Figure S3. Phylogenetic tree based on Ets domain sequences of Ets transcription factor superfamily members. ETS domain sequences of squid Ets-4 homolog was aligned with those of representative ETS members and analyzed by the neighbor-joining method using Clustal W. Spi1 were taken as out-group. The squid Ets-4 homolog is most closely related to Drosophila D-ets-3 (a homolog of human Fli-1). Close relationships among the squid Ets-4, D-ets-3, and human Fli-1/Erg is supported with high bootstrap value (91%). The accession number corresponding to each ETS member is Spi1_human P17947 as an outgroup, C. elegans protein C33A11.4 CAB01862.4, Elf3_human P78545, Etv6_human P41212, POK_Drosophila Q01842, Elf-1_human NP_758961, D-ets-4_Drosophila P29775, Etv5_human P41161, Elk-1_human P19419, Erg_human P11308, Erf_human P50548, GABPA_human Q06546, Ets-1_human P14921, ETS-family transcription factor_bivalve [Chlamys farreri] AAU11487, Ets at 98B_Drosophila AAF56746, D-ets-6/Ets at 21C_Drosophila AAF51484, D-ets-3/Ets at 65A_Drosophila AAF50697, D-ets-2/pointed_Drosophila NP_524461, Fev_human NP_059991, and Fli-1_human AAH10115. Numbers at nodes indicate bootstrap values obtained 1,000 repetition. Bar indicates kimura distance calculated by Clustal W.