

Development of High Throughput KASP SNP Markers for Wheat Curl Mite **Resistance and their Application in Marker-assisted Breeding**



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INTRODUCTION

- Biotic stresses coupled with abiotic stresses are the major limiting factor for crop production feeding the 9 billion people by 2050. Wheat curl mite (*Aceria tosichella* Keifer, WCM) have been a
- persistent concern to farmers and researchers for at least six decades
- impacting forage and grain yields, and crop water use efficiency.
- At least 90 plant species hosts have been reported across 34 countries around the world for WCM.
- WCM transmit Wheat streak mosaic virus (WSMV), Wheat mosaic virus (WMoV), Triticum mosaic virus (TriMV), and Brome streak mosaic virus (BrSMV) (Navia et al, 2013).
- Yield loss by mite-virus complexes have been reported up to 100% at

Genotyping and Mapping:

- **DNA** was extracted from all the lines including parental and check using CTAB protocol.
- DNA samples were sent to the USDA genotyping center, Fargo, ND for 90K SNP genotyping.
- Illumina's iSelect infinium assay platform was used to genotype SNP.
- Fluorescent intensities of A and B allele captured by iScan were analyzed using Genome studio software.
- A total of 90,900 markers including SNPs , DArT, SSR and STS, were screened.
- 7985 polymorphic markers were utilized to construct the genetic

Screening of Resistance source of WCM present in hard red

winter wheat

- Out of 43 resistant lines, 28 have 1AL.1RS translocation containing
 - *Cmc3* gene, while eight lines derived from TAM112 showed
- resistance even when no 1AL.1RS translocation was present.
- TAM 112 is in the pedigree of 20 resistant lines (Table 2).
- Results suggest that, in addition to 1AL.1RS translocation, there is
- another gene governing WCM resistance in TAM 112.
- In general line with either 1AL.1RS or TAM112 in its pedigree
- exhibited a high degree of resistance to WCM.

II. Mapping of the WCM resistant gene in TAM 112

field level and up to 7% state wide (Appel et al, 2001, Slykhuis, 1976).

- TAM 112, most widely grown hard red winter wheat (HRWW), is noted for resistance to WCM.
- With the application of molecular marker, we are trying to predict the genotype performance and at the same time help to accelerate the breeding process.

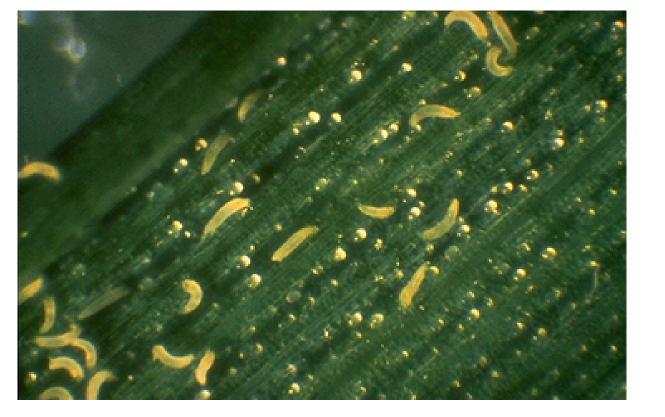






Figure 2: (Left to Right): **Susceptible and Resistant line**

OBJECTIVES

Screen the currently grown winter wheat to know their resistance to

map.

JoinMap version 4.0 was used to create the linkage map.

RESULTS AND DISCUSSIONS

0.0 CmcTAM112

13.0 ++- SNP1

17.4 SNP2 17.5 SNP3 19.3 SNP4

28.3 + SNP5

39.4 🕂 SN P6 41.8 ++ SNP7

45.8 +++ SNP8

64.3 U SNP13

Figure 3: Wheat curl

mite resistance gene

mapped

No

No

50.9 SN P9 51.3 SN P10 51.8 SN P11 SNP12

	NRP				
WCM reaction	Ν	SRPN	TXE	CEB	Total lines
R	3	9	16	17	45
S	34	30	21	33	118
Н	3	1	3	2	9
Total	40	40	40	52	172

 Table 1: Response of wheat lines across
the four tests to wheat curl mite

WCM reaction	SRPN	TXE	CEB	Total lines
R	6	10	4	20
S	0	1	0	1
Н	0	2	0	2
Total	6	13	4	23

 Table 2: Response of wheat lines with TAM
112 in their pedigree across the four tests to wheat curl mite

- 99 homozygous resistant: 5 heterozygous: 20 homozygous susceptible.
- TAM 112 was consistently resistant to the WCM colony from Texas collection.
- In addition to the 1AL.1RS translocation, another WCM resistant

gene in TAM 112 was mapped (Figure 3).

- The resistance gene present in TAM 112 was designated as **CmcTAM112.**
- SNP1 was at 13.0 cM away from the *Cmc_{TAM112}* proximally, among 13 SNP markers mapped onto the same linkage group.

III. Development of KASPar SNP for Cmc_{TAM112}

- Two closely linked array SNPs were converted into Kompetitive allele specific PCR (KASP) SNP for high throughput screening (Figure 2 & 3).
- The KASP SNP s were mapped back onto the regions.
- These two KASP SNP were used to screen a collection of cultivars and elite lines, only two out of 33 did not match between genotypes and phenotypes (Table 3).

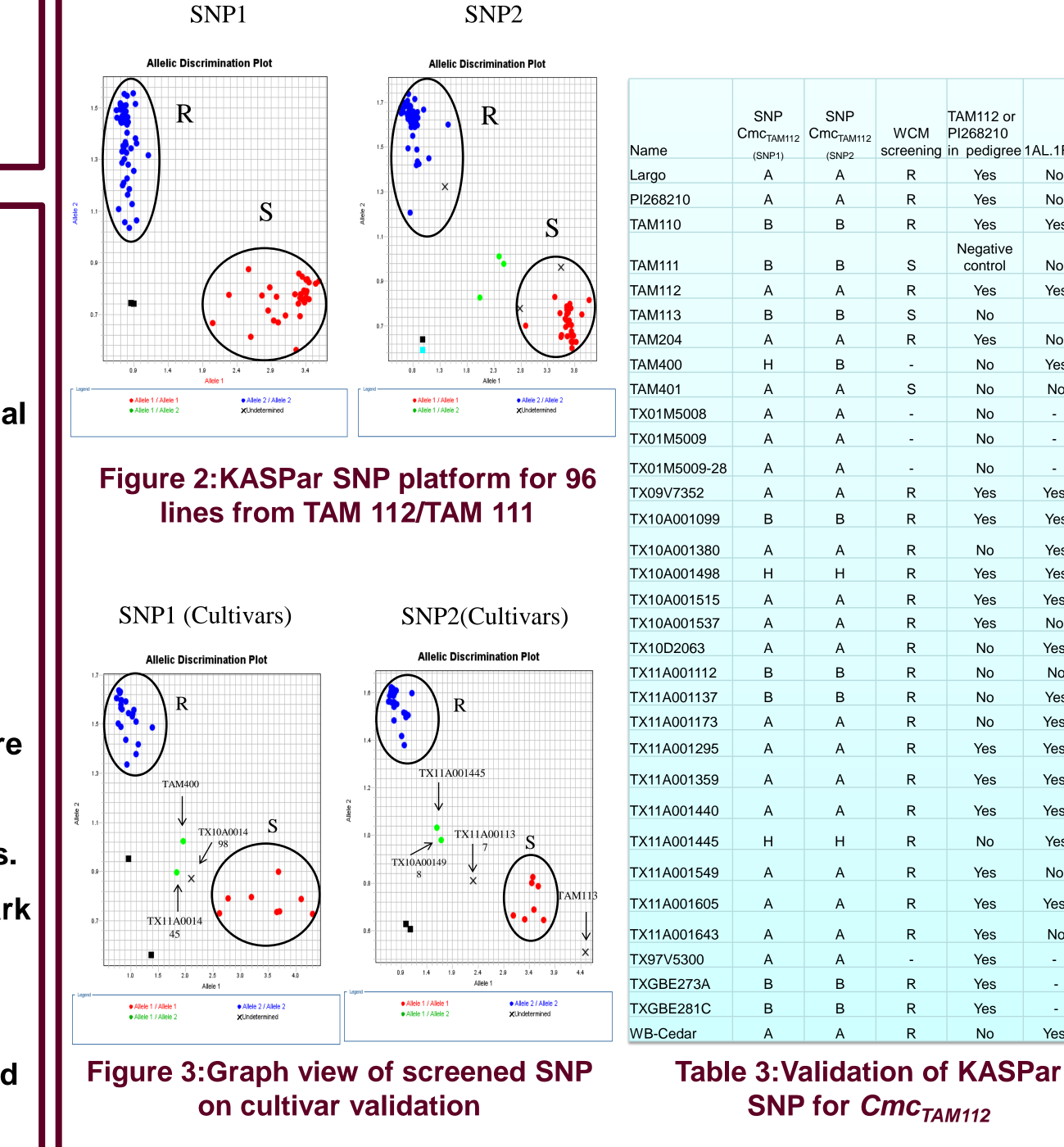
WCM.

- Map the Wheat curl mite resistance gene in the TAM 112.
- Develop a diagnostic KASP SNP markers for wheat curl mite resistance.

MATERIALS AND METHODS

Phenotyping:

- For screening objective, 40 wheat lines each from Northern Regional Performance Nursery (NRPN), Southern Regional Performance Nursery (SRPN), Texas Elite (TXE) lines, and 52 Cultivar and Elite **Breeding (CEB) lines were screened.**
- For mapping objective, 124 F7 recombinant inbreed lines from TAM 112/TAM 111 were screened.
- TAM 112 (resistant check), TAM 111, Karl 92 (susceptible check) were also screened.
- A total of 15 plants per line were screened in replicated experiments.



These SNPs are promising for marker-assisted breeding as wheat

curl mite screening is very time and money consuming.

CONCLUSIONS

Resistance in TAM 112 is governed by at least two genes, One is

originated from wheat-rye translocation, while the other from Ae. tauschii.

Because of the two resistance genes, TAM 112 was consistently

resistant in the screening.

> As both of the genes are single dominant in nature, the resistance

can be easily transferred to other adapted wheat cultivars in wheat

breeding program.

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REFERENCES



period were maintained inside the growth chamber.

Each plant was infested with WCM at the two-leaf stage.

Scoring on resistant (R) or susceptible (S), was done on the first and

second week after double infestation (Figure 2)

Appel, J., E. DeWolf, W. Bockus, and T. Todd. 2011. Preliminary 2011 Kansas wheat disease

loss estimates. Kansas Dep. Of Agric., Topeka, KS.

Navia, D., R. S. Mendonca and A. Skoracka et al. 2013. Wheat curl mite, Aceria tosichella,

and transmitted viruses: an expanding pest complex affecting cereals crops.

Slykhuis, J. T. 1976. Virus and virus like diseases of cereal crops. Annu. Rev. Phytopathol.



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