui 1 | –                 | Guinea              | D. cayenensis-rotundata | AJ305455     | unpublished	a |
| G1      | –                 | India               | D. alata | KJ463416           | unpublished	a |
| Guad 1  | 1998              | Guadeloupe          | D. alata | AF548500           | (Bousalem et al. 2003) |
| Guad 2  | 1998              | Guadeloupe          | D. alata | AF548501           | (Bousalem et al. 2003) |
| Guad 3  | 1998              | Guadeloupe          | D. alata | AF548502           | (Bousalem et al. 2003) |
| Guad 4  | 1998              | Guadeloupe          | D. trifida | AF548503      | (Bousalem et al. 2003) |
| Guad 5  | 1998              | Guadeloupe          | D. trifida | AF548504      | (Bousalem et al. 2003) |
| Guad 6  | 1998              | Guadeloupe          | D. esculenta | AF548505   | (Bousalem et al. 2003) |
| Guy 1   | 1998              | French Guyana       | D. trifida | AF548506      | (Bousalem et al. 2003) |
| Guy 2   | 1998              | French Guyana       | D. trifida | AF548507      | (Bousalem et al. 2003) |
| Guy 3   | 1998              | French Guyana       | D. trifida | AF548508      | (Bousalem et al. 2003) |
| Guy 4   | 1998              | French Guyana       | D. trifida | AF548509      | (Bousalem et al. 2003) |
| Guy 5   | 1998              | French Guyana       | D. trifida | AF548510      | (Bousalem et al. 2003) |
| Guy 6   | 1998              | French Guyana       | D. trifida | AF548511      | (Bousalem et al. 2003) |
| Guy 7   | 1998              | French Guyana       | D. trifida | AF548512      | (Bousalem et al. 2003) |
| Guy 8   | 1998              | French Guyana       | D. trifida | AF548513      | (Bousalem et al. 2003) |
| JPN 1   | 1997              | Japan               | D. alata | AF548519      | (Bousalem et al. 2003) |
| JPN 2   | 1998              | Japan               | D. alata | AF548520      | (Bousalem et al. 2003) |
| JPN 3   | 1998              | Japan               | D. alata | AF548521      | (Bousalem et al. 2003) |
| Mart 1  | 1998              | Martinique          | D. cayenensis-rotundata | AF548493    | (Bousalem et al. 2003) |
| Mart 2  | 1998              | Martinique          | D. alata | AF548494      | (Bousalem et al. 2003) |
| Mart 3  | 1998              | Martinique          | D. alata | AF548495      | (Bousalem et al. 2003) |
| Mart 4  | 1998              | Martinique          | D. alata | AF548496      | (Bousalem et al. 2003) |
| Mart 5  | 1998              | Martinique          | D. cayenensis-rotundata | AF548497    | (Bousalem et al. 2003) |
| Mart 6  | 1998              | Martinique          | D. alata | AF548528      | (Bousalem et al. 2003) |
| PNG 1   | 1993              | Papua New Guinea    | D. alata | AF548523      | (Bousalem et al. 2003) |
| PNG 2   | 1993              | Papua New Guinea    | D. alata | AF548524      | (Bousalem et al. 2003) |
| PNG 4   | 1994              | Papua New Guinea    | D. alata | AF548526      | (Bousalem et al. 2003) |
| Togo 1  | 1999              | Togo                | D. alata | AF548514      | (Bousalem et al. 2003) |
| Togo 2  | 1999              | Togo                | D. dumetorum | AF548527    | (Bousalem et al. 2003) |
| India 3-Da | 2008              | India               | D. alata | FJ390024      | (Bousalem et al. 2003) |
| India 4-Da | 2009              | India               | D. alata | HQ231894      | (Bousalem et al. 2003) |
| MS_StoDgo | 2014              | Cuba                | D. esculenta | MK780009    | unpublished	a |
| M4_StoDgo | 2014              | Cuba                | D. rotundata subsp. rotundata | MK780008  | unpublished	a |
| M3_StoDgo | 2014              | Cuba                | D. alata | MK780007      | unpublished	a |
| 10_StoDgo | 2011              | Cuba                | D. esculenta | MK780006   | unpublished	a |
| 9_StoDgo | 2011              | Cuba                | D. esculenta | MK780005   | unpublished	a |
| 4_StoDgo | 2011              | Cuba                | D. alata | MK780004      | unpublished	a |
VII and Group X, intra-group identity percentage of functional gene/protein and UTR at nucleotide and amino acid levels were calculated and compared with the Brazilian strain, the representative of Group IV (Additional file 3: Table S3 and Additional file 4: Table S4). As summarized in Table 2, at the whole genome level, identities were 87.8–99.7% for isolates within Group VII and 86.1–97.6% for isolates within Group X. However, identity between Group VII and Group X was only 84.9–87.5%. Even lower levels of identity were found between Group X and Group VII (82.8–83.6%) or between Group VII and Group IV (83.9–84.4%). It was noticed that 5′-UTRs were the most divergent, 65.1–96.2% within groups and 58.0–82.0% between groups. As to the Brazilian strain (group IV), it shares up to 84.4% identity to either Group X or Group VII, lower than that between Group X and Group VII. Less variation was found at amino acid level for the viral proteins of the isolates, both within and among groups of isolates, with the highest identity of 93.7–99.0% for isolates from Group X and 94.6–99.9% for isolates from Group VII at polyprotein level. Among the mature proteins, 6 K1 was the most conserved with an identity of 96.1–100% for isolates in Group X and 98.0–100% for isolates in Group VII, while P1 was the most variable, 77.8–99.4% for isolates in Group VII. An identity of 71.2–79.1% for P1 between Group X and Group VII was the lowest among all processed proteins. Brazilian strain shares an identity up to 91.3–92.0% to Group X or Group VII at polyprotein level, lower than the 92.6–93.9% between Group X and Group VII.

**Genome recombination events in YMMV**

Among the 12 genomes of YMMV isolates (ten from China, one from Brazil and one from Korea), three isolates (NN1, FX1 and NC1) seemed to have a "pure" genome, i.e. without any apparent chimeric genome fragment from the known isolates (Fig. 5). Six clear recombination signals representing four recombination events involving four YMMV genomes were detected with P-value < 1.0 × 10⁻⁶ (Table 3). CP was the hottest region for recombination, involving five isolates, and P3 and CI each occurred once in one of the isolates. As shown in Fig. 5, one recombination event was found in NC1, CN1, NC2 and CN20, respectively. And there were two recombination regions in CN1 and CN20, which were located at both ends of the genome, respectively. There was also a recombination event in NC1 and NC2, but only one recombination region existed in each recombination event. The region of recombination event I was located at the 5′ end of NC1 genome, while the region of recombination event III was located at the middle and rear of NC2 genome.

**Discussion**

**Geographical distribution of YMMV isolates and movement of yam germplasm**

YMMV is a world-wide dispersed potyvirus in yams, but not much is known about its natural history and evolutionary relation. A powerful tool for tracing the origin and evolution of a virus is phylogenetic and phylodynamic analysis of viral sequences (Ren et al. 2013). Data obtained using such methods have contributed to the surveillance of viral spread and drug resistance as well as the identification of strains as vaccine candidates (Lam et al. 2010; Norström et al. 2012).

As the 3′-terminal region of CP and 5′-terminal region of 3′-NCR of YMMV has been used for genotype classification in YMMV to investigate the diversity of YMMV (Bousalem et al. 2003), these sequences were used in this study to establish evolutionary relationship of YMMV isolates. As shown in Fig. 2, the 14 distinct groups clustered were strongly associated with geographical distribution, among which Group X was from South China, Group VII from North China. Most of the isolates from other Asia regions were closer to Chinese isolates than to those from the Central America and West Africa, indicating that the differentiation of YMMV was resulted from geographic isolation. Of particular note is that Group VII isolates from North China represent the most distant relation to the Group X from South China, and these two groups are different from those outside China, suggesting that YMMV isolates from China may have undergone further differentiation. The phylogenetic tree constructed from this study was partially in accordance with the assumption that Asian-Pacific origin of YMMV was likely from *D. alata* species (Bousalem et al. 2003). But our data also showed that YMMV isolates from China and India share a common ancient ancestor, different from those YMMV isolates from Central America and West Africa which share the same common ancient ancestor. Distinct geographical distribution of YMMV groups also suggests that germplasm exchange of yams has been infrequent between South China and North China, as well as among countries world-wide.
Phylogenetic relation and viral recombination at whole-genome scale among YMMV isolates

To address the genetic relation among different genotypes further, we employed 12 YMMV complete genome sequences to analyze the phylogenetic relationship. The predicted sizes of the coding regions were identical among YMMV isolates except CN20 (Additional file 2: Table S2). The extent of genetic diversity, reflected in percentage identity, varies within and among proteins, in the order NIa-VPg > HC-Pro > Nla-Pro > 6 K1 > CI > Nlb > CP > 6 K2 > PIPO > P3 > P1 (Table 2). The phylogenetic tree constructed based on the whole genomes (Fig. 3) matches basically well with the tree constructed based on the partial CP core region and the 3′-NCR sequences (Fig. 2), with isolates from South China (NN1, NC1, NC2, NC3, FX1 and CN1) in Group X, isolates from North China (CN20, XZ1, WX1 and WX3) in group VII, and Brazilian isolate in group IV. The only exception is the isolate CN20, the placing of which is inconsistent in the two trees, suggesting a mixed genome. YMMV genome recombination was first observed by comparison of the partial 3′-terminal genome sequences of the isolates collected from different geographical locations (Bousalem et al. 2003). Indeed, in the current study, a detailed examination revealed that 4 out of the 12 YMMV isolates may have gone through genome recombination events (Fig. 5). While recombination events were spotted in the genome regions encoding P3, CI and CP, the most frequent recombination events were found in the CP-encoding 3′-end region. Viral recombination is a powerful contributor to genetic variation, adaptation to new hosts, escape from the host immune response, and emergence of newly infectious agents (Becher et al. 2001; Simon-Loriere and Holmes 2011). Although recombinants did not seem to have a significant impact on symptoms (Fig. 1), recombination in CP between YMMV isolates may provide a selective advantage for virus dissemination by a vector to adapt to the local environment, although the molecular mechanisms of this hypothesis needs to be clarified.

Conclusions

Data presented in this study demonstrated that YMMV infection was prevalent in the main yam producing areas in China, and there was a significant genetic diversity and a clear correlation between the coat protein gene sequence and the geographical origin of YMMV isolates. Four chimeric genome patterns were identified in 12 isolates, suggesting a high frequency of genome recombination event. Therefore, due cautions should be taken in the exchange of germplasms between the north and the south of China, as well as among nations in the world, to prevent possible occurrence of new virulent isolates through genome recombination between and among YMMV isolate types.

Methods

Sample collection and virus identification

During a survey of viral diseases on yam in 2010–2015, a total of 365 yam leaf samples were collected from...
Guangxi Province, Henan Province, Jiangsu Province, Jiangxi Province and Shandong Province in China, and 105 samples were identified as YMMV positive by IC-RT-PCR using polyclonal antibodies developed in our laboratory (Zou et al. 2011).

Nucleic acid extraction and analysis
Total RNA was extracted from an amount of 100 mg of yam leaf tissue using an RNAprep pure Plant Kit (Tiangen Inc., Beijing, China) and quantified by agarose gel electrophoresis and Qubit® 2.0 Fluorometer quantitation assay. Viral cDNA was synthesized with the total RNA as template using Transcriptor High Fidelity cDNA Synthesis Kit (Roche Applied Science, Mannhelm, Germany) and PCR amplification of YMMV genome fragments was performed using AmpliTaq DNA polymerase and the Expand high-fidelity PCR system (Roche Applied Science, Mannhelm, Germany) with virus-specific primers (Additional file 5: Table S5).

Deep sequencing, RACE and assembly of viral genome from total yam RNA
An amount of 5 μg of the total RNA was used for cDNA library construction using TruSeq Illumina mRNA library construction kit (Illumina Inc., San Diego, CA). Deep sequencing was performed on an Illumina Solexa GAIIx platform. The CLC Genomics Workbench V6.0.1 software was used for deep sequencing data analysis. Raw reads from the Illumina RNA-Seq were trimmed to remove low quality reads and sequencing adaptor. The clean reads were assembled into contigs using the De novo Assembly algorithm. Contigs were then mapped to

Table 2 Identity percentage at nucleotide and amino acid levels within and between YMMV isolate groups

| Nucleotide level | Isolate group | 5′-UTR Coding region 3′-UTR | Full-length |
|------------------|---------------|-----------------------------|------------|
| Within Group X   | 72.4–94.9     | 86.1–97.7                   | 86.1–97.6 |
| Within Group VII  | 65.1–96.2     | 81.7–99.3                   | 87.8–99.7 |
| Group X / Group VII | 58.0–82.0   | 80.9–90.2                   | 84.9–87.5 |
| Group X / Group IV | 70.7–76.6   | 80.3–85.9                   | 82.8–83.6 |
| Group VII / Group IV | 65.9–74.2   | 78.9–88.7                   | 83.9–84.4 |

| Amino acid level | Isolate group | P1 | HC-Pro | P3 | PIPO | 6K1 | CI | 6K2 | Nla-Vpg | Nla-Pro | Nib | CP | Polyprotein |
|------------------|---------------|----|--------|----|------|-----|----|-----|---------|---------|-----|----|-------------|
| Within Group X   | 81.2–95.6     | 83.6–98.3 | 86.6–97.0 | 96.1–100 | 95.8–99.2 | 90.4–100 | 95.2–100 | 95.4–99.6 | 94.0–99.6 | 95.1–99.2 | 93.7–99.0 |
| Within Group VII  | 77.8–99.4     | 94.3–100 | 86.6–100 | 98.0–100 | 97.0–100 | 96.2–100 | 98.4–100 | 97.5–100 | 95.9–99.8 | 94.7–100 | 94.6–99.9 |
| Group X / Group VII | 71.2–79.1   | 85.1–928 | 83.6–940 | 94.1–100 | 95.5–975 | 92.3–981 | 96.3–990 | 95.0–983 | 940–967 | 93.2–966 | 92.6–93.9 |
| Group X / Group IV | 76.6–80.3    | 82.2–87.1 | 88.1–91.0 | 96.1–100 | 93.8–949 | 88.5–90.4 | 95.2–96.9 | 94.2–95.8 | 930–946 | 91.4–925 | 91.3–920 |
| Group VII/Group IV | 73.8–77.8    | 86.8–90.2 | 85.1–88.1 | 94.1–96.1 | 94.1–94.9 | 90.4–92.3 | 95.2–96.3 | 94.6–95.4 | 938–944 | 91.0–91.0 | 91.7–91.8 |
the YMMV reference sequence (Genbank accession No. JX470965).

RACE amplification was performed using the SMARTer® RACE cDNA Amplification Kit (Clontech Laboratories, Inc.). PCRs were carried out on cDNA with Phusion® High-Fidelity PCR Master Mix with HF Buffer (Thermo Fisher Scientific, Inc.). PCR fragments were purified from agarose gels and cloned into pJET1.2/blunt Cloning Vector (Thermo Fisher Scientific, Inc.). Sequences from each isolate were confirmed by analysis of at least three overlapping independent RT-PCR products on an Applied Biosystems 3730XL DNA Sequencer. Overlapped sequences were assembled and analyzed with Vector NTI Advance™ version 10 software (Invitrogen Inc., Carlsbad, CA).

Alignment and phylogenetic analysis
Nucleotide sequences of YMMV isolates were identified by BLAST search against NCBI (http://blast.ncbi.nlm.nih.gov/Blast.cgi) and conserved domain database (CDD) were used for conserved domain identification and structure analysis. DNASTAR (DNASTAR, USA) was used to analyze nucleotide and amino acid sequence divergence. Alignment of the nucleotide or amino acid sequences was performed using the Clustal W program (Thompson et al. 1994). Phylogenetic analysis was performed with MEGA6 software using the maximum likelihood method with bootstrap of 1000 times repeat (Tamura et al. 2011), using CP/3′-UTR region (Mumford and Seal 1997) or the whole genome of the virus.

Genome recombination analysis
A collection of isolate genomes was first aligned using Clustal W program, then scanned by Recombination Detection Program version 4 (RDP4) with default settings for the different detection methods and a Bonferroni corrected \( P \) value cut-off of 0.05. (Martin et al. 2015). And the detected site with a Bonferroni corrected \( P \) values of less than 1.0 \( \times 10^{-6} \) was considered a clear recombination site, otherwise it was considered as a tentative recombination sites (Ohshima et al. 2007).

### Supplementary information

**Supplementary information**

Supplementary information accompanies this paper at https://doi.org/10.1186/s42483-020-00051-0.

**Additional file 1: Table S1.** Symptom-showing samples collected and YMMV infection rate detected by IC-RT-PCR.

**Additional file 2: Table S2.** Amino acid identity within YMMV isolate groups.

**Additional file 3: Table S3.** Nucleotide identity within YMMV isolate groups.

**Additional file 4: Table S4.** Amino acid identity within YMMV isolate groups.

**Additional file 5.** Table S5. Primers used in this work.

**Abbreviations**

CP: Coat protein; HTS: High throughput sequencing; IC-RT-PCR: Immunocapture Reverse transcription polymerase chain reaction; NCR: Nocoding region; RT-PCR: Reverse transcription polymerase chain reaction; YMMV: Yam mild mosaic virus

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**Authors’ contributions**

BSC conceived the study and revised the manuscript. CWZ and LZ performed the experiments. CWZ, JRM and ZTY drafted the manuscript. JRM, ZQW and BHW contributed reagents/materials/analysis tools. All authors read and approved the final manuscript.

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**Availability of data and materials**

Not applicable.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

### Table 3 Recombination crossover sites in YMMV genomes detected by RDP4

| Event No. | Found in | Recombinant | Major | Minor | Detecting methods | \( P \) values |
|-----------|----------|-------------|-------|-------|-------------------|--------------|
| I         | 1        | NC1         | NN1   | NC2   | R G B M C         | 7.741 \( \times 10^{-13} \), 8.558 \( \times 10^{-14} \), 6.872 \( \times 10^{-13} \), 2.670 \( \times 10^{-3} \), 2.442 \( \times 10^{-3} \) |
| II        | 1        | CN1         | WX1   | NC3   | R B C S          | 1.066 \( \times 10^{-6} \), 6.842 \( \times 10^{-10} \), 1.148 \( \times 10^{-10} \), 3.226 \( \times 10^{-6} \) |
| III       | 1        | NC2         | FX1   | WX3   | M C S 3      | 3.367 \( \times 10^{-11} \), 4.606 \( \times 10^{-13} \), 2.116 \( \times 10^{-6} \), 5.180 \( \times 10^{-7} \) |
| IV        | 1        | CN20        | WX1   | NC3   | R B M C         | 4.927 \( \times 10^{-3} \), 1.287 \( \times 10^{-12} \), 6.484 \( \times 10^{-6} \), 5.180 \( \times 10^{-7} \) |

*Recombinant isolates identified by the recombinating detection programs, R (RDP), G (Geneconv), B (Bootscan), M (Maxchi), C (Chimaera), S (Siscan) and 3 (3seq) programs in RDP4 v.4.56. The analysis was carried out with default settings for the different detection methods and a Bonferroni corrected \( P \) values cut-off of 0.05. And the detected site with a Bonferroni corrected \( P \) values of less than 1 \( \times 10^{-6} \) was considered a clear recombination site whilst a detected site with \( P \) values of grater than 1 \( \times 10^{-6} \) as a tentative recombination sites. The \( P \) values obtained by different detection methods are shown in the last column of the table, where the bold font indicates that the \( P \) value of less than 1 \( \times 10^{-6} \).
Competing interests
The authors declare that they have no competing interests.

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