

A target protein for nalidixic and oxolinic acids in *Escherichia coli*, the *nalA* gene product (Pnal), was purified to homogeneity as judged by gel electrophoresis, using an in vitro complementation assay. The nicking-closing activity is distinct from *E. coli* omega protein in several properties, including the ability to relax positively super-twisted DNA. DNA gyrase from a strain with a *nalA* mutation conferring drug resistance (*nalA*(r)) is 1/100 as sensitive to oxolinic and nalidixic acids with respect to inhibition of super-twisting and induction of the pre-linearization complex. A polypeptide of this molecular weight is uniquely induced by a lambda *nalA* transducing phage, thereby showing that the purified Pnal is a product of the *nalA* gene. DNA gyrase preparations and Pnal catalyze a third reaction sensitive to .nalidixic and oxolinic acids, the ATP-independent relaxation of super-twister DNA