

1 **Genetic Structure and Diversity of Banana Bunchy Top Virus (BBTV) in the**
2 **Philippines**

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17 structure

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26 **Abstract**

27 Banana bunchy top virus (BBTV) is an important disease of banana in the Philippines and in
28 other banana-producing countries. This study was conducted to investigate the genetic structure
29 and diversity of Philippine BBTV isolates which remain unexplored in the country. BBTV-
30 infected plant tissues were sampled from banana-growing provinces (i.e., Cagayan, Isabela,
31 Quirino, Batangas, Laguna, Rizal, Quezon, Palawan, Cebu, Leyte, and Davao del Sur) and the
32 partial DNA-R gene of BBTV was sequenced. Analysis of all local BBTV isolates showed a
33 nucleotide diversity (π) of 0.00721, average number of nucleotide differences (k) of 5.51984,
34 and haplotype diversity (hd) of 0.971. Neutrality tests using Fu's F_s and Tajima's D showed
35 significant and highly negative values which suggest an excess number of rare alleles due to
36 recent population expansion or from genetic hitchhiking. Haplotype network and phylogenetic
37 analyses revealed that the local BBTV isolates were closely related to Southeast Asian (SEA)
38 group and exhibited a monophyletic clade with distinct haplotype grouping from other SEA
39 sequences. However, some Indonesian and Indian reference sequences were also clustered
40 within the Philippine BBTV group suggesting sequence homology. Results also showed that
41 the local BBTV isolates may be categorized into three major haplotype groups (HA, HB, and
42 HC) but only the HC group remained distinct upon comparison with other Philippine and SEA
43 reference sequences. BBTV isolates from Quezon were the most diverse while isolates from
44 Palawan displayed low genetic diversity indices and belonged only in the HC group. The
45 assessment of the degree of variability among Philippine BBTV isolates will provide a
46 reference towards the development of high-throughput BBTV detection systems as well as
47 enable to devise plant breeding strategies to manage the current BBTV spread and variations.

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51 **Introduction**

52 Banana bunchy top disease (BBTD) is one of the major threats in banana producing
53 countries such as India, China, Taiwan, Indonesia and the Philippines (Debbarma et al., 2019;
54 Qazi, 2016). In the Philippines, BBTD is present in almost all banana-growing areas. The
55 disease affects both smallholders and large banana plantation growers. Spread of the disease
56 can be minimized through elimination of infected materials and use of virus-free tissue culture-
57 derived planting materials. These management strategies are usually employed by big
58 commercial growers instead of the smallholders. In the 1990s, smallholder production of the
59 popular dessert cv. Lakatan in the northern Philippines was virtually eliminated by severe
60 BBTD infection (Molina et al., 2009). Initial symptoms of the disease involve the appearance
61 of dark green streaks in the veins which becomes apparent with a combination of marginal
62 chlorosis or yellowing of the leaves. Dashes and dots creating a “Morse code” pattern in the
63 leaves and petiole may also be observed. This will lead into small emerging bunchy leaves
64 forming a rosette pattern. In serious cases, disease plants were observed to be severely stunted
65 and unable to bear fruits (Dale 1987; Hooks et al., 2008).

66 The disease is known to be caused by Banana bunchy top virus (BBTV), a single-
67 stranded DNA virus belonging to the Babuvirus genus, which can bring catastrophic loss to a
68 banana plantation. The systemic virus can easily be transmitted by an aphid (*Pentalonia*
69 *nigronevosa*; Magee, 1927), a vector with a wide host range including *Musa textilis* and other
70 members of the family Musaceae. The disease transmission is of the persistent, circulative,
71 non-propagative type (Anhalt and Almeida, 2008), with efficiency ranging from 46-67%
72 (Magee, 1927; Wu and Su., 1990; Hu et al., 1996). The virus is made up of 6 genetic
73 components, namely, DNA-C coding for the cell cycle link protein, DNA-S coding for the
74 capsid protein, DNA-M coding for the movement protein, DNA-N coding for the nuclear
75 shuttle protein, DNA-U3 coding for potential protein with unknown function, and DNA-R

76 coding for the replication initiation protein (Amin et al., 2008, Kumar et al., 2017;
77 Wickramaarachchi et al., 2016).

78 The phylogenetic relationship among BBTV DNA-R sequences revealed that the virus
79 can be categorized into two different lineages based on geographical distribution: the South
80 Pacific/ Pacific-Indian Oceans (PIO) group and the Asian/ Southeast Asian (SEA) group (Yu
81 et al., 2012; Karan et al., 1994). BBTV isolates obtained from Australia, Egypt, Hawaii, India,
82 Myanmar, Pakistan, Sri Lanka and Tonga belong under the PIO group, while the isolates
83 collected from China, Indonesia, Japan, Philippines, Taiwan and Vietnam are considered
84 members of the SEA group (Yu et al., 2012).

85 In-depth studies on the diversity of BBTV from various countries were already
86 reported, such as in Democratic Republic of Congo (Mukwa et al., 2016), Pakistan (Amin et
87 al., 2007), sub-Saharan Africa (Kumar et al., 2011), Indonesia (Chiaki et al., 2015), Japan
88 (Furuya et al., 2005) and India (Banerjee et al., 2014). The information on the degree of genetic
89 diversity and distribution of BBTV in these countries provided useful and fundamental
90 information to control BBTD through various pest management approaches. Unfortunately, the
91 reported BBTV sequences from the Philippines have only been made available through foreign
92 efforts, which generally aims to provide insights into BBTV diversity and population structure
93 at the global level. Local and intensive reports regarding the diversity and spread of this
94 important banana disease will be vital towards devising a specific management system in the
95 country. Thus, a detailed assessment of the current Philippine BBTV diversity and genetic
96 structure using the DNA-R region will be discussed in this paper.

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101 **Materials and Methods**

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103 **Survey and collection**

104 A survey was done in banana growing areas in the Philippines (Figure 1). Symptoms depicting
105 classic BBTV infection such as leaf chlorosis, dash-dot pattern, rosetting, and stunting were
106 observed and recorded. Leaf samples from representative symptomatic and asymptomatic
107 samples were collected and processed for DNA extraction.

108

109 **DNA extraction**

110 Samples were processed using a CTAB DNA extraction protocol adapted from Doyle and
111 Doyle (1990) with a few modifications. Approximately, 300 mg of fresh leaf tissue was ground
112 into fine powder with liquid nitrogen using a sterilized mortar and pestle. Ground tissue was
113 then transferred into a sterilized 1.5 ml microcentrifuge tube. Exactly 700 µl of extraction
114 buffer with 2% PVP was added to each sample and then incubated at 65 °C for an hour. One
115 volume CH₃Cl-isoamyl alcohol (24:1) was added followed by centrifugation at 10,000 rpm for
116 10 minutes at 23 °C. Aqueous phase was transferred into a new sterile 1.5 ml microcentrifuge
117 tube. DNA was precipitated by adding 0.8 volume of cold isopropanol and incubated at -20 °C
118 for 30 minutes, followed by centrifugation at 10,000 rpm for 15 minutes. The DNA pellet was
119 washed using 1 mL of Wash 1 (0.2 M sodium acetate, 76% ethanol; filter sterilized) for 10
120 minutes followed by 1 ml of Wash 2 (10 mM ammonium acetate, 76% ethanol; filter sterilized)
121 for 5 minutes, and the pellet was air-dried for 30 minutes. The DNA pellet was resuspended in
122 Tris-EDTA buffer (10 mM Tris-HCl, 1 mM disodium EDTA, pH 8.0) and purified by
123 incubation with 0.1 mg/ml RNase at 37 °C for 1 hour, and centrifugation at 10,000 rpm for 5
124 minutes. DNA was collected into individual sterile 1.5 ml microcentrifuge tubes and stored at
125 -20 °C.

126 **PCR detection**

127 The presence of BBTV was confirmed by performing PCR detection. Each 15 µL reaction
128 mixture is consisting of 1X PCR buffer (10 mM Tris pH 9.1 at 20 °C, 50 mM KCl, 0.01%
129 Triton™ X-100; Vivantis Technologies, Malaysia), 1.76 mM MgCl₂, 0.2mM dNTPs, 2 µM of
130 BBT1 (5'-CTC GTC ATG TGC AAG GTT ATG TCG-3') and BBT2 (5'-GAA GTT CTC
131 CAG CTA TTC ATC GCC-3') primers (Thompson and Dietzgen, 1995; Harding et al., 1993;
132 Integrated DNA Technologies Pte. Ltd., Singapore), 1U of Taq Polymerase (Vivantis
133 Technologies, Malaysia), and 20 ng DNA. The PCR mixture was then run in a T100 thermal
134 cyclor (BioRad, USA) with initial denaturation at 94 °C for 10 minutes, followed by 30 cycles
135 of 94 °C for 1 minute, 53 C for 1 minute, 72 °C for 2 minutes and a final extension of 72 °C
136 for 10 minutes. PCR products were viewed with electrophoresis using 1% agarose gels in 1X
137 TBE buffer at 100 V for 40 min and visualized using 0.5 ug/ml ethidium bromide staining and
138 UV illumination using the Enduro GDS Touch Imaging System (Labnet International, Inc,
139 Edison, New Jersey, USA).

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141 **Outsourced sequencing**

142 Confirmed BBTV isolates were further processed for the partial sequencing of the DNA-R
143 region of the virus (Table 1). PCR amplification was done using previously established primers
144 for assessing genetic similarity (Islam et al., 2010): BBTVREP-F (5'- ATG GCG CGA TAT
145 GTG GTA TGC -3') and BBTVREP-R (5'-TCA GCA AGA AAC CAA CTT TAT TCG - 3').
146 The DNA-R primer was optimized using the same PCR conditions as BBT1 and BBT2 primers.
147 PCR products were then sent for outsourced capillary sequencing (Apical Scientific, Malaysia).

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151 **Sequence analysis**

152 Raw paired sequences (forward and reverse) were quality trimmed and analyzed using
153 Geneious Prime® (version 2019.0.4). Trimmed and assembled sequences were then aligned
154 using ClustalW (Thompson et al., 2003) at default settings. Resulting alignment was then used
155 for phylogenetic analysis. The partial DNA-R sequences from collected BBTV samples were
156 compared with published reference sequences in NCBI (Appendix Table 1) to determine its
157 relation to the South Pacific group and Asian group and to confirm the identity of the virus.
158 Two (2) Abaca bunchy top virus (ABTV) reference sequences (accession numbers:
159 EF546813.1, EF546807.1) served as outgroups in the analysis.

160

161 **Genetic diversity and demographic analyses**

162 Parameters of genetic diversity and demographic analysis using the partial DNA-R gene of
163 BBTV populations isolated in the Philippines were computed using DNA Sequence
164 Polymorphism (DnaSP) (Rozas et al., 2017). The estimates of evolutionary divergence (genetic
165 distance) over sequence pairs between and within population (inter- and intra-population,
166 respectively) of BBTV isolates were computed using Molecular Evolutionary Genetics
167 Analysis (MEGA X) (Kumar et al., 2018) based on T93 nucleotide substitution model (Tamura
168 and Nei 1993). The rate variation among sites was modeled with a gamma distribution (+G)
169 and discreet evolutionary invariable sites (+I).

170

171 **Haplotype network and phylogenetic analyses**

172 Based on DNA-R sequence of BBTV, two haplotype networks were constructed using
173 Population Analysis with Reticulate Trees (PopART) (Leigh and Bryant, 2015): (1) median
174 joining network of collected Philippine isolates; and (2) minimum spanning network of
175 collected Philippine isolates with other published reference sequences from the Philippines and

176 SEA. The BBTV phylogenetic tree was reconstructed using the maximum likelihood statistical
177 method implemented in IQ-TREE (Nguyen et al., 2015) with best-fit substitution model
178 selected based on Bayesian information criterion (BIC) through ModelFinder
179 (Kalyaanamoorthy et al., 2017). The tree was generated using TIM2 model (AC=AT, CG=GT
180 and unequal base frequency; Posada 2008) with empirical base frequencies (+F) and FreeRate
181 heterogeneity across sites model (+R3) (Yang, 1995; Soubrier et al., 2012). The resulting
182 phylogenetic tree was validated with 1,000 replicates of ultrafast bootstrapping (Hoang et al.,
183 2018) and visualized using FigTree (Rambaut, 2018).

184

185 **Results**

186

187 **Genetic diversity and demographic analysis**

188 Banana leaves showing characteristic symptoms of banana bunchy top disease (BBTD) were
189 collected from 11 banana growing areas in the country, namely, Cagayan, Isabela, Quirino,
190 Batangas, Laguna, Rizal, Quezon, Palawan, Cebu, Leyte, and Davao del Sur (Figure 1). Among
191 the BBTV populations with partial DNA-R sequences (Table 1), the highest number of
192 segregating sites (S) was observed in Quezon and Batangas (S=25 and 28, respectively) while
193 the lowest was observed in Rizal and Cebu (S=6) (Table 2). The nucleotide diversity (π) was
194 highest in Quezon ($\pi=0.00893$) and lowest in Rizal ($\pi=0.00385$). The average number of
195 nucleotide differences (k) was highest in Quezon, Cebu, and Leyte (ranging from 6 to 6.897)
196 and lowest in Rizal, Palawan, and Leyte (ranging from 3 to 3.2). The number of haplotype (h)
197 was highest in Batangas and Quezon (h=12) with relatively high haplotype diversity (hd) of
198 0.967 and 0.987, respectively. A high haplotype diversity (hd) of 1 was observed in Cebu (h=2),
199 Laguna (h=8), and Davao del Sur (h=5) as the haplotype number in these locations
200 corresponded to the number of samples obtained. Lowest haplotype diversity (hd) was

201 observed in Palawan (0.644). Analysis of all Philippine BBTv populations showed 59 total
202 segregating sites (S) with nucleotide diversity (π) of 0.0021 and average number of nucleotide
203 differences (k) of 5.51984, while the haplotype number (h) was 41 with haplotype diversity
204 (hd) of 0.971.

205

206 For the test of neutrality (Table 2), significantly ($P < 0.02$) negative Fu's F_s was observed in
207 BBTv population from Batangas (-5.478), Laguna (-4.309), and Quezon (-5.026) (Table 2),
208 while only Batangas has significantly ($P < 0.05$) negative Tajima's D (-1.80970). Meanwhile,
209 Tajima's D was not computed for Cebu samples due to small sample size. Analysis of all
210 Philippine BBTv isolates showed a significant and highly negative Fu's F_s (-33.210) and
211 Tajima's D (-1.98369).

212

213 BBTv isolates from Cebu, Leyte, Davao del Sur, Quezon, Laguna, and Batangas showed the
214 highest inter-population genetic distance with Isabela isolates (ranging from 0.009 to 0.011),
215 while BBTv isolates from Quirino, Cagayan, Palawan, and Rizal showed the highest inter-
216 population genetic distance with Batangas isolates (ranging from 0.007 to 0.010) (Table 3).
217 Intra-population genetic distance was highest in Quezon isolates, while Rizal isolates showed
218 the lowest.

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220 The data on genetic diversity, demographic analysis, and intra-population genetic distance
221 were not computed for BBTv isolates from Cagayan, Isabela, and Quirino as only one
222 representative isolate was obtained from each location.

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226 **Haplotype analysis**

227 Haplotype network of BBTv samples isolated from the Philippines revealed three major
228 haplotype groups (HA, HB, and HC) using the partial DNA-R gene (Figure 2a). The first
229 haplotype group (HA) includes isolates from Laguna, Batangas, and Quezon which are
230 provinces from the Luzon region. The second haplotype group (HB) includes isolates from
231 Laguna and Quezon (Luzon region), Cebu and Leyte (Visayas region), and Davao del Sur
232 (Mindanao region). The third haplotype group (HC) includes isolates from all sampled
233 provinces (except Laguna and Batangas). Interestingly, all Palawan isolates were included in
234 this haplotype group only. Additional BBTv samples and genes will be analyzed to confirm
235 the haplotype groupings observed.

236

237 Using the DNA-R gene, a SEA haplotype network consisting of collected Philippine isolates
238 and reference sequences from the Philippines and SEA was constructed (Figure 2b). It showed
239 distinct grouping of Philippine isolates from its neighboring Asian countries (such as
240 Indonesia, Taiwan, Japan), while China and Vietnam showed the most distant haplotype
241 grouping. Here, the HC group remains distinct wherein BBTv sequences from Indonesia (n=2)
242 and India (n=1) were also included.

243

244 **Phylogenetic analysis**

245 Phylogenetic analysis showed two broad clades/groups of BBTv, namely, the SEA and PIO
246 groups, with high bootstrap support values of 88% and 90%, respectively (Figure 3). All
247 Philippine BBTv sequences were found in the SEA clade. The Philippine reference sequences
248 and Philippine BBTv isolates in this study clustered together wherein sequences from
249 Indonesia (n=8) and India (n=1) were also included. BBTv sequences from India and Egypt
250 were both found in SEA and PIO clades. Notably, the clustering of Philippine BBTv isolates

251 appeared to follow the three haplotype groupings observed in this study (HA, HB, and HC).
252 Philippine reference sequences were found among the HA and HB isolates but not on HC
253 isolates. On the other hand, Indonesian sequences were found among HB and HC isolates,
254 while an Indian BBTV sequence was found among HC isolates. In the SEA clade, China and
255 Vietnam sequences formed a separate cluster from the rest of SEA sequences with a well-
256 supported bootstrap value of 82%.

257

258 **Discussion**

259 Banana bunchy top disease has been an important disease of the banana crop in the Philippines
260 (Molina et al., 2009). Molecular information regarding the Philippine BBTV isolates has been
261 lacking and remains unexplored. Thus, this study was performed to investigate the genetic
262 structure and diversity of the BBTV isolates in the Philippines. Here, the partial DNA-R region
263 of BBTV was sequenced due to its wide application in assessing genetic diversity and other
264 molecular analyses (Bell et al., 2002; Furuya et al., 2005; Amin et al., 2007; Kumar et al., 2011;
265 Shekhawat et al., 2012; Banerjee et al., 2014; Chiaki et al., 2015; Mukwa et al., 2016). Survey
266 and sample collection were conducted in banana growing areas in various regions in the
267 Philippines for BBTV detection and diversity analysis. The banana cultivars wherein BBTV
268 was isolated include Saba, Lakatan, Latundan, and other unknown varieties.

269

270 Among the sampling sites, Quezon appear to have the most diverse BBTV population as shown
271 by high nucleotide segregating sites (S), nucleotide diversity (π), average number of nucleotide
272 differences (k), haplotype number (h), haplotype diversity (hd), intra-population genetic
273 distance, and a significant negative Fu's F_s value. Significant and negative Fu's F_s was
274 observed also in Laguna BBTV population, while significant and negative Fu's F_s and
275 Tajima's D values were observed in BBTV population from Batangas. Fu's F_s is regarded as

276 a more sensitive indicator of population expansion and a more powerful test of neutrality than
277 the Tajima's D , which probably contributed to the inconsistent results (Zeng et al., 2006). More
278 BBTV samples should be collected from different provinces for genetic diversity and
279 demographic analyses to confirm the results obtained. Overall, significant and highly negative
280 Fu's F_s and Tajima's D were observed using all Philippine BBTV isolates. These results
281 suggest that there is an excess number of rare alleles in Philippine BBTV isolates, probably
282 due to its recent population expansion (or from genetic hitchhiking) as evidenced also by
283 overall high haplotype diversity with relatively low overall nucleotide diversity.

284

285 Haplotype network and phylogenetic analyses of partial DNA-R of combined SEA and
286 Philippine sequences suggest that geographic location heavily affects the distribution of BBTV
287 as indicated by geographically proximate haplotypes in each group. Viruses may have evolved
288 independently mainly because countries are separated by sea, and host movement could have
289 been limited within the haplotype groupings. The complex haplotype network of BBTV
290 isolates suggests that the Philippines, as part of SEA, is a hotspot of an on-going BBTV
291 diversification (Stainton et al., 2015). Furthermore, the Philippines, along with other
292 neighboring countries such as New Guinea and Indonesia, are believed to be the center of origin
293 of domesticated bananas (Perrier et al., 2011). This may indicate that the intensive
294 domestication of bananas within the region might have been a possible driver for the
295 diversification of BBTV in the country. Meanwhile, the haplotype network analysis revealed
296 three major haplotype groups (HA, HB, and HC) of BBTV isolates collected in the Philippines.
297 Interestingly, BBTV isolates from Palawan were only found in the HC group. This province
298 also has very low haplotype number (h) and haplotype diversity (hd), and a relatively low
299 nucleotide diversity (π). These results could be probably caused by recent population
300 bottleneck and recent introduction of BBTV in the area due to the movement of planting

301 materials (e.g., Lakatan variety) from other provinces in the Philippines. Due to the strict
302 quarantine implementation in the province, BBTV has been reported only recently in Palawan.
303 It appears that the HC group is widespread in the country and was also introduced in Palawan.
304 Upon inclusion of Philippine and SEA reference sequences in the haplotype network, however,
305 only the HC group remained distinct wherein few BBTV reference sequences from India and
306 Indonesia were also included.

307

308 As expected, phylogenetic analysis based on partial DNA-R showed that the collected local
309 isolates were more closely related with the SEA group (where the Philippines is geographically
310 classified) than the PIO group (Karan et al., 1994). This could also mean that plant and virus
311 movement is limited within the SEA region (Karan et al., 1994; Wickramaarachchi 2016).
312 Philippine BBTV isolates formed a monophyletic clade which suggests a monophyletic origin
313 of the majority of local isolates from a common SEA ancestor. The collected local isolates also
314 clustered with Philippine reference sequences which confirms their identity as BBTV and may
315 indicate that virus movement could be limited in the country (Stainton et al., 2015). In the
316 phylogenetic tree, the clustering of Philippine BBTV isolates seemed to follow the observed
317 three haplotype groupings (HA, HB, and HC). However, as shown in the tree, no Philippine
318 reference sequences appeared to cluster with HC isolates. A more exhaustive survey of
319 reference sequences in the Philippines will be performed and more BBTV samples in the
320 Philippines will be sequenced to verify the findings on haplotype groupings and phylogenetic
321 analysis.

322

323 Some BBTV sequences from Indonesia and India were also clustered within the Philippine
324 clade (the former clustered with HB and HC isolates, while the latter clustered with HC
325 isolates), suggestive of sequence homology and possible BBTV movement in these countries.

326 It was previously inferred that the Indian subcontinent is a major contributor to the long-
327 distance dispersal of BBTv, both as donor and recipient. For instance, the introduction events
328 of SEA isolates were recently detected between 1976 and 1991 in India (Stainton et al., 2015).
329 Thus, it may be deduced that there is a probable dispersal event of Philippine BBTv isolates
330 to India. Outside the Philippine clade, BBTv sequences from Egypt and India (which are
331 known to be closely related with PIO group) were also clustered in the larger SEA clade,
332 indicating the presence of isolates that are related with SEA group. On the other hand, reference
333 sequences from Vietnam and China formed a monophyletic clade and appeared to be separated
334 from the larger SEA group. Similar observation was reported by Rao (2017) wherein the
335 constructed DNA-R phylogenetic tree depicts a further separation of BBTv isolates from
336 China and Vietnam into sub-groups 2, 3, and 4; while the rest of the members of the Asian
337 group exclusively formed the sub-group 1.

338

339 In summary, the results of this study showed that BBTv is widespread and diverse in the
340 Philippines and undergoing population expansion. However, more samples and genes should
341 be analyzed to confirm the results obtained especially at the province level. Additional
342 reference sequences from the Philippines and other countries with reported BBTv occurrence
343 will be included in the analysis. Recombination analysis shall be also performed to provide
344 further understanding regarding the evolutionary history of Philippine BBTv isolates.
345 Nevertheless, the insights drawn from this research endeavor will provide a framework in the
346 development of improved BBTv-specific detection marker systems in the country as well
347 enable the strategic BBTv-resistant variety deployment across various regions in the
348 Philippines.

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350

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362

363 **Conflict of interest**

364 The authors declare no conflict of interests.

365

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536 **Table 1.** Collected BBTV Philippine isolates sequenced in this study.
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Isolate	Location	Collection Year	Host variety
Balam_Cebu_B22	Balamban, Cebu	2015	Unknown
Baybay_Ley_B34	Baybay, Leyte	2015	Unknown
Baybay_Ley_B35	Baybay, Leyte	2015	Unknown
BenSol_Isab_B18	Benin Soledad, Isabela	2015	Unknown
Calin_Dav_B12	Calinan, Davao City	2015	Unknown
Calin_Dav_B14	Calinan, Davao City	2015	Unknown
Can_Que3	Candelaria, Quezon	2019	Unknown
Can_Que_07	Candelaria, Quezon	2019	Tordan and Saba
Can_Que_08	Candelaria, Quezon	2019	Tordan and Saba
Can_Que_11	Candelaria, Quezon	2019	Tordan and Saba
Can_Que_12	Candelaria, Quezon	2019	Tordan and Saba
Can_Que13	Candelaria, Quezon	2019	Tordan and Saba
Can_Que15	Candelaria, Quezon	2019	Tordan and Saba
Diffun_QUI_B19	Diffun, Quirino	2015	Unknown
Luc_Que_01	Lucban, Quezon	2019	Unknown
Luc_Que_02	Lucban, Quezon	2019	Unknown
Luc_Que_05	Lucban, Quezon	2019	Lakatan Tagalog
Luc_Que4	Lucban, Quezon	2019	Lakatan Tagalog
Luc_Que10	Lucban, Quezon	2019	Lakatan Tagalog
Luc_Que6	Lucban, Quezon	2019	Lakatan Tagalog
Luis_Lag_04	Luisiana, Laguna	2019	Lakatan
Luis_Lag_07	Luisiana, Laguna	2019	Lakatan
Luis_Lag_08	Luisiana, Laguna	2019	Lakatan
Luis_Lag6	Luisiana, Laguna	2019	Lakatan
Luis_Lag7	Luisiana, Laguna	2019	Lakatan
Magda_Lag_B10	Magdalena, Laguna	2015	Unknown
Naga_Cebu_B24	Naga, Cebu	2015	Unknown
Orm_Ley_B31	Ormoc, Leyte	2015	Unknown
Orm_Ley_B32	Ormoc, Leyte	2015	Unknown
Orm_Ley_B32b	Ormoc, Leyte	2015	Unknown
Pagsan_Lag_01	Pagsanjan, Laguna	2019	Saba
Pagsan_Laguna2	Pagsanjan, Laguna	2019	Saba
Rizal_Cag_B16	Rizal, Cagayan	2015	Unknown
Roxas_Pal_01	Roxas, Palawan	2019	Lakatan and Saba
Roxas_Pal_03	Roxas, Palawan	2019	Lakatan and Saba
Roxas_Pal_04	Roxas, Palawan	2019	Lakatan and Saba
Roxas_Pal_05	Roxas, Palawan	2019	Lakatan and Saba
Roxas_Pal_07	Roxas, Palawan	2019	Lakatan
Roxas_Pal12	Roxas, Palawan	2019	Lakatan
Roxas_Pal15	Roxas, Palawan	2019	Lakatan
Roxas_16	Roxas, Palawan	2019	Lakatan

Roxas_Pal2	Roxas, Palawan	2019	Lakatan
Roxas_Pal5	Roxas, Palawan	2019	Lakatan
Talisay_Bat_08	Talisay, Batangas	2019	Saba
Talisay_Bat_09	Talisay, Batangas	2019	Saba
Talisay_Bat_10	Talisay, Batangas	2019	Saba
Talisay_Bat_10b	Talisay, Batangas	2019	Saba
Talisay_Bat3	Talisay, Batangas	2019	Saba
Talisay_Bat4	Talisay, Batangas	2019	Saba
Talisay_Bat6	Talisay, Batangas	2019	Saba
Tanauan_Bat_11	Tanauan, Batangas	2019	Latundan
Tanauan_Bat_16	Tanauan, Batangas	2019	Latundan
Tanauan_Bat_17	Tanauan, Batangas	2019	Latundan
Tanauan_Bat_18	Tanauan, Batangas	2019	Latundan
Tanauan_Bat_20	Tanauan, Batangas	2019	Latundan
Tanauan_Bat19	Tanauan, Batangas	2019	Latundan
Tanauan_Bat11	Tanauan, Batangas	2019	Latundan
Tanay_Riz_B01	Tanay, Rizal	2015	Unknown
Tanay_Riz_B02	Tanay, Rizal	2015	Unknown
Tanay_Riz_B03	Tanay, Rizal	2015	Unknown
Tanay_Riz_B04	Tanay, Rizal	2015	Unknown
Tugbok_Dav_B11	Tugbok, Davao City	2015	Unknown
Tugbok_Dav_B13	Tugbok, Davao City	2015	Unknown
Tugbok_Dav_B15	Tugbok, Davao City	2015	Unknown

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Table 2. Parameters of genetic diversity and demographic analysis using DNA-R gene of BBTV populations isolated in the Philippines.

Location	n	S			π (SD)	k	h	hd (SD)	Fu's F_s		Tajima's D	
		SS	PIS	Total S					F_s	P value ^a	D	Significance
Cagayan	1	-	-	-	-	-	-	-	-	-	-	-
Isabela	1	-	-	-	-	-	-	-	-	-	-	-
Quirino	1	-	-	-	-	-	-	-	-	-	-	-
Batangas	14	23	5	28	0.00653 (0.00133)	5.07692	12	0.967 (0.044)	-5.478	0.00400	-1.80970	$P < 0.05^*$
Laguna	8	7	5	12	0.00567 (0.00057)	4.39286	8	1.000 (0.063)	-4.309	0.00602	-0.25574	$P > 0.10^{ns}$
Rizal	4	6	0	6	0.00385 (0.00167)	3.00000	3	0.833 (0.222)	0.731	0.62381	-0.80861	$P > 0.10^{ns}$
Quezon	13	15	10	25	0.00893 (0.00162)	6.89744	12	0.987 (0.035)	-5.026	0.01200	-0.90389	$P > 0.10^{ns}$
Palawan	10	8	3	11	0.00416 (0.00158)	3.22222	3	0.644 (0.101)	3.321	0.94790	-0.76710	$P > 0.10^{ns}$
Cebu	2	6	0	6	0.00772 (0.00386)	6.00000	2	1.000 (0.500)	1.792	0.53944	-	-
Leyte	5	8	0	8	0.00411 (0.00162)	3.20000	3	0.700 (0.218)	1.458	0.75358	-1.17432	$P > 0.10^{ns}$
Davao del Sur	5	5	7	12	0.00798 (0.00164)	6.20000	5	1.000 (0.126)	-1.011	0.15015	0.55247	$P > 0.10^{ns}$
ALL	64	37	22	59	0.00721 (0.00053)	5.51984	41	0.971 (0.011)	-33.210	0.00000	-1.98369	$P < 0.05^*$

n=no. of BBTV sequences; S=segregating sites; SS=singleton sites; PIS=parsimony informative sites; π =nucleotide diversity; k=average no. of nucleotide differences, h=no. of haplotypes; hd=haplotype diversity; SD=standard deviation; *=significant; ^a should be regarded as significant (5% level) if $P < 0.02$.

Table 3. Estimates of evolutionary divergence (inter- and intra-population genetic distance) of BBTV populations collected in the Philippines.

Location	Inter-population genetic distance										Intra-population genetic distance	
	Cebu	Leyte	Isabela	Davao del Sur	Quezon	Quirino	Laguna	Cagayan	Palawan	Batangas	Distance	SE
Cebu											0.00513	0.00259
Leyte	0.00644										0.00360	0.00142
Isabela	0.01037	0.01011									-	-
Davao del Sur	0.00592	0.00618	0.00959								0.00566	0.00196
Quezon	0.00700	0.00769	0.01017	0.00705							0.00809	0.00201
Quirino	0.00644	0.00618	0.00385	0.00566	0.00645						-	-
Laguna	0.00499	0.00628	0.01020	0.00596	0.00711	0.00628					0.00392	0.00140
Cagayan	0.00514	0.00489	0.00516	0.00437	0.00555	0.00128	0.00627				-	-
Palawan	0.00789	0.00763	0.00790	0.00680	0.00769	0.00400	0.00885	0.00271			0.00370	0.00126
Batangas	0.00707	0.00865	0.01112	0.00781	0.00838	0.00737	0.00570	0.00831	0.01022		0.00619	0.00143
Rizal	0.00595	0.00534	0.00614	0.00489	0.00613	0.00225	0.00676	0.00096	0.00368	0.00894	0.00192	0.00107
Mean distance (SE): 0.00722 (0.00162)												

SE=standard error

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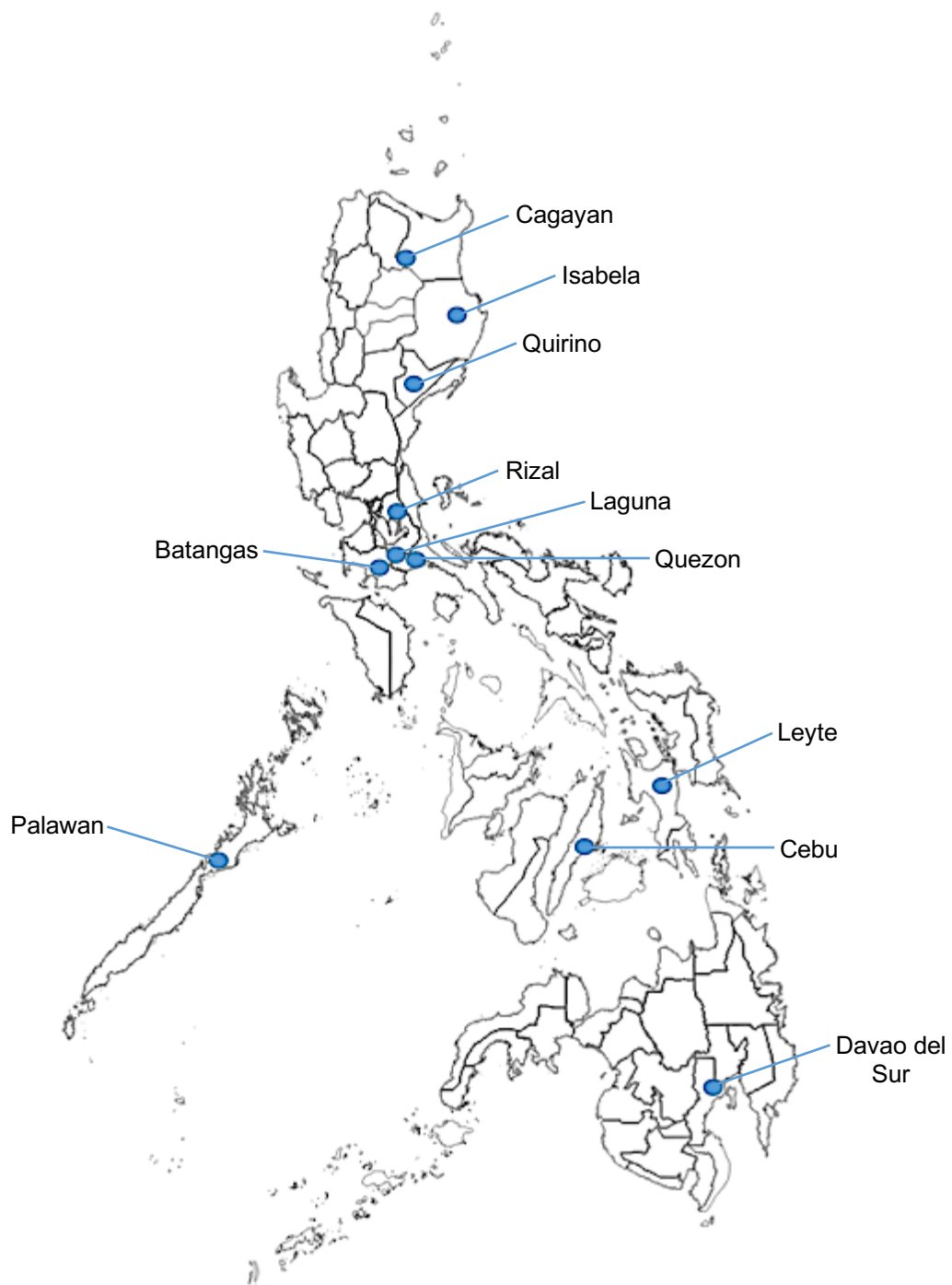


Figure 1. Geographical distribution of BBTB collected from banana-growing areas in the Philippines.

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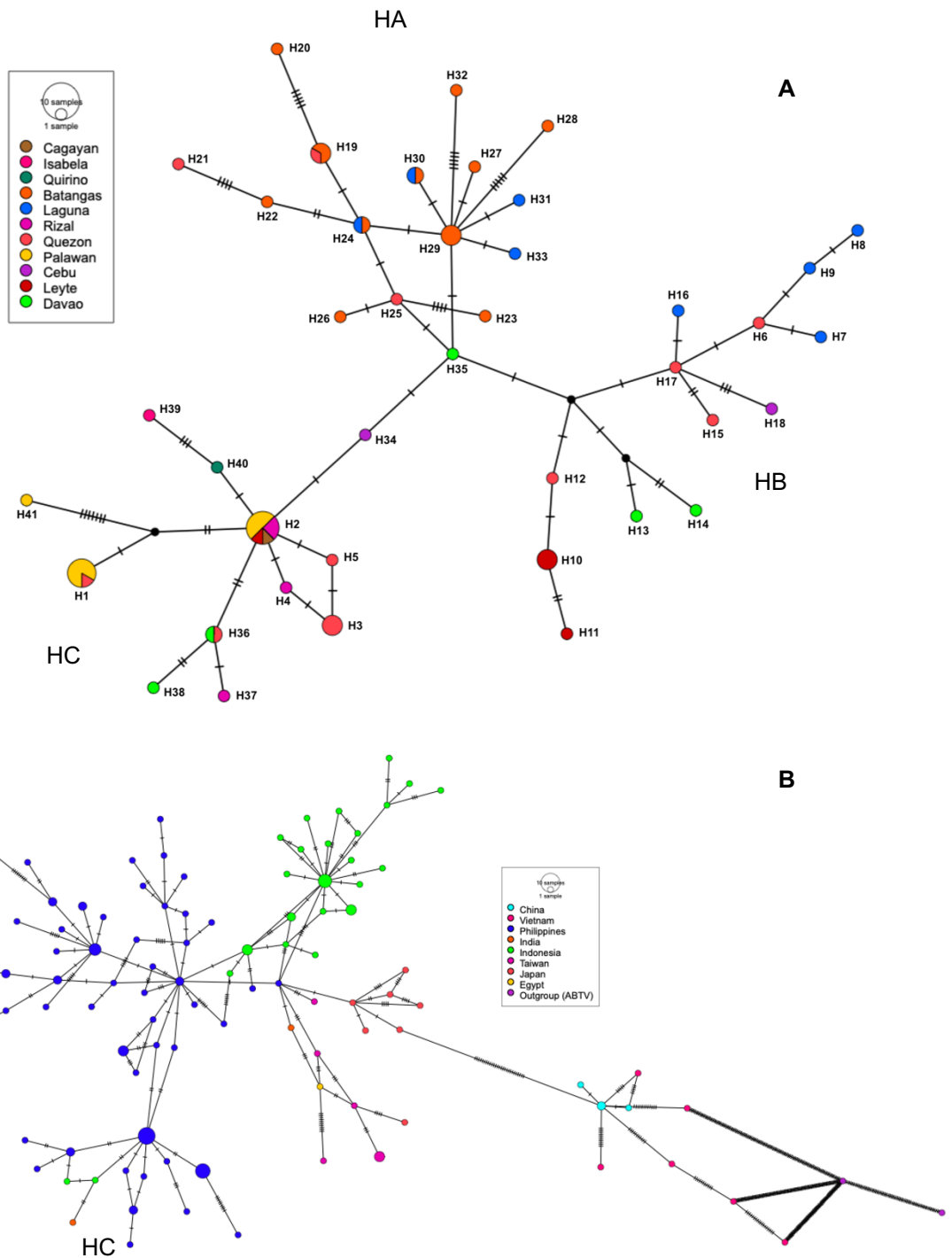
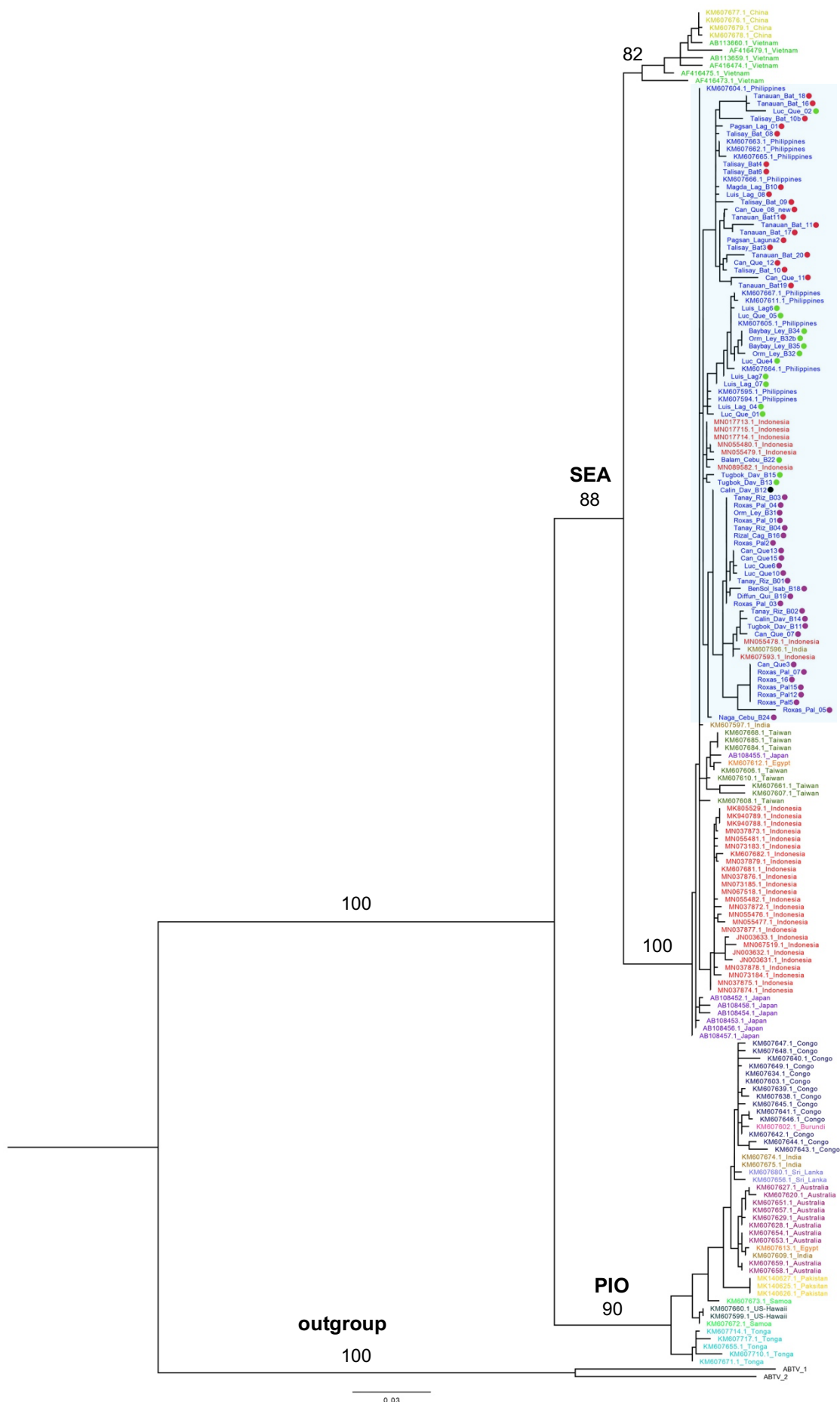


Figure 2. Haplotype network based on partial DNA-R sequence of BBTV constructed using PopART (Leigh and Bryant, 2015): median joining network of the collected Philippine isolates (A) and minimum spanning network of collected isolates with Philippine and other SEA reference sequences (B). Haplotype Groups: HA, HB, HC.

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151 **Figure 3.** Maximum likelihood phylogenetic tree constructed using IQ-TREE from the partial
152 DNA-R sequence alignment of collected Philippine BBTV isolates and published Philippine,
153 SEA, and PIO reference sequences. Best-fit model was selected according to BIC using
154 ModelFinder (Kalyaanamoorthy et al., 2017). The tree was generated using TIM 2 model
155 (AC=AT, CG=GT and unequal base frequency; Posada 2008) with empirical base frequencies
156 (+F) and FreeRate heterogeneity across sites model (+R3) (Yang, 1995; Soubrier et al., 2012).
157 The tree was tested with 1,000 replicates of ultrafast bootstrapping (Hoang et al., 2018) and
158 visualized using FigTree (v1.4.4) (Rambaut 2018). The numbers in the branches are bootstrap
159 support values. Clade highlighted in blue contains the collected Philippine BBTV isolates with
160 colored dots corresponding to haplotype grouping: red dot = HA, green dot = HB, and purple
161 dot = HC. SEA = Southeast Asian group, PIO = Pacific-Indian Oceans group. ABTV sequences
162 served as outgroups.
163