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PROJECT 3 ABSTRACT

(1 Page Limit)

We will characterize the common set of 49 elite lines and 100 RILs from the Purdue program that have type I and/or type II FHB resistance (the same lines that were characterized in 2011). Plots will be a 1m row. The study will consist of 3 replications, statistical RCBD, grown in the misted nursery, seeded in disced corn residue. We will record data for FHB incidence, severity, index, FDK and submit composite samples of each line for DON analysis. Genetics studies in SWW suggest that there are several unique sources of FHB resistance that are controlled by several OTL with moderate to small effects, thus complicating traditional MAS approaches. Consequently, recurrent selection is likely to be an effective breeding tool to accumulate favorable alleles. We propose to develop knowledge of the types of resistance, the genetics of this resistance, and efficient breeding methodologies for improving FHB resistance in SWW. Specifically, we will determine the genetic structure of FHB resistance in SWW, develop models to implement genomic selection (GS) for multiple FHB traits in SWW. Our approach will be to phenotype and genotype a set of 49 elite SWW lines by each of the 7 collaborators, plus each collaborator will phenotype 100 RILs developed in their respective programs that have good FHB resistance; a total of 749 lines. The populations will be phenotyped for multiple FHB traits (INC, SEV, IND, FDK, DON) in field trials. The data will be used in an association analysis (AA) to determine the genetics of resistance in SWW to estimate the effect of QTL on multiple mechanisms of FHB resistance and the frequency of favorable alleles in the SWW. The data will also be used to develop a GS model that predicts the breeding value of individuals using estimated gene effects from the entire genome. The model can be used in subsequent selection cycles to choose superior parents with little or no phenotyping. Collectively, AA and GS will allow us to 1) select parents that are fixed for the same major genes, 2) design crosses that facilitate combining different genes and multiple mechanisms of FHB resistance, and 3) allow selection of superior individuals without phenotyping thereby reducing time per breeding cycle.