

Weng peer-reviewed journal publications

- 1) **Weng Y**, Liu DJ. Morphology, scab resistance and cytogenetics of Inter-generic hybrids between wheat and two *Roegneria* species. *Scientia Agricultura Sinica* 22: 1-7. 1989.
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- 6) **Weng Y**, Liu DJ. Advances in RFLP mapping of wheat, barley and rye (I). *Journal of World Agriculture* 11: 24-26. 1993.
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- 13) **Weng Y**, Lazar MD, Michels Jr. GJ, Rudd JC. Phenotypic mechanisms of host resistance against greenbug revealed by near isogenic lines of wheat. *Journal of Economical Entomology* 97: 654-660. 2004. (Corresponding author)
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