

Introduction

Colombia is one of the world's main producers and exporters of tropical fruits, however, it lacks a robust preventive management programme for the control of plant viruses¹.

Recent Next Generation Sequencing (NGS) according to Massart et al. (2017)⁴ revealed 57% identity to the movement protein (*MP*) of *Lilac ring spot virus* (*Jambava*), and 65% identity to the replicase of the *Tobacco mosaic virus* (*Jambava*) which mean drastic reduction of crop yields and major economic losses. For this purpose, molecular biological tools such as nucleic acid isolation, polymerase chain reaction (PCR), and NGS are being used to measure the frequency and distribution of this virus in Colombia, to characterize which symptoms are associated with it, and to identify the pathways for its transmission.



Figure 1. Maps of Colombia, where the new *Santivania* in purple passion fruit was found, and the region where the samples were taken from (a,b). Purple passionfruit field in Chocó and Cauca, Colombia (c).

Materials and Methods

- Samples of *R. solani* plant material with virus-suspected symptoms were collected in 2016 and 2017 from 5 different farms in the Altiplano Cundiboyacense Andean plateau (Cundinamarca and Boyaca departments, Colombia). 97 samples from these 5 farms, were used in this study.
- RNA isolation according to Goorn et al. (1990) from leaf material
- cDNA synthesis with pMLMV RTase (beach material) or Premium ReverseAid RTase (fresh material) and random hexamer primers
- nested-PCR for quality control of RNA and cDNA synthesis according to Menzel et al. (2002)²¹
- Diagnostic RT-PCR for the detection of the novel luteivirus with primers developed for its RNA1, RNA2, and RNA3
- RT-PCR for viral-specific detection of the CP coding region of RNA3 of the luteivirus

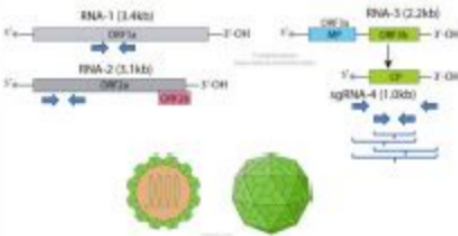


Figure 1. Genome structure of the *Saccharomyces* genus, with a separate tree with *S. (C)* genomes. The positions of the developed primers are shown by blue arrows. For the *RGA5* primers, the different combinations for amplification of the CP region by PCR are shown (a). Morphology of the heterokaryons (b). Figure adapted from www.mycosynth.com.

Results



Figure 3. Healthy carriers of three asymptomatic symptoms associated with the new hepatitis isolate by RT-PCR.

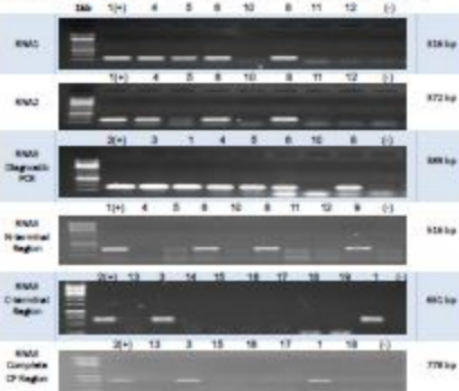


Figure 3. Detection of the new variants by means of RT-PCR, using designed primers based on the previously obtained NGS analysis. RNP1, RNP2 and RNP3 diagnostic PCR, and primers targeting RNP3 5' terminal region, RNP3-5'UTR-terminal region, and RNP3-5'UTR-TPR domain. In all PCR, one positive control (1), one negative control (2), one no-template control (3) and water (4) were included.

Table 1. Results from the detection of the new SARS-CoV by means of diagnostic RT-PCRs, using RNA1, RNA2, and RNA3 primers.

Symptom	Bleeding	Hematuria	Cramps	Waking	Maternal Deformities	Maternal/Infantile Cramps	Bleeding/Waking	Healthy
Gynecological symptoms with irregularities	15	7	3	18	6	21	11	6
RNA1	1	0	0	1	4	2	1	0
RNA2	1	0	0	0	4	2	1	0
RNA3	1	0	0	1	4	2	1	0
Diagnosis								
RNA2-B-terminal Region	1	0	0	1	4	2	1	0
RNA2-C-terminal Region	1	0	0	0	3	2	1	0
RNA2-C-terminal CP Region	1	0	0	0	3	1	0	0

Conclusions

The novel lavirus found in purple passion fruit (*Passiflora edulis Sims*) in Colombia can be detected by RT-PCR, using different types of primer combinations of the 3 RNAs of its genome. Therefore, this could be a useful tool to detect the presence of the virus in the passion fruit.

The virus could be detected in samples showing blistering, mottling, blistering/deformation, and blistering/deformation/ichthiosis symptoms, as seen in Table 1. The coat protein (CP) gene is generally more variable. Therefore, the characterisation of the virus, is best done by amplification of the N-terminal and C-terminal regions of the CP gene (Invitrogen, PE-RT-CP).

References