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Development and validation of a one-step reverse transcription real-time PCR assay for simultaneous detection and identification of tomato mottle mosaic virus and tomato brown rugose fruit virus

Antonio Tiberini; Ariana Manglli; Anna Taglienti; Ana Vučurović; Jakob Brodarič; Luca Ferretti; Marta Luigi; Andrea Gentili; Nataša Mehle

1 CREA – Research Centre for Plant Protection and Certification, Via C.G. Bertero 22 - 00156 Roma, Italy.
2 National Institute of Biology, Department of Biotechnology and Systems Biology, Večna pot 111, SI-1000 Ljubljana, Slovenia
3 University of Nova Gorica, School for Viticulture and Enology, Dvorec Lanthieri, Glavni trg 8, SI-5271 Vipava, Slovenia
* Correspondence: antonio.tiberini@crea.gov.it

Abstract: Tobamovirus species represent a threat to solanaceous crops worldwide, due to their extreme stability and being seed-borne. In particular, recent outbreaks of tomato brown rugose fruit virus in tomato and pepper crops led to the establishment of prompt control measures, and the need for reliable diagnosis was urged. Another member of the genus, tomato mottle mosaic virus, has recently risen attention due to reports in different continents and its common features with tomato brown rugose fruit virus. In this study, a new real-time RT-PCR detection system was developed for tomato brown rugose fruit virus and tomato mottle mosaic virus on tomato leaves and seeds using TaqMan chemistry. This test was designed to detect tomato mottle mosaic virus by amplifying the movement protein gene in a duplex assay with tomato brown rugose fruit virus target on the CP-3’NTR region, which was already validated as a single assay. The performance of this test was evaluated, displaying analytical sensitivity 10^{-5}-10^{-6}-fold dilution for seeds and leaves, respectively, and good analytical specificity, repeatability, and reproducibility. Using the newly developed and validated test, tomato brown rugose fruit virus detection was 100% concordant with previously performed analyses on 106 official samples collected in 2021 from different continents.

Keywords: ToBRFV, ToMMV, Solanum lycopersicum, Capsicum annum, leaves detection, seeds detections, performance criteria

1. Introduction

Tomato (Solanum lycopersicum L.) is one of the most important crops worldwide, with an annual production of around 180 million tonnes of fresh weight [1]. Viral diseases are major causes of yield losses in tomato production, the more representative and important species that could seriously affect this crop belong to Crinivirus, Cucumovirus, Begomovirus, Potexvirus, Potyvirus, Tospovirus, Polerovirus, and Tobamovirus [2,3]. The latter genus has long been considered a threat for agricultural crops; tobacco mosaic virus (TMV) and tomato mosaic virus (ToMV) have been reported to infect tomato and other important solanaceous crops for more than a century, representing the type species of the genus and most noticeable and economically important viruses up to the development of resistance genes [4,5]. Tobamoviruses are characterized to have stable virions with a mechanical
transmission pathway. In most cases, virions contaminate the seed coat and infectivity is preserved in seeds for up to several years [6]. In nurseries, during plantlets emersion, the tobamovirus on the seed coat may be transmitted to wounded roots [6]. Furthermore, even though low seed-to-seedling transmission rate is reported, due to the large-scale usage of seeds and seedlings in farming, the contribution of a single infected seedling to an outbreak, or long-distance transmission, may become significant [7]. Due to mechanical transmission, the primary source can be easily spread by contacts, hands, tools, greenhouse structure and bumblebees [8]. Tobamovirus infectious particles are reported to be stable in plant debris and in contaminated soil for years [9]. In recent years, a new member of the Tobamovirus genus, tomato brown rugose fruit virus (ToBRFV) was characterized and reported to induce symptoms of mosaic, discoloration and deformation on leaves, and size reduction, discoloration, brown rugosity, malformation on fruits, making them unmarketable; reduction of plant vigour was also observed in diseased plants. Such severe symptomatology was also observed in tomato cultivars harbouring resistance genes (Tm-1, Tm-2, Tm-22) against tobamoviruses [10]. After emerging in Israel and Jordan in 2014 [10,11], the virus was reported in several European countries [12] (e.g. Italy [13], United Kingdom [14], The Netherlands [15], Greece [16] and Spain [17]). In North America ToBRFV was first reported in Mexico in 2018 [18], then it spread also in California [19]. In Asia it was reported first in Turkey [20] and then in China in 2019 [21]. Due to its high transmissibility and resistance breaking, ToBRFV was included in the A2 list of the European and Mediterranean Plant Protection Organization (EPPO) and subjected to prompt measures (Commission Implementing Decision (EU) 2020/1191 and subsequent amendments) to prevent its spread and establishment in the European Union. Validated molecular tests for the detection and identification of ToBRFV are available in EPPO Standard PM 7/146 (1) [22]; for seed testing, only real-time reverse-transcription polymerase chain reaction (real-time RT-PCR) tests are recommended.

A member of subgroup I of tobamoviruses, which also includes TMV, ToMV and ToBRFV, is tomato mottle mosaic virus (ToMMV). It was first reported in Mexico in 2013, on a sample collected in 2009 [23]; the associated disease is characterized by symptoms of plant stunting, severe mosaic, mottling, distortion and necrosis on leaves, fruit necrosis in tomato [23–25]; in pepper symptoms as apical yellowing and necrosis were observed [24,26]. After the first report, ToMMV has been detected in the United States [27], Spain[26], Israel [28], China [29], Brasil [30] and Czech Republic [31]. Pepper seed lots imported in Australia were also found infected by ToMMV [32]. Moreover, Mut Bertome [33] reported that the presence of ToMMV was confirmed in batches of tomato seeds imported by Spanish companies between 2019 and 2021. In their study 130 tomato samples and 85 pepper samples from different countries were analysed by end point RT-PCR, and ToMMV was detected in 5 tomato seed samples from China (3), the USA (1) and Israel (1).

In addition, ToMMV, that shares a close phylogenetic relationship with ToBRFV [34], has proven to partially break resistance of tomato resistant to ToMV [25]. According to Nagai et al. [35], Tm-2i gene is involved in the resistance against ToMMV; this study reported that three out of seven tomato varieties and hybrid lines infected with ToMMV tested...
positive for the virus, suggesting an hypothetical ability to overcome resistance genes [35].
Additional resistance tests were run on a certified ToMV-resistant variety by Sui et al.: around 10% of plants were successfully infected with ToMMV [25].

In view of the above, the EPPO Panel on Phytosanitary Measures recommended the inclusion of ToMMV in the EPPO Alert List (2020); Australia has implemented emergency measures for ToMMV, requiring testing for imported tomato and pepper seed lots; finally, express Pest Risk Analyses were recently performed in Germany [36] and the Netherlands [37]. In a context of envisaged phytosanitary measures against ToMMV, proper diagnostic tests for its detection and identification, as stated by EPPO PM 7/76 (5) [38], are critical for the appropriate application of such measures. In particular, because of the high risk of cross-reactivity that serological methods present in the detection of tobamoviruses, molecular methods represent the gold standard for the specific detection of viral species within this genus. To date, there are few published protocols for the detection and identification of ToMMV. These are Sanger sequencing of generic to bromovirus RT-PCR or nested RT-PCR products (e.g. [10, 39, 40]) and multiplex endpoint RT-PCR for TMV, ToMV, and ToMMV [25]. In this work, we report the development and validation, according to EPPO PM 7/98 (4) guidelines [41], of a duplex real-time RT-PCR assay for the detection and identification of ToMMV and ToBRFV in leaves and seeds. To our knowledge, this is the first molecular method developed for the simultaneous detection and identification of the two emerging viruses mentioned above, which currently pose the greatest threat to solanaceous crops worldwide.

2. Results

2.1 Primers/probe design and evaluation
ToMMV specific primers/probe sets obtained targeting CP and MP regions are summarized in Table 1. The set 1 and 2 were labeled in HEX whereas sets 3 and 4 in Texas Red. Preliminary reactions performed including the ToMMV-positive/ToBRFV-positive/negative controls showed that all the ToMMV primers/probe sets were specific and allowed to correctly identify ToMMV, with good Cq values ranging from 6.8 to 13.6, and discriminate it from ToBRFV without cross-reactions (Figure 1). The best annealing temperature was set at 60 °C with primers and probe concentration at 0.3 and 0.2 μM respectively. For all the sets, no amplification signals were obtained for negative controls. The primes set 1 and 2, targeting CP region and labeled in HEX, showed the lowest level of relative fluorescence units (RPU), with a value of 2500 and 1000, respectively whereas sets 3 and 4 showed RPU values above 3500 (Figure 1).

| Set | Primer/probe ID | 5’-3’ Sequence | Labelling | Genome Position (nt) | Coding region |
|-----|----------------|----------------|-----------|----------------------|---------------|
| 1   | ToMMV MWAT Fw  | GGAACAGGTTTCTACAACCAGAG | HEX       | 6124-6146            | CP            |
|     | ToMMV MWAT Rv  | CGTACGCACCACGTGTTT   | Texas Red | 6235-6254            |               |
All four primers/probe sets were tested versus some other tobamovirus species (TMV, ToMV, TGMV, BPeMV, PMMoV, CGMMV). None of the set cross-react with TMV, ToMV, TGMV, BPeMV and PMMoV. However, all three primers/probe sets targeting CP resulted to cross-react with CGMMV, whereas no signal was observed for the set targeting MP region (set 3) (Figure 1).

**Figure 1.** Amplification curves obtained testing each ToMMV primers/probe set versus ToMMV and non-target tobamovirus species (ToFBRFV, TMV, ToMV, TGMV, BPeMV, PMMoV, CGMMV). Cross-reaction of some primers/probe sets were observed only with CGMMV. (Probe labelling: green for HEX, red for TexasRed).

| Set       | Forward Primer | Reverse Primer | Probe/Labelling | RPU Range |
|-----------|----------------|----------------|----------------|-----------|
| 1         | TCTGCGCCACCGTGCTAAGTAAT | GGAACAGGTTCATAACACGAG | HEX           | 6180-6203 |
| 2         | TCTGCGCCACCGTGCTAAGTAAT | GGAACAGGTTCATAACACGAG | HEX           | 6124-6146 |
| 3         | TCTGCGCCACCGTGCTAAGTAAT | GGAACAGGTTCATAACACGAG | HEX           | 6180-6203 |
| 4         | TCTGCGCCACCGTGCTAAGTAAT | GGAACAGGTTCATAACACGAG | HEX           | 6124-6146 |

In view of these results, primers/probe set 2, targeting CP region and showing the lowest level of RPU was excluded from further evaluation. The remaining primers/probe sets 1, 3 and 4 were tested in duplex/triplex assay with ToBRFV assays. Considering the targeting
region/labelling matches, a total of four combinations were obtained and tested as reported in table 2.

**Table 2.** Quantification cycle (Cq) values obtained for each ToMMV and ToBRFV duplex/triplex assay analyzing the same sample set. Results considered as positive are marked in black bold.

|                | Duplex A | Duplex B | Triplex A | Triplex B |
|----------------|----------|----------|-----------|-----------|
|                | ToBFRV M&W set 1 | ToMMV | ToBFRV M&W set 3 | ToBFRV ISHI-Veg set 3 | ToBFRV ISHI-Veg set 4 | ToMMV set 4 |
| Healthy leaf   | NA       | NA       | NA        | NA        | NA        | NA        | NA        |
| Healthy seeds  | 37.8*    | NA       | NA        | 36.0*     | 36.5*     | NA        | 35.4*     | 36.1*     | NA |
| ToBRFV infected leaves | 16.2    | NA       | 17.5      | NA        | 18.2      | 18.4      | NA        | 18.7      | 18.8 |
| ToBRFV infected seeds | 27.9    | NA       | 25.8      | NA        | 28.4      | 28.2      | NA        | 28.5      | 28.1 |
| ToMMV infected leaves | NA     | 9.2      | NA        | 12.3      | NA        | 13.6      | NA        | NA        | 8.8 |
| Mixed leaf infected sample | 15.2    | 10.1     | 16.6      | 12.3      | 18.6      | 19.2      | 14.4      | 18.2      | 18.8 |
| NAC1           | NA       | NA       | NA        | NA        | NA        | NA        | NA        | NA        | NA |
| NAC2           | NA       | NA       | NA        | NA        | NA        | NA        | NA        | NA        | NA |

*Cq above the Cq threshold considered as negative; NAC-negative amplification control; NA-no exponential amplification curve

The results of duplex/triplex assays showed expected amplification curves with Cq values ranging from 8.8 to 19.2 for leaf samples and from 25.8 to 28.5 for seed samples (Table 2). All the duplex/triplex assays were able to discriminate ToMMV from ToBRFV infected samples and to detect both targets in the mixed infected sample. In all the assays, except for duplex B, some high Cq values, ranging from 35.4 to 37.8, were observed for ToBRFV signal in healthy seed sample. No assay displayed any reaction in healthy leaf and non-amplification controls (NAC 1, 2). Based on these results, the duplex assay using ToMMV set 3 targeting the MP region and the ToBRFV M&W assay (targeting the CP region)-Duplex B, were selected for full validation.

2.2. Duplex assay validation

The selected duplex assay (Duplex B) was further evaluated and included in the validation process according to EPPO PM 7/ 98 (4) guidelines [41].
2.2.1. Analytical sensitivity

The LOD was determined using the 10-fold dilutions series (leaves and seeds) obtained from ToMMV and ToBRFV mixed infected samples (Table 3). In leaves, the LOD were $10^{-5}$ for ToMMV and $10^{-6}$ for ToBRFV, whereas in seeds both viruses could be detected up to the $10^{-5}$ dilution (Figure 2a and 2b). In addition, the E (%) and $R^2$ values were determined for each primers/probe set included in the duplex assay on the basis of the standard curves obtained in leaves and seed dilutions series (Figure 3). Both sets resulted to have a similar efficiency in both matrices, with $R^2$ close to the optimum value of 1 and a comparable efficiency above 95% (Table 3).

**Figure 2.** Amplification curves obtained testing the 10-fold dilutions series of leaf material (panel a) and seeds material (panel b) in mixed infection ToMMV and ToBRFV, with the selected duplex assay (Duplex B: ToMMV set 3 (probe labelled with TexasRed, in red) and ToBRFV M&W (probe labelled with FAM, in blue)).
Figure 3. Duplex assay efficiency. ToBRFV and ToMMV TaqMan® assay standard curves in leaf and seed samples obtained from tenfold serial dilutions. The quantification cycle (Cq) value was plotted against the log concentration of RNA. The curve formula in the graph reports slope and R² values.
In addition, for each virus and matrix LOD was evaluated and the efficiency of the selected ToMMV and ToBRFV primers/probe sets was also calculated in a single assay (Table 3). In both, ToBRFV and ToMMV single assays, the LOD was 10-fold lower than in duplex assay (except for ToBRFV in leaves confirming a LOD of $10^{-6}$), and the efficiency was 1-2% lower in duplex than in single assays. However, the standard curves overlapped sharing common slope values (Figure 4a and 4b).

Table 3. Quantification cycle values obtained for the 10-fold dilutions series (in both leaf and seeds) tested for ToBRFV (in single or duplex assay) and for ToMMV (in single or duplex assay). In addition, for each assay the $E$ (%) and $R^2$ values were determined and reported in table.

|          | Leaves |          |          |          | Seeds |          |          |
|----------|-------|----------|----------|----------|-------|----------|----------|
|          | ToBRFV | ToBRFVd  | ToMMV    | ToMMVd   | ToBRFV | ToBRFVd  | ToMMV   | ToMMVd   |
| $10^8$   | 14.1  | 13.7     | 12.2     | 12.5     | 15.2   | 14.9     | 12.0     | 12.3     |
| $10^4$   | 17.4  | 17.2     | 16.0     | 16.1     | 18.3   | 18.2     | 15.5     | 15.9     |
| $10^2$   | 21.1  | 20.8     | 19.5     | 19.8     | 21.8   | 21.8     | 19.4     | 19.5     |
| $10^3$   | 24.4  | 24.2     | 22.7     | 23.2     | 25.4   | 25.2     | 22.9     | 23.0     |
| $10^4$   | 27.8  | 27.4     | 26.1     | 26.3     | 28.8   | 28.8     | 26.3     | 26.3     |
| $10^5$   | 30.8  | 31.0     | 29.1     | 30.2     | 32.2   | 31.9     | 29.3     | 29.3     |
| $10^6$   | 34.5  | 34.1     | 32.8     | NA       | 34.5   | NA       | 31.9     | NA       |
|          |       |          |          |          |       |          |          |          |
|          | $10^7$ | 38.8*    | NA       | NA       | NA     | NA       | NA       | NA       |
|          | $10^8$ | 39.4*    | NA       | NA       | NA     | NA       | NA       | NA       |
|          | $R^2$  | 0.999    | 0.999    | 0.999    | 0.999  | 0.999    | 0.999    | 0.999    |
|          | $E$    | 97.63    | 95.67    | 98.03    | 95.20  | 95.83    | 95.42    | 97.33    | 96.57    |

s—single assay; d—duplex assay; NA—no exponential amplification curve; *Cq above the Cq threshold and considered as negative (no clear exponential amplification curve)

Figure 4. Comparison of single vs duplex assay efficiencies. In the graphs are reported the TaqMan® assay standard curves obtained in the single and in the duplex assay (for ToMMV on the right and ToBRFV on the left), in leaves (panel a) and seeds (panel b) samples using tenfold serial dilutions.
The quantification cycle (Cq) value was plotted against the log concentration of RNA. The curve formula in the graph reports slope and $R^2$ values.

2.2.2. Analytical specificity
The non-target tobamovirus species were tested by duplex assay at CREA and at NIB (Supplementary Table S2 and Table 4). In general, no important cross-reaction was observed, except for PaMMV and RMV; among them, only the cross-reaction with PaMMV could represent a threat for identification of ToBRFV and ToMMV, in particular in pepper, which is the chief host of PaMMV.

Table 4. Results of testing different isolates of non-target tobamoviruses with duplex ToBRFV&ToMMV assay and with single ToBRFV M&W assay.

|          | NIB | CREA |
|----------|-----|------|
| Virus  | Collection | ID         | Single assay ToBRFV M&W | Duplex ToBRFV (Cq) | Single assay ToMMV M&W | Duplex ToBRFV (Cq) | ToMMV (Cq) |
|--------|------------|------------|-------------------------|--------------------|-------------------------|--------------------|------------|
| BPeMV  | DSMZ       | BN-4708    | nt                      | nt                 | NA                      | NA                 | NA         |
|        |            |            | nt                      | nt                 | NA                      | NA                 | NA         |
|        |            |            | NA                      | NA                 | NA                      | NA                 | NA         |
|        |            |            | NA                      | NA                 | NA                      | NA                 | NA         |
| CGMMV  | NIB        | NIB V 271  | NA                      | NA                 | NA                      | NA                 | NA         |
|        |            |            | NA                      | NA                 | NA                      | NA                 | NA         |
|        |            |            | NA                      | NA                 | NA                      | NA                 | NA         |
| CGMMV  | NIB        | NIB V 320  | NA                      | NA                 | NA                      | NA                 | NA         |
|        |            |            | NA                      | NA                 | NA                      | NA                 | NA         |
|        |            |            | NA                      | NA                 | NA                      | NA                 | NA         |
| CGMMV  | DSMZ       | PV-0375    | nt                      | nt                 | NA                      | NA                 | NA         |
|        |            |            | nt                      | nt                 | NA                      | NA                 | NA         |
| ObPV   | DSMZ       | PV-1176    | 38.4*                   | NA                 | NA                      | NA                 | NA         |
|        |            |            | NA                      | NA                 | NA                      | NA                 | NA         |
| ORSV   | DSMZ       | PV-1048    | NA                      | NA                 | NA                      | NA                 | NA         |
|        |            |            | NA                      | NA                 | NA                      | NA                 | NA         |
| PaMMV  | DSMZ       | PV-0606 (2020)** | 33.9 | 33.5 | 31.5 | nt | nt | nt |
|        |            |            | 33.8 | 33.5 | 31.4 | nt | nt | nt |
| PaMMV  | DSMZ       | PV-0606 (2021)** | 33.8 | 34.2 | 29.6 | nt | nt | nt |
|        |            |            | 34.0 | 33.7 | 29.3 | nt | nt | nt |
| PMMoV  | CREA       | CREA-552   | nt                      | nt                 | nt                      | NA                 | NA         |
|        |            |            | nt                      | nt                 | nt                      | NA                 | NA         |
| PMMoV  | DSMZ       | PV-0165    | nt                      | nt                 | nt                      | NA                 | NA         |
|        |            |            | nt                      | nt                 | nt                      | NA                 | NA         |
| RMV    | DSMZ       | PV-0145    | 33.9 | 36.9 | NA   | nt | nt | nt |
|        |            |            | 34.4 | 35.2 | NA   | nt | nt | nt |
| SFBV   | DSMZ       | PV-1058    | NA                      | NA                 | NA                      | 38.2*              | NA         |
|        |            |            | NA                      | NA                 | NA                      | nt                 | nt         |
|        |            |            | NA                      | NA                 | NA                      | nt                 | nt         |
| Strain  | Source | Code     | Cq   | Cq   | Cq   | Cq   |
|---------|--------|----------|------|------|------|------|
| SHMV    | DSMZ   | PV-0156  | NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
| ToMV    | NIB    | NIB V 036| NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
| ToMV    | NIB    | NIB V 049| NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
| ToMV    | NIB    | NIB V 072| NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
| ToMV    | NIB    | NIB V 104| NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
| ToMV    | DSMZ   | PV-0141  | nt   | nt   | nt   | NA   |
|         |        |          | nt   | nt   | nt   | NA   |
|         |        |          | nt   | nt   | nt   | NA   |
| TMGMV   | DSMZ   | PV-0124  | NA   | NA   | NA   | NA   |
|         |        |          | NA   | NA   | NA   | NA   |
|         |        |          | NA   | NA   | NA   | NA   |
| TMV     | NIB    | NIB V 037| NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
|         |        |          | NA   | NA   | NA   | nt   |
| TMV     | DSMZ   | PV-1252  | nt   | nt   | nt   | NA   |
|         |        |          | nt   | nt   | nt   | NA   |
|         |        |          | nt   | nt   | nt   | NA   |
| TMV     | DSMZ   | PV-0137  | NA   | NA   | NA   | NA   |
|         |        |          | 38.5*| NA   | NA   | NA   |
|         |        |          | NA   | NA   | NA   | NA   |
| TMV     | DSMZ   | PV-0943  | NA   | NA   | NA   | NA   |
|         |        |          | NA   | NA   | NA   | NA   |
| YMoV    | DSMZ   | PV-0527  | NA   | NA   | NA   | NA   |
|         |        |          | NA   | NA   | NA   | NA   |

NA-no exponential amplification curve; nt-not tested; *Cq above the Cq threshold and considered as negative (no clear exponential amplification curve); **two batches of isolate were tested (one purchased from DSMZ in 2020 and one in 2021).

The inclusivity of duplex assay was assessed using different ToBRFV and ToMMV isolates (Supplementary Table S1 and Table 5). In all cases, the identification of ToBRFV and
ToMMV was successful. In addition, the \textit{in-silico} analysis versus all the full-genome sequences of ToMMV available in GenBank highlighted only up to two nucleotide polymorphisms in forward and/or reverse primer (Supplementary Figure S1). However, these single point mutations were observed in a group of isolates including the DSMZ isolate PV-1267 that was shown to be successfully detected (Table 5).

Table 5. Results of testing different isolates of ToMMV and ToBRFV with duplex ToBRFV&ToMMV assay and with single ToBRFV M&W assay

| virus          | collection         | ID      | ToBRFV M&W (Cq) | ToBRFV M&W (Cq) | ToMMV (Cq) |
|----------------|--------------------|---------|-----------------|-----------------|------------|
| ToMMV         | DSMZ               | PV-1267 | 36.9*           | NA              | 8.4        |
|                |                    |         | 38.9*           | NA              | 8.0        |
|                |                    |         | 37.6*           | NA              | 8.1        |
| ToMMV         | IBMCP; Spain       | S1      | nt              | NA              | 12.7       |
|                |                    |         | nt              | NA              | 12.8       |
|                |                    |         | nt              | NA              | 12.9       |
| ToMMV         | IBMCP; Spain       | S2      | nt              | NA              | 15.9       |
|                |                    |         | nt              | NA              | 15.7       |
|                |                    |         | nt              | NA              | 15.7       |
| ToBRFV        | CREA               | MR50 (100,000 x dilution) | 25.5 | 25.7 | NA |
|                |                    |         | 25.3 | 26.0 | NA |
| ToBRFV        | Volcani center; Israel | S21    | nt              | 6.7             | NA         |
|                |                    |         | nt              | 6.8             | NA         |
|                |                    |         | nt              | 5.1             | NA         |
| ToBRFV        | Volcani center; Israel | S22    | nt              | 6.8             | NA         |
|                |                    |         | nt              | 4.7             | NA         |
|                |                    |         | nt              | 4.6             | NA         |

NA-no exponential amplification curve; nt-not tested; *Cq above the Cq threshold and considered as negative (no clear exponential amplification curve)

2.2.3. Selectivity
We have shown that the duplex assay developed in this study can be used to detect and identify ToBRFV and ToMMV in leaf and seed matrices (Table 3). In addition, we have shown that this assay can reliably detect ToBRFV and ToMMV in seeds of different cultivars (see section 2.3).

2.2.4. Repeatability and reproducibility
The repeatability of the duplex assay was evaluated by analyzing three replicates of RNA samples containing various concentrations of ToBRFV or ToMMV in the same run. Repeatability within each sample was 100%, with standard deviation (SD) of the mean Cq values obtained always less than 1.5 Cq for the high, medium, and low quantities of target
RNA (data not shown). Reproducibility also proved to be 100% (SD below 1.5 Cq; Table 6). Reproducibility was analyzed for two dilutions of a ToBRFV- and ToMMV-positive RNA samples, with medium and low target concentrations. The repeatability and reproducibility were assessed at both CREA and NIB, analyzing different target RNA samples and using different reagents (see Materials and Methods section). In each laboratory, different real-time RT-PCR runs, 4 (CREA) and 6 (NIB), were performed on different days. In addition, at NIB two different instruments were used (see Materials and Methods section).

**Table 6.** Results of testing medium and low concentration of targets and negative amplification controls with duplex ToBRFV&ToMMV assay at NIB and at CREA on different days using different instruments, technicians and reagents.

| run ID samples | ToBRFV (Cq) | ToMMV (Cq) | ToBRFV (Cq) | ToMMV (Cq) | ToBRFV (Cq) | ToMMV (Cq) | ToBRFV (Cq) | ToMMV (Cq) | ToBRFV (Cq) | ToMMV (Cq) |
|----------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|-------------|
| NIB            |             |             |             |             |             |             |             |             |             |             |
| ToBRFV PK1     | 25.6 NA     | 26.7 NA     | 24.7 NA     | 25.6 NA     | 25.3 NA     | 25.5 NA     |             |             |             |             |
| ToBRFV PK2     | 31.8 NA     | 32.5 NA     | 31.2 NA     | 30.3 NA     | 31.0 NA     | 30.4 NA     |             |             |             |             |
| ToMMV PK1      | NA 26.2 NA  | NA 26.6 NA  | NA 25.7 NA  | NA 27.8 NA  | NA 27.2 NA  | NA 27.3 NA  |             |             |             |             |
| ToMMV PK2      | NA 32.9 NA  | NA 33.0 NA  | NA 32.9 NA  | NA 35.4 NA  | NA 34.2 NA  | NA 35.6 NA  |             |             |             |             |
| NAC1           | NA NA NA NA | NA NA NA NA | NA NA NA NA | NA NA NA NA | NA NA NA NA |             |             |             |             |
| NAC2           | NA NA NA NA | NA NA NA NA | NA NA NA NA | NA NA NA NA | NA NA NA NA |             |             |             |             |
| CREA           |             |             |             |             |             |             |             |             |             |             |
| ToBRFV MR50(1) | 23.4 NA     | 23.6 NA     | 23.3 NA     | 23.4 NA     | nt nt nt nt |             |             |             |             |
| ToBRFV MR50(2) | 30.8 NA     | 30.8 NA     | 31.1 NA     | 31.0 NA     | nt nt nt nt |             |             |             |             |
| ToMMV-1        | NA 24.4 NA  | NA 24.9 NA  | NA 24.8 NA  | NA 25.2 NA  | nt nt nt nt |             |             |             |             |
| ToMMV-2        | NA 31.4 NA  | NA 31.2 NA  | NA 31.2 NA  | NA 32.0 NA  | nt nt nt nt |             |             |             |             |
| NAC1           | NA NA NA NA | NA NA NA NA | NA NA NA NA | NA NA NA NA | nt nt nt nt |             |             |             |             |
| NAC2           | NA NA NA NA | NA NA NA NA | NA NA NA NA | NA NA NA NA | nt nt nt nt |             |             |             |             |

NA-no exponential amplification curve; nt-not tested; NAC-negative amplification control; ToBRFV PK1 – MR50 (1) 10^5 dilution; ToBRFV PK2 – MR50(2) 10^6 dilution; ToMMV PK1 – PV-1267 10^6 dilution; ToMMV PK2 – PV-1267 10^4 dilution; ToMMV-1 – PV-1267 10^4 dilution; ToMMV-2 – PV-1267 10^4 dilution.

2.3 Use of the validated duplex assay in diagnosis of seed samples

To further assess the feasibility of the duplex assay developed and validated in this study for a rapid and sensitive detection of ToBRFV and ToMMV in seeds, a wide number of RNA samples previously assayed for ToBRFV was tested with the duplex assay. A total of 106 tomato and pepper official seed samples from 2021 (RNA extracted from 62 tomato seed samples and from 44 pepper seed samples) from different origins (China, Italy, Brazil, India, Slovenia, Serbia) and previously analyzed for ToBRFV at CREA or at NIB, were included (Supplementary Table 2). In all these samples, the absence/presence of ToBRFV was confirmed with 100% concordant results; all 3 tomato and all 5 pepper seed samples, previously found to be ToBRFV positive, were also positive confirmed with the duplex ToBRFV & ToMMV assay. Regarding ToMMV, fifteen seeds official samples
resulted with Cq values in a range from 23.1 to 37.7 for ToMMV (Supplementary Table 2). All these 15 samples were further analyzed by generic tobamovirus nested PCR [40] and by Sanger sequencing of the nested PCR products; and the presence of ToMMV was clearly confirmed in five samples in which the Cq values of ToMMV ranged from 23.1 to 29.0. In samples with Cq values for ToMMV of 29.7 to 37.7, no nested PCR product or pure sequence was obtained, or the presence of another tobamovirus (ToMV) was confirmed (Supplementary Table 2). The origin of all seed samples in which the presence of ToMV and ToMMV have been confirmed is China.

3. Discussion

The development of diagnostic techniques that are as rapid, effective, and sensitive as possible is one of the most important issues in the controlling of plant viruses. In particular, regulated and/or emerging plant viruses, could indeed pose a major threat to most crops, and the information gained from rapid detection and identification is important for determining any measures needed to prevent the introduction of these pathogens and/or limit their spread. In view of this, it is essential to have reliable diagnostic tests. Recently, ToBRFV highlighted how easily viruses belonging to tobamovirus genus can pose a phytosanitary risk, especially for tomato and pepper crops. For this reason a prompt test performance studies (TPSs) organized under several national and transnational initiatives/European projects, as well as the publication of the EPPO standard PM7/146 (1), has been providing valid tools for the establishment of ToBRFV containment measures, included also in Commission Implementing Decision (EU) 2020/1191 and subsequent amendments.

Recently, the increasing number of reports about the presence of ToMMV raised the attention to this tobamovirus, currently included in EPPO alert list, from 2020. The experience gained with ToBRFV should provide the guidelines for managing the threat of this emerging virus. The lack of a method such as real-time RT-PCR test for ToMMV, is the major diagnostic gap, which this study aims to address. In particular, in this study, a real-time RT-PCR test was developed for the detection and identification of ToMMV, which can also be used in a duplex assay with primers/probe set of a test already validated and recommended in the EPPO standard PM7/146(1) for ToBRFV detection (i.e., M&W) [22].

The ToMMV assay developed in this study targeting the MP region confirmed that this region could be a suitable target for primers/probe set design. This region, provides sufficient variability to distinguish between species of the genus Tobamovirus, which are known to have a high degree of homology [42]. In fact, the other pre-tested sets developed in this study and targeting CP region, reacted unspecifically with CGMMV, a tobamovirus that infects Cucurbitaceae plant species, and a very limited number of Solanaceae species [43].

The ToBRFV&ToMMV duplex assay (Duplex B) included in the validation process according to EPPO standard PM 7/98 (4) was evaluated in terms of analytical sensitivity and specificity, selectivity, repeatability, and reproducibility. The LODs obtained for both leaf
(10^4/10^5) and seed (10^5) matrices and for both target viruses were in line with those of other real-time RT-PCR tests for the detection of tobamovirus species [22,44,45]. In addition, the performance criteria when testing dilution series (leaf or seeds samples) showed that there was no relevant reduction in LOD and/or primer efficiency (E) for the duplex assay compared to the single assays. The standard curves obtained in leaf and seed samples are similar, indicating equal slope and efficiency. The effectiveness of ToBRFV detection and identification in leaf and seed samples is not affected by either the ToBRFV M&W test alone or the duplex assay. This comparison is essential as the ToBRFV M&W test has already been validated (TPS under the H2020 Valitest and Euphresco 2019-A-327 projects) and is one of the two mandatory assays for ToBRFV detection in seeds (Commission Implementing Decision (EU) 2020/1191).

In addition, the duplex assay showed optimal analytical specificity on both, inclusivity, and exclusivity. Twenty-two non-target isolates of tobamovirus species were tested and the only relevant cross-reaction was obtained for paprika mild mottle virus – PaMMV (Table 5). Indeed, the high Cq values (29.3-35.0) for both ToMMV and ToBRFV (the latter in the single and duplex assay), could represent an unspecific reaction that should be considered, since PaMMV shares common hosts (i.e., peppers) with them. Nevertheless, in silico analysis (data not shown) did not reveal sequence identity with both primers/probe sets that would warrant a cross-reaction. Moreover, PaMMV is reported to have sequence homology of about 64% with different ToMMV isolates [42]; in view of this, cross-reaction between the two targets does not seem likely to occur. Similar to PaMMV, cross-reactivity of RMV with ToBRFV in single and duplex assays is not expected due to sequence homology. Nevertheless, the samples of non-target tobamoviruses included in this study are from artificially inoculated test plants. A high virus concentration is expected in these samples; such a virus titer is hardly found in naturally infected leaf samples and not at all in seed samples, but further analysis and investigation are required.

The results obtained for inclusivity show that the duplex assay is able to detect all known ToBRFV and ToMMV isolates. In support of the inclusivity data (tested on few isolates from different sources), in silico analysis showed that the 14 different ToMMV whole genome sequences had very low variability, suggesting low selection pressure for this virus. In addition, point mutations were found in regions targeting forward and reverse primers (not the probe). However, these mutations occurred at the 5’/3’ ends of the primers, and in the DSMZ PV-1267 ToMMV isolate used throughout the development of this assay. This confirms that the proposed test is capable to identify ToMMV and detecting all its isolates.

Results of repeatability and reproducibility, showing a standard deviation between tested samples less than 1.5 Cq, confirm that this duplex assay can be used for the detection and identification of ToBRFV and ToMMV with a high degree of confidence. As these results were obtained independently by two laboratories (CREA and NIB) on different days and with different reagents, operators and equipment, the associated validation data can be considered robust.

As mentioned above, ToMMV is added in the EPPO alert list and is considered an emerging pathogen. As with ToBRFV, seed testing is important to prevent the spread of plant
viruses, especially viruses that, like tobramoviruses, arise through mechanical/contact transmission. Recent reports of ToMMV occurrence, particularly in pepper seeds imported to Australia in 2020 [32] and in batches of tomato seeds imported by companies in Spain between 2019 and 2021 [33], confirm the importance of seed testing to ensure free movement without phytosanitary risks. Since all these reports used an endpoint RT-PCR, it is likely that the presence of ToMMV could be underestimated.

In view of the above, this study focused on the detection of ToBRFV and ToMMV on seeds. One hundred and six tomato/pepper official seed samples previously tested for ToBRFV at CREA or NIB were assayed using the duplex assay developed in this study. Regarding ToBRFV, data showed 100% concordance with previous analysis. The results of the 106 official seed samples showed that 15 samples taken during the inspection of seed lots imported from China gave some positive signals for ToMMV, where we could confirm the presence of ToMMV in five samples by Sanger sequencing of amplicons belonging to the region RdRp (Supplementary Table S2) and in addition, one of these samples was tested and the presence of ToMMV was also confirmed by Sanger sequencing of amplicon belonging to region CP (data not shown). These results suggest that the spreading of ToMMV is underestimated, especially on seeds, confirming what reported by [33], and that more accurate testing should be done on seeds, especially those imported from third countries.

Samples confirmed positive for ToMMV by Sanger sequencing of amplicons yielded duplex real-time RT-PCR Cq values ranging from 23.1 to 29.0 (supplemental Table S2). One sample with a Cq value of 29.7 is thought to have a mixture of ToMMV and ToMV, whereas all other nine samples that did not show ToMMV on Sanger sequencing had Cq values between 31.7 and 37.7. Due to the expected lower sensitivity of nested RT-PCR compared to the sensitivity of the real-time RT-PCR and since the nested RT-PCR used is based on degenerate primer sets targeting the tobramovirus genus, and the species titers in mixed infections could be different, the presence of a low ToMMV titer in such samples is also possible, including hypothetical contaminations. Taking all results together, including the exclusivity tests, signals with high Cq values for ToMMV and ToBRFV (in both, the single and duplex ToBRFV (M&W) assays) are signals that should be further evaluated.

4. Material and methods

4.1. Plant Material and RNA extraction

The samples (leaf and seed matrices) used in this study are listed in Supplementary Table 1 including: i) isolates already characterized and included in CREA (Council for Agricultural Research and Economics, Research Centre for Plant Protection and Certification) and NIB (National Institute of Biology) collections; ii) isolates purchased from DSMZ (German Collection of Microorganisms and Cell Cultures GmbH) or kindly provided by other institutions; iii) RNAs extracted from seeds official samples previously analyzed for ToBRFV at CREA or at NIB (Supplementary Table 2).
The total RNAs used for the assay set-up and validation were extracted according to the RNA extraction methods for leaves and seeds reported in the ToBRFV EPPO Standard PM 7/146 (1) [22]. Total RNAs yield and quality were assessed by NanoDrop ND1000 spectrophotometer (Thermo Fisher, Italy), or real-time RT-PCR amplifying nadS [46] or cytochrome oxidase [47] were used as controls to evaluate the quality of the RNA in the extractions (data not shown). Buffer controls were included with all isolations (i.e. negative isolation control), to monitor potential contamination through the extraction procedures.

4.2 Primer design and reaction set up

The reference full genome sequence of ToMMV (NC_022230) was retrieved from GenBank and used as template for TaqMan® probe and primer set design by Primer Express® Software Version 3.0 (Applied Biosystems, USA) with default parameters. In particular, two different regions coding for coat protein (CP) and movement protein (MP), were included in the design since they represent target regions for the two validated real-time RT-PCR tests available for ToBRFV detection [22]; Menzel and Winter test (M&W) targeting a fragment from the end of the CP gene to the middle of 3-NTR ) [22], and International Seed Federation-International Seed Health Initiative for Vegetable Crops (ISHI-Veg) test being a duplex assay involving both CP and MP regions [22].

The designed TaqMan® probe and primers sets were synthesized by Eurofins genomics (Kolhn, Germany) and labelled in HEX and/or Texas Red according to the correct match with the fluorophore used in ToBRFV detection tests based on real-time RT-PCR (i.e. M&W, FAM-labelled, and ISHI-Veg, FAM/HEX-labelled). Real-time RT-PCR reactions were performed using the CFX96 optical reaction module with C1000 Thermalcycler (BioRad, Milan, Italy). Preliminary reactions were performed including two negative (healthy leaf and seed samples), one ToMMV-positive (ToMMV isolate PV-1267 from DSMZ), one ToBRFV-positive (ToBRFV isolate MR50 – leaf), and one negative amplification control, in two technical repetitions. These samples were tested at different primers and probe concentrations as well as annealing temperature ranging from 55 °C to 62 °C (data not shown). Once determined the optimized amplifications conditions, reactions were carried out in 10 µl reaction volumes containing 5 µl of 2× Mastermix, 0.25 µl of 40X RT Enzyme (both from TaqMan® RNA-to-Ct™ 1-Step Kit, Thermo Fisher,Milan, Italy), 0.3 µM of each primer, 0.2 µM of labeled TaqMan® probe, and 1 µl of template RNA (0.4–43 ng). The optimized one step real-time RT-PCR cycling conditions included a RT step at 48 °C for 30 min, an initial denaturation step at 95 °C for 10 min followed by 40 cycles of denaturation and annealing/elongation for 15 s at 95 °C and 1 min at 60 °C, respectively. Analyses were performed using the BIORAD CFX Maestro v2.2. For assessing exclusivity, each ToMMV primers/probe set was tested including the following non-target isolates from DSMZ collection: TMV (PV-0137), ToMV, tobacco mild green mosaic virus - TMGMV, bell-pepper mosaic virus - BPeMV, pepper mild mottle virus - PMMoV, cucumber green mottle mosaic virus – CGMMV (PV-0375), retrieved from DSMZ collection (Supplementary Table 1). Further, ToMMV primers/probe sets were evaluated in duplex/triplex assay with the corresponding ToBRFV test (M&W or ISHI-Veg), according to targeting region and/or labelling. The duplex/triplex assays were evaluated on a sample set composed of healthy leaves/seeds controls, leaves/seeds samples infected by ToBRFV (ascertained to
be in single infection), ToMMV leaves sample and a mixed ToMMV and ToBRFV infected sample (prepared by mixing the same amount (0.05g) of freeze-dried material of ToBRFV MR50 isolate and ToMMV PV1276 isolate. Based on the general quality assessment, i.e., amplification curves patterns, value of quantification cycle (Cq), concordance of relevant controls, amplification background signals and performance in the duplex/triplex with ToBRFV tests taking into major account the different primers/probe set targeting region, the best ToMMV primers/probe set was selected to be further validated as a duplex assay at CREA.

Selected duplex (ToMMV and ToBRFV M&W) real-time RT-PCR was also tested by NIB according to the following mixture reaction (final volume of 10 µl): 2 µl of sample RNA, 0.3 µM of each primer, 0.2 µM of each labeled TaqMan® probe] using the AgPath-ID™ One-Step RT-qPCR mix (Applied Biosystems, Italy). Real-time RT-PCR was carried out in 384-well plates (Applied Biosystems, USA), and the reactions were run on an QuantStudio™ 7 Pro System or Viia7™ System sequence detection (Applied Biosystems, USA). One-step real-time RT-PCR cycling conditions included a RT step at 48 °C for 10 min, an initial denaturation step at 95 °C for 10 min followed by 45 cycles of denaturation and annealing/elongation for 15 s at 95 °C and 1 min at 60 °C, respectively.

The Design & Analysis Software v2.5.1 (Applied Biosystems, USA) was used for fluorescence acquisition and determination of Cq values. Before analysis, ROX™ reference dye was excluded to prevent its interfering with Texas RED dye, and the baseline was set manually. The fluorescence threshold was also set manually, i.e., to a level that was above the baseline and sufficiently low to be within the exponential increase region of the amplification curve. If no exponential amplification curve was produced, a sample was considered negative. If an exponential amplification curve was produced, the Cq were determined. In addition, especially for ToBRFV, a Cq threshold value was determined if needed according to what reported in [22].

4.3. Duplex assay validation

The ToBRFV&ToMMV duplex assay was validated according to EPPO PM 7/98 (4) guidelines. To determine all the criteria included in the validation process, a sample set summarized in Supplementary Table S1 was used.

4.3.1. Analytical sensitivity, limit of detections and efficiencies

Eight ten-fold serial dilutions using both leaf and seed matrices were prepared. To assess the analytical sensitivity in leaf samples, artificially mixed infected samples were created by mixing the same amount of freeze-dried material (0.05g each) derived from leaves infected with ToBRFV (isolate MR50) and ToMMV (isolate PV-1276 from DSMZ). The sample was homogenized with PO4 buffer (pH 7.2) and 100 µl were serially diluted in 900 µl of healthy crude extract (tomato plant), to obtain a total eight 10-fold serial dilutions. Then, for each aliquot of the dilutions series, total RNA was extracted by RNeasy Plant Mini Kit (Qiagen), adding 100 µl to 380 µl of RLT buffer (included in the kit) and following manufacturers’ instructions.

To assess the analytical sensitivity in seed samples, two ToBRFV infected seeds (CREA collection) and 5 mg of freeze-dried leaf material infected by ToMMV (PV-1276 from DSMZ) were spiked in 998 tomato healthy seeds. The material was transferred in a
grinding bag (BIOREBA XL), soaked in GH+ buffer, as reported in [22], for 1h at room temperature, and homogenized with an Interscience BagMixer (position 4) for 90s. Eight ten-fold serial dilution were prepared (100 µl of homogenised infected sample in 900 µl of homogenised healthy tomato seeds) and following instructions reported in [22]. All the aliquots from the serial dilution for each matrix were assayed by the duplex assay for ToMMV and ToBRFV.

For both matrices (leaf and seeds) the limit of detection (LOD) was determined, the amplification efficiencies (E) calculated, and all the results obtained compared. In detail, standard curves were obtained by plotting mean Cq values of the ten-fold serial dilutions (leaves and seeds) versus minus the logarithm of the RNAs dilution factors. The Cq values and the following equation were used to determine efficiency (E) of each primers/probe set with the slope of a linear regression model; the linear correlation coefficient (R²) was also reported.

\[ E(\%) = (10^{\frac{1}{slope}} - 1) \times 100 \]

The same dilution series have been tested and the efficiency calculated also for cases where selected ToMMV primers/probe set and ToBRFV assay were used as single assay.

4.3.2. Analytical specificity

A total of 13 different tobamovirus species were assayed at CREA and NIB as relevant non-target (Supplementary Table S1); for some tobamovirus species (TMV, ToMV) commonly present in tomato and/or pepper, different isolates were included. The inclusivity was assessed using several ToBRFV and ToMMV isolates according to different geographic origins and hosts. In addition, the inclusivity results were further widened by in silico analysis, by aligning (Clustal W supported by MEGA X software [48]) the duplex assay primers/probe sets versus all the full genome sequences of ToMMV and ToBRFV retrieved in GenBank (Supplementary Table S3).

4.3.3. Selectivity

To determine whether variations of the sample material could affect the test performance or not, different cultivars of host plant and different matrices (leaf and seeds) were tested.

4.3.4. Repeatability, reproducibility

These performance parameters were tested by performing technical replicates and repeating the assay in two laboratories (i.e., CREA and NIB), with different operators, in different days and on different instruments; deviations due to the use of different reagents were also checked.

5. Conclusion

The ToBRFV&ToMMV real-time RT-PCR duplex assay developed in this study was found to be reliable for use in routine analyses, according to international validation guidelines. Moreover, this aspect appears to be of fundamental importance for seed testing, where the requirements for a sensitive and reliable diagnostic tool are crucial given the phytosanitary risk posed by the free movement of seeds worldwide. Since testing of seeds for ToBRFV is mandatory in most countries, the development of a duplex assay for two
tobamoviruses as presented in this study could provide an effective diagnostic tool for testing imported seeds, especially for those countries where ToMMV must also be tested (e.g., Australia). A rapid and sensitive diagnostic test is essential for the introduction of containment measures to protect tomato and pepper crops.

**Supplementary material**

**Supplementary figure S1.** *In-silico* analysis (alignment by Clustal X) of the ToMMV selected primers/probe set (set 3) versus all the full-genome sequences of ToMMV available in GenBank. Highlighted are nucleotide polymorphisms observed in forward and reverse primer sequence.

**Supplementary Table S1.** List of viral isolates/species included in this study

**Supplementary Table S2.** List of official samples assayed by the duplex assay developed and validated in this study.

**Supplementary Table S3.** List of tomato mottle mosaic virus isolates which genome sequences were retrieved in GenBank and included in the *in-silico* inclusivity analysis.

**Author Contributions**

A.Ti., conceived and designed the research. A.Ti., A.M., N.M., and J.B. conducted experiments. A.Ti, A.Ta., and N.M. wrote the manuscript. M.L., L.F., and A.G. revised the manuscript. All authors approved the publication. All authors have read and agreed to the published version of the manuscript.

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**Conflicts of Interest**

The authors declare no conflict of interest - Ethical statement. This research did not involve any animal and/or human participant. The authors declare that they have no conflict of interest.

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