Cassava brown streak disease and *Ugandan cassava brown streak virus* reported for the first time in Zambia

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**Abstract**

A diagnostic survey was conducted in July 2017 in two northern districts of Zambia to investigate presence or absence of cassava brown streak disease (CBSD) and its causal viruses. Twenty-nine (29) cassava fields were surveyed and cassava leaf samples collected from 116 plants (92 symptomatic and 24 non-symptomatic). CBSD prevalence was ~79% (23/29) across fields. Mean CBSD incidence varied across fields but averaged 32.3% while mean disease severity was 2.3 on a 1-5 rating scale. RT-PCR screening of all 116 samples with one generic and two species-specific primer pairs yielded DNA bands of the expected sizes from all symptomatic plants with the generic (785 bp) and Ugandan cassava brown streak virus (UCBSV)-specific (440 bp) primers. All 24 non-symptomatic samples were negative for UCBSV and all samples tested negative with primers targeting *Cassava brown streak virus*. The complete genome of a representative isolate of UCBSV (WP282) was determined to be 9,050 nucleotide (nt) in length, minus the poly A tail. A comparative analysis of this isolate with global virus isolates revealed its nature as a sequence variant of UCBSV sharing 94/96% maximum complete polyprotein nt/ amino acid identities with isolates from Malawi (MF379362) and Tanzania (FJ039520). This is the first report of CBSD and UCBSV in Zambia thus expanding the
geographical distribution of the disease and its causal virus and further reinforcing the need to
strengthen national and regional phytosanitary programs in Africa.

Introduction
Cassava brown streak disease (CBSD) and cassava mosaic disease (CMD) are major constraints
to cassava (*Manihot esculenta* Crantz) production in sub-Saharan Africa. Both diseases produce
distinct symptoms in cassava. CMD and CBSD causal viruses are transmitted by the whitefly
*Bemisia tabaci* (Gennadius) in persistent and semi-persistent manners, respectively. However, in
a recent study, the DAG motif associated with aphid transmission of potyviruses was found
within coat protein genes of cassava brown streak virus (CBSV) genomes and two other
ipomoviruses suggestive of their potential aphid transmissibility (Ateka et al., 2017). Whereas
CMD and its causal viruses have been subject of intensive studies (Thottappilly et al. 2003 and
references therein; Alabi et al. 2015 and references therein), the body of knowledge on CBSD
and its causal viruses is just beginning to evolve (Legg et al. 2015 and references therein).
First described in 1936 in the coastal area of Tanzania (Storey 1936), CBSD is now known to be
caused by two positive-sense ssRNA viruses namely CBSV and Ugandan cassava brown streak
virus (UCBSV) that belong to the genus *Ipomovirus* in the family *Potyviridae* (Mbanzibwa et al.
2009; Winter et al. 2010). The genome organization of both viruses is similar comprising a large
polyprotein that encodes 10 genes but lack the helper component proteinase (HC-Pro) (Winter et
al. 2010) present in most members of *Potyviridae* including in *Sweet potato mild mottle virus*,
the typical *Ipomovirus* (Tugume et al. 2010). Rather, both viruses encode a unique HAM1
protein thought to be of cellular origin (Mbanzibwa et al. 2009). In cassava, CBSD symptoms
due to either virus are largely indistinguishable from each other although foliar symptoms due to
UCBSV are reported to be milder than those induced by CBSV (Winter et al. 2010; Patil et al.
2011). It is unclear if such phenotypic differences are due to virus- or host-dependent factors,
the environment, or their interactions. However, CBSV isolates are more genetically diverse than
UCBSV isolates and intraspecies, but not interspecies, recombination has been reported for both
viruses (Mbanzibwa et al. 2009; Ndunguru et al. 2015; Alicai et al. 2016). Country-wide surveys
have shown that UCBSV is more widespread than CBSV with confirmed reports of the former
virus in eight countries compared to that of the latter in four countries (Legg et al. 2015).
Additional reports exist of the presence of CBSD in several other sub-Saharan African countries
such as Angola, Gabon and Madagascar but the association of either of both viruses with these occurrences is yet to be determined (Legg et al. 2015). In addition, CBSV has been shown to have a faster rate of evolution compared to UCBSV (Alicai et al. 2016).

The landlocked country of Zambia is bordered to the north by the Democratic Republic of Congo (DRC), to the west by Angola, to the east by Malawi and Mozambique and to the south by Zimbabwe, Botswana and Namibia. So far, there are confirmed reports of UCBSV in the Democratic Republic of Congo (Mulimbi et al. 2012) and Malawi (Nichols 1950; Mbewe et al. 2015) and of CBSV in Mozambique (Hillocks et al. 2002) and Malawi (Nichols 1950; Mbewe et al. 2015). These reports heightened concerns of CBSD spread into Zambia due to the risks of inadvertent introductions via trans border movement of contaminated cassava cuttings. Hence, survey efforts were initiated in 2009 to determine the incidence and distribution of CBSD, CMD, and their causal viruses in the seven major cassava growing provinces of Zambia. These efforts led to the detection and characterization of CMD causal viruses in farmers’ fields across the seven cassava growing provinces (Chikoti et al. 2013; 2015; Mulenga et al. 2016) but CBSD was reported to be absent (Chikoti et al. 2013; 2015). While areas along the long border of Zambia with Malawi were extensively surveyed, the extreme north of Luapula and Northern provinces covering Chienge and Kaputa districts that border the Democratic Republic of Congo were not covered. Villages and other human settlements in both districts are populated with people of mixed citizenry and it is not uncommon for cassava farmers across these largely unmanned border areas to exchange vegetative cuttings. Therefore, an intensive survey was conducted in farmers fields located in the extreme north of Luapula and Northern provinces to determine the presence or absence of CBSD and its causal viruses.

**Materials and Methods**

**Survey and sample collection.** The survey was conducted in July 2017 in the extreme northern part of Zambia covering the districts of Chienge in Luapula province and Kaputa in Northern province (Fig. 1). The two districts lie on the border with the Democratic Republic of Congo with no physical barrier between the two countries. Both districts were not covered in previous surveys (Chikoti et al. 2013; 2015; Mulenga et al. 2016) due to challenges associated with their accessibility. Surveys were conducted in the agriculture camps of Mwabu (4 fields), Chienge central (5 fields) and Chipungu (4 fields) in Chienge district (13 fields) and Chocha (2 fields),
Kaputa central (7 fields), Chipili (3 fields) and Kasepa (4 fields) in Kaputa district (16 fields).

Cassava fields were surveyed along secondary and feeder roads encompassing areas along Lakes Mweru and Mweru Wantipa that lie at altitudes ca. 940 m above sea level (Fig. 1). Twenty-nine (29) cassava fields were assessed across both districts at 5 to 10 km intervals and each field was georeferenced using a handheld global positioning systems (GPS) device (Garmin International Inc., Olathe, KS, USA). CBSD incidence was assessed for each field as a proportion of cassava plants showing characteristic foliar and/or stem brown streak symptoms (Fig. 2) out of a total of 30 visually inspected plants along two diagonals (15 plants per diagonal) (Sseruwagi et al. 2004).

Disease severity was assessed on a scale of 1-5 (Rwegasira et al. 2011). Mean incidence and severity were calculated along field diagonals for each camp and district regardless of the cultivar. To confirm root and stem diagnostic symptoms, four plants per field with foliar symptoms were uprooted, sliced and inspected for presence of root necrosis (Fig. 2G) while stems with brown streak-like symptoms (Fig. 2F) were sliced from the node upward. Symptomatic leaf samples (Figs. 2A-2E) were collected individually from four plants per field and preserved dry as per Aloyce (2013). A total of 116 (symptomatic = 92 and non-symptomatic = 24) samples were collected during the survey. All samples were packaged in labeled envelopes and transported to the Mount Makulu Central Research Station, Chilanga, Lusaka, Zambia for analysis.

**RNA extraction.** RNA was extracted from each dried cassava leaf sample using a previously described CTAB protocol (Alicai et al. 2016). Briefly, 50 mg of leaf tissue per sample were ground in 1 ml CTAB extraction buffer containing 2% (w/v) CTAB, 2% PVP-40, 100 mM Tris-HCl pH8.0, 25 mM EDTA, 2M NaCl, and 2% mercaptoethanol (added immediately before use). Approximately 750 µl of the extract were pipetted into clean 2 ml microcentrifuge tubes and incubated in a water bath at 65°C for 15 min, then cooled at room temperature. Equal volume of chloroform:isoamyl alcohol (24:1 v/v) was added to the extract, tubes vortexed until content turned cloudy followed by centrifugation at 12,000 rpm for 15 min. This step was repeated twice. Finally, 600 µl of extract were transferred into a 1.5 ml microcentrifuge tube to which one-third volume ice cold isopropanol was added followed by incubation at -20°C for 1 hr. The tubes were then centrifuged at 13000 rpm for 30 min at 4°C. The pellets were washed twice with 70% ethanol and air dried at room temperature. The dried pellet was dissolved in 50 µl DEPC-treated
water, analyzed on a NanoDrop 2000 series spectrophotometer (ThermoFisher Scientific Inc., Waltham, MA, USA) and stored at -20°C until use.

**cDNA synthesis and PCR.** Two micrograms of total RNA per sample were subjected to cDNA synthesis using the Revert Aid First Strand cDNA synthesis kit (ThermoFisher Scientific Inc.) according to the manufacturer’s protocol. Approximately 400 ng cDNA per sample were used in a 25 µl PCR reaction volume containing 1X DreamTaq Buffer (Life Technologies, Carlsbad, CA, USA), 0.2 mM dNTPs, 0.2 mM each sense and antisense primers, and 1 U of DreamTaq DNA Polymerase. Each of the 116 samples was screened with generic primers CPBSV1 and CPBSV2 (Aloyce 2013) and species-specific primers, CBSVF2/CBSVR7 and CBSVF2/CBSVR8 (Abarshi et al. 2012) capable of discriminating between CBSV and UCBSV, respectively. Cycling conditions were initial denaturation at 94°C for 3 min, followed by 35 cycles of 94°C for 30 s, 52°C for 30 s, 72°C for 45 s; with a final extension step of 72°C for 5 min. Additionally, each sample was screened for the presence of African cassava mosaic virus (ACMV) and East African cassava mosaic virus (EACMV), two CMD causal viruses known to be prevalent across cassava fields in Zambia (Mulenga et al. 2016; Chikoti et al. 2015). The ACMV-specific primers JSP01/02 and EACMV-specific primers EAB555F/R (Pita et al., 2001) were employed for this purpose. The PCR products were electrophoresed in 1% agarose gel pre-stained with 10 mg/ml ethidium bromide in 1X Tris-acetate-EDTA buffer and visualized using a Gel Doc XR System (Bio-Rad Laboratories, Hercules, CA, USA). An aliquot of the O’GeneRuler 1-kb DNA Ladder (ThermoFisher Scientific, Inc.) was loaded alongside PCR amplicons for size verification.

**Complete genome amplification and characterization of UCBSV.** Based on results of initial screening of the 116 field collected samples, dried leaf tissues from a representative isolate (WP282) that gave DNA amplicons of the expected size with one of the primer pairs was shipped under a USDA-APHIS-PPQ permit to the Texas A&M AgriLife Research and Extension, Weslaco, Texas, USA facility for further molecular characterization. Total nucleic acid was extracted from isolate WP282 using a Spectrum Plant total RNA Kit (Sigma-Aldrich, St. Louis, MO USA) and used for cDNA synthesis with the cDNA synthesis Kit, ThermoScript™ RT-PCR System (ThermoFisher Scientific, Inc.) as per manufacturer’s instructions. The genome of the CBSD causal virus present in isolate WP282 was amplified as four pieces of overlapping DNA fragments using newly designed primers (Table 1) derived
based on consensus sequence of virus isolates available in GenBank (Supplementary Table 1). The 25 µl PCR reaction volume for each primer pair consisted of 1 x PrimeSTAR GXL buffer, 0.2 mM dNTPs, primers 0.2 µM each, 1.5 U PrimeSTAR GXL DNA polymerase (Takara-Bio USA, Inc., Mountain View, CA, USA) and 2 µl of a 1:10 dilution of the template cDNA. Thermal cycling conditions were 35 cycles of 98°C for 10 s, annealing 55°C for 15 s and extension 68°C for 1 min./Kb. Correct size amplicons on a pre-stained (10 mg/ml ethidium bromide) 1% agarose gel were excised and gel eluted using Zymoclean™ Gel DNA recovery kit (Zymo Research Corp., Irvine, CA). The eluate was A-tailed and cloned into a pCR2.1 TOPO-TA vector then transformed into One Shot TOPO10 Escherichia coli chemically competent cells, according to manufacturer’s protocol (Life Technologies, Carlsbad, CA). Two plasmids with the correct size inserts per DNA amplicon were recovered using a GenElute™ Plasmid Miniprep Kit (Sigma-Aldrich, St. Louis, MO) and initially sequenced in both directions with the M13F/R primers. Additional primers were used to walk each plasmid DNA sample (data not shown). A consensus sequence was obtained for each PCR fragment and the putative full-length viral genome was assembled using the CAP contig assembly program of the BioEdit software (Hall 1999). The genome sequence obtained from isolate WP282 was annotated based on homologies with corresponding sequences in GenBank.

**Sequence analysis.** The complete polyprotein and gene-specific sequences of isolate WP282 were compared with corresponding GenBank sequences (Supplementary Table 1). Sequence alignments were generated with the MUSCLE alignment program (http://www.ebi.ac.uk/Tools/msa/muscle/) and the alignment files employed for calculating pairwise sequence identity matrices with the BioEdit program (Hall 1999). The Simplot v.3.5.1 program (Lole et al. 1999) was used to depict a graphical representation of genome-wide sequence comparisons between isolate WP282 and selected GenBank virus isolates. The phylogenetic relationship between isolate WP282 and publicly available complete genome GenBank isolates of CBSV (14) and UCBSV (19), including infectious clones of UCBSV, was inferred using Bayesian analyses conducted with ExaBayes v.1.4.1 (Aberer et al. 2014) as described in Ndunguru et al. (2015). The polyprotein alignment file involving only UCBSV isolates was scanned for the presence of reticulate phylogenetic networks using the Neighbor-Net method as implemented in SplitsTree v4.10 (Huson and Bryant 2006). The detection of reticulate networks involving isolate WP282 rather than a tree-like topology will indicate that its
evolutionary history has been shaped by recombination (Alabi et al. 2011; Huson and Bryant 2006) as reported for CBSD causal viruses (Mbanzibwa et al. 2011; Ndunguru et al. 2015; Alicai et al. 2016). The exact nature of any potential recombination event was then investigated using the RDP4 software (Martin et al. 2015).

Results

CBSD symptoms and prevalence. Characteristic CBSD symptoms were observed in farmers’ fields in the seven agriculture camps located in Chienge (n = 3) and Kaputa (n = 4) districts. The cassava landraces commonly encountered across the 29 surveyed fields were Nshile, Kalulundyongo, Ndelekulwa, Chibungabunga and Kalilanshindo. In addition, a cassava cultivar referred to as “Research”, presumably denoting that it was received from a research institute, was encountered in one field in Chienge district. Although diverse CBSD symptoms were observed on these cultivars (Fig. 2), leaf chlorotic blotches (Figs. 2A-2B) was the most common symptom on landraces Nshile, Kalulundyongo and Ndelekulwa whereas Chibungabunga and Kalilanshindo displayed feathery veinal yellowing (Figs. 2D-2E). Of the 13 fields surveyed in Chienge district, ~77% (10/13) had CBSD with disease incidence recorded ranging from 10% to 100%; and district-wide mean incidence of ~35% (Table 2). Similarly, ~81% (13/16) of fields surveyed in Kaputa district had CBSD with disease incidence recorded ranging from 3% to 67%; and district-wide mean incidence of 30% (Table 2). Overall CBSD prevalence across fields visited in both districts was ~79% (23/29). Moderate CBSD symptoms were observed across the 23 CBSD-affected fields regardless of the cassava cultivar with a mean severity score of 2.3 (range = 2-3.5). Characteristic CBSD root (Fig. 2G) and stem (Fig. 2F) symptoms were also observed on few plants in a limited number of locations (Table 2). In these locations, incidence of typical CBSD stem symptoms (Fig. 2F) ranged from 2.2% to 10% (low) and mean severity of root necrosis ranged from 2 to 2.3 (mild to moderate) (Table 2). Interestingly, a few plants (Chienge = 2 and Kaputa = 3) showed symptoms of both CBSD and CMD (Fig. 2E) suggesting mixed infections of causal viruses of both diseases in these samples. Based on these results, leaf tissue samples taken from all 116 plants (symptomatic = 92; non-symptomatic = 24) were screened for presence of CBSD and CMD causal viruses.

RT-PCR analyses of CBSVs. The pair of generic CPBSV1/2 primers (Aloyce 2013) produced the expected 785 bp amplicon from all 92 symptomatic samples, while all 24 non-symptomatic
samples were negative. This indicated that each of the 92 symptomatic samples was positive for at least one of the two CBSD causal viruses. Further analysis of these 92 samples with virus-specific primers resulted in the amplification of the expected 440 bp UCBSV-specific DNA fragment from all 92 samples with the primer pair CBSVF2/CBSVR8. No DNA band was obtained from all 92 samples with the CBSV-specific primers CBSVF2/CBSVR7 indicating that UCBSV is the only CBSD causal virus present in these samples. Additionally, five (5.4%) of the 92 UCBSV-positive samples also tested positive for ACMV (4/5) and EACMV (1/5) indicating mixed infections of UCBSV with either of both cassava mosaic begomoviruses (CMBs) in these samples. Cassava plants bearing mixed infections of UCBSV and these CMBs showed more severe foliar symptoms (Fig. 2E) relative to plants with single infection of UCBSV (Figs. 2A-2D).

**Determination and comparative analysis of complete genome sequence of UCBSV from Zambia.** The genome of UCBSV isolate WP286 from Zambia was amplified as four pieces of overlapping DNA fragments using four pairs of newly designed primers (Table 1). Assembly of these sequence fragments resulted in a genome length of 9,050 nucleotides (nt) (GenBank Accession number MG257787) representing the complete genome of Zambian UCBSV isolate WP282, minus the poly A tail. The WP286 sequence has a 5’ untranslated region (UTR) of 134 nt and a 3’ UTR of 207 nt (excluding the poly A tail). The genome encodes a 2,902 amino acid (aa) long polyprotein and homology-based prediction of its proteolytic cleavage sites revealed the expected 10 individual proteins for UCBSV isolates (Fig. 3). The SimPlot analysis of an alignment file of isolate WP286 (MG257787) together with 16 GenBank isolates of UCBSV revealed that it is more closely related to isolate MLB3 (FJ039520) from Tanzania and isolate Malawi (MF379362) from Malawi relative to other GenBank isolates (Fig. 3). In pairwise comparisons, the complete polyprotein sequence of isolate WP286 shared 94% nt and 96% aa identity levels with isolates MLB3 and Malawi, its closest relatives (Table 3). In contrast, polyprotein nt/aa identity levels between isolate WP286 and the remaining GenBank isolates ranged from 86-88/92-94% (Table 3). The levels of nt/aa identities for the 10 encoded genes were similar to those of the polyprotein with a few exceptions (Table 3). For instance, whereas isolate WP282 shared similar levels of nt/aa identities with isolates MLB3 and Malawi with respect to P1, P3, CI, NIa and NIb proteins, contrasting results were obtained for the VPg, Ham1 and CP genes (Table 3). While the VPg and CP genes of isolate WP282 were more identical to
those of isolate MLB3, its Ham1 showed greater resemblance to that of isolate Malawi (Table 3). Overall, higher levels of Ham1 and CP nt/aa identities were obtained for comparisons between isolate WP282 and GenBank isolates compared to other genes indicating greater similarities between the 3' termini genes of UCBSV relative to the 5' termini genes.

**Phylogenetic analysis and test for recombination.** The Bayesian phylogenetic analyses confirmed the clustering of global isolates of CBSV/UCBSV into two major clades, one for each virus with isolate WP282 segregating into the UCBSV clade as expected (Fig. 4). The same five UCBSV-specific subclades identified by Alicai et al. (2016) were recovered in this study with isolate WP282 segregating into clade C along with isolates UCBSV_TZ_MLB3_(FJ039520), UCBSV_MA_43_(FN433933), and UCBSV_TZ_Tan_23_(KR108839) (Fig. 4). These results confirm the results of Alicai et al. (2016) that members of UCBSV clade C (WP282, MLB3, MA_43 and Tan_23) represent a unique UCBSV lineage. A reticulate (non-tree-like) network was detected when the UCBSV sequence alignment file was subjected to analysis with the Neighbor-Net method of the SplitsTree v4.10 program (Fig. 5). Further analysis with the different RDP4-implemented programs revealed two putative recombination events involving isolate WP282 (Table 4) besides events involving other UCBSV isolates as previously reported (Mbanzibwa et al. 2011; Ndunguru et al. 2015; Alicai et al. 2016). Results from this analysis suggest that isolate WP282 arose potentially as a consequence of genetic recombination.

**Discussion**

CBSD and CMD are no doubt the most serious constraints to sustainable production of cassava in sub-Saharan African countries. Unlike CMD which is endemic in all sub-Saharan African cassava-producing countries, the geographical distribution of CBSD and its causal viruses have so far been limited largely to the southeastern axis of the African continent. At present, there are confirmed reports of CBSD occurrence in nine countries including Uganda, Kenya, Malawi, Tanzania, Mozambique, Rwanda, Burundi, Democratic Republic of Congo and the Indian Oceanic island of Mayotte (Legg et al. 2015 and references therein). Given the proximity of Zambia to some of these countries including the fact that it shares extensive land borders with Malawi and the Democratic Republic of Congo (Fig. 1) it seemed inevitable that CBSD would eventually spread into Zambia. Consequently, CBSD and its causal viruses are considered priorities in surveillance efforts by scientists from the Zambian Agricultural Research Institute.
and the International Institute of Tropical Agriculture. Interestingly, despite extensive surveys
(Chikoti et al. 2013; 2015), CBSD and its causal viruses have not been found in Zambia prior to
this study thus prompting the current survey effort. With the mean incidences of ~32% across
~79% (23/29) farmer’s fields in the surveyed districts, it is evident that CBSD is relatively
widespread in both Chienge (Luapula province) and Kaputa (Northern province) districts of
Zambia. In these fields, CBSD was documented on commonly grown cassava landraces and on
the cultivar that farmers referred to as “research”. Due to the annual growth pattern of cassava, it
is difficult to estimate how long CBSD had been present but a ~79% (23/29) prevalence and
~32% mean disease incidence across surveyed fields and all six encountered cassava cultivars
would suggest that local spread has occurred over time albeit at low rates. The only logical
explanation for CBSD incursion into Zambia is that the disease was probably initially introduced
inadvertently via farmer-to-farmer exchange of contaminated cassava planting material rather
than by the whitefly vector. Subsequently, the combination of low whitefly population in most of
the cassava growing provinces in Zambia (Chikoti et al. 2015) and low B. tabaci transmission
efficiency of CBSVs (Maruthi et al. 2005) contributed to a limited CBSD spread encountered
during the survey. This is in addition to the universally accepted view that long distance spread
of whitefly transmitted viruses is caused by human movement of infected planting materials
from regions where disease occurs to new areas (Legg 1999; Mbewe et al. 2015).

The detection of UCBSV alone from all 92 symptomatic plants sampled in this study support the
view that it is more widespread than CBSV (Bigirimana et al. 2011; Mbanzibwa et al. 2011;
Mulimbi et al. 2012; Mbewe et al. 2015; Michel et al. 2016). The exact reasons for the
widespread nature of UCBSV relative to CBSV remains unclear especially against the
background that isolates of the latter virus are more virulent than those of the former (Ogwok et
al. 2015; Alicai et al. 2016). While initial reports point to an altitude dependent prevalence of
CBSV and UCBSV (Alicai et al. 2007; Mbanzibwa et al. 2009), an altitude independent
occurrence of both viruses has also been reported recently (Ndunguru et al. 2015; Alicai et al.
2016). Hence, studies of the comparative epidemiology of both viruses and the relative ability of
the whitefly vector to acquire and transmit them are warranted in order to better understand
factors responsible for the widespread nature of UCBSV than CBSV.

The majority (87/92) of the symptomatic UCBSV-positive cassava plants were singly infected
with the virus and exhibited mild symptoms phenotype (Figs. 2A-2D) in agreement with
previous reports (Mbanzibwa et al. 2009; Legg et al. 2015; Ogwok et al. 2015). These results indicated prevalence of single infection of UCBSV in the surveyed fields. However, five symptomatic plants of the landraces Chibungabunga and Kalilanshindo from three cassava fields located in Kaputa district showed severe symptoms (Fig. 2E). Further evaluation of these plants showed mixed infections of UCBSV with ACMV or EACMV thus suggestive of synergism between the co-infecting viruses. Root and stem symptoms (Figs. 2F-2G) were also observed in some of the disease-affected plants in the two districts, Chienge and Kaputa consistent with symptoms documented in previous studies (Nichols 1950; Mbewe et al. 2014; Patil and Fauquet 2014) and indicating that these phenotypes directly correlated to negative impacts of CBSD are also present under Zambian growing conditions. However, foliar symptoms predominated in all 23 CBSD-affected fields in agreement with previous reports which indicated that foliar CBSD symptoms are the most common and streaks on stems and root necrosis are irregular in cassava plants naturally infected with CBSD causal viruses (Ogwok et al. 2010; Mbewe et al. 2014; Ndunguru et al. 2015; Alicai et al. 2016).

The genome characterization of isolate WP286 from Zambia and its clustering pattern (Fig. 4) further support the distinct nature of the UCBSV clade C isolates identified in a recent study (Ndunguru et al. 2015; Alicai et al. 2016). In this clade, isolate WP286 showed contrasting patterns of gene-specific sequence similarities and differences with other members indicating its distinctness as a sequence variant of UCBSV. Interestingly, the observed contrasting patterns of sequence similarities between WP282 and its closest relatives occurred in the 3’ distal Ham1 and CP genes whereas similar levels of sequence similarities were observed for other regions of the genome (Table 3; Fig. 3). These observations point to potential genetic exchanges occurring among natural populations of UCBSV perhaps as a consequence of an expansion in its geographical distribution (Mbandizwa et al. 2011). Notably the sequence of recently generated UCBSV infectious clones Kikombe (KX753356) and Kikombe-IC (KX753357) (Nanyiti, Unpublished) differ considerably from those of clade C isolates (Table 3; Figs. 3 and 4) hence there may be need to generate additional clade-specific infectious clones for use by breeders and scientists in breeding and reverse genetics studies.

A pertinent question of interest to national and regional regulatory agencies is regarding the possible route of spread of CBSD into Zambia. The relatively high level of similarity of isolate WP282 with UCBSV isolates from Malawi and Tanzania (Table 3, Figs. 3 and 4) suggests
possible incursion of the disease from either country into Zambia. However, recent intensive and
extensive surveys conducted especially in farmer’s fields along the long border of Zambia with
Malawi reported absence of CBSD in these areas (Chikoti et al. 2013; 2015). Therefore, it is
logical to propose that isolate WP282 was probably introduced into Zambia from Tanzania. The
support for this inference is found in higher levels of sequence similarities obtained between
isolate WP282 and Tanzanian isolate MLB3 in nine of ten genes compared to eight of ten genes
with isolate Malawi (Table 3). Interestingly, isolate MLB3 was sourced from the Kagera region
of Northwestern Tanzania which is bordered to the west by Lake Tanganyika, a freshwater lake
that supports commerce and human movement across countries Tanzania, Democratic Republic
of the Congo, Burundi, and Zambia. It is also conceivable that UCBSV introduction into Zambia
came from the Democratic Republic of Congo with whom the surveyed districts share land
border (Fig. 1). However, no genomic data is available from the Democratic Republic of Congo
hence future studies are needed to help us track the true origin of UCBSV introduction into
Zambia. Regardless, there exists very high risk of CBSD spread post-introduction to other
cassava-growing provinces of Zambia via movement of infected stems and/or whiteflies if urgent
exclusionary measures are not taken by relevant regulatory agencies in the country. Therefore,
our results point to the need to strengthen phytosanitary programs and intensify farmer extension
and outreach efforts in the region to stem the tide of disease spread via movement of plant
germplasm.

In conclusion, we have detected the occurrence of CBSD for the first time in Zambia and the
association of a sequence variant of UCBSV with the disease in affected fields. Efforts have been
initiated by the Zambian Agriculture Research Institute (ZARI) to destroy cassava plants in all
affected fields to avert further spread inland to cassava fields in the seven major cassava
producing provinces of the country.

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of cassava brown streak viruses reveals the presence of distinct virus species causing
Table 1. Oligonucleotides used for amplification of complete genome of Ugandan cassava brown streak virus (UCBSV) isolate WP282 from Zambia.

Table 2. Incidences of foliar, stem and root symptoms of cassava brown streak disease (CBSD) recorded in Zambia based on a diagnostic survey conducted in July 2017 in two northern districts of Luapula and Northern provinces of the country.

Table 3. Percent nucleotide (nt)/amino acid (aa) sequence identities between polyprotein and encoded genes of Zambian isolate WP282 of *Ugandan cassava brown streak virus* and global isolates of the virus.

Table 4. Putative recombination events involving *Ugandan cassava brown streak virus* isolate WP282.

Supplementary Table 1. Sequences assembled from the NCBI database and used to design primers UCBSVv502/UCBSVc3182 and UCBSVv6456/UCBSVc9055 (Table 1).

Figure 1. A partial map of Africa showing Zambia and adjoining countries and survey locations targeted in this study; Chienge and Kaputa districts of Luapula and Northern provinces of Zambia. Red and black dots denote cassava fields with and without cassava brown streak disease (CBSD) symptoms, respectively. LP = Luapula province, LSP = Lusaka province, EP = Eastern province, NP = Northern province, CBP = Copperbelt province, CP = Central province, WP = Western province, SP = Southern province, and NWP = Northwestern province.

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**Figure 4.** Bayesian analysis depicting evolutionary relationships between Ugandan cassava brown streak virus (UCBSV) isolate WP282 from Zambia and global isolates of the virus and Cassava brown streak virus (CBSV). The analysis was conducted using ExaBayes version 1.4.1 (Aberer et al., 2014) essentially as described in Ndunguru et al. (2015).

**Figure 5.** SplitsTree reticulate (non-tree-like) networks analyses of complete polyprotein sequences of *Ugandan cassava brown streak virus*. The network analyses were performed using the Neighbor-Net method as implemented in SplitsTree v4.10 (Huson and Bryant, 2006). GenBank accession numbers of the isolates are provided on Table 3.
Table 1. Oligonucleotides used for amplification of complete genome of Ugandan cassava brown streak virus (UCBSV) isolate WP282 from Zambia (MG257787).

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<tr>
<th>Name</th>
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<th>Size (bp)</th>
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*aSmall case letters v and c indicate virion and complementary sense primers, respectively.
Numerals following both symbols indicate genome position of the starting nucleotide for each primer based on isolate WP282.

bY = C+T; R = A+G; D = G+A+T; M = A+C; W =A+T. Primers UCBSVv502/ UCBSVc3182 and UCBSVv6456/ UCBSVc9055 were designed based on consensus sequence of alignments derived from 17 GenBank isolates (Supplementary Table 1) while the remaining two primer pairs are specific to isolate WP282.
Table 2. Incidences of foliar, stem and root symptoms of cassava brown streak disease (CBSD) recorded in Zambia based on a diagnostic survey conducted in July 2017 in the northern districts of Chienge (Luapula province) and Kaputa (Northern province) of the country.

<table>
<thead>
<tr>
<th>District</th>
<th>Camp</th>
<th>Mean altitude (masl)</th>
<th>No. of fields (CBSD+ve/CBSD-ve)</th>
<th>Mean CBSD incidence&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Mean root CBSD&lt;sup&gt;c&lt;/sup&gt;</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Foliar (%</td>
<td>Stem (%)</td>
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<sup>a</sup>CBSD+ve = presence of cassava plants showing CBSD symptoms; CBSD-ve = absence of cassava plants showing CBSD symptoms

<sup>b</sup>% incidence = (number of symptomatic plants/total number of plants evaluated) x 100

<sup>c</sup>Mean root necrosis was calculated using the formula (x/n), where x = number of roots with necrosis, n = total number of tubers inspected. Only four plants were sampled per field and number of root tubers per plant ranged from 2 to 6. NS = No CBSD root necrosis symptoms. Mean root CBSD severity range was calculated as previously described by Rwegasira et al. (2011).
Table 3. Percent nucleotide (nt)/amino acid (aa) sequence identities between polyprotein and encoded genes of Zambian isolate WP282 of *Ugandan cassava brown streak virus* (MG257787) and global isolates of the virus.

<table>
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<th>C1</th>
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Descriptions of Ugandan cassava brown streak virus (UCBSV) encoded proteins can be found in Mbanzibwa et al. (2011).
Table 4. Putative recombination events involving *Ugandan cassava brown streak virus* isolate WP282 (MG257787).

<table>
<thead>
<tr>
<th>Putative recombinant isolate</th>
<th>GenBank accession no.(^a)</th>
<th>‘Parent-like’ isolates(^b)</th>
<th>Method(^c)</th>
<th>P-value(^d)</th>
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<tr>
<td>WP282</td>
<td>6919 (NIb), 7772 (CP)</td>
<td>Unknown Ma43 (90%)</td>
<td>M, C, 3S</td>
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<td>Malawi</td>
<td>7796 (CP), 31 (P1)</td>
<td>WP282 (95%) Ma43 (98%)</td>
<td>R, G, B, M, C, S, 3S</td>
<td>5.645 x 10(^{-35})</td>
</tr>
</tbody>
</table>

\(^a\)Genomic region corresponding to the putative recombination breakpoints are shown in parenthesis.

\(^b\)The percent similarity value shared by the putative recombinant with each parent-like isolate is shown in parenthesis.

\(^c\)The different recombination detection used in the RDP3 analyses are R, RDP; G, GENECONV; B, BOOTSCAN; M, MAXCHI; C, CHIMAERA; S, SISCAN; and 3S, 3SEQ. Only events supported by at least three of the seven RDP4-implemented methods (Martin et al. 2005) are reported.

\(^d\)The support probability (P-value) for the method in bold font is shown.
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199x121mm (131 x 131 DPI)
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<table>
<thead>
<tr>
<th>Virus</th>
<th>GenBank accession no.</th>
<th>Reference</th>
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