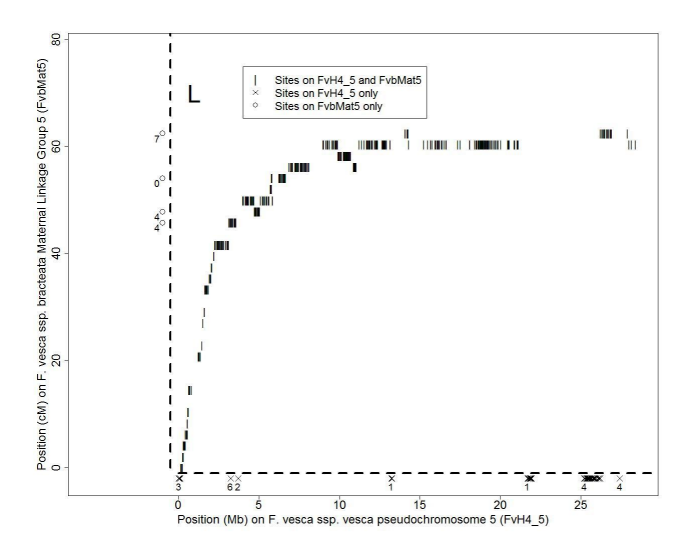
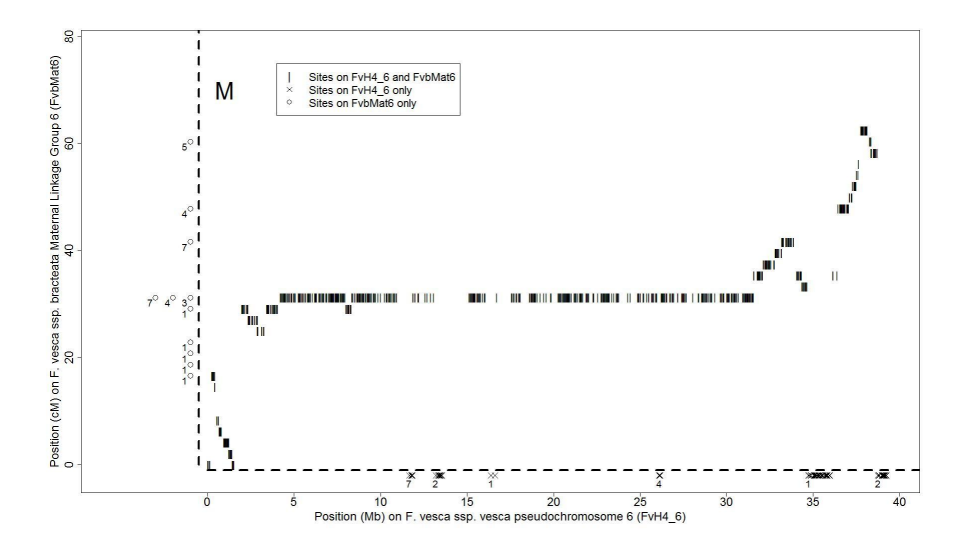


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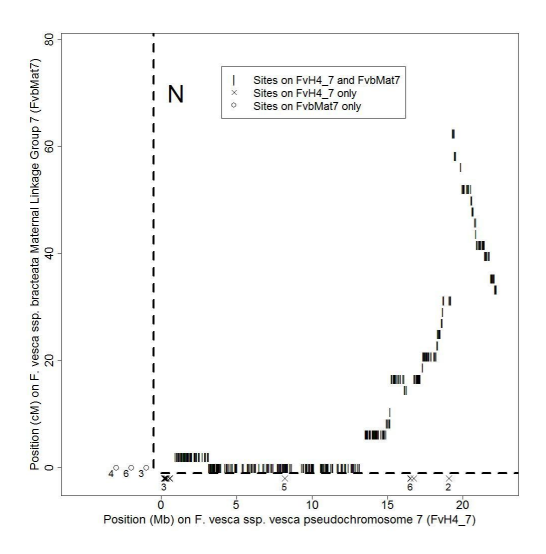


Figure S2 (A-N) Linkage groups in *F. vesca* ssp. *bracteata* plotted against the *F. vesca* ssp. *vesca* reference genome (FvH4). Each short vertical line represents an informative marker in our linkage map. Each linkage group is plotted against the FvH4 pseudochromosome with which it shares the most makers (the main pseudochromosome; Table 1). Markers that aligned somewhere in FvH4 other than the main pseudochromosome are indicated as circles on along left side of the figure, numbered according to FvH4 pseudochromosome. Markers that aligned to the main pseudochromosome but mapped to a different Fvb linkage group are indicated as x marks along the bottom of the figure, numbered according to Fvb linkage group. Markers that map to one part of the linkage group but align to a different part of the main pseudochromosome are within-chromosome translocations.