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We identified a 116-kD protein, termed p116^{Rip}, that binds to activated RhoA under two-hybrid conditions in yeast. Pull-down experiments in mouse N1E-115 neuroblastoma cells revealed an interaction between GST-RhoA and a protein of that comigrates with p116Rip and is recognized by anti-p116Rip antibody (Fig. 5 D). While this protein does not interact with GST alone, our recent experiments indicate that it is not p116^{Rip} but an unidentified bacterial protein that has the same apparent molecular size as p116^{Rip} and cross-reacts with the anti-p116^{Rip} antibody used. Therefore, the conclusion that p116^{Rip} interacts with RhoA in N1E-115 cells is premature.

Overexpression of p116^{Rip} in N1E-115 cells mimics dominant-negative RhoA in stimulating cell flattening and neurite outgrowth. We are currently characterizing p116^{Rip} in further biochemical detail and evaluating the relationship between p116^{Rip} and RhoA action.

We apologize for any additional work that our error may have caused other investigators.
