## Supplemental material

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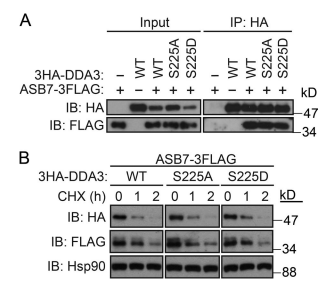


Figure S1. Phosphorylation of DDA3 has no effect on the interaction with ASB7 or stabilization of DDA3. (A) Interaction between  $3\times$ HA-DDA3 (WT, S225A, or S225D) and  $3\times$ FLAG-ASB7.  $3\times$ HA-DDA3 (WT, S225A, or S225D) and  $3\times$ FLAG-ASB7 (as indicated) were expressed in HEK293T cells in the presence of the proteasome inhibitor MG132 (10  $\mu$ M for 6 h), immunoprecipitated (IP) with anti-HA antibody, and immunoblotted (IB) with anti-HA or anti-FLAG antibody. (B) DDA3 WT, S225A, and S225D are comparably stable. HEK293T cells expressing  $3\times$ HA-DDA3 (WT, S225A, or S225D) and  $3\times$ FLAG-ASB7 were exposed to 50  $\mu$ g/ml CHX for 1 or 2 h. The lysates were subjected to Western blotting with antibodies against HA, FLAG, or Hsp90. Hsp90 is shown as a loading control.