

## AR50756PU-S

## Human XPA / XPAC (1-273, His-tag) - Purified

### Alternate names:

DNA repair protein complementing XP-A cells, Xeroderma pigmentosum group A-complementing protein

### Quantity:

0.1 mg

### Concentration:

1 mg/ml (determined by Bradford assay)

### Background:

XPA, also known as DNA repair protein complementing XP-A cells, belong to the XPA family. This protein is involved in DNA excision repair. It Initiates repair by binding to damaged sites with various affinities, depending on the photoproduct and the transcriptional state of the region. Defects in XPA are a cause of xeroderma pigmentosum complementation group A (XP-A), which is a rare human autosomal recessive disease characterized by solar sensitivity, high predisposition for developing cancers on areas exposed to sunlight and, in some cases, neurological abnormalities.

### Uniprot ID:

[P23025](#)

### NCBI:

[NP\\_000371](#)

### GenelD:

[7507](#)

### Species:

Human

### Source:

E. coli

### Format:

**State:** Liquid purified protein

**Purity:** >85% by SDS - PAGE

**Buffer System:** 20 mM Tris-HCl buffer (pH 8.0) containing 0.4M Urea, 10% glycerol

### Description:

Recombinant human XPA protein, fused to His-tag at N-terminus, was expressed in E.coli.

#### AA Sequence:

MGSSHHHHH SSGLVPRGSH MGSMAADGA LPEAAALEQP AELPASVRAS IERKRQRALM  
LRQARLAARP YSATAAAATG GMANVKAAPK IIDTGGGFIL EEEEEEEQKI GKVVHQPGPV  
MEFDYICEE CGKEFMDSYL MNHFDLPTCD NCRDADDKHK LITKTEAKQE YLLKDCDLEK  
REPPLKFIVK KNPHHSQWGD MKLYLKLQIV KRSLEVWGSQ EALEEAKEVR QENREKMKQK  
KFDKKVKELR RAVRSSVWKR ETIVHQHEYG PEENLEDDMY RKTCTMCGHE LTYEKM

**Molecular weight:** 33.8 kDa (296aa)

### Storage:

Store undiluted at 2-8°C for one week or (in aliquots) at -20°C to -80°C for longer. Avoid repeated freezing and thawing. Shelf life: one year from despatch.

### General Readings:

Pan Y.R., et al. (2009) Cell Cycle 8:655-664 James, et al. (2005) Andrews' Diseases of the Skin: Clinical Dermatology. (10th ed.). Saunders.