## Chromosome composition in an F<sub>2</sub> hexaploid x durum cross analyzed by DArT markers and MCFISH

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A major constraint to tetraploid durum wheat production in Australia is widespread susceptibility to crown rot, due to infection by Fusarium pseudograminearum. Several sources of partial resistance to this disease are available in hexaploid bread wheats and genetic markers for quantitative trait loci conditioning this resistance have been identified. We are currently attempting to transfer crown rot resistance from these hexaploid sources into susceptible tetraploid wheats. However, knowledge of the fate of D-genome material in hexaploid/tetraploid crosses is incomplete, while the degree of recombination between the A- and B-genomes of the parents in these crosses is also of critical interest. Diversity Array Technology (DArT) markers and multicolour fluorescence in situ hybridisation (MCFISH) were employed to investigate parental inheritance in the  $F_2$  progeny from a cross between the hexaploid bread wheat line '2-49' and the tetraploid durum variety 'Bellaroi'. Of the 83 F<sub>2</sub> progeny analyzed with DArT, 82 contained one or more D-genome chromosomes, either complete or partial. The marker profiles indicated that all lines possessed recombined A- and B-genome loci derived from both parents, indicating the absence of parental selfs. The majority of Aand B-genome chromosomes showed a random re-assortment of parental genes. MCFISH analysis was conducted on 28 additional plants from the same F<sub>2</sub> population. All lines contained varying numbers of D-genome chromosomes, while two plants carried A-D translocations. Investigations of F<sub>3</sub> plants from an independent 249/Bellaroi cross indicated that only 16 out of 33 plants still contained D genome material. Two F3 plants with translocations of D genome material were identified.