# CIDER-Seq: unbiased virus enrichment and single-read, full length genome sequencing 

Devang Mehta*, Matthias Hirsch-Hoffmann, Andrea Patrignani ${ }^{1}$, Wilhelm Gruissem and Hervé Vanderschuren**<br>Institute of Molecular Plant Biology, Department of Biology, and ${ }^{1}$ Functional Genomics Center Zurich, ETH Zurich, 8092 Zurich, Switzerland<br>\#Current address: AgroBioChem Department, University of Liège, Gembloux, Belgium<br>*Corresponding authors: devang@ethz.ch \& herve.vanderschuren@ulg.ac.be


#### Abstract

Deep-sequencing of virus isolates using short-read sequencing technologies is problematic because the viruses are often present in populations with high sequence similarity. We present a new method for generating single-read, full-length virus genomes by combining an improved Random Circular Amplification (RCA)-based virus enrichment protocol with SMRT-sequencing and a new sequence de-concatenation method. We demonstrate that CIDER-Seq produces more than 250 full-length geminivirus genomes from symptomatic field-grown plants.


Advances in sequencing technologies have produced extensive data about viral diversity and facilitated the identification of previously unknown viruses. The abundance of new virus sequence data recently led to the consensus that virus taxonomy should be revised by incorporating metagenomic sequence data ${ }^{1}$. However, using high-throughput sequencing technologies for virus detection and accurate identification remains difficult because of risking the assembly of artificial chimeric viral genomes from short sequence reads.

Sequence-bias introduced during virus enrichment prior to sequencing is equally problematic for deep-sequencing of virus genomes. Amplification methods rely on either Polymerase Chain Reaction (PCR) with primers designed to bind conserved sequences in the genome, or random circular amplification (RCA) utilizing the unique properties of Phi29 DNA polymerase and randomnucleotide primers ${ }^{2}$. PCR-based methods can result in a biased amplification of viral templates because of differing primer-template affinities ${ }^{3}$. Random primer-based RCA is less prone to primer complementarity bias but results in hyper-branched, high molecular weight concatenated products ${ }^{4}$ that must be linearized with restriction enzymes (REs) ${ }^{5}$ or mechanical shearing prior to Sanger or next generations sequencing (NGS). Using REs requires prior information on conserved RE sites and therefore results in the loss of viral sequences that may have none or multiple recognition sites for the selected REs. An alternative method called "polymerase cloning" or "ploning" employs endonucleases and DNA repair enzymes to linearize RCA products (hence voiding the need for virus sequence information or specific restriction enzyme cut sites), followed by shotgun sequencing and whole genome assembly.

CIDER-Seq (Circular DNA Enrichment Sequencing) is a sequence-independent viral nucleic acid enrichment method utilizing assembly-free Single Molecule Real Time (SMRT; Pacific Biosciences Inc.) long-read sequencing (Fig. 1a). Our enrichment method requires only an estimate of virus genome size and, optionally, a single reference sequence. CIDER-Seq also enables the direct single molecule sequencing of RCA-derived products. We have also developed a new algorithm, DeConcat, to parse concatenated DNA strands that are generated by RCA reactions, allowing us
a.


1. Initial size-selection

2. Random Circular Amplification
r



3. SMRT Library prep (barcoded)

4. Library size-selection
5. Branch Release

6. Debranching

1
b.


Figure 1: Circular DNA enrichment scheme and validation. (a) Enrichment of circular DNA based on automated size selection, non-denaturing random circular amplification (RCA) and linearization and repair of the RCA product followed by Single Molecule Real Time (SMRT) library creation. SMRT libraries were optionally size-selected prior to sequencing on a PacBio RS II instrument. (b) Semi-quantitative PCR on a sample testing permutations of the three enrichment steps. Manihot esculenta PROTEIN PHOSPHATASE 2A (MePP2A)-specific primers were used to determine the amount of linear cassava genomic DNA compared to enriched circular viral DNA amplified with cassava geminivirus-specific primers.
to sequence complete virus genomes without reference-based or de novo assembly. To validate our approach, we sequenced the cassava mosaic geminivirus (CMG), which causes cassava mosaic disease (CMD) and severe economic losses for farmers, particularly in sub-Saharan Africa. To date, nine CMG species have been identified that share $68 \%$ to $90 \%$ sequence identity based on Sanger sequencing of PCR and RCA products ${ }^{7}$. CMGs are bipartite viruses of the genus

Begomovirus in the family Geminiviridae - the most populous family of eukaryote-infecting viruses ${ }^{8}$. They are comprised of two separate genomes (designated DNA A and DNA B) ${ }^{9}$, and thus serve as a good model to validate the functionality of our method for segmented viruses (Fig. 1a, Steps 18).

Using samples from cassava plants with CMD symptoms we first enriched complete DNA genomes of CMGs by automated size selection of DNA molecules in the 2.8 kb size range (Step 1). Phi29 DNA polymerase RCA of the selected DNAs was carried out using random hexameric primers (Step 2). Using a 'ploning' based protocol, we amplified the products of the initial amplification in an additional primer-free Phi29 DNA polymerase reaction to further "de-branch" the hyper-branched DNA ${ }^{6}$ (Step 3). Next, the hyper-branched structure was resolved using ssDNA-digesting S1 Nuclease (Step 4) and repaired with DNA Polymerase I and T4 DNA Polymerase (Step 5). The linear DNA fragments were purified using magnetic beads (Step 6), which excluded small DNA fragments produced during the RCA steps.

To optimize the enrichment, we performed a semi-quantitative PCR (semi-qPCR) with 15,25 and 40 cycles to quantify viral DNA titers relative to host DNA before and after Steps 1-3 (Fig. 1b). The semi-qPCR revealed viral and host amplicon DNA bands at the expected size but also longer RCAspecific DNA bands. Maximum enrichment was obtained after Steps 1-3 (Fig. 1b, highlighted) and avoidance of the denaturation step common to most RCA protocols. The debranching step catalyzed by S1 nuclease also reduced the abundance of genomic DNA in favor of a more distinct viral RCA DNA product.


Figure 2: Data analysis method and results. (a) Overview of the data analysis method including a simplified depiction of the DeConcat algorithm. (b) Distribution of the BLAST-based binned sequencing reads from the non-size selected (NSS) and size selected (SS) libraries. (c) Length distribution of sequencing reads before and after DeConcat for NSS and SS libraries.

The enrichment was applied to total DNA extracts from six independent leaf samples collected from a cassava field trial in Kenya. The six samples were barcoded and pooled to build a SMRT sequencing library. We sequenced the library in two independent sequencing runs: an intact library without size selection (NSS), and a size-selected (SS) library (to maximize the number of reads obtained from inserts greater than 3 kb ) (Step 8, Fig. 1a). We then filtered the raw sequencing reads for high-quality insert sequences (predicted minimum quality of 99.9 and minimum insert length of 3000 bp ). The SS library produced 275 high quality reads of insert (ROIs) of the expected size and quality range, while the NSS library produced only 112 . As expected, a majority of ROIs were greater than the 3 kb size cut-off, indicating that the inserts were comprised of concatenated RCA products.

We developed a custom data analysis pipeline to resolve the concatenated sequences into their component parts (Fig. 2a). First, in a basic filtering step all ROIs were binned into non-viral, CMG DNA A and CMG DNA B sequences. The results from this step indicated that approximately 99\% of the ROIs in both libraries were found in the CMG bins (Fig. 2b). Next, we trimmed non-viral DNA from the ends of each ROI by performing a circular DNA-aware multiple sequence alignment (see Methods), which did not significantly affect the size distribution of the dataset. These trimmed ROIs were next passed through the de-concatenation algorithm we termed 'DeConcat' (Supplementary Fig. 1, see Methods). The resulting de-concatenated ROIs had a greatly reduced size distribution (Fig. 2c), with a clear peak at 2.8 kb . Thus, DeConcat was able to reduce RCA-generated SMRTsequenced ROIs of a wide size-distribution into expected CMG-length virus sequences. This result validated the parameters and scoring formula applied in DeConcat and also demonstrated that the Phi29 DNA polymerase enrichment step amplified circular DNA in the 3 kb range with high fidelity.

We performed tBLASTn using the final DeConcat reads against virus protein reference sequences to annotate reads belonging to viral ORFs. The results revealed frameshift mutations in one or more ORFs in several reads. Frameshift mutations are caused by insertions or deletions, which is the most frequent type of error in SMRT sequencing ${ }^{10}$. To distinguish if these frameshifts were of biological origin or SMRT-sequencing errors, we relaxed the minimum quality threshold of $99.9 \%$ for ROIs in the initial filtering step to $99.5 \%$ and $99 \%$. When comparing the frequency and number of frameshift mutations between the three quality thresholds we found that the number of frameshifts per sequence increased with decreasing quality thresholds (Supplementary Fig. 2a). and the frequency of frameshifts in each viral protein increased linearly with protein length (Supplementary Fig. 2b), indicating that they are likely caused by SMRT sequencing errors. Between 5 to $8 \%$ of reads in the $>99.99$ quality threshold dataset have no frameshift errors (Supplementary Fig. 2c).

We next selected DNA A sequences from the NSS and SS datasets that were longer than 2.7 kb (and therefore likely complete virus sequences) for pairwise alignments using the SDT virus classification software ${ }^{11}$ to analyze the diversity of the CMG sequences. Most DNA sequences belonged to three categories with approximately $75 \%$ identity (Fig. 3a, arrows), while sequences within each category were $>91 \%$ identical. This was also reflected in the frequency distribution of pairwise sequence identities calculated by SDT (Fig. 3b). Since the recently revised Begomovirus taxonomy guideline ${ }^{12}$ classifies sequences with less than $91 \%$ identity as distinct species, we conclude that CIDER-Seq can distinguish two distinct CMG species. We constructed a Maximum Likelihood phylogenetic tree including published sequences of the nine reported CMG species found on the African continent ${ }^{7}$ to determine the species that most closely match our dataset. The cladogram shows that the two major species we detected in the field samples using SDT belong to
the African cassava mosaic virus (ACMV) and the East African cassava mosaic virus (EACMV) clades (Fig. 3c).


Figure 3: Sequence demarcation and phylogenetic analysis of the cassava mosaic geminivirus (CMG) DNA A dataset. (a) Pairwise identity matrix of all full length DNA A sequences ( $>2700 \mathrm{bp}$ ) generated using CIDER-seq. Three species groups are marked by arrows. (b) Frequency plot of pairwise identity (DNA A sequences) showing the taxonomic cut-offs for species and isolate demarcation. (c) Unrooted cladogram constructed from a maximum likelihood tree of all DNA A sequences with reference DNA A genomes of the 9 CMG species found on the African continent. Coloured clades mark sequences closest to respective reference species. Nodes marked in blue are supported by $>70$ bootstraps and the node sizes are scaled from 70-100 bootstrap values.

Together, CIDER-seq effectively enriches circular DNA molecules and produces single-read, fulllength DNA sequence data from the RCA reaction products. The DeConcat algorithm parses the concatenated reads of the RCA products into individual component DNA sequences of the appropriate size range without training the algorithm with prior information of desired sequence length. Considering the popularity of Phi29 DNA polymerase-based amplification methods such as MDA or MALBAC in single-cell genome sequencing protocols ${ }^{13}$, DeConcat can increase analysis quality of concatenated sequences from single-cell DNA samples as well.

We have validated CIDER-seq for deep-sequencing of Geminiviruses, which is an important family of plant-infecting viruses that is causing increasing economic losses to farmers ${ }^{14}$. Using infected field material we could generate 270 high-quality, non-chimeric full-length virus genome
sequences, representing nearly $50 \%$ of all CMG sequences deposited in GenBank to date. CIDERSeq can also be applied to other important viruses with similar genome sizes and topology, including e.g. Porcine circoviruses, Chicken anemia virus, and the recently discovered, ubiquitous, human-infecting Torque teno virus. The circular dsDNA Human papillomavirus is another important target for future CIDER-Seq experiments. The sequencing of non-viral circular DNA templates such as plasmids is another potential application of CIDER-Seq, particularly in the context of clinical deep-sequencing of antibiotic resistance gene carrying plasmids ${ }^{15}$. In addition to circular DNA templates, single molecule sequencing of linear DNA and RNA viruses could also be facilitated by DeConcat in combination with size selection, Phi29 polymerase amplification and SMRT sequencing. Based on the current accuracy of long-read sequencing technologies CIDER-Seq produces results with an estimated error rate that is far below virus species and isolate demarcation thresholds. We also note that an error rate of $0.1 \%$ is comparable to the Q30 threshold of Illumina short reads. Further improvements of either SMRT read-length or single-read quality will increase the number of full-length viral genomes and expand our method to larger viruses. Considering recent calls to incorporate metagenomics data in virus classification ${ }^{1}$, CIDER-Seq is a superior method for high-quality full-genome sequencing of viruses infecting plants, animals and bacteria that will facilitate building accurate sequence datasets for virus taxonomy and evolutionary studies.

## Methods

## Sampling and DNA extraction

Mature symptomatic cassava leaves were harvested from infected plants grown for nine months in a confined field trial in Alupe, Kenya. The plants included cassava genotypes 60444 and TME14 as well as transgenic lines in the same genotypic backgrounds. Total nucleic acid was extracted from leaf samples pooled from three plants of each genotype. Extraction was performed using a CTAB (cetyl trimethylammonium bromide) protocol ${ }^{16}$ combined with an ethanol precipitation step. Total nucleic acid was quantified using a Qubit dsDNA BR Assay Kit (Q32850, Thermo Scientific).

## Size Selection

For the pre-enrichment size-selection step, $5 \mu \mathrm{~g}$ of total nucleic acid was loaded on a $0.75 \%$ agarose gel cassette and separated on a BluePippin instrument (SAGE Science). DNA fragments between $0.8-5 \mathrm{~kb}$ were extracted. This size range was selected because geminivirus DNA is present as dsDNA replicative intermediates (running between 2 and 3 kb ) and ssDNA mature forms, which migrate at lower size ranges on agarose gels. Post-enrichment size-selection was similarly performed and fragments $>3 \mathrm{~kb}$ were extracted.

## Random Circle Amplification

Random rolling circle amplification was performed as previously described ${ }^{2}$ with some modifications. A $20 \mu \mathrm{l}$ reaction was set up using $5 \mu \mathrm{l}$ of size-selected template DNA, 1 mM dNTPs, $10 U$ Phi29 DNA polymerase (EP0092, Thermo Scientific), $50 \mu \mathrm{M}$ Exo-resistant random primer (SO181, Thermo Scientific), 0.02 U inorganic pyrophosphatase (EF0221, Thermo Scientific) and 1X Phi29 DNA polymerase buffer (supplied with enzyme). The reaction was run at $30^{\circ} \mathrm{C}$ for 18 hours and stopped by heating to $65{ }^{\circ} \mathrm{C}$ for 2 minutes. Product DNA was purified by sodium acetate/ethanol precipitation. We also used the illustra TempliPhi 100 amplification kit (25640010, GE Life Sciences) and obtained similar amplification results.

## Phi29 debranching

$10 \mu \mathrm{~g}$ of amplified DNA was used in a debranching reaction with 5 U of Phi29 DNA polymerase without a primer at $30^{\circ} \mathrm{C}$ for 2 hours and stopped by heating at $65^{\circ} \mathrm{C}$ for 2 minutes. The product was precipitated with sodium acetate/ethanol. The purified product was treated with 50 U S1 nuclease (ENO321, Thermo Scientific) in a $20 \mu \mathrm{l}$ reaction at $37{ }^{\circ} \mathrm{C}$ for 30 minutes and stopped by adding $3.3 \mu \mathrm{l}$ of 0.5 M EDTA and heating at $70^{\circ} \mathrm{C}$ for 10 minutes. DNA was purified by sodium acetate/ethanol precipitation.

## DNA repair

De-branched DNA was treated with 3U T4 DNA polymerase (M0203L, New England Biolabs) and 10U E. coli DNA polymerase I (M0209L, New England Biolabs) with 1X NEBuffer 2 and 1mM dNTPs in a $50 \mu \mathrm{l}$ reaction. The reaction was incubated at $25^{\circ} \mathrm{C}$ for 1 hour and stopped by heating at $75^{\circ} \mathrm{C}$ for 20 minutes. After cooling, 5 U of Alkaline Phosphatase (EF0651, Thermo Scientific) was added. Dephosphorylation was conducted at $37^{\circ} \mathrm{C}$ for 10 minutes and stopped by heating to $75^{\circ} \mathrm{C}$ for 5 minutes. The repaired DNA was purified using KAPA Pure Beads (KK8000, Kapa Biosystems) at a 1.5 X volumetric ratio and quantified using a Qubit dsDNA BR Assay Kit (32850, Thermo Scientific).

## Semi-quantitative PCR

Semi-quantitative PCR was performed using the primers described in Supplementary Table 2. DreamTaq polymerase (EP0705, Thermo Scientific) was used to amplify 10 ng of template in $50 \mu \mathrm{l}$ reactions set for 15,25 and 40 cycles each. PCR products were separated using a $1 \%$ agarose gel in 1X sodium borate acetate buffer and visualised by staining with ethidium iodide.

## SMRT barcoding, library preparation and sequencing

A Bioanalyzer 2100 12K DNA Chip assay (5067-1508, Agilent) was used to assess fragment size distribution of the enriched DNA samples. The sequencing libraries were produced using the SMRTBell ${ }^{\text {TM }}$ Barcoded Adapter Complete Prep Kit - 96, following manufacturer's instructions (100-514-900. Pacific Biosciences). Approximately 200 ng of each DNA sample was end-repaired using T4 DNA Polymerase and T4 Polynucleotide Kinase according to the protocol supplied by Pacific Biosciences. A PacBio barcoded adapter was added to each sample via a blunt end ligation reaction. The 9 samples were then pooled together and treated with exonucleases in order to create a SMRT bell template. A Blue Pippin device (Sage Science) was used to size select one aliquot of each barcoded library to enrich the larger fragments $>3 \mathrm{~kb}$. Both the non-size selected and the size selected library fractions were quality inspected and quantified on the Agilent Bioanalyzer 12Kb DNA Chip and on a Qubit Fluorimeter respectively. A ready-to-sequence SMRTBell-Polymerase Complex was created using the P6 DNA/Polymerase binding kit 2.0 (100-236-500, Pacific Biosciences) according to the manufacturer instructions. The Pacific Biosciences RS2 instrument was programmed to load and sequence the samples on " $n$ " SMRT cells v3.0 (100-171-800, Pacific Biosciences), taking 1 movie of 360 minutes each per SMRT cell. A MagBead loading (100-133600 , Pacific Biosciences) method was chosen to improve the enrichment of longer DNA fragments. After the run, a sequencing report was generated for every cell via the SMRT portal to assess the adapter dimer contamination, sample loading efficiency, the obtained average read-length and the number of filtered sub-reads.

## Data analysis

Following SMRT sequencing and the generation of barcode-separated subreads ${ }^{17}$ we followed a custom data analysis method depicted in Fig. 2a. First, we implemented the RS_ReadsOfInsert. 1 program using the SMRTPipe command line utility (Pacific Biosciences) using the following filtering
criteria: Minimum Predicted Accuracy=99.9, and Minimum Read Length of Insert (in bases) $=3,000$. (For the error analysis, the same analysis was repeated by changing the Minimum Predicted Accuracy to 99.5 and 99.0 respectively). Resulting high quality ROls were binned into three categories, virus DNA A, virus DNA B or non-viral DNA based on BLAST results (expect value threshold $=1.0$ ) against a database comprised of the full-length East African Cassava mosaic virus (EACMV) DNA A (AM502329) and DNA B (AM502341).

## Sequence Trimming

Next, to identify the putative viral DNA sequence start and end points in each sub-read, the binned ROIs were aligned against two modified EACMV DNA sequences (DNA A and DNA B, for sequences in their respective bins) using MUSCLE ${ }^{18}$. The modified sequences consisted of: a) a three times concatenated full-length genome sequence and b) a genome sequence flanked on either side by two half sequences. This was done to simulate the linearization of the circular genome and allow for the best alignment of the generated sub-read. A 10 nt sliding window was then run from both ends of the alignment. The first window (at both sequence ends) to detect a $90 \%$ sequence identity (i.e. 9 out of 10 nucleotides in the sliding window are identical) between the read and the two modified reference sequences was designated as the start and end point of the viral sequence respectively (Fig. 2, Step 4). The sequence between these two points (called the trimmed ROI) was further analysed using DeConcat (Supplementary Fig. 1).

## DeConcat Algorithm Description

DeConcat begins (Step 1, Supplementary Fig. 1) by cleaving the trimmed ROI (A-B') at the 30nt position to produce two segments (A-A' and B-B') and aligning them using MUSCLE (Step 2a). (This fixed distance of 30 nt was determined by benchmarking a range of values from 500 nt to 30 nt. The benchmarking results are shown in Supplementary Fig. 3) Using the alignment consensus, a score is calculated by dividing the total consensus length by the number of consensus fragments separated by gaps (Step 3). The algorithm iterates back (Step 4) to step 1, increases the cleavage position by 30 , proceeds to step 2 a and step 3 and iterates back to step 1 . This proceeds until all possible cleavage positions have been used. The algorithm retains the alignment with the highest score in step 3. A second iteration now aligns the reverse complement sequence of the first segment (i.e. converts A-A' to A'-A) (Step 2b), calculates the score, and if the score is higher than the previously retained alignment, the reverse complement alignment is used. If the computed score of the two best aligned segments is $>20$, a 10 nt sliding window on both ends is applied and the first windows with $>90 \%$ identity are used to determine start and end positions of the alignment fragments.

The final retained alignment for each ROI can take one of eight possible overlap patterns (Step 5, Supplementary Fig.1). If the smaller segment lies completely within the larger one (cases 1a-1d) the algorithm re-starts with the larger segment (Step 6) and eliminates the smaller one. If the second segment ( $B-B^{\prime}$ ) overlaps the end of the first segment ( $A-A^{\prime}$ ) (case 2), the algorithm restarts (step 6) with both segments as independent ROIs. If the B-B' overlaps with the front of $A-A^{\prime}$ (case 3 ), the overlapping part of $B-B^{\prime}$ is eliminated and the remaining segment ( $B-B^{\prime \prime}$ ) is reattached in its original position at the end of $A-A^{\prime}$. If $B-B^{\prime}$ overlaps the end of $A^{\prime}-A$ (i.e. the reverse complement of $A-A$ ' produced in step $2 b$ ) (case 4), the overlapping part of $A^{\prime}-A$ is eliminated, the remaining segment ( $A^{\prime}-A^{\prime \prime}$ ) is reverse complemented and $B-B^{\prime}$ is re-attached to the end of $A^{\prime \prime}-A^{\prime}$. In case $5, B-B^{\prime}$ overlaps with the start of $A^{\prime}-A$. Here, the overlapping part of $B-B^{\prime}$ is eliminated to produce $B-B^{\prime \prime}$. $A^{\prime}-A$ reverts back to its original configuration $A-A^{\prime}$ and the two segments ( $A-A^{\prime}$ \& $B-B^{\prime \prime}$ ) are reattached. Once case-resolution (step 5) is completed, the resulting sequence is entered back into
the algorithm at step 1 for another round of de-concatenation. The case resolutions have been designed so as to maintain the integrity of the initial fragment, i.e. the order in which bases are produced by the sequencer is never changed. Effectively, case-resolution simply results in sequence reduction from the ends.

## DeConcat Performance

After running DeConcat on our two libraries, we found that in both NSS and SS data sets a majority of sequences had scores in the range of 4-6 at the end of the DeConcat program (Supplementary Fig. 4a), indicating that for these sequences the final de-concatenation round had indeed resolved most repeat sequences. Interestingly, most reads in the SS data set required just 1-3 rounds of deconcatenation to resolve their sequences (Supplementary Fig. 4b).

We also found that of the eight possible alignment cases in DeConcat (see Methods), cases 1a, 1b and 3 were the most frequent (Supplementary Fig. 4c) in both data sets. It should be noted however that cases are assigned by DeConcat iteratively and hence do not reflect overall sequence topology. It is thus likely that the frequency of cases 1 a and 1 b is simply due to alignments of fragments differing greatly in size during DeConcat rounds. The relatively high frequency of case 3, however, does suggest that RCA-derived sequences are often direct sequence concatemers. Overall, based on frequencies of cases 1c, 1d, 4 and 5 it appears that cases where concatemers are formed between sequences and their opposite strands are rare.

To test the hypothesis that more DeConcat processing rounds usually result in shorter sequences we plotted the number of DeConcat rounds per sequence against its length (Supplementary Fig. $4 d)$. However, in many cases, several processing rounds still resulted in sequences $\sim 2800 \mathrm{bp}$ in length-further demonstrating the ability of the system to resolve RCA products into single molecules.

We also tested the ability of DeConcat to process RCA-derived long sequencing reads in the absence of a reference sequence that is used in the trimming step (Step 4, Fig. 2). When comparing results with and without the trimming step, we found that the lengths of only 8.8-15.3\% of sequences were affected by omitting the trimming step. Further, the average change in length of these sequences was only 15-30 bp (Supplementary Fig. 5). We conclude that DeConcat can effectively resolve RCA-derived sequences even in the absence of a reference sequence and does not require prior information on expected monomer length distributions. However, the trimming step is recommended to improve sequencing in cases where multiple viral molecules may be joined together due to the action of the Phi29 DNA polymerase.

## Sequence Annotation and Phasing

The de-concatenated ROIs were annotated using tBLASTn against virus protein reference sequences (Supplementary File 1) with an e-value of 0.01. The high-scoring pairs (HSPs) were summarized and sequences with protein-annotations that had contradicting and incorrect coding strands (according to the reference annotation) or lacked the full set of geminivirus proteins were eliminated. Sequences were replaced by their reverse complements where necessary to maintain all sequences in the same strand (i.e. +strand relative to the geminivirus AV1/AV2 genes). Annotation positions were also adjusted accordingly. Next, since geminiviruses have circular genomes, for comparative analyses we phased all sequences to the same start position (minus an offset) of a selected reference protein (in our case AV1). The proteins were annotated by using the start and end codon positions derived from the tBLASTn HSPs. The results were saved in FASTA (sequences only) and GenBank (sequences with ORF annotations) formats. Frameshift errors were
detected using the HSP results from tBLASTn to identify cases where the same protein annotation had a break or overlap in the alignment results. The error/sequence statistic was calculated by dividing the total number of frameshifts detected by the total number of sequences in each dataset.

## Phylogenetic analyses

Phased sequence reads and reference sequences (also phased) (Supplementary File 2) were aligned using MUSCLE implemented in the CLC Genomics Workbench 10.0 using default parameters. The alignment was used to create a Maximum Likelihood phylogeny using a General Time Reversible model with 100 bootstraps. The phylogeny was used to create an unrooted cladogram and clades were defined based on the position of the 9 reference sequences used. Sequence demarcation analysis was performed using the SDT-MPI solftware ${ }^{11}$ with the MUSCLE alignment option.

## Code Availability

Python packages along with installation and usage guidelines are available at: www.github.com/hirschhm/ciderseq.

## Data Availability

Full length annotated genome sequences, raw sequence data and data generated during intermediate CIDER-Seq steps are freely available at: www.dx.doi.org/10.5281/zenodo. 830530

## References

1. Simmonds, P. et al. Consensus statement: Virus taxonomy in the age of metagenomics. Nat. Rev. Microbiol. 15, 161-168 (2017).
2. Dean, F., Nelson, J., Giesler, T. \& Lasken, R. Rapid amplification of plasmid and phage DNA using Phi29 polymerase and a multiply-pimed rolling circle amplification. Genome Res. 11, 1095-1099 (2001).
3. Sipos, R., Szekely, A., Revesz, S. \& Marialigeti, K. in Bioremediation, Methods in Molecular Biology (ed. Cummings, S. P.) 599, (Humana Press, 2010).
4. Lasken, R. S. \& Stockwell, T. B. Mechanism of chimera formation during the Multiple Displacement Amplification reaction. BMC Biotechnol. 7, 19 (2007).
5. Inoue-Nagata, A. K., Albuquerque, L. C., Rocha, W. B. \& Nagata, T. A simple method for cloning the complete begomovirus genome using the bacteriophage Phi29 DNA polymerase. J. Virol. Methods 116, 209-211 (2004).
6. Zhang, K. et al. Sequencing genomes from single cells by polymerase cloning. Nat. Biotechnol. 24, 680-6 (2006).
7. De Bruyn, A. et al. Divergent evolutionary and epidemiological dynamics of Cassava Mosaic Geminiviruses in Madagascar. BMC Evol. Biol. 16, (2016).
8. International Committee on Taxonomy of Viruses. ICTV Master Species List. (2016). doi:10.15468/i4jnfv
9. Hanley-Bowdoin, L., Bejarano, E. R., Robertson, D. \& Mansoor, S. Geminiviruses: masters at redirecting and reprogramming plant processes. Nat. Rev. Microbiol. 11, 777-88 (2013).
10. Laehnemann, D., Borkhardt, A. \& Mchardy, A. C. Denoising DNA deep sequencing data -high-throughput sequencing errors and their correction. Brief. Bioinform. 17, 154-179 (2016).
11. Muhire, B. M., Varsani, A. \& Martin, D. P. SDT: a virus classification tool based on pairwise
sequence alignment and identity calculation. PLoS One 9, e108277 (2014).
12. Brown, J. K. et al. Revision of Begomovirus taxonomy based on pairwise sequence comparisons. Arch. Virol. 160, 1593-619 (2015).
13. Gawad, C., Koh, W. \& Quake, S. R. Single-cell genome sequencing: current state of the science. Nat. Rev. Genet. 17, 175-188 (2016).
14. Mansoor, S., Zafar, Y. \& Briddon, R. W. Geminivirus disease complexes: the threat is spreading. Trends Plant Sci. 11, (2006).
15. Conlan, S. et al. Single-molecule sequencing to track plasmid diversity of hospital-associated carbapenemase-producing Enterobacteriaceae. Sci. Transl. Med. 6, (2014).
16. Chang, S., Puryear, J. \& Cairney, J. A simple and efficient method for isolating RNA from pine trees. Plant Mol. Biol. Report. 11, 113-116 (1993).
17. Pacific Biosciences Inc. Pacific Biosciences Glossary of Terms. 1-8 (2015).
18. Edgar, R. C. MUSCLE: Multiple sequence alignment with high accuracy and high throughput. Nucleic Acids Res. 32, 1792-1797 (2004).

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## Author Contributions

Conceptualization: DM, WG and HV; Methodology, Investigation, Formal Analysis: DM, MHH \& AP; Software: MHH; Visualisation, Writing-original draft: DM; Funding acquisition, Resources, Writingreview \& editing, Supervision: WG \& HV. All authors agree with the final version of the manuscript.

## Competing Financial Interests

The authors declare that they have no competing financial interest.

## Materials and Correspondence

Devang Mehta (devang@ethz.ch ) \& Herve Vanderschuren (herve.vanderschuren@ulg.ac.be)

## Supplementary Information

Supplementary Table 1: Primer sequences

| Primer Name | Sequence | Use |
| :--- | :--- | :--- |
| mePP2A_genomic_F | CGC TGT GGA AAT ATG GCA TCA | Cassava |
| reference gene |  |  |



Supplementary Figure 1: DeConcat workflow (see Methods for details)
a.



d.


Supplementary Figure 2: Performance of the DeConcat algorithm. (a) Frequency curve of the number of DeConcat rounds required to process each sequencing read. (b) Frequency curve of the final DeConcat alignment score obtained for the CMG datasets. (c) Frequency of different alignment forms used during DeConcat. (d) Correlation plot of the number of DeConcat rounds required to completely resolve sequences versus the length of the final de-concatenated sequence.

## a. Non-size selected library


b. Size selected library


Supplementary Figure 3: DeConcat benchmarking with differently sized cuts in step 1. (a) nonsize selected library, (b) size-selected library (>3 kb).


Supplementary Figure 4: Analysis of frameshift mutations in de-concatenated CMG sequences using different SMRT sequence quality thresholds. (a) The number of frameshifts decreases with increasing quality thresholds. (b) Percentage of frameshift-free sequences. (c) Total number of CMG sequences produced using different quality thresholds for SMRT analysis. (d) Number of frameshifts per protein correlates with the amino acid length of the respective protein.
a.

b.

| Effect | Non-size selected <br> library | Size-selected library |
| :--- | :--- | :--- |
| No. of sequences decreased in length | 6 | 9 |
| No. of sequences increased in length | 11 | 15 |
| No. of sequences changed | 17 | 24 |
| Average change in length | 28.9 | 18.0 |

Supplementary Figure 5: The effect of applying DeConcat directly on sequencing reads without reference-based trimming. (a) Number of sequences ( $y$-axis) versus the change in bp ( $x$-axis) without trimming. (b) Summary of changes without trimming.

Supplementary File 1: Reference protein sequences from the East African cassava mosaic virus.
$>A V 2$
MWDPLVNEFPDSVHGLRCMLAIKYLQALEDTYEPSTLGHDLVRDLVSVIRARNYVEATRRYHHFHSRLEGSS KAELRQPIQEPCYCPHCPRHKSKTGLDKQAHVQKAHNVQDV*
>AV1
MSKRPGDIIISTPGSKVRRRLNFDSPYRNRATAPTVHVTNRKRAWINRPMYRKPTMYRMYRSPDIPRGCEGP CKVQSYEQRDDVKHLGICKVISDVTRGPGLTHRVGKRFCIKSIYILGKIWMDENIKKQNHTNNVMFYLLRDR RPYGNAPQDFGQIFNMFDNEPSTATIKNDLRDRFQVLRKFHATVIGGPSGMKEQALVKRFYRLNHHVTYNHQ EAGKYENHTENALLLYMACTHASNPVYATLKIRIYFYDSIGN*
>AC4
MPFRDTYIMSPINRRRCSRVSLVECLQLTWSTCELRVLEFKPRMSFSHTQSVLYPKNTSCHSFKHYLSHQTL SSLKSVESCIRMGNLTCMPSSNSRGKSRLRTIVSSIVYTQPVAPISTPTFKVPNQAQMSSPIWIRTETPSNG DNFRSMDDLLEAVNNQRMMLTPKALTAAVSQKLLMSLGN*
>AC1
MRTPRFRVQAKNVFLTYPKCSIPKEHQLSFIQTLSLPSSNPKFIKICRELHQNGEPHLHALIQFEGKITITNN RLFDCVHPTCSTNFHPNIQGAKSSSDVVKSYLDKDGDTVEWGQFQIDGRSARGGQQSANDAYAKGLNSGSKSE ALNVIRELVPKDFVLQFHNLNSNLDRIFQEPPAPYVSPFPCSSFDQVPVEIEEWVADNVRDSAARPWRPNSI VIEGASRTGKTIWARSLGPHNYLCGHLDLSPKVFNNAAWYNVIDDVDPHYLKHFKEFMGSQRDWQSNTKYGK PVQIKGGIPTIFLCNPGPTSSYKEFLDEEKQEALKAWALKNAIFITLTEPLYSGSNQSQSQTIQEASHPA*
>AC2
MQSSSPSQNHSTQVPIKVSHRQFKKRAIRRRRVDLVCGCSYYLHINCSNHGFTHRGTHHCSSSSNEWRVYLGN KQSPVFHNHQAPTTTIPAEPGHHNSPGSIQSQPEEGAGDSQMFSQLQDLDDLTASDWSFLKGL*
>AC3
MDLRTGELITAPQAMNGVYTWEINNPLYFTITRHQQRPFLLNQDIITVQVRFNHNLRKELGIHKCFLNFRIW TTLRPQTGLFLRVFRYQVLKYLDNIGVISINDVIRAADHVLFNVIAKTIECQLTHEIKFNVY*
>AC5
MSTCHVQKQSILCVILILPCLLMIICHVMIQPVKPFNQSLLLHARWTTNNSGMKFPQYLKPIPQIVLNCCST GLIIKHIKYLPKVVLGRIAIRPSIPKQIKHHIIRVILLLNIFIHPDLTKYVNGLDTKPLSDPVCQPRPTCHIT NHLTDTKVLHIIPLLIRLDLTWAFTAPRYVWASIHPVHCGLSVHGPVYPGPFSICDVDSGGSSTVPVWAVEV QPSTNLRAWSGNDDISWSLRHNYEP*
>BV1
MYSIRKQPRNFQRKCNSNTTNRFPIRRKYVGGHTRPSVRRRLSYEPVERPLVYNVLCEKQHGDVFNLQQNTS YTSFVTYPSRGPSGDGRSRDYIKLQSMSVSGVIHAKANCNDDPMEVSHVVNGVFVFSLIMDTKPYLPAGVQA LPTFEELFGSYSASYVNLRLLNNQQHRYRVLHSVKRFVSSAGDTKVSQFRFTKRLSTRRYNIWASFHDGDLV NAGGNYRNISKNAILVSYAFVSEHSMSCKPFVQIETSYVG*
$>B C 1$
MDTSVPVISSDYIQSARTEYKLTNDESPITLQFPSTIERTRVRIMGKCMKVDHVVIEYRNQVPFNAQGSVIV TIRDTRLSDEQQDQAQFTFPIGCNVDLHYFSASYFSIDDNVPWQLLYKVEDSNVKNGVTFAQIKAKLKLSAA KHSTDIRFKQPTIKILSKDYGPDCVDFWSVGKPKPIRRLIQNEPGTDYDTGPRYRPITVQPGETWATKSTIG RSTSMRYTGPKHIDIDDSSSKQYASEAEFPLRGLHQLPEASLDPGDSVSQTQSMSKKDIESIIEQTVNKCLI AHRGSSHKDL*

## Supplementary File 2: Reference genome sequences of nine species of cassava geminiviruses.

>EACMVCV, East African cassava mosaic virus Cameroon virus
TATTTACACATATGCCATTGGGGGACATCCTATATAATGCCCCCCATTCCACCGTTTCCC CTGGAGTTTTGAGTGTCCCCCGATCCAAAACGACAGCCAATATGCCGAGAGCCGGTCGTT TTCAAATAAATGCCAAAAATTATTTCATAACCTATCCCAGATGCTCATTAACAAAGGCAG AGGCCCTTTTCCCAATTAAAAGCCCTTTTCTTACCCGACGAATATCAAATTCATTAGGGTTT GCAGAGAACTACATCAGGATGGGGTGCCTCATCTCCATGTTCTCATCCAATTCGAAGGCA AGTTCCAATGTACCAACCCCAGATTCTTCGATCTCATTTCCCCATCCCGATCAACACATT TCCATCCGAACATTCAGGGAGCTAAATCATCGTCCGATGTCAAAGCTTACATTGAAAAGG GAGGGGAATTTCTTGACGATGGAATTTTCCAAGTCGATGCCAGAAGTGCCAGGGGGGAGG GCCAGCATTTTAGCTCAGGTATATGCAGAAGCGTTGAATGCTTCTTCTAAATCAGAAGCTC TTCAAATTATCAAAGAAAAAGATCCAAAGTCCTTTTTTTTTACAGTTCCATAACATATCTG CTAACGCGGATCGAATCTTCCAGGCTCCGCCACAAACTTACGTTAGTCCGTTCTTATCAT CCTCATTTACGCAAGTCCCAGAGGACATAGAAGTCTGGGTATCCGAAAATATATGCCGTC CCGCTGCGCGGCCATGGAGACCGATCAGTATTGTTCTAGAAGGTGATAGCCGAACCGGCA AAACAATGTGGGCTCGTTCACTGGGACCCCCATAATTATCTTTGTGGACACCTGGATCTGT CTCCCAAGATATATTCAAACGACGCATGGTACAACGTCATTGACGACGTAGACCCGCATT ATCTAAAGCATTTCAAAGAGTTCATGGGGGCCCCAACGAGATTGGCAATCAAACACAAAAT ACGGAAAGCCCATTCAAATTAAAGGTGGGATTCCCACCATCTTCTTATGCAATCCGGGCC CCAATTCGTCCTATAAAGAATACCTAGACGAGGACAAGAATTCCAATCTAAGGAATTGGG CGCTCAAGAATGCGCTCTTTCATCTCCCTCACCGAGCCACTCTTCTCCTCCACCGATCAAA GCCAGGCACAGGCAAGCTAAGATCAGAGCACCTAGACGTCGACGCATCGACCTAACTTGT GGCTGTTCCATCTATCGCAGCATTAATTGTCACAACCATGGATTTACGCACAGGGGAAGA CATTGGTGCTCTTCAATGGAGGAATGGCGCCTTTTATCTGGGAGATTCCAAATCCCCTATA TTTCACAATCCTCAACCACGACAGCATGCCGTTCAACATGAATCACGACATAATCACTGT CCAGATTCGGTTCAACTACAACCTGAGGAAAGCACTGGGGATGCACAAGTGTTTTCTCAA CTTCCGGATCTGGACTCGTTTTACATCCTCAGACTTGGCGTTTTCTTCAAAACCTTTAGGAC ACAAGTCATGAAATATCTCAATAATTTAGGTGTTATTTCGATTTCAACTGTTATAGATGC AGTGCATCATGTATTGAATATAGTTTTTGTGGGAACCCTATATGTATCACAAGATCATGC AATCAAATTTAATATTTATTAATTTGTCACTGAATCATAGAAATAGATGCGTATTTTCAG CGTAGCGTACACAGGATTTGAGGCATGTGTACATGCCATATACAATAACAACGCATTCTC GGTATGATTCTCATACTTGGCCTGTTCCTGATGATTTATACACTACATGATTATTGATCCT AAAAAACCTCTTAACCAGAGCTTGTTCCTTCATCCCAGAGGGTCCACCAACAACAGTCGC

ATGGAATTTACGCAACACCTGATACCGGTCCCTAAGATCATTCTTCACAGTTGCAGTCGT AGGTTCATTATCAAACATGTTGAACACTTGTCCAAAATCTTGAGGACTCGGCCCATAAGG CСTTCTATCTCGCACGAGGAAAAACATAACATGGTTCGTATGATTTTGCTTCTTGATATT CTCATCCATCCAGATCTTGCCCAATATATATATGGACTTCACACACAACCTCTTCCCAAC TCTATGGGTAATGCCTGGCCCACGAGTAACTTCACTAACACATCGGACCATACCCGTATG CTTCACATCATCCCTCTGCTCAAACGACTGGACCTTACATGGGCCTTCACAGCCCTTAGG GACGTCTGGGCTTCGATACATCCTGTACATCTTGGGCTTCCGATACATGGGCCTGTTGGC CCATATTCTGCTTCTGGTGACGCGGACAGTGGGGGCAACCACACGGTTCGTGTATGGGCT GTCGAAGTTCAGCCTCCTCCGCACCTTGGATACGGGTGTTGAGATTATTATATCTCCTGG TCGCTTCGCCATAACTCTTACACCGTAGCACCCCTATTAGATCACGTATATATTCAGCCC CGACAGTACCGCGATCATATTCCTGTTCCAAATGTAACAGGTATTTAACAGCAAGCATCG AACGGAAACCGTGCACGGTTTCCGGAAAATCGTTAACTAACGGATTCCACATTTTGACGC GCTCCACTACTTCGTGACGAAGTATTTAAAGTCAAAAATCCATATCTAGCATTCAAGGCG CAAATATTATTGGCCGACAAACATGCGTGCGCGGGGACCACTTTCTTTTACGGGCGCGGA CTATTATGGGGCCCATCTGCTTTTTTCGGGCGCGGCCATCCGGTAATATTAATCGGATGG CCGCCAATATTCGCAATTCGAATTTTGAATGGAGTCTACC
>ACMV, African cassava mosaic virus
ATTACAAGAATGCCATTTAGAGACACCTATATAATGTCTCCAATTAACAGGAGGAGATGC TCAAGAGTGTCTCTAGTTGAGTGTCTCCAATTGACTTGGTCAACATGCGAACTCCGCGTT TTAGAGTTCAAGCCAAGAATGTCTTTCTCACATACCCAAAGTGTTCTATACCCAAAGAAC ACCAGCTGTCATTCATTCAAACACTATCTCTCCCATCAAACCCTAAGTTCATTAAAATCT GTAGAGAGCTGCATCAGAATGGGGAACCTCACTTGCATGCCCTCATCCAATTCGAGGGGA AAATCACGATTACGAACAATCGTCTCTTCGATTGTGTACACCCAACCTGTAGCACCAATT TCCACCCCAACATTCAAGGTGCCAAATCAAGCTCAGATGTCAAGTCCTATCTGGATAAGG ACGGAGACACCGTCGAATGGGGACAATTTCAGATCGATGGACGATCTGCTAGAGGCGGTC AACAATCAGCGAATGATGCTTACGCCAAAGGCCTTAACAGCGGCAGTAAGTCAGAAGCTC TTAATGTCATTAGGGAATTAGTCCCAAAGGACTTTGTACTTCAATTTCATAATCTCAATA GTAATTTAGATAGGATTTTCCAGGAGCCACCAGCTCCTTATGTTTCTCCCTTCCCATGTT CTTCCTTTGACCAAGTTCCTGTTGAAATTGAAGAATGGGTCGCTGATAATGTTAGGGATT CCGCTGCGCGGCCATGGAGACCCAATAGTATTGTAATAGAAGGTGCTAGCAGAACAGGGA AGACGATATGGGCCAGATCTTTAGGCCCACACAATTACCTGTGTGGACACCTGGACCTTA GTCCAAAGGTCTTCAATAATGCTGCCTGGTACAACGTCATTGATGACGTCGATCCCCACT ACCTAAAGCACTTTAAAGAATTCATGGGGTCCCAGAGGGACTGGCAGTCCAACACGAAAT

ACGGGAAACCCGTTCAAATTAAAGGTGGCATTCCCACTATCTTCCTCTGCAATCCAGGAC CTACCTCGTCCTATAAAGAGTTCCTAGACGAGGAAAAGCAAGAAGCGCTAAAGGCCTGGG CATTAAAGAATGCAATCTTCATCACCCTCACAGAACCACTCTACTCAGGTTCCAATCAAA GTCAGTCACAGACAATTCAAGAAGCGAGCCATCCGGCGTAGGAGAGTGGATCTTGTCTGT GGCTGTTCATATTACCTCCATATCAACTGCTCCAATCATGGATTTACGCACAGGGGAACT CATCACTGCTCCTCAAGCAATGAATGGCGTGTATACCTGGGAAATAAACAATCCCCTGTA TTTCACAATCACCAGGCACCAACAACGACCATTCCTGCTGAACCAGGACATCATAACAGT CCAGGTTCGATTCAATCACAACCTGAGGAAGGAGCTGGGGATTCACAAATGTTTTCTCAA CTTCAGGATCTGGACGACCTTACGGCCTCAGACTGGTCTTTTCTTAAGGGTCTTTAGATA CCAAGTACTTAAATACTTGGATAATATAGGTGTTATCTCAATTAACGATGTCATTAGAGC TGCTGATCATGTTTTGTTCAATGTAATTGCCAAAACTATTGAGTGTCAGTTGACTCATGA AATAAAATTCAATGTTTATTAATTGCCAATACTGTCATAGAAGTATATACGTATTTTCAA CGTAGCATATACAGGATTGGAGGCATGAGTACATGCCATGTACAGAAGCAAAGCATTCTC TGTGTGATTCTCATACTTCCCTGCCTCCTGATGATTATATGTCACGTGATGATTCAACCT GTAAAACCTTTTAACCAAAGCCTGCTCCTTCATGCCAGATGGACCACCAATAACAGTGGC ATGAAATTTCCTCAATACCTGAAACCTATCCCTCAAATCGTTCTTAATTGTTGCAGTACT GGGCTCATTATCAAACATATTAAATATCTGCCCAAAGTCTTGGGGCGCATTGCCATAAGG CCTTCTATCCCTAAGCAGATAAAACATCACATTATTCGTGTGATTTTGCTTCTTAATATT TTCATCCATCCAGATCTTACCAAGTATGTAAATGGACTTGATACAAAACCTCTTTCCGAC CCTGTGTGTCAGCCCAGGCCCACGTGTCACATCACTAATCACCTTACAGATACCAAGGTG CTTCACATCATCCCTCTGCTCATACGACTGGACCTTACATGGGCCTTCACAGCCCCTAGG TATGTCTGGGCTTCTATACATCCTGTACATTGTGGGCTTTCTGTACATGGGCCTGTTTAT CCAGGCCCGTTTTCGATTTGTGACGTGGACAGTGGGGGCAGTAGCACGGTTCCTGTATGG GCTGTCGAAGTTCAGCCTTCGACGAACCTTCGAGCCTGGAGTGGAAATGATGATATCTCC TGGTCGCTTCGACATAATTACGAGCCCTGATAACTGAGACTAGATCTCTAACCAAATCGT GGCCCAAAGTACTGGGCTCGTATGTATCCTCTAAGGCCTGCAAATATTTAATTGCAAGCA TACACCTAAGCCCATGCACCGAGTCTGGAAACTCATTCACCAGTGGATCCCACATTGCGC ACTAGCAACAACTTCGCCTGTAAGTATATAGTGGTCCACCACTAATACATAACCTTTAAA GCTACAGCATGATTGGCCGACATAAGTAGTGCGCGGGGACCACGTTTAAAGGGGGGCGGG GCCAACCGGTAATATTATACGGTTGGCCCCTTGGGTGTTCTGCGCCTTTTGAGCCTTTTA ATTCAAATTAAAGTTCAACTT
>EACMKV, East African mosaic Kenya virus
TATTACATAAATGCCATTTAGGGGGCATCATATAAATTGCCCCCCTTTCCCCCGATTGCT

## ACAGACTTTGAGTGCCCCCCGATTGCTATACGACAGCGAAAATGCCAAGGGCTGGTCGTT

 TTAGCATCAAAGCCAAAAACTATTTCCTAACATATCCCAAATGCTCTCTATCCAAAGAGG AGGCATTGGATCAAATCCGACAACTCCAAACCCCAACAAATAAATTGTTCATCAAGATCT GCAGAGAACTCCATGAAAATGGGGAACCTCATCTGCATGCCCTCATTCAGTTCGAGGGCA AGTACAATTGTACCAACCATCGATTCTTCGACCTCATATCCCCATCCCGGTCAGCACATT TCCATCCAAACATTCAGGGAGCTAAATCAAGCTCCGACGTCCAGTCCTATATGGACAAGG ACGGAGACACCATCCAATGGGGCACGTTTCAGATCGACGGACGATCTGCTCGAGGAGGAC AACAATCAGCCAATGACGCTTACGCCAAGGCTCTTAACTCAGCAAATAAGTCAGAGGCTC TTAATGTAATACGCGAACTAGCTCCAAAAGATTTTGTTTTACAGTTTCATAATTTACATA GCAATTTAGATAGGATTTTTCAAGAGCCTCTGACTCCTTATGTTTCTCCATTTCTTTCAT CTTCTTTCACTAACGTTCCTGAGGAACTTGAAGATTGGGTTTCCGAGAACGTGATGGGTT CCGCTGCGCGGCCATGGAGACCTACTAGTATCGTCATCGAGGGCGATAGTAGGACGGGGA AGACGATGTGGGCCCGCTCTTTGGGTCCACACAACTACTTGTGTGGACACCTGGATCTTA GTCCAAAGGTCTACAGCAACGACGCCTGGTACAACGTCATTGATGACGTCGACCCCCACT ACCTCAAACACTTCAAAGAATTCATGGGGGCCCAAAGGGACTGGCAAAGCAATACAAAGT ACGGGAAGCCGATTCAAATTAAAGGCGGCATTCCCACTATCTTCCTCTGCAATCCGGGCC CAACATCATCATATAAAGAGTTTCTGGACGAGGAAAAGAACCAGTCCCTTAAAGCCTGGG CTTTAAAGAATGCCACCTTCATCACCCTCCACGAGCCATTGTTCTCTAGTGCCCATCAAA GTCCAACACCGCACAGCGAAGACCAGGGCCGTCAGACGTAGGCGGGTAGACCTCGAATGC GGCTGCTCGTTCTATCTCCATATCGACTGCATCAACCATGGATTCTCGCACAGGGGAACT CATCACTGCGCCTCAAGCAAGGAATGGCGTTTTTACCTGGGACATAACAAATCCCCTCTA TTTCGAAATCACCAACCACGACAAGAGGCCAGGGAACATGAACCACGACATCATCACACT CCAGATACGGTTCAACCACAACCTCCGGAAGGCATTGGGGATTCACAAGTGTTTTCTCAA CTTCAGGGTCTGGACGACCTTACGGCCTCAGACTGGTCTTTTCTTAAGAGTATTTAGATA TCAAATGCTCAAGTATTTGGATATGATAGGCGTTATTTCCATTAACACTGTACTTCAAGC TGTTGATCATGTTATGTACGATGTATTACTAAACACGCTCCAAGTTACGGAGCAACATGC AATAAAATTCAACCTTTATTAATTTGTCACTGCATCATAAAAATAGATGCGTATTTTCAG CGTAGCGTACACAGGATTAGCGGCATGTGTACATGCCATATACAATAACAACGCATTCTC AGTATGATTCTCATACTTGGCCTGTTCCTGATGATTATACACTACATGATTATTGATCCT AAAAAACCTCTTAACCAGAGCTTGTTCCTTCATCCCAGAGGGTCCACCAACCACAGTCGC ATAGAATTTACGTAACACCTGATACCGGTCCCTAAGATCATTCTTCACAGTTGCAGTAGT TGGTTCATTATCAAACATGTTGAACACTTGCCCAAAATCTTGAGGACTCGGACCATACGG CСTTCTATCTCGAACGAGGAAGAACATCACATGGTTCGTGTGATTTTGCTTCTTGACATTCTCATCCATCCAGATCTTGCCCAATATATATATGGACTTAACACAAAACCTCTTCCCGAC TCTATGGGTAATGCCTGACCCACGAGTAACATCACTGACACATCGGACCATACCAGTGTG CTTAACATCATCCCTCTGTTCATAGGACTGAACCTTACATGGGCCTTCACAGCCCTTAGG GACATCTGGGCTTCGATACATTCTGTACATCTTGGGCTTCCGATACATGGGTCTGTTGGC CCATATTTTGCTTCTGGTGACGCGGACAGTGGGGGCAACAACACGGTTCGTGTATGGGCT GTCGAAGTTCAGCCTCCTCCGCACCTTGGATACGGGTGTTGAGATTATTATATCTCCTGG TCGCTTCGACATAACTCTTACACCGTAGAACCCCTATTAGATCCCGTATATACTCAGCCC CGACAGTACCGCGGTCGTATTCCTGTTCCAGATGTAACAGGTATTTAACAGCAAGCATAG AACGGAAACCGTGAACGGTTTCGGGAAAGTCGTTCAACAATGGATCCCACATGTTGACGC GCTCCACTACTTCGCGACGAAGTCTATAAAGACAAACAACATATATCTAGCCTTTCACGC GTGAATATGACTGGCCGACCAAAACAAGTGCGCAGGGACCACTTTCTTTCACGGGCGCGG CCATCCTGTGGGGTCCACCTGCTTTTTCGGGCGCGGCCATCCGGTAATATTATACGGATG GCCGCTTTCCACGTTTGAATTTCAAATTCAATGAGGA
>SACMV, South African cassava mosaic virus
AATTACACATATGCCATTTGGGGGGCATCATATAAATTGCCCCCCATTCCCCCGATTGCT AGGAACTTTGAGTGCCCCCCGATTGCTATACGACAGCGAAAATGCCGAGGGCTGGTCGTT TTAGCATAAAAGCCAAAAATTATTTCCTCACGTATCCGAAATGCACTCTCTCGAAAGAAG CGGCATTAGATCAACTCCGACAACTCCAAACCCCAACAAATAAATTGTTCATCAAGATCT GCAGAGAACTCCATGAAAATGGGGAACCTCATTTGCATGCCCTCATTCAGTTCGAGGGCA AGTACAATTGTACCAACCAACGATTCTTCGACCTCATATCCCCTTCCAGGTCAACACATT TCCATCCAAACATTCAGGGAGCTAAATCCAGTTCTGACGTCAAGTCCTATTTGGACAAGG ACGGAGACACCATCCAATGGGGCGAGTTTCAGATCGACGGACGATCTGCTCGCGGCGGAC AACAATCCGCCAATGACGCTTACGCCAAGGCTCTTAACGCAGCAAGTAAAACAGAGGCTC TTAATGTAATCCGGGAACTAGCCCCAAAGGATTTTGTTTTACAGTTTCATAATTTAAATA GCAATTTAGATAGGATTTTTCAGGAGCCTCCGATTCCTTATATTTCTCCCTTTCTTTCTT CTTCTTTCACTCATGTTCCTGAGGAACTTGAAGACTGGGTTTCCGAGAACGTGATGGGTT TCGCTGCGCGGCCATGGAGACCGAGTAGTATCGTCATCGAGGGCGATAGTAGGACAGGGA AGACGATGTGGGCCCGATCTCTGGGACCACACAACTACTTATGTGGACATTTGGATCTCA GTCCAAAGGTTTACAGCAACGACGCATGGTACAACGTCATTGATGACGTCGACCCCCATT ACCTCAAGCACTTCAAAGAATTCATGGGGGCCCAAAGGGACTGGCAAAGCAATACCAAGT ACGGGAAGCCGATTCAAATTAAAGGCGGCATTCCCACTATCTTCCTATGCAATCCAGGAC CGACATCATCATATAAAGAGTTTCTGGACGAGGAAAAGAACCAGTCCCTTAAAGCCTGGG CTTTAAAGAATGCAACCTTCATCACCCTCCACGAGCCATTGTTCTCAAGTGCCCATCAAA

GTCCAACACCGCACCGCGAAGACTAGGGCCCTCAGACGTAGGAGGGTAGACCTCGAATGC GGCTGCTCGTTCTATCTCCATATCGACTGCATCAACCATGGATTCTCGCACAGGGGAACT CATCACTGCGCCTCAAGCAAAGAATGGCGTTTTTTACCTGGGAAATAACAAATCCCCTCTA TTTCGACATCACCAACCACGACAAGCGGCCAGGGAACATGAACCACGACATCATCACCCT CCAGATACGGTTCAACCACAACATCAGGAAGGCATTGGGGATTCACAAGTGTTTTCTCAA CTTCAAGGTCTGGACGACCTTACGGCCTCCGACTGGTCTTTTCTTAAGAGTATTTAAATA TCAAGTGCTCAAGTATTTTAAATATGATAGGCGTTATTTCCCATTAACACTGTACTCAGAGC TGTTGATCATGTTCTGTACGATGTATTACTAAACACACTCCAAGTTACGGAGCAACATGC AATAAAATTCAACCTTTATTAATTTGTTACTGCATCATAAAAATAGATGCGTATTTTAAG CGTAGCATACACTGGATTAGAGGCATGCGTACATGCCATATACAACAATAACGCATTCTC TGTATGATTCTCATACTTAGCTGCCTCCTGGTGATTATACACAACATGATTATTTATCCT AAAAAATCTCCTCACCAAAGCCTGCTCCTTCATTCCAGAAGGACCCCCAACAACGGTGGC ATGAAACTTCCGCATAACTCGATACCTATCCCTAAGATCGTTCTTCACAGTGGCTGTACT GGGCTCATTATCAAACATATTAAAAACCTGTCCAAAGTCCATGGGGCTATTGCCATAGGG CCTTCTGTCACGGACTAAGAAGAACATGACCTGGTTTGTATGGTTCTGCTTCTTGATGTT TTCATCCATCCATATCTTACCTAACACATATATAGACTTGATACAGAACCTTTTACCTAC TCTATGTGTAATTCCCGAACCACGCGTGACATCACTAACACAACGAACACTGCCAGTATG CTTAACGTCATCTCGCTGTTCATAAGATTGAACCTTACATGGGCCTTCACAGCCACGCGG AACATCAGGGCTTTTTGAACATTCTGTACATTCTGGGCTTTTCGGTACATGGGCCGGAACGT CCATGATCGACGCTTGTTTGTGCCTTGGACAATGGGGACAGCAGCACGGCTGCTGAACGG GCTGTCGAAGTTCAGCCTTCGACGCACCTTCGAGACGGGAGTGGAAATGATTATATCGGC GGGACGCTTCGACATAATCACGGGCTCGGATCACACCGATGAGATCACGGACTAGATCGT GGCCCAAAGTATTGGGCTCGTAGGTTTCCTCCAAGGCCTGCAAATATTTTAATAGCAAGCA TACAGCGAAAACCGTGCACAGACTCGGGGAACTCATTCAACAATGGATCCCACATGTTGA CACGCTCCACTACTTCGCGACGAAGTCTATAACGACATAAAACAAATATCTAGGCTTTCA CGCGTGAATATGACTGGCCGACAGCAACACGTGCGTGGGGACCACTTTCTTTTACGGGCG CGGTCATCCAATGGGGTCCACCTACTTTGTCGGGCGCGGCCATCCGGTAATATTAGACGG ATGGCCGCTTTCCACGTTCGAAATTCAAATTTCATCACAA
>EACMZV, East African cassava mosaic Zanzibar virus
AATTACCATTATGCCATTTGGAGACACTATATATTGTCTCCAATTCCGTGGAGACATCAA CCGAGGTGTCTCCAATTGCTCTCGCATAAATGACTCCCCCCAAGCGTTTTAAAATACAGG CCAAAAACTATTTTTCTCACATATCCCAAATGTTCTCTATCTAAACACGACGCATTATCCC AAATATTAAACCTCCCAACTCCCACAAACAAGAAATACATCAAAGTGTGCAGAGAACTTC

ACGACGATGGGCAACCTCATCTCCACATGCTTATTCAGTTCGAAGGCAAATTCTCATGCA CAAATAAGCGATTCTTCGACCTGGTATCCCCCACACGATCAACACATTTCCATCCGAACA TTCAAGGAGCTAAATCCAGCTCCGACGTCAAGTCCTACATCGACAAAGATGGGGATACCA CTGAGTGGGGCGAATTCCAGATCGACGCCAGATCGGCTAGAGGCGGCTGCCACAATGCTA ATGACGCATGTGCCGAAGCATTAAACTCCGGTTCCAAGGCAGCAGCACTTCTAATTATTA AGGAGAAACTCCCAAAAGAATTTATTTTTCAATATCATAATTTAAGTAGTAATTTAGATA GGATTTTTCAAGAGCCACCAGCTCCTTATGTTTCTCCATTTCTGTCTTCTTCTTTTACTA ACGTTCCTGAGGAACTTGAAGTCTGGGTTTCCGAGAACGTGATGGGTTCCGCTGCGCGGC CTTGGAGACCTAATAGTATTGTTATTGAGGGTGATAGTCGTACAGGGAAGACAATGTGGG CCAGATCATTGGGACCACATAATTATTTGTGTGGACACCTGGATCTCAGTCCAAAGGTCT ACAGCAACGACGCCTGGTACAACGTCATTGATGACGTCGACCCCCACTACCTCAAACACT TCAAAGAATTCATGGGGGCCCAACGGGACTGGCAAAGCAATACAAAGTACGGGAAGCCAA TTCAAATTAAAGGCGGCATTCCCACTATCTTCCTATGCAATCCAGGACCAACATCATCAT ATAAAGAGTTTCTGGAAGAGGAAAAGCACCAATCCCTTAAAGCCTGGGCTTTAAAGAATG CCACCTTCGTCACCCTCCACGAGCCATTGTTCTCAAGTGCCCATCAAAGTCCAACACCGC ACAGCGAAGACCAGGGCCATCAGACGTAGGCGCGTAGACCTCGAATGCGGCTGCTCGTTT TATATCCATATTAACTGCATCAACCATGGATTCTCGCACAGGGGAACTCATCACTGCGCC TCAAGCAACGAATGGCGTTTTTACCTGGGAAATAACAAATCCCCTATATTTCGCAATCAC CACCCACGACAAGAGACCCGGGAACATGAACCACGACATCATCACAATCCAGATACGGTT CAACCACAACATCCGGAAGGCATTGGGGATTCACAAGTGTTTTCTCAACTTCAAGATCTG GACGACCTTACGGCCTCCGACTGGTCTTTTCTTAAGAGTATTTAGATCTCAAGTGCTCAA GTATTTAGACATGATAGGCGTTATTTCCATTAACACTGTACTTCGATCTGTTGATCATGT TCTGTACGATGTATTACTAAACACGCTCCAAGTTACGGAGCAACATGCAATAAAATTCAA CCTTTATTAATTTGTCACTGCATCATAGAAATAGATGCGTATTTTCAGAGTCGCATACAC AGGATTTGAGGCATGTGTACATGCCATATACAATAACAACGCATTCTCAGTATGATTCTC ATACTTGGCCTGTTCCTGATGATTATACACTACATGATTATTAATCTTAAAAAACCTCTT AACCAGCGCTTGTTCCTTCATCCCAGAGGGTCCACCAACAACAGTCGCATAGAATTTACG CAGCACTTGATACCGGTCCCTAAGATCATTCTTCACAGTTGCAGTCGTAGGTTCATTATC AAACATGTTGAACACTTGTCCAAAATCTTGCGGACTCGGACCATAAGGCCTTCTATCTCG CACGAGGAAGAACATAACATGGTTCGTATGGTTTTGCTTCTTAATATTCTCATCCATCCA AATCTTGCCCAATATATATATGGACTTCACACAGAACCTCTTCCCGACTCTATGGGTAAT GCCTGACCCACGAGTAACCTCACTGACACATCGGACCCTACCAGTGTGCTTAACATCATC CCTCTGTTCATACGACTGAACCTTACATGGGCCTTCACAGCCCTTAGGAACATCTGGGCT

TCGATACATTCTGTACATCTTGGGCTTCCGGTACATGGGCCTGTTAGCCCATATTTTGCT TCTGGTGACGCGGACAGTGGGGGCAACAACACGGTTCGTGTATGGGCTGTCGAAGTTCAG CCTCCTCCGCACCTTGGATACGGGTGTTGAGATTATTATATCTCCTGGTCGCTTCGACAT AACTCTTACACCGTAGAACCCCTATTAGATCCCGTATATACTCAGCCCCGACAGTACCGC GATCGTATTCCTGTTCAAGATGTAACAGGTATTTAACCGCAAGCATGGAACGGAAACCGT GAACGGTTTCAGGGAAATCGTTTAACAATGGATCCCACATGTTGACGCGCTCCACTACTT CGCGACGAAGTCTATAAAGACAAACAACAAATATCTAGACTTCCACGCGTGAATATGACT GGCTGACCGAAACAAGTGCGCGGGGACCACTTTCTTTCACAGGCGCGGCCATCCTGTGGG GTCCACCTGCTTTTTTGGGCGCGGCCATCCGGTAATATTATACGGATGGCCGCTTTTGGT TCACATTTTGAAAATAGATCTACCT
>EACMV, East African cassava mosaic virus
CTATTTACACATATGCCATTGGGGGACCTCATATATAATGCCCCCCATTCCACCGATTGC TATAGACTTTGAGTGTCCCCCGATCCAAAACGACAACCAATATGCCAAGAGCCGGTCGTT TTCAAATAAATGCCAAAAATTATTTCATAACCTATCCCCGATGCTCCTTAACAAAAGAAG AGGCCCTTTTCCCAATTACAAGCCCTTTCGTACCCGACGAATATCAAATTCATTAGGGTTT GCAGAGAACTACATCAGGATGGGGTGCCTCATCTCCATGTTCTCATCCAATTCGAAGGCA AGTTCCAATGTACCAACCCGAGATTCTTCGATCTCATTTCCCCATCCCGATCAACACATT TCCATCCGAACATTCAGGGAGCTAAATCATCGTCCGATGTCAAAGCTTACATTGAAAAGG GAGGGGAATTTCTTGACGCTGGACTTTTCCAAGTCGATGCCAGAAGTGCAAGAGGGGAGG GCCAGCATTTAGCTCAGGTATATGCAGACGCGTTGAATGCTTCGTCTAAATCCGAGGCTC TTCAAATTATTAAAGAAAAAGATCCAAAGTCCTTTTTTTTTACAGTTCCATAACATATCTG CTAACGCAGATAGAATCTTCCAGGCTCCGCCACAAACTTACGTTAGTCCGTTCTTATCAT CATCTTTTTACACAAGTCCCAGAAGACATAGAGGTATGGGTATCCGAAAATATATGCAGTC CCGCTGCGCGGCCATGGAGACCGATCAGTATTGTTCTAGAAGGTGATAGCCGAACCGGCA AAACAATGTGGGCTCGTTCACTGGGACCCCATAATTATCTTTGTGGACACCTGGATCTGT CTCCCAAGGTATATTCAAACGACGCCTGGTACAACGTCATTGATGACGTCGACCCCCACT ACCTCAAACACTTCAAAGAATTCATGGGGGCCCAAAGAGACTGGCAAAGCAATACAAAAT ACGGGAAGCCAATTCAAATTAAAGGCGGCATTCCCACTATCTTCCTCTGCAATCCAGGAC CAACATCATCATATAAAGAGTTTCTGGACGAGGAAAAGAACCAATCCCTTAAAGCCTGGG CTTTAAAGAATGCCACCTTCATCACCCTCCACGAGCCATTGTTCTCTAGTGCCCATCAAA GTCCAACACCGCACAGCGAAGACCAGGGCCGTCCGACGTAGGCGGGTAGACCTCGAATGC GGCTGCTCGTTCTATCTCCATATCGACTGCATCAACCATGGATTCTCGCACAGGGGAACT CATCACTGCGCCTCAAGCAAGGAATGGCGTTTTTACCTGGGACATAACAAATCCCCTCTA

TTTCGAAATCACCGACCACGACAAGAGGCCAGGGAACATGAACCACGACATCATCACACT CCAGATACGGTTCAACCACAACCTTCGGAAGGCATTGGGGATTCACAAGTGTTTTCTCAA CTTCAAGGTCTGGACGACCTTACGGCCTCAGACTGGTCGTTTCTTAAGAGTATTTAAATA TCAAGTGCTCAAGTATTTAGATATGATAGGCGTTATTTCCATTAACACTGTCCTTCAAGC TGTTGATCATGTGGTGTACGATGTATTACTAAACACACTCCAAGTTACGGAGCAACATGT AATAAAATTCAACCTTTATTAATTTGTCACTGCATCATAAAAATAGATGCGTATTTTCAG CGTAGCATACACAGGATTCGAGGCATGTGTACATGCCATATACAATAACAACGCATTCTC TGTGTGATTCTCATACTTCCCTGCCTCTTGATGATTATATGTCACGTGATGATTCAGCTT GTAAAACCTTTTCACCAACGCCTGCTCCTTCACGCCAGATGGACCACCAACAACAGTGGC ATGAAATTTCCTCAACACCTGAAACCTATCCCTCAAATCGTTCTTAATTGTTGCAGTACT GGGCTCATTATCAAACATGTTAAATATCTGCCCAAAGCCTTGGGGCGCATTGCCATACGG CCTTCTATCCCTAAGCAGGTAAAACATCACATTATTAGTGTGATTCTGCTTCTTAATATT TTCATCCATCCAGATCTTACCAAGAATGTAAATGGACTTGATACAAAACCTCTTTCCGAC CCTGTGTGTCAGCCCAGGCCCACGCGTCACATCACTAATCACCTTACAGATACCAAGGTG CTTCACATCATCCCTCTGCTCAAACGACTGGACCTTACATGGGCCTTCACAGCCCTTAGG GACATCTGGGCTTCGATACATTCTGTACATCTTGGGTTTCCGATACATGGGCCTGTTGGC CCATATTTTGCTTCTGGTGACGCGGACAGTGGGGGCAACAACACGGTTCGTGTATGGGCT GTCGAAGTTCAGCCTCCTCCGCACCTTGGATACGGGTGTTGAGATTATTATATCTCCTGG TCGCTTCGACATAATTCTTACACCGTAGAACCCCTATTAGATCCCGTATATACTCAGCCC CGACAGTACCGCGATCGTATTCCTGTTCCAGATGTAACAGGTATTTTAACCGCAAGCATAG AACGGAAACCGTGAACGGTTTCGGGAAAATCGTTCACCAATGGATCCCACATGTTGACGC GCTCTACTACTTCGCGACGAAGTCTATAAAGACAAACAACAAATATCTAGACTTTCACGC GTGAATATGACTGGCCGACCAAAACAAGTGCGCAGGGACCACTTTCTTTCACGAACGCGG TCATTGTGGGATCCACCTGCTTTTTTCGGGCGCGGCCATCCGGTAATATTAATCGGATGGC CGCCAATATTCGGAATTCAAATTTTGAATGGAAGTCTAC
>ACMBFV, African cassava mosaic Burkina Faso virus
GTTGGGATTATTACGACAATGCCATTTGGTGCTAAGTATATATAGAACCCCAATACACTA GGGATAAATCACAGAAGAGTCAATCGGTGCTAGTTGACCAAATGGCTCCTCCTCGAAAAT TCAGAATTAACTCCAAAAATTATTTCCTCACATATCCCAAATGCTCTCTCACAAAAGACG AAGCACTTTCCCAAATCAGAAACTTAGAAACACCAACAAACAAAAAATACATCAAAATCT GCAAAGAATTACACGACGATGGGGAGCCTCATCTCCATGTGCTTATTCAGTTCGAAGGAA AATACAACTGCCAAAATCAACGATTCTTCGACCTGGTATCCCCAACCAGGTCAGCACATT TCCATCCAAACATTCAAGGAGCTAAATCAAGCTCCGATGTCAAGTCCTACATCGACAAGG

## ACGGAGACACCCTTGAATGGGGAGAATTCCAGGTCGACGGAAGAAGTGCTAGGGGAGGCT

 GCCAGAACGCTAACGACGCTGCAGCAGAGGCCTTTAAATGCAGGTTCCGCTGACGCCGCTA TGGCTATTATTAGGGAGAAACTCCCTAAAGAATTTATTTTTTCAATATCATAATTTAAAAA ATAATTTAGATAGGATTTTTCATGGCCCCTCCAGAGCCATATATTTCTCCTTTTTTTATCTT CTTCTTTTACTCACGTTCCGGAAGAACTTGAAGACTGGGTTTCTGAGAACGTCATGAGTT CCGCTGCGCGGCCTTGGAGACCGACTAGTATAGTTATTGAAGGTGATAGTCGTACAGGCA AGACAATGTGGGCCAGGTCATTAGGTCCACACAATTACTTGTGTGGGCATCTAGATCTGA GTCCAAAGGTGTATTCAAATGATGCTTGGTATAACGTCATTGATGACGTCGATCCCCACT ATCTCAAACACTTTAAAGAGTTCATGGGGGCCCCAAAGAGACTGGCAAAGCAACACCAAAT ACGGGAAGCCCATTCAAATTAAAGGTGGAATTCCCACTATCTTCCTCTGCAATCCAGGAC CTACCTCGTCCTATAAAGAGTTCCTAGACGAGGAAAAGCAACAAGCGCTAAAGGCTTGGG CATTAAAGAATGCAATCTTCATCACCCTCACAGAACCACTCTACTCCGGTTCCAATCAAA GTCAGTCACAGACAATTCAAGACGCGAGCCATCCGGCGTAGGAGAGTGGATCTTGTTTGT GGCTGTTCATATTACCTCCATATCAACTGCTCCAATCATGGATTTACGCACAGGGGAACT CATCACTGCTCCTCAAGCAATGAATGGCGTGTATACCTGGGAAATAAACAATCCCCTGTA TTTCACAATAACCGAGCACCAACAACGCCCATTCCTGATGAACCAGGACATCATAACCGT CCAAGTTCGATTCAATCACAACCTGAGGAAGGAGTTGGGGATTCACAAATGTTTTTCTAAC CTTCAGGATCTGGACGACCTTACGGCCTCAGACTGGTCTTTTCTTAAGGGTCTTTAGATA CCAAGTACTTAAATACTTGGATAATATAGGTGTTATCTCAATTAACAGTGTAATTAGAGC TGCTGATCATGTTTTTGTTCAATGTAATTGAAAAAACTATTGAGTGTCAGTTGACTCATGA AATAAAATTCAATGTTTATTAATTCCCAATACTGTCATAGAAGTATATACGTATTTTCAA CGTAGCATATACAGGATTGGAGGCATGAGTACATGCCATATACAAAAGCAATGCATTCTC TGTGTGATTCTCGTATTTCCCTGCCTCTTGATGATTATATGTCACGTGATGATTCAACCT GTAAAACCTTTTCACCAACGCCTGCTCCTTCATGCCAGATGGACCACCAATAACAGTGGC ATGAAATTTCCTCAAAACCTGAAACCTATCCCTCAAATCGTTCTTAATTGTTGCAGTACT GGGCTCATTATCAAACATGTTAAATATCTGCCCAAAGTCTTGGGGCGTATTGCCATAAGG CCTTCTATCCCTAAGCAGATAAAACATTACATTATTCGTGTGATTCTGCTTCTTAATATT TTCATCCATCCAGATCTTACCAAGAATGTAAATGGACTTGATACAAAACCTCTTACCGAC TCTGTGTGTCAGCCCAGGCCCACGTGTCACATCACTAATCACCTTACAGATACCAAGGTG CTTCACATCATCCCTCTGCTCAAACGACTGGACCTTACATGGACCTTCACAGCCCCTAGG TATGTCTGGGCTTCTATACATCCTGTACATCATGGGCTTTTCTGTACATGGGCCTGTTCAT CCAGGCCCGTTTTCGATTTGTGACGTGGACAGTGGGGGCAGTAGCACGGTTCCTGTATGG GCTGTCGAAGTTCAGCCTTCGACGAACCTTGGATCCTGGCGTGGAAATGATGATATCTCCTGGACGCTTCGACATAATTACGAGCCCTGATAACCGAGATTAGATCTCTAACCAGATCGT GGCCCAAAGTACTGGGCTCGTATGTATCCTCTAAGGCCTGCAAATATTTAATTGCAAGCA TACACCTCAGCCCATGCACCGAGTCTGGAAACTCATTCACCAGTGGATCCCACATTGCGC ACTAGCAACAACTTCGCCTGTAAGTATATAGTGGTCCACCACTATTAAATGACCTTTAAC GCTACAGCATGATTGGCCGACATACATAGTGCGCGGGGACCACGTTTAAAGGGGGGAGGG GACCACTTTTTTTTTTTTCGCGCCCACCGGTAATATTAGAACGGTGGGCGCTATGGGGGT TCAAATTT
>CMMGV, Cassava mosaic Madagascar virus
AATATTATACGGATGGCCGCTTTTGGTCCAATTTTTGAATTTTGAACATAGCTTTAACTA ATTACCATAATGCCATTTGGAGACACTATATATTGTCTCCAATTCAGTAGAGACATCAAC CAAGGTGTCTCCAATTGCTCTCGCATAAATGCCTCCTCCCAAGCGTTTTAAAATACAAGC CAAAAACTATTTCCTCACATATCCCAAATGCTCTCTATCTAAACACGACGCATTATCCCA AATCTTAAACATCCCAACTCCAACTAATAAGAAATACATCAAAGTGTGCCGAGAACTTCA CGAAGATGGGCAACCTCATCTCCACATGCTTATTCAATTCGAAGGCAAATTCTCATGCAC AAATAAGCGATTATTCGACCTGGTATCCCCAACAACGTCAACCCATTTCCATCCAAACAT TCAAGGAGCCAAATCCAGCTCCGACGTCAAGTCCTACATCGACAAAGATGGGGATACAAC TGAGTGGGGCGAATTCCAGATCGACGCAAGATCTGCTAGAGGCGGCTGCCAAAATGCTAA TGACGCATGTGCCGAAGCCTTAAACTCACGTTCGAAGGCAGCTGCACTTCTAATTATTAA GGAGAAACTCCCCAAAGAATTTATTTTTCAATATCATAACTTAAGTAGTAATTTAGATAG GATTTTTCAAGAGCCACCAGCTCCCTATGTTTCTCCATTTCTGTCTTCTTCATTCGACCA AGTTCCTGACGACCTTGAGGTCTGGGTGTCAGAAAACATTATGCATCCCGCTGCGCGGCC TTGGAGACCGAATAGTATTGTTATTGAGGGTGATAGTCGTACAGGGAAGACAATGTGGGC CAGATCATTGGGACCACATAATTATTTATGTGGTCATCTCGACCTCAGTCCCAAAGTCTT CACTAATGATGCATGGTACAACATTATTGATGATGTCGATCCGCACTATCTAAAGCACTT TAAAGAGTTCATGGGTGCACAACGAGACTGGCAAAGCAACACAAAATACGGAAAGCCAAT TCAAATTAAAGGCGGAATTCCCACTATCTTCCTCTGCAATCCGGGGCCGACTTCTTCATA TAAAGAATATCTCGATGAAGAAAAGAATGCATCTCTCAAAGCGTGGGCACTGAAAAATGC AACCTTCGTCACCCTCAGCGAGCCATTGTTCACAGGTTCCCATCAAGGTCCTACACCGCA TAGCCAAGACGAGACGCATCAGACGTAGACGTATCGACCTAAGCTGCGGCTGTTCATATT ATCTCCACATCGACTGCATCAATCATGGATTCACGCACAGGGGCACTCATCACTGCTCCT CAAGCGCAGAATGGCGTTTTTACCTGGGAGATAAACAATCCCCTTTATTTCACGATACCC AGACACGACTCGAGACCGTCCCACCTGAACCACGACATCATCACCATCCAAATACGCTTC AACCACAACATCCGGAAGGAATTGGGGATTCACAAATGTTTTCTGAACTTCCAGGTCTGG

## ACGACCTTACACCCTCAGACTGGTCATTTCTTAAGCGTATTTAAGCGTCAAGTTCTTAAG

 TATTTAGATAATGTAGGCGTTATTTCAATAAACACTGTAATTCGCGCTGTTGATCACGTA TTGTACAATGTACTTGTAAACACACTCCAAGTTATGGAGTCCCACGAAATAAAATTTAAT TTGTATTAATTTGTTACTGCATCATAAAAATAGATGCGTATTTTAAGCGTAGCATACACT GGATTAGAGGCATGCGTACATGCCATATACAACAATAACGCATTCTCTGTATGATTCTCA TACTTAGCTGCCTCCTGGTGATTATACACCACATGATTATTAATCCTAAAAAATCTCCTA ACCAAAGCTGCTCCTTCATCCCAGAAGGACCCCCAACAACGGTGGCATGAAACTTCCGCA TAACTTGATACCTATCCCTAAGATCGTTCTTCACAGTTGCTGTACTGGGCTCATTATCAA ACATATTAAAAACCTGTCCAAAGTCCATGGGGCTAGTGCCATAGGGCCTTCTGTCACGGA CTAAGAAGAACATGACCTGGTTAGTATGGTTCTGCTTCTTGATATTTTCATCCATCCATA TCTTACCTAACACATATATAGACTTGATACAAAACCTTTTACCCACTCTATGCGTAATTC CCGGACCACGCGTGACATCACTCACACAACGAACAGTGCCAGTATGCTTGACGTCATCTC GCTGTTCATATGATTGAACCTTACATGGGCCTTCACAGCCACGAGGAACATCAGGGCTTT TGAACATCCTGTACATCCTGGGCTTCCGATACATGGGTCTGAACTTCCATGATTTGAACT TGCTTGTGCCTGGGACAATGGGGACAACAGCACGGCTGCTGTATGGGCTGTCGAAGTTCA GCTTCCGACGCACCTTCGAGGCGGGAGTGGCAATTATTATATCGGCGGGACGCTTCGACA TAGTCACGAGCCCTAATAACAGAAATGAGATCCCGAATTAAATCGCGGCCCAAAGTATTG GACTCGTAGGTTACCTCCACAGCCTGCAAATATTTTTATAGCTAGCATACAACGAAATCCA TGAACCGAGTCAGGAAATTCATTTTAACAATGGATCCCACATTTTGAAATGCAAAACTACT TGTTGGACAAGCATATAAAGACCATTATAATTATGTAGTCTTCCCAGACACAATGTGATT GGTCGACATAAATTAGTGCGTGGGGACCATTGTGGGGCCCACTTACTTTTTCGGGCGCGG CCATCCGGT