Genome-wide association mapping and genomic prediction unrayels CBSD resistance in a

2 Manihot esculenta breeding population

- 3 Siraj Ismail Kayondo^{1,2}, Dunia Pino Del Carpio³, Roberto Lozano³, Alfred Ozimati^{1,3}, Marnin Wolfe³, Yona
- 4 Baguma¹, Vernon Gracen^{2,3}, Offei Samuel², Morag Ferguson⁵, Robert Kawuki¹ and Jean-Luc Jannink^{3,4}
- 5 1) National Crop Resources Research Institute, NaCRRI, P.O. Box, 7084 Kampala, Uganda,
- 6 2) West Africa Center for Crop Improvement, (WACCI), University of Ghana,
- 7 8 3) School of Integrative Plant Sciences, Section on Plant breeding and Genetics, Cornell University, Ithaca, New York,
- 4) US Department of Agriculture Agricultural Research Service (USDA-ARS)
- 9 5) International Institute for Tropical Agriculture (IITA), Nairobi
- 10 Corresponding author: kawukisezirobert@gmail.com

ABSTRACT

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- 14 Cassava (Manihot esculenta Crantz), a key carbohydrate dietary source for millions of people in
- 15 Africa, faces severe yield loses due to two yiral diseases: cassava brown streak disease (CBSD)
- 16 and cassava mosaic disease (CMD). The completion of the cassava genome sequence and the
- of 17 marker profiling clones from African breeding genome
- 18 (www.nextgencassava.org) provides cassava breeders the opportunity to deploy additional
- 19 breeding strategies and develop superior varieties with both farmer and industry preferred traits.
- 20 Here the identification of genomic segments associated with resistance to CBSD foliar symptoms
- 21 and root necrosis as measured in two breeding panels at different growth stages and locations is
- 22 reported. Using genome-wide association mapping and genomic prediction models we describe
- 23 the genetic architecture for CBSD severity and identify loci strongly associated on chromosomes
- 4 and 11. Moreover, the significantly associated region on chromosome 4 colocalises with a 24
- 25 Manihot glaziovii introgression segment and the significant SNP markers on chromosome 11 are
- 26 situated within a cluster of nucleotide-binding site leucine-rich repeat (NBS-LRR) genes
- 27 previously described in cassava. Overall, predictive accuracy values found in this study varied
- 28 between CBSD severity traits and across GS models with Random Forest and RKHS showing
- 29 the highest predictive accuracies for foliar and root CBSD severity scores.
- Key words: Genome-wide association studies (GWAS), virus severity, augmented designs, de-31
- 32 regressed best linear unbiased Predictions (drg-BLUPs), NBS-LRR proteins

33 34 **INTRODUCTION** 35 Cassava (Manihot esculenta Crantz), is a major source of income and dietary calories for more 36 than 800 million people across the globe especially in Sub Saharan Africa (SSA) and recently, 37 due to the unique starch qualities of the storage roots cassava is also turning into an industrial 38 crop (Pérez et al., 2011). Although cassava is a resilient crop, its production is threatened by 39 viral diseases such as Cassava brown streak virus disease (CBSD), which causes major yield 40 losses to poor farming families (ASARECA:, 2013; Ndunguru et al., 2015; Patil et al., 2015). 41 CBSD is caused by two major strains; Cassava brown streak virus (CBSV) and Ugandan 42 cassava brown streak virus (UCBSV) both CBSVs have successfully colonized the lowland and 43 highland altitudes across East Africa and new strains are emerging (Winter et al., 2010; 44 Ndunguru et al., 2015; Alicai et al., 2016). In Uganda, because of CBSVs and agronomical 45 practices, cassava yields were recorded to be eight times lower than the yield potential for this 46 crop (ASARECA:, 2013). 47 In addition to the uncontrolled exchange of infected cassava stakes among farmers across 48 borders, CBSVs are transmitted by the African whitefly (Besimia tobaci) in a semi-persistent 49 manner (Legg, Sseruwagi, et al., 2014; McQuaid et al., 2016). Upon infection, the viruses use 50 the transport system of the plant and cause yellow chlorotic vein patterns along minor veins of 51 leaves in susceptible cassava clones (Ogwok et al., 2010; Maruthi et al., 2016; Anjanappa et al., 52 2016). On the stem, prominent brown elongated lesions commonly referred to as "brown 53 streaks" are formed and in the storage roots, necrotic hard-corky layers are formed in the root 54 cortex of the most susceptible cassava clones (Hillocks et al., 1996; Legg, Somado, et al., 2014; 55 Ndyetabula et al., 2016). 56 Earlier, CBSD resistance breeding initiatives have highlighted the polygenic nature of 57 inheritance in both intraspecific and interspecific cassava hybrids (Nichols, 1947; Hillocks and 58 Jennings, 2003; Munga, 2008; Kulembeka, 2010). In view of the rapid virus evolution and the 59 insufficiency of dependable virus diagnostic tools (Alicai et al., 2016) breeding for durable 60 CBSD resistance, has been the main strategy to control CBSD spread in Eastern Africa. Most of 61 the available elite cassava lines have exhibited some level of sensitivity to CBSVs ranging from

62 mild sensitivity to total susceptibility. Moreover, clones classified as resistant and tolerant show 63 diverse symptom expression, restricted virus accumulation or recovery after clonal propagation (Hillocks and Jennings, 2003; Alicai et al., 2016). 64 Overall, in cassava for many traits the rate of genetic improvement following a traditional 65 breeding pipeline has been slower due to the combination of several biology-related issues such 66 67 as: poor flowering, length of breeding cycle, limited genetic diversity and slow rate of multiplication of planting materials. 68 69 Recently, using genotypic and phenotypic information genome wide association mapping 70 (GWAS) has been used to unravel the genetic architecture of cassava mosaic disease (CMD) 71 (Wolfe et al., 2016) and beta carotene content (Esuma et al., 2016). Both studies have been 72 successful in identifying associated loci with traits of interest. In addition, the performance of 73 genomic prediction for different traits was previously evaluated using historical phenotypic and 74 genotyping by sequencing (GBS) datasets from the International Institute of Tropical Agriculture 75 in Nigeria (Elshire et al., 2011; Ly, Hamblin, Rabbi, Melaku, Bakare, Gauch, et al., 2013). 76 Genomic Selection (GS) is a breeding method alternative to marker assisted selection and 77 conventional phenotypic selection which can accelerate genetic gains through the use of 78 phenotypic and genotypic data from a training population (Meuwissen et al., 2001; Jannink et 79 al., 2010; Lorenz et al., 2011). The performance of different GS models has been evaluated in 80 various species and in many traits (Resende et al., 2012; Gouy et al., 2013; Heslot et al., 2014; 81 Charmet et al., 2014; Cros et al., 2015). Recently the potential of GS for CMD resistance has been reported with predictive accuracies ranging from 0.53 to 0.58 (Wolfe et al., 2016). 82 83 In the present study we followed a GWAS approach in combination with genomic prediction to 84 unravel the genetic architecture of CBSD in two Ugandan breeding populations. While one of 85 our main objectives was to assess the current predictive accuracy for CBSD we also aimed to 86 identify the most promising genomic prediction models that can account for CBSD genetic 87 architecture. 88 GWAS identified loci strongly associated with CBSVs resistance to foliar symptoms which co-89 locate with an introgression block from a cassava wild progenitor, M. glaziovii (Bredeson et al., 90 2016) and with root necrosis which were close to a cluster of plant defence response-related

genes annotated in the cassava genome (Lozano et al., 2015). The presence of introgressions segments from the wild progenitors into the elite breeding lines is the result of cassava improvement programs at the Amani Research Station throughout the 1940s and 1950s (Jennings and Iglesias, 2002; Hillocks and Jennings, 2003). Here we demonstrated with the synergistic implementation of GWAS and GS that GWAS could be used as a prioritization tool to identify markers for genomic prediction for CBSD resistance in cassava. In addition to unravelling the genetics of CBSD resistance these findings may help in the identification of significant causal polymorphisms to guide marker-assisted breeding for CBSD severity that may greatly improve cassava breeding in the face of increasing disease threats to agricultural production.

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(Supplementary Figure 1)

MATERIALS AND METHODS Plant material Phenotypic data was collected from two GWAS panels (Supplementary table 1), GWAS panel 1 composed of 429 clones and GWAS panel 2 which was composed of 872 clones. The combined dataset of 1281 cassava clones were developed through three cycles of genetic recombination between cassava introductions and local elite lines by the National root crops breeding program at NaCRRI. These cassava clones have a diverse genetic background whose pedigree could be traced back to introductions from the International Institute of Tropical Agriculture (IITA), International Center for Tropical Agriculture (CIAT) and the Tanzania national cassava breeding program (Supplementary table 1). **Phenotyping** The GWAS panel trials were conducted in five locations; Namulonge, Kamuli, Serere, Ngetta and Kasese in Uganda. GWAS panel 1 data was collected in two years across three locations, each trial was designed and laid out as a 6 by 30 alpha-lattice design with two-row plots of five plants each at a spacing of 1 meter by 1 meter. GWAS panel 2 was evaluated in three locations, on each location, five rows of test clones were bordered by two CBSD susceptible clones in order to increase CBSD disease pressure (TME204). Clones from GWAS panel 2 were evaluated as single entries per location being connected by six common checks in an augmented completely randomized block design with 38 blocks per site (Federer et al., 2002; Federer and Crossa, 2012). CBSD severity was scored at 3 (CBSD3S), 6 (CBSD6S), and 9 (CBSD9S) months after planting (MAP) for foliar and 12 MAP (CBSDRS) for root symptoms respectively. The CBSD9S scores were not available for GWAS panel 1. CBSD severity was measured based on a 5-point scale with a score of 1 implying asymptomatic conditions and a score 5 implying over 50% leaf vein clearing under foliar symptoms. However, at 12 MAP a score of 5 implies over 50% of root-core being covered by a necrotic corky layer.

Clones were classified with a score of 5 if pronounced vein clearing at major leaf veins were jointly displayed with brown streaks on the stems and shoot die-back that appeared as a candle-stick. Clones with 31 – 40% leaf vein clearing together with brown steaks at the stems were classified under score 4. A Score of 3 was assigned to clones with 21 – 30% leaf vein clearing with emerging brown streaks on the stems. While a score of 2 was assigned to clones that only displayed 1 – 20% leaf vein clearing without any visible brown streak symptoms on the stems. Plants classified with a score of 1 showed no visible sign of leaf necrosis and brown streaks on the stems. On the other hand, root symptoms were also classified into 5 different categories based on a 5 – point standard scale (Jennings and Iglesias, 2002; Hillocks and Jennings, 2003).

Two-stage genomic analyses

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- 160 For the two stage analyses, the first stage involved accounting for trial-design using a linear
- mixed model to obtained de-regressed BLUPs (drgBLUPs) and the second stage involved the use
- of de-regressed BLUPs in GWAS and Genomic prediction.
- For the panel 1 we fitted the model: $= \mathbf{X}\beta + \mathbf{Z}_{clone}c + \mathbf{Z}_{range(loc)}r + \mathbf{Z}_{block(range)}b + \varepsilon$, using
- 164 the *lmer* function from the *lme4* R package (Bates et al., 2015). In this model, β included a fixed
- effect for the population mean and location. The incidence matrix $\mathbf{Z}_{\text{clone}}$ and the vector c
- represent a random effect for clone $c \sim N(0, I\sigma_c^2)$ and I represent the identity matrix. The range
- variable, which is the row or column along which plots are arrayed, is nested in location-rep and
- is represented by the incidence matrix $\mathbf{Z}_{\text{range(loc.)}}$ and random effects vector $r \sim N(0, \mathbf{I}\sigma_r^2)$. Block
- effects were nested in ranges and incorporated as random with incidence matrix $Z_{block(range)}$ and
- effects vector $b \sim N(0, \mathbf{I}\sigma_b^2)$. Residuals ε were fit as random, with $\varepsilon \sim N(0, \mathbf{I}\sigma_{\varepsilon}^2)$.
- For panel 2 we fitted the model $y = X\beta + Z_{clone}c + Z_{block}b + \varepsilon$ Where y was the vector of raw
- phenotypes, β included a fixed effect for the population mean and location with checks included
- as a covariate. The incidence matrix \mathbf{Z}_{clone} and the vector c are the same as the aforementioned
- model and the blocks were also modeled with incidence matrix \mathbf{Z}_{block} and \mathbf{b} represents the
- random effect for block. The best linear predictors (BLUPs) of the clone effect (\hat{c}) were extracted
- as de-regressed BLUPS following the formula (Garrick *et al.*, 2009):

$$deregressed \ BLUP \ = \ \frac{BLUP}{1 - \frac{PEV}{\sigma_c^2}}$$

Where PEV is the prediction error variance of the BLUP and σ_c^2 is the clonal variance component.

DNA preparation and Genotyping by sequencing (GBS)

Total genomic DNA was extracted from young tender leaves of all cassava clones included in the phenotyping trials according to standard procedures using the DNAeasy plant mini extraction kit (QIAGEN, 2012). Genotyping-by-sequencing (GBS)(Elshire *et al.*, 2011) libraries were constructed using the ApeKI restriction enzyme (Hamblin and Rabbi, 2014). Marker genotypes were called using TASSEL GBS pipeline V4 (Glaubitz *et al.*, 2014) after aligning the reads to the Cassava v6 reference genome (Prochnik *et al.*, 2012; Goodstein *et al.*, 2014). Variant Calling Format (VCF) files were generated for each chromosome. Markers with more than 60% missing calls were removed. Genotypes with less than five reads were masked before imputation.

191 Additionally, only biallelic SNP markers were considered for further processing.

The marker dataset consisted of a total of 173,647 bi-allelic SNP markers called for 986 individuals. This initial dataset was imputed using Beagle 4.1 (Browning and Browning, 2016). After timputation, 63,016 SNPs had an AR2 (Estimated Allelic r-squared) higher than 0.3 and were kept for analysis; from these, 41,530 had a minor allele frequency (MAF) higher than 0.01 in our population. Dosage files for this final dataset were generated and used for both GWAS and GS analyses.

Genetic correlations and heritability estimates

Correlation across CBSD traits was estimated using pairwise correlations for each location using the drgBLUPs values obtained after fitting the aforementioned linear mixed model. Broad sense

203 heritabilities (plot-mean basis) were calculated using the estimated variance components from 204

the first step of the two-step genomic analysis as explained previously.

In addition, SNP-based heritabilities were calculated for each GWAS panel by fitting a singlestep mixed-effects model, the full models which specified clone as a random effect were fitted using the emmreml function from the EMMREML R package (Akdemir and Okeke, 2015). The random effect was modeled as having co-variance proportional to the kinship matrix, which was

calculated using the A.mat function from the rrBLUP R package (Endelman, 2011).

Genome-wide association analysis for CBSV severity

- 213 Although pedigree records indicate the two GWAS panels to be closely related a principal
- 214 component analysis (PCA) was performed in order to characterize these panels and to identify
- 215 any population stratification between the two GWAS panels. We used the imputed dataset of
- 216 63,016 SNP markers to calculate the PCs with the function *princomp* in R.
- 217 With the imputed dataset of 63,016 SNP markers and 986 individuals genome wide association
- 218 was performed using a mixed linear model association analysis (MLMA) accounting for kinship
- 219 as implemented in GCTA (v 1.26.0) (Yang et al., 2011). Specifically, we followed a leave one
- 220 chromosome out approach, with this approach the chromosome on which the candidate SNP
- 221 markers are tested gets excluded from the genomic relationship (GRM) calculation. Bonferroni
- 222 correction (reference) was used to correct for multiple testing with a significance threshold set at
- 223 5.9. Manhattan plots with transformed -log₁₀(P-value) were generated using R package *qqman*
- 224 (Turner, 2014).

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Genomic prediction models

- 227 To assess the potential of implementing genomic selection for CBSD, seven genomic prediction
- 228 models were keenly examined; genomic best linear unbiased prediction (GBLUP), reproducing
- 229 kernel Hilbert spaces (RKHS), BayesCpi, Bayesian LASSO, BayesA, BayesB and Random
- 230 forest (RF).

231 **GBLUP**. In this prediction model, the GEBVs are obtained after fitting a linear mixed model 232 where the genomic realized relationship matrix is based on SNP marker dosages. Accordingly, 233 the genomic relationship matrix was constructed using the function A.mat in the R package 234 rrBLUP (Endelman, 2011) and follows the formula of VanRaden (2008), method two. GBLUP 235 predictions were made with the function emmreml in the R package EMMREML (Akdemir and 236 Okeke, 2015). 237 Multi-kernel GBLUP. Because the most significant QTLs for foliar severity 3 and 6 MAP were 238 mapped on chromosomes 4 and 11 (this paper) we followed a multi-kernel approach by fitting 239 three kernels with genomic relationship matrices constructed with SNP markers from 240 chromosomes 4 (G_{chr4}), 11 (G_{chr11}) and SNPs from the other chromosomes (G_{allchr-[4,11]}). Multikernel GBLUP predictions were made with the function emmremlMultiKernel in the R package 241 242 EMMREML (Akdemir and Okeke, 2015). **RKHS**. Unlike GBLUP for RKHS we use a Gaussian kernel function: $K_{ij} = \exp(-(d_{ij}\theta))$, 243 where K_{ij} is the measured relationship between two individuals, d_{ij} is their Euclidean genetic 244 245 distance based on marker dosages and θ is a tuning ("bandwidth") parameter that determines the 246 rate of decay of correlation among individuals. This function is nonlinear and therefore the 247 kernels used for RKHS can capture non-additive as well as additive genetic variation. To fit a 248 multiple-kernel model with six covariance matrices we used the emmremlMultiKernel function 249 in the EMMREML package, with the following bandwidth parameters: 0.0000005, 0.00005, 250 0.0005, 0.005, 0.01, 0.05 (Multi-kernel RKHS) and allowed REML to find optimal weights for 251 each kernel. 252 Bayesian maker regressions. We tested four Bayesian prediction models: BayesCpi (Habier et 253 al., 2011), the Bayesian LASSO (Park and Casella, 2008), BayesA, and BayesB (Meuwissen et 254 al., 2001). The Bayesian models we tested allow for alternative genetic architectures by way of 255 differential shrinkage of marker effects. We performed Bayesian predictions with the R package 256 BGLR (Pérez and De Los Campos, 2014) 257 Random Forest. Random forest (RF) is a machine learning method used for regression and 258 classification (Breiman, 2001; Strobl et al., 2009; Charmet and Storlie, 2012). Random forest 259 regression with marker data has been shown to capture epistatic effects and has been successfully

260 used for prediction (Breiman, 2001; Motsinger-Reif et al., 2008; Heslot et al., 2012; Charmet et 261 al., 2014; Spindel et al., 2015). We implemented RF using the random Forest package in R 262 (Liaw and Wiener, 2002) with the parameter, ntree set to 500 and the number of variables 263 sampled at each split (*mtry*) equal to 300. 264 265 **Introgression Segment Detection** 266 To identify the genome segments in the two GWAS panels, we followed the approach described 267 in Bredeson et al., (Bredeson et al., 2016). We used the M. glaziovii diagnostic markers 268 identified in Supplementary Dataset 2 of Bredeson et al. (Bredeson et al., 2016), these ancestry 269 diagnostic (AI) SNPs were identified as being fixed for different alleles in a sample of two pure 270 M. esculenta (Albert and CM33064) and two pure M. glaziovii. 271 Out of 173,647 SNP in our imputed dataset, 12,502 matched published AI SNPs. For these AI 272 SNPs, we divided each chromosome into non-overlapping windows of 20 SNP. Within each 273 window, for each individual, we calculated the proportion of genotypes that were homozygous 274 (G/G) or heterozygous (G/E) for M. glaziovii allele and the proportion that were homozygous for 275 the *M. esculenta* allele (E/E). We assigned G/G, G/E or E/E ancestry to each window, for each 276 individual only when the proportion of the most common genotype in that window was at least

We also used this approach on six whole-genome sequenced samples from the cassava HapMap

II (Ramu et al., 2016). These included the two "pure cassava" and M. glaziovii (S) from

twice the proportion of the second most common genotype. We assigned windows a "No Call"

Bredeson et al. (Bredeson et al., 2016), plus an additional M. glaziovii, and two samples labeled

Namikonga. Because these samples came from a different source from most our samples, we

could find only 11,686 SNPs that matched both the sites in the rest of our study sample and the

list of ancestry informative sites for analysis.

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status otherwise.

Linkage disequilibrium plots

To confirm whether a large haplotype block present on chromosome 4 colocate with a GWAS QTL identified on this chromosome we calculated LD scores of every SNP marker on chromosome 4 in a 1Mb window using GCTA (Yang *et al.*, 2011). Briefly, LD score for a given marker is calculated as the sum of R² adjusted between the index marker and all markers within a specified window. The adjusted R² is an unbiased measure of LD:

$$R_{adj}^2 = R^2 - \frac{(1-R^2)}{(n-2)}$$

Where "n" is the population size and R² is the usual estimator of the squared Pearson's correlation (Bulik-Sullivan *et al.*, 2015). The resulting LD scores were then plotted against the GWAS log₁₀ (Pvalue) of every marker on chromosome 4.

To highlight the importance of the associated markers on chromosome 11 we calculated pairwise squared Pearson's correlation coefficient (*r*²) between the top significant GWAS SNP hit on this chromosome and neighboring markers in a window of 2Mb (1Mb upstream and 1Mb

Candidate gene identification

downstream). (plink ref)

We used the mlma GCTA output to filter out SNP markers based on -log10 (P-value) values higher than the Bonferroni threshold (~ 5.9). The resulting significant SNP markers were then mapped onto genes using the SNP location and gene description from the M.esculenta_305_v6.1.gene.gff3 available in Phytozome 11(Goodstein et al., 2014) for *Manihot esculenta* v6.1 using the intersect function from bedtools (Quinlan and Hall, 2010).

RESULTS

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Phenotypic variability for severity to cassava brown streak virus infection

- In the present study field disease scoring was done based on a standard CBSD scoring scale that
- ranges from 1 to 5 for both foliar and root symptoms (Supplementary figure 1).
- 319 Datasets for CBSD foliar and root severities of the evaluated germplasm are presented in
- 320 Supplementary figures 2 and 3, both GWAS panels exhibited differential response to CBSVs at
- three, six, nine and twelve months as revealed in the great variability of the deregressed BLUPs.
- 322 Interestingly, clones which displayed an intermediate response were by far more abundant than
- 323 clones with susceptible or resistance response.
- Phenotypic correlations for foliar and root severities (CBSD3S, CBSD6S and CBSDRS) within
- panels and within and across locations are presented in Supplementary figure 4, Supplementary
- tables 2 and 3 with clear differences in CBSD severity scores.
- For panel 1, results varied across locations and CBSD severity traits the lowest correlation value
- was between Ngetta and Kasese (0.09) and the highest between Namulonge and Kasese (0.60)
- both values correspond root severity scoring (Supplementary table 2A).
- For panel 2 the results varied across locations and CBSD severity traits with correlation values
- ranging between -0.08 for CBSD9S (Namulonge-Kamuli) and 0.51 CBSD3S (Kamuli-Serere)
- 332 (Supplementary table 2B).
- Within locations across traits the highest correlation values were found in panel 1 for foliar
- scorings CBSD3S and CBSD6S ($r^2 > 0.5$) (Supplementary table 3A). For panel 2, correlation
- across traits varied depending on the location, nonetheless correlations across foliar traits were
- generally higher than those between foliar and root severity (Supplementary table 3B).
- Heritability estimate values for CBSD3S, CBSD6S and CBDRS were low to intermediate with
- broad-sense heritability (H²) estimates spanning a wide range (11% to 73%) for both panels
- across locations (Table 1). For GWAS panel 2, broad-sense heritability (H²) estimates ranged
- between 56% and 63% for CBSD3S and between 60% and 62% for CBSD6S; while for GWAS
- panel 1 ranged between 11% and 51%.
- Narrow-sense heritability (h²), also referred to as SNP heritability, was estimated using the
- variance components obtained as a result of fitting a one step model using the genetic

- relationship matrix (GRM) for each panel. For panel 1, the broad- and narrow-sense heritability
- values were comparable across locations except for the multi-location model. For panel 2, for
- 346 most locations the broad-sense heritability estimates were larger than the narrow-sense
- 347 heritability estimates. The high variability observed within and across GWAS panels reflects
- differences in population composition, field design and environmental effects.

Genome wide association mapping for CBSV severity in cassava

- 351 The extent of subpopulation structure between the two GWAS panels was examined by PCA,
- which showed no distinct clusters: clones from both panels had mixed distribution. Overall, the
- 353 first three PCAs accounted for 60% of the genetic variation observed in the data (Figure 1). The
- 354 first PC accounted for 30% of the observed variation while the second and third PCs contributed
- 355 20% and 10% respectively.

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- 356 Genotype-phenotype associations for CBSD severity traits based on the combination of multi-
- 357 location data and 986 individuals are presented in Figure 2. Additional GWAS analyses
- performed on each panel individually are presented in Supplementary tables 4 and 5 and
- 359 Supplementary figures 5-12.
- We characterized SNP markers with a -log10 (P-value) above the Bonferroni threshold > 5.9 as
- 361 significant marker-trait associations and further annotated those into candidate genes
- 362 (Supplementary table 4).
- For the combined dataset, we identified 83 significant SNP markers associated to CBSD3S; the
- markers mapped to chromosome 11 with 61 markers located within genes (Supplementary Table
- 365 4). The QTL on chromosome 11, top hit reference SNP - \log_{10} (P-value) = 9.38, explained 6% of
- the observed phenotypic variation.
- 367 On the other hand, for CBSD6S, we identified significant SNPs on chromosome 11,
- 368 chromosome 4 and chromosome 12. On chromosome 11, 33 SNPs surpassed the Bonferroni
- threshold with 27 SNP markers located within genes. The QTL on chromosome 11 is located on
- 370 the same region as the QTL identified for CBSD3S and explained 5% of the observed
- 371 phenotypic variation (Figure 3A).

372 It suffices to note that although several SNPs on chromosome 11 for CBSD6S exceeded the Bonferroni threshold, six SNPs were in linkage disequilibrium ($r^2 > 0.6$) with the top reference 373 SNP hit. The SNP markers, with an $r^2 > 0.2$ to the reference SNP, were annotated into candidate 374 375 genes: Manes11G130500, a gene that is known to encode glycine-rich protein. 376 Manes11G130000 gene that encodes Leucine-rich repeat (LRR) containing protein, 377 Manes11G130200 gene that encodes the trigger factor chaperone and peptidyl-prolyl trans and 378 Manes 11G131100 that encodes a protein kinase (Figure 3B). 379 Since several SNPs on the chromosome 4 QTL region are in high LD, no single locus can be 380 highlighted as candidate gene(s) to be associated with CBSD severity (Figure 4A). The large 381 haplotype on chromosome 4 is an introgression block from the a wild relative of cassava (M. 382 glaziovii) (Jennings, 1959; Bredeson et al., 2016). We further confirmed the presence and 383 segregation of the introgressed genome segment in both panels using a set of diagnostic markers 384 from *M. glaziovii* (Figure 4B, supplementary figure 13 and 14). 385 The significant QTL on chromosome 12 has been previously identified for CMD resistance in cassava (Wolfe et al., 2016) Accordingly, after correction for CMD scoring in the first step 386 387 calculation of CBSD deregressed BLUPs, the QTL on chromosome 12 was no longer significant 388 and only QTLs on chromosomes 4 and 11 remained (supplementary Figure 15). 389 For CBSDRS we could not identify SNPs surpassing the Bonferroni correction partly to the 390 complexity of this trait with apparently several small effect genes and low heritability. However,

Genome-wide prediction for CBSV severity in cassava

phenotypic variance respectively.

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An important objective within this study was to assess the accuracy of prediction in cassava for CBSD-related traits. Using the combined dataset, we compared the performance of seven genomic prediction models with contrasting assumptions on trait genetic architecture. Some model predictions represent genomic estimated breeding values (GEBV) in that they are sums of additive effects of markers, while other model predictions represent genomic estimated total

the results of the analysis of CBSDRS multi-location data of panel 1 identified significant

regions on chromosomes 5, 11 and 18 (-log10 (P-value) > 6.5), which explained 8, 6 and 10%

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genetic value (GETGV) because they include non-additive effects. Predictive accuracy for CBSD related traits had mean values across methods of 0.29 (CBSD3S), 0.40 (CBSD6S) and 0.34 (CBSDRS) (Figure 5 and Supplementary table 6). Predictive accuracies for CBSD3S varied in the range of 0.27 (BayesB and GBLUP) and 0.32 (RF), for CBSD6S we obtained a predictive value of 0.40 for most methods except for RKHS (0.42) and RF (0.41) and for CBSD root severity scores varied from 0.31 (BayesA, B, C and GBLUP) to 0.42 (RF and RKHS). It is clear from the results that higher predictive accuracies were consistently achieved when using Random forest and RKHS for the prediction of both foliar and root CBSD resistance traits. Although for foliar symptoms the increase in predictive accuracy using those methods is modest, for CBSDRS the increase in predictive accuracy was 0.10. Based on the GWAS results, we identified for CBSD3S, CBSD6S and CBSDRS the strongest marker associations on chromosomes 4 and 11. Markers from chromosomes 4, 11 and markers on other chromosomes were used independently to construct covariance matrices that were fitted in a multikernel GBLUP model (Supplementary figure 16). For all CBSD traits the mean predictive accuracy values from the single-kernel GBLUP model were similar to the mean total predictive accuracy following the multi-kernel approach (Supplementary table 6). Differences were found on the contribution of the individual kernels to the total predictive accuracies. For example, the multikernel GBLUP model for CBSD3S had the lowest total predictive accuracy (0.27) with the highest contribution coming from chromosome 11 and the rest of the genome (0.19). In contrast, the multikernel GBLUP model for CBSD6S gave the highest predictive accuracy (0.40) and most of the accuracy came from chromosome 4 (0.29). The multikernel GBLUP approach for CBSDRS had a total predictive accuracy of 0.30 with the rest of the genome (0.29) contributing the most to the total predictive accuracy (Supplementary figure 16).

430 **DISCUSSION** 431 Cassava brown streak disease has been identified as one of the most serious threats to food 432 security (Pennisi, 2010) owing to the significant loses it imparts in cassava wherever it occurs. 433 Host plant resistance, that is obtained through breeding efforts has been so far the most effective 434 approach. However, this is only achievable when the host-pathogen behaviour and interaction is 435 well understood and/or when the genetics of resistance to CBSD are clearly known. 436 In the present study, ~1200 cassava clones from the NaCRRI breeding program in Uganda were 437 evaluated for CBSD severity scores in leaves and root. Specifically, this paper sought to provide 438 fundamental information on the genetics of resistance to CBSD which was previously unknown. 439 From our analyses it was evident that correlation among foliar CBSD severities were higher than 440 correlation between foliar and root severities. 441 Selection of resistant clones has been hampered by the fact that some clones do not show 442 symptoms on leaves or storage roots, while other varieties may only express symptoms on leaves 443 and not on roots and still others do not show symptoms on leaves but instead on roots only 444 (ASARECA:, 2013). Moreover, a lack of correlation between virus load and symptom 445 expression in a field evaluation of selected cassava genotypes has been reported (Kaweesi et al., 446 2014). Previous studies have also reported that 79% plants with above- ground symptoms of 447 CBSD also exhibited root necrosis and 18% of plants had no visible symptoms of CBSV 448 (Hillocks et al., 1996) 449 Recently, efforts to understand CBSD have focused on CBSD resistance population development 450 and preliminary insights into chromosomal regions and genes involved in resistance (Kawuki et 451 al., 2016; Anjanappa et al., 2016, 2017). These studies have highlighted the existence of a QTL 452 on chromosome 11 for CBSD root necrosis among cassava clones of Tanzanian origin (Kawuki 453 et al., 2016). 454 In our study, based on foliar CBSD severity scoring using a multi-location dataset we identified 455 significant QTL regions on chromosome 4 and 11, though these associations were not always 456 consistent when the panels were analyzed separately and per location. Overall, these results 457 highlight the advantage of using a large GWAS panel and a multi-location approach were plants

are exposed to different disease pressures to identify additional genomic regions.

459 On chromosome 11, a cluster of genes underlies the significant OTL; candidate genes for further 460 study are: Manes11G131100, Manes11G130500, Manes11G130200 and Manes11G130000. 461 Lozano et al. 2015 previously reported Manes11G130000 when studying the distribution of 462 NBS-LRR in cassava. Furthermore, a recent study on early transcriptome response to brown 463 streak virus infection in susceptible and resistant cassava varieties identified Manes.11G130000 464 among the differentially expressed genes in the susceptible line 60444 from the ETH cassava 465 germplasm collection (Anjanappa et al., 2017). The QTL on chromosome 11 is particularly 466 unstable across locations, which may be related to NBS-LRR genes conferring resistance to a 467 particular strain, UCBSV exhibits a lower mutation rate, while CBSV is more aggressive and 468 mutates faster. 469 Throughout the 1940s and 1950s at the Amani Research Station, Manihot glaziovii and cassava 470 varieties of Brazilian origin were used for crosses to obtain CBSD resistant varieties (Jennings 471 and Iglesias, 2002). One of the introgression segments from these wild relatives has been 472 reported to be located on chromosome 4, however the level of linkage disequilibrium in that 473 region remains as a major constraint for the identification of the gene or genes that are 474 responsible for CBSD resistance (Bredeson et al., 2016). Current on-going research efforts are 475 focused on dissecting the extent of the effects of wild introgressions on cassava traits (Marnin 476 Wolfe personal communication). 477 One important objective of the present study was to test our ability to predict CBSD severity in 478 cassava, which is, particularly relevant in two situations. First, when the objective is the 479 introduction of germplasm from Latin america and/or from West Africa to East Africa and for 480 early seedling or clonal selection of resistant lines. 481 Thus, using a cross-validation approach, we evaluated the suitability of seven GS models with 482 the expectation that the results may differ due to differences in genetics of foliar and root CBSD 483 severity traits (B. J. Hayes et al., 2009; Grattapaglia et al., 2011). 484 In cassava, previous genomic prediction studies have evaluated the predictive ability of GBLUP 485 using historical phenotypic data from the International Institute of Tropical Agriculture (IITA) 486 and GBS markers and in a small training population with relatively low-density markers (de 487 Oliveira et al., 2012; Ly, Hamblin, Rabbi, Melaku, Bakare, Okechukwu, et al., 2013).

488 Principally, the GS models evaluated have varying underlying assumptions genomic-BLUP 489 (GBLUP) model assume an infinitesimal genetic architecture; Bayesian methods such as BayesA 490 and BayesB relax the assumption of common variance across marker effects (De Los Campos et 491 al., 2009; Habier et al., 2011; Legarra et al., 2014), RKHS and random forest methods can model 492 epistatic and other non-additive effects. 493 A first assessment of predictive accuracy of CBSD foliar and root traits in cassava indicate that 494 the use of genomic selection is a promising breeding method for resistance to Cassava brown 495 streak virus. We found moderate to high predictive accuracies for these traits in relation to results 496 from other traits in cassava (Ly, Hamblin, Rabbi, Melaku, Bakare, Okechukwu, et al., 2013). 497 However, predictive accuracy values are lower in comparison to the values reported for cassava 498 mosaic virus (Ly, Hamblin, Rabbi, Melaku, Bakare, Okechukwu, et al., 2013) possibly due the 499 presence of a large effect GWAS QTL (CMD2) for CMD. 500 Although, a priori knowledge of the loci affecting a trait is not needed for GS, we also tested a 501 multiple kernel approach using GWAS results as a reference to construct covariance matrices. 502 GWAS results have been incorporated in genome-wide prediction models to increase predictive 503 accuracy through de-novo GWAS or using previously published GWAS results (Zhang et al., 504 2014; Spindel et al., 2016). 505 In our study, to avoid a correlation effect across covariance matrices we partitioned SNP markers 506 into three sets: markers on GWAS QTLs chromosomes (chr 4 and 11) and markers on rest of the 507 genome to built genomic relationship matrices (G_{chr4},G_{chr11},G_{allchr-[4,11]}) Remarkably, the 508 predictive accuracy of each kernel modeled the genetic architecture found though GWA 509 analyses. Our GWAS and GS results indicate that resistance to CBSD root necrosis severity is 510 polygenic in nature, which is in accordance to Kawuki et al.'s (2016) results. 511 Our results suggest that non-additive effects are likely to play a role shaping CBSD resistance 512 particularly root necrosis. This conclusion derives from GS results using Random Forest and 513 RKHS, which gave the highest predictive accuracies, and from the observed differences in broad 514 sense and narrow sense heritability values. 515 CBSD is a disease that has devastating consequences in cassava production and poses a risk

particularly to countries in Central and West Africa where CBSD is not currently present. Our

study provides, through GWAS and genomic prediction, an insight into the genetic regulation of CBSD severity in leaves and roots. Although we were able to identify a candidate NBS-LRR gene on chromosome 11, the function of this gene in CBSD resistance requires further validation and more importantly, there is a risk that this gene might not be a source of durable resistance to CBSVs. Within this context, genomic selection arises as a promising tool that can accelerate breeding, though the average predictive accuracy is lower than CMD, this is highly variable across locations and the breeding panel evaluated. Further work will require screening of large diversity panels in multiple environments, identification of QTLs specific to viral strains and the introgression of genomic regions conferring resistance to CBSD from wild relatives and Latin American accessions.

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Figures and supplementary legends

Figure 1. Principal components analysis of panel 1 and panel 2 clones. The top two panels and the lower left panel show the distribution of clones in PC1-PC3. The lower right panel shows the variance explained by the first ten principal components. Green color shows the distribution of panel 1 clones and the orange color shows the distribution of panel 2 clones.

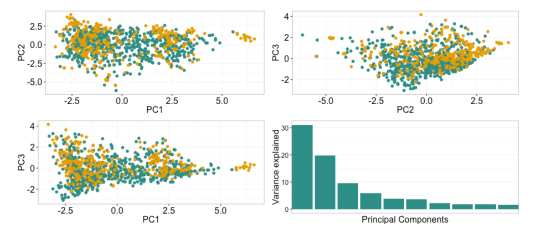


Figure 2. **GWAS results for CBSD severity** .Analysis was performed with a multilocation combined dataset of panels 1 and 2.(a) scoring 3 MAP (b) 6 MAP and (c) root necrosis severity. Red line indicates Bonferroni threshold.

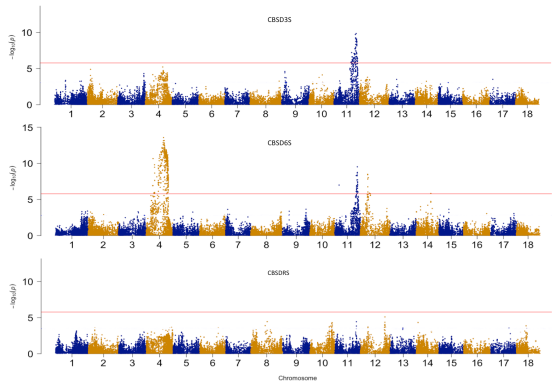
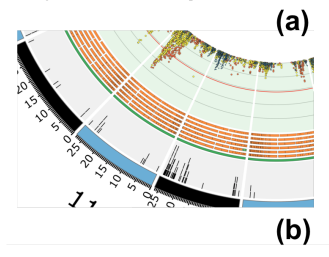


Figure 3. **Chromosome 11 region with QTL for CBSD severity** (a) 3 MAP (yellow), 6 MAP and root necrosis (blue). Outer ring black lines indicate clusters of NBS-LRR genes (Lozano et al 2015). Intermediate ring indicate regions homozygous (G/G)(blue) or heterozygous (G/E)(green) for *M. glaziovii* allele and the proportion that were homozygous for the *M. esculenta* allele (E/E)(orange) on seven clones. (b) LD association plot, 2 Mb region in chromosome 11, top SNP indicated in red, annotated genes within that region are indicated in the panel below.



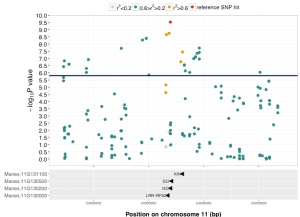


Figure 4. Chromosome 4 region with QTL for CBSD severity with introgression segment (a) 3 MAP (yellow), 6 MAP and root necrosis (blue). Outer ring black lines indicate clusters of NBS-LRR genes (Lozano et al 2015). Intermediate ring indicate regions homozygous (G/G)(blue) or heterozygous (G/E)(green) for *M. glaziovii* allele and the proportion that were homozygous for the *M. esculenta* allele (E/E)(orange) on seven clones. (b) Introgression region on chromosome 4 (colors description) are the same as the aforementioned),Nam: Namikonga,w: wild *M. glaziovii*, cm: CM330645,Alb:Albert, P1: panel 1 clones and P2 panel 2 clones.

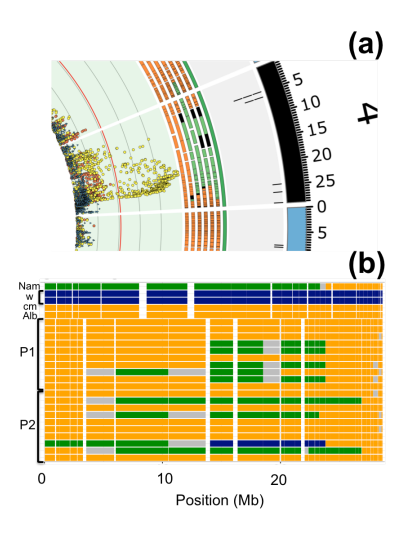


Figure 5. **Cross validation results for CBSD severity**. 3 MAP (CBSD3S), 6 MAP (CBSD6S) and Root necrosis (CBSDRS). x-axis: predictive accuracy and y-axis: genomic prediction model.

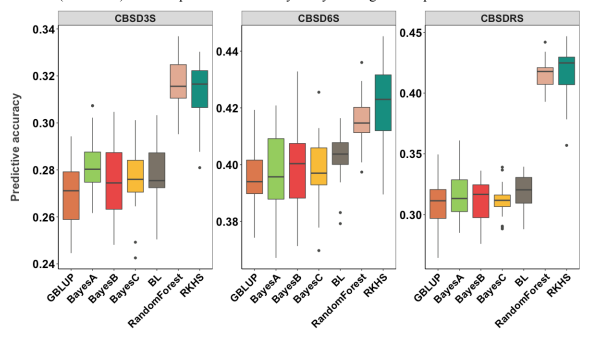


Table 1. Broad sense heritability (H²) and SNP heritability (h²) of foliar and root CBSD severity. Broad-sense heritability (H²) values were calculated using the variance components obtained from a model fitted using the *lmer* function from the lme4 R package. SNP heritability values were calculated using the variance components obtained obtained from a model fitted using the EMMREML R package. Heritability values estimates were calculated for sets 1 and 2 separately.

Trait	H^2	h^2	LOCATION-YEAR	Panel
CBSD3S	0.11	0.32	NAMULONGE	1
CBSD6S	0.31	0.39	NAMULONGE	1
CBSDRS	0.55	0.59	NAMULONGE	1
CBSD3S	0.43	0.48	NGETTA	1
CBSD6S	0.51	0.53	NGETTA	1
CBSDRS	0.73	0.72	NGETTA	1
CBSD3S	0.27	0.29	KASESE	1
CBSD6S	0.21	0.27	KASESE	1
CBSDRS	0.39	0.47	KASESE	1
CBSD3S	0.61	0.17	MULTI LOCATION	1
CBSD6S	0.35	0.31	MULTI LOCATION	1
CBSDRS	0.37	0.34	MULTI LOCATION	1
CBSD3S	0.60	0.37	NAMULONGE	2
CBSD6S	0.60	0.32	NAMULONGE	2
CBSD9S	0.68	0.34	NAMULONGE	2
CBSDRS	0.24	0.53	NAMULONGE	2
CBSD3S	0.63	0.28	SERERE	2
CBSD6S	0.60	0.28	SERERE	2
CBSD9S	0.73	0.34	SERERE	2
CBSDRS	0.15	0.48	SERERE	2
CBSD3S	0.56	0.27	KAMULI	2
CBSD6S	0.62	0.29	KAMULI	2
CBSD9S	0.75	0.34	KAMULI	2
CBSDRS	0.28	0.44	KAMULI	2
CBSD3S	0.42	0.28	MULTI LOCATION	2
CBSD6S	0.47	0.34	MULTI LOCATION	2
CBSD9S	0.56	0.38	MULTI LOCATION	2
CBSDRS	0.25	0.33	MULTI LOCATION	2

- 785 Supplementary figure 1. Cassava brown streak disease symptoms on leaves and roots of 786 sampled plants; Severity Score from 1 (no visible symptoms) to 5 (severely disease plants. (a) 787 leaf veins chlorosis severity progresses with severity score, (b) dark brown necrotic areas within 788 storage roots severity scale. 789 790 **Supplementary figure 2.**Panel 1 phenotypic distribution of CBSD severity traits. 791 (A) deregressed BLUPs distribution of CBSD 3 months foliar severity, (B) deregressed BLUPs 792 distribution of CBSD 6 months foliar severity, (C) deregressed BLUPs distribution of CBSD 12 793 months root severity
- 795 **Supplementary figure 3.**Panel 2 phenotypic distribution of CBSD severity traits.

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- 796 (A) deregressed BLUPs distribution of CBSD 3 months foliar severity, (B) deregressed BLUPs
- 797 distribution of CBSD 6 months foliar severity, (C) deregressed BLUPs distribution of CBSD 9
- 798 months foliar severity, (D) deregressed BLUPs distribution of CBSD 12 months root severity
- 800 Supplementary figure 4. Correlation plots between de-regressed BLUPs for foliar and root
- symptoms. De-regressed BLUPs were calculated for different locations in panel 1 and panel 2.
- 803 **Supplementary figure 5**. GWAS results for CBSD severity in panel 1 measure at Kasese.(a)
- scoring CBSD 3 months foliar severity (b) 6 CBSD 6 months foliar severity and (c) root necrosis
- severity. Red line Bonferroni correction. Blue line log_{10} P-value = 3.8.
- 807 **Supplementary figure 6.** GWAS results for CBSD severity in panel 1 measure at Ngetta.(a)
- scoring CBSD 3 months foliar severity (b) 6 CBSD 6 months foliar severity and (c) root necrosis
- severity . Red line Bonferroni correction. Blue line log_{10} P-value = 3.8.
- Supplementary figure 7. GWAS results for CBSD severity in panel 1 measure at Namulonge.
- 812 (a) scoring CBSD 3 months foliar severity (b) 6 CBSD 6 months foliar severity and (c) root
- 813 necrosis severity. Red line Bonferroni correction. Blue line \log_{10} P-value = 3.8.

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Supplementary figure 8. GWAS results for CBSD severity in with a multilocation dataset of panel 1 (a) scoring CBSD 3 months foliar severity (b) 6 CBSD 6 months foliar severity and (c) root necrosis severity. Red line Bonferroni correction. Blue line \log_{10} P-value = 3.8. **Supplementary figure 9.** GWAS results for CBSD severity in panel 2 measure at Kamuli. (a) scoring CBSD 3 months foliar severity (b) 6 CBSD 6 months foliar severity (c) 9 CBSD 9 months foliar and (c) root necrosis severity. Red line Bonferroni correction. Blue line log₁₀ Pvalue = 3.8. **Supplementary figure 10**. GWAS results for CBSD severity in panel 2 measure at Namulonge. (a) Scoring CBSD 3 months foliar severity (b) 6 CBSD 6 months foliar severity (c) 9 CBSD 9 months foliar and (c) root necrosis severity. Red line Bonferroni correction. Blue line log₁₀ Pvalue = 3.8. **Supplementary figure 11.** GWAS results for CBSD severity in panel 2 at Serere. (a) Scoring CBSD 3 months foliar severity (b) 6 CBSD 6 months foliar severity (c) 9 CBSD 9 months foliar and (c) root necrosis severity. Red line Bonferroni correction. Blue line log_{10} P-value = 3.8. Supplementary figure 12. GWAS results for CBSD severity in with a multilocation dataset of panel 2 (a) scoring CBSD 3 months foliar severity (b) 6 CBSD 6 months foliar severity (c) 9 CBSD 9 months foliar and (c) root necrosis severity. Red line Bonferroni correction. Blue line log_{10} P-value = 3.8. Supplementary figure 13. local LD in chromosome 4. Plot of the mean LD score for each marker .With a smooth line representing a relative measure of the local LD in chromosome 4. Dots are colored with the -log₁₀ P-value for the association test for CBSD severity six months after planting.

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Supplementary figure 14. Introgressions segment detection. For each clone of the two GWAS panels we calculated the proportion of genotypes that were homozygous (G/G) or heterozygous (G/E) for M. glaziovii allele and the proportion that were homozygous for the M. esculenta allele (E/E). Supplementary figure 15. (a) GWAS results for 6MAP CBSD severity panels 1 and 2 (b) GWAS Results after correction including markers in chromosome 12 as a covariate. Supplementary figure 16. Multi-kernel GBLUP approach by fitting three kernels constructed with non-overlapping SNPs (MAF> 0.01) from chromosomes 4, 11 and SNPs from the other chromosomes. Crossvalidation GS predictive accuracies results for CBSD severity were calculated using the multilocation dataset of the combined panels. Scoring CBSD 3 months foliar severity (CBSD3S), CBSD 6 months foliar severity (CBSD6S) and root necrosis severity (CBSDRS). **Supplementary Table 1.**Pedigree information from GWAS panels 1 and 2. Details are shown on the parental lines per clone and selected traits that came from the maternal side. Supplementary table 2. Correlation values across locations in panel 1 and panel 2. (A) Correlation of deregressed BLUPs across locations within traits in panel 1 measured in three locations.(B) Correlation of deregressed BLUPs across locations within traits in panel 2 measured in three locations **Supplementary table 3**. Correlation values across locations and traits in panel 1 and panel 2. (A) Correlation of deregressed BLUPs across locations and traits in panel 1 measured in three locations.(B) Correlation of deregressed BLUPs across locations and four traits in panel 2 measured in three locations

Supplementary table 4. Panel 1 and 2 and combined panels GWAS results. Gene annotation is only shown for significant SNPs.

Supplementary table 5. Explained variance of phenotypic traits. Details are shown of the reference SNP, the -log10(pval)(score),chromosome and explained variance.

Supplementary table 6.Genomic prediction accuracy values. (A) Cross validation results using 7 GS models for CBSD severity prediction of 3 MAP CBSD3S, 6 MAP CBSD6S and Root necrosis (CBSDRS) (B) Multi-kernel GBLUP crossvalidation by fitting three kernels constructed with non-overlapping SNPs (MAF> 0.01) from chromosomes 4, 11 and SNPs from the other chromosomes. RKHS = Reproducing kernel Hilbert spaces regression, Total accuracy is the accuracy obtained by following the GBLUP multikernel approach.