

binant), resulted in linkage with the mutation. The SSR marker *umc1962*, tested on the same segregating population, resulted in complete linkage with E3443_23, confirming the mapping data previously described. The region containing *de18* was edged by the SSR markers *umc1367* and *umc2069* (Pasini et al., Mol. Breed. 22:527-541, 2008). Furthermore, a QTL for kernel weight was localized in the same bin 10.03 (Austin and Lee, Theor. Appl. Genet. 92: 817-826, 1996). The mutant *orange pericarp 2* (*orp2*) was placed on chromosome 10 bin 10.03, near the SSR marker *bnlg1712*, at least 38 cM from *de18*. Moreover, it is also known that other candidate genes involved in auxin efflux transports, such as *ZmPIN1a*, *ZmPIN1b* and *ZmPIN1c* (Gallavotti et al., Plant Physiol. 147:1913-1923, 2008), were localized on different chromosomes (chromosome 9, 5 and 4 respectively). In the same manner, the mutant *orange pericarp1* (*orp1*), involved in tryptophan biosynthesis, was mapped to chromosome 4.

In order to build a high resolution map around the *de18* mutant, a large F3 population was developed from the cross A69Y x *de18*. This population, consisting of 391 homozygous wild type, 52 homozygous mutant and 188 heterozygous F3 families, will be screened for recombinants in the interval of flanking SSR markers. Recombinant pools will be screened with AFLP markers to saturate the region.

Maize long oligonucleotide microarrays (Maize Oligonucleotide Array Project, version 1, 45k) were used to determine the differential gene expression between the mutant *de18* and its wild type B37. mRNAs were extracted from seeds at four different stages of development, 7, 14, 21 and 28 days after pollination, and used to perform the hybridization. After normalization and statistical analysis of data groups, differentially expressed genes were detected. We identified many genes involved in the process of endosperm development and linked to auxin metabolism. To confirm the accuracy and reproducibility of the microarray results, 4 differentially expressed genes were selected for confirmation by real-time PCR. The genes selected were: anthranilate phosphoribosyltransferase, auxin response factor 8, *dull2* starch synthase and zein alpha precursor 19kDa. The real-time PCR results showed that the expression trends of these genes were partially consistent with those derived from the microarray analysis. In addition, *DR5*, another gene involved in auxin transport, was tested in quantitative PCR, but no significant differences between *de18* and B37 were detected.

Mutator-induced alleles at the *reduced grain filling1* locus of maize

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The defective kernel mutant *rgf1*, *reduced grain filling1*, induces alterations in both pedicel and transfer layer development, resulting in reduced starch accumulation and a final grain weight of 30% that of the wild type (Maitz et al., Plant J. 23:29-42, 2000). The mutant is dominant, with gene dosage-dependent expression of the phenotype. Selfing of heterozygous kernels gave rise to F2 cobs on which four classes of kernel size were segregating. *rgf1* was mapped to chromosome 2 bin 2.04 within a 0.1 cM interval with the aid of recombinant pools derived from an F3 population of 1406 lines. We report here the isolation of new mutants at the *rgf1* locus that we have obtained by tagging using the *Mutator* (*Mu*)

transposable element, in collaboration with the Dipartimento Produzione Vegetale, University of Milan, Italy (F. Salamini).

A *rgf1/Rgf1* strain containing *Mu* was used as the female parent in crossing with a wild type *Rgf1/Rgf1* line. About 2,500 F1 ears were obtained, for a total of 650,000 seeds; we expected to obtain ears bearing kernels with dosage-dependent reduced size, half of them with the allelic composition *rgf1/rgf1/Rgf1* and the other half *rgf1/Rgf1/Rgf1*. Of the 2,500 cobs, 13 carried wild type kernels, probably arising from *Mu* insertion within the *rgf1* locus: 10 ears harboured one wild type kernel, while sectors of wild type kernels were observed on the remaining 3 cobs (Figure 1). Plants



Figure 1. Phenotypes of three F₁ ears derived from the tagging experiment with the putative wild type *Mu*-induced kernels indicated by arrows. The new mutants are named 11508-3, 11508-7 and 11508-10.

were grown from wild type kernels and selfed. In six cases (11508-1, 11508-3, 11508-5, 11508-6, 11508-7 and 11508-10), the segregation data were compatible with the presence of *Mu*-induced alleles. The frequency of these *rgf1-Mu* dependent alleles was 9.2×10^{-6} . In the F₂ generation, the 11508-1 *rgf1-Mu* allele yielded a 3 wild type:1 *rgf1* segregation, which is compatible with the presence of a recessive *rgf1* allele generated by *Mu*. This hypothesis will be tested by crossing 11508-1 to the wild type. Another group of four putative mutants (11508-3, 11508-5, 11508-6 and 11508-10) produced F₂ cobs with all wild type kernels, suggesting that the insertion of *Mu* into the *rgf1* locus resulted in a wild type allele. F₂ progenies will be developed for each mutant and wild type individuals will be screened with the two SSR markers flanking *rgf1*. It is expected to find the dominant pattern at the SSR loci in some wild type kernels. For a third type of mutant, 11508-7, the F₂ segregation was compatible with a lethal recessive allele caused by *Mu* insertion. Heterozygous *rgf1/rgf1-lethal* individuals and their progenies will be analyzed for germination capability.