Scab epidemics were widespread in Virginia in 1998 and resulted in significant losses in yield and quality. In 2003 and 2009, scab epidemics devastated much of the wheat crop in Virginia. Scab epidemics in Virginia also were significant in 2013 and 2017. Currently, production of cultivars having moderate FHB resistance derived predominantly from native sources, and fungicide applications offer the primary means of disease control. However, neither control strategy provides optimal protection in years of severe epidemics. Enhanced efforts are needed to develop cultivars and superior parental lines having gene Fhb1 combined with other validated genes and QTL derived from both exotic and native sources. Simultaneous use of both marker-assisted selection (MAS) and Doubled Haploid (DH) methods offers great potential to accelerate the development of cultivars having FHB resistance superior to that conferred solely by gene Fhb1 or QTL from native sources.

Research will continue to focus on enhanced MAS breeding efforts in selection of parents, designing crosses, gene introgression and pyramiding, population enrichment, and selection of pure lines. Marker haplotypes of parents for validated FHB resistance QTL and other traits of importance such as dwarfing genes, disease and insect resistance, rye translocations, and quality are assessed and utilized to enhance breeding efficiency. Markers linked to 15 scab resistance genes located on wheat chromosomes 2D, 3B (Fhb1), and 5A of Ning 7840, 1B of Jamestown, 1A and 2A of Tribute, 3B and 4B of Ernie, 2B and 3B of Bess, 3B of Massey, and 1A, 4A, and 6A of Neuse are being used to screen, characterize and select parents and their progeny for FHB resistance genes. Plants from top cross populations developed in 2015 (13), 2016 (13), 2017 (15), and 2019 (6) comprised of parents having FHB resistance QTL derived from Ning 7840 and other native sources have been enriched via MAS for FHB resistance and will be advanced using either the DH and/or pedigree method to derived pure lines.

Research will also focus on Genomic Selection to provide information on genomic estimated breeding values (GEBV) to aid in selection of superior elite lines, parents and crosses. Parental lines will be selected and crosses made on the basis of phenotypic and genotypic data, and GEBV derived from GS models including both agronomic and disease data from the Gulf Atlantic and Mason Dixon regional nurseries that are evaluated in uniform yield trials and for FHB resistance in scab nurseries. Genotypic data is routinely generated and available for wheat entries in regional, uniform, and scab nurseries by the Genotyping Center at Raleigh. New elite lines derived from the GS project will be genotyped and evaluated in observation yield tests at two locations and in the scab nursery at Mt. Holly. MAS breeding efforts and GS in conjunction with doubled haploid (DH) technology will aid in increasing breeding efficiency and development of FHB resistant varieties.