

Molecular Biology

Entry Clone Accession: BC008716

Entry Clone Source: MGC

SGC Construct ID: DDR1A-c002

Protein Region: P601-V913

Vector: pFB-LIC-Bse

Tag: N-6HIS;N-TEV

Host: DH10Bac

Sequence (with tag(s)):

MGHHHHHSSGVLDGTENLYFQSMPRVDFPRSRLFKEKLGEQFGEVHLCEVDSPQDL
VSLDFPLNVRKGHPLLAVKILRPDATKNARNDLKEVKIMSRLKDPNIIRLLGVCVQDDP
LCMITDYMENGDLNQFLSAHQLEDKAAEGAPGDQAAQGPTISYPMLLHVAAQIASGM
RYLATLNFVHRDLATRNCLVGENFTIKIADFGMSRNLYAGDYYRVQGRAVLPIRWMAWE
CILMGKFTTASDVWAFGVTLWEVLMCRAQPGQLTDEQVIENAGEFFRDQGRQVYLSR
PPACPQGLYELMLRCWSRESEQRPPFSQLHRFLAEDALNTV

Sequence after tag cleavage:

SMPRVDFPRSRLFKEKLGEQFGEVHLCEVDSPQDLVSLDFPLNVRKGHPLLAVKILRP
DATKNARNDLKEVKIMSRLKDPNIIRLLGVCVQDDPLCMITDYMENGDLNQFLSAHQL
EDKAAEGAPGDQAAQGPTISYPMLLHVAAQIASGMRYLATLNFVHRDLATRNCLVGEN
FTIKIADFGMSRNLYAGDYYRVQGRAVLPIRWMAWE CILMGKFTTASDVWAFGVTLWEV
LMCRAQPGQLTDEQVIENAGEFFRDQGRQVYLSRPPACPQGLYELMLRCWSRESEQR
PPFSQLHRFLAEDALNTV

DNA Sequence:

CCATGGGCCACCATCATCATCATTCTCTGGTAGATCTGGTACCGAGAACCTG
TACTTCCAATCCATGCCAGAGTGGATTCCCTCGATCTGACTCCGCTTCAAGGAGA
AGCTTGGCGAGGCCAGTTGGGAGGTGCACCTGTGAGGTCGACAGCCCTCAA
GATCTGGTTAGTCTGATTCCCCCTTAATGTGCGTAAGGGACACCCCTTGCTGGTAGC
TGTCAAGATCTACGCCAGATGCCACCAAGAATGCCAGGAATGATTCTGAAAGA
GGTGAAGATCATGTCGAGGCTCAAGGACCCAAACATCATTGGCTGCTGGCGTGTG
TGTGCAGGACGACCCCCCTCTGCATGATTACTGACTACATGGAGAACGGGACCTCAAC
CAGTCCTCAGTGCCACCAAGCTGGAGGACAAGGCAGCCGAGGGGCCCTGGGA
CGGGCAGGCTGCGCAGGGGCCACCATCAGCTACCAATGCTGCTGCATGTGGCAGC
CCAGATCGCCTCCGGCATGCCTATCTGCCACACTCAACTTGTACATCGGACCTG
GCCACGCGGAACTGCCTAGTTGGGAAAATTCAACATCAAATCGCAGACTTGGC
ATGAGCCGGAACCTCTATGCTGGGACTATTACCGTGTGCAGGGCCGGCAGTGTG
CCATCCGCTGGATGGCCTGGAGTGCATCCTCATGGGAAGTTCACGACTGCGAGTG
ACGTGTGGCCTTGGTGTGACCTGTGGAGGTGCTGATGCTCTGTAGGGCCCAGC
CCTTGGCAGCTCACCGACGAGCAGGTATCGAGAACGCCGGGGAGTTCTCCGGG
ACCAGGGCCGGCAGGTGTACCTGTCCCAGGCCCTGCCTGCCGCAGGGCTATATG
AGCTGATGCTCGGTGCTGGAGCCGGAGTCTGAGCAGCGACCACCCCTTCCCAGC
TGCATCGGTTCTGGCAGAGGATGCACTAACACGGTGTGACAGTAAAGGTGGATAC
GGATCCGAATTGAGCTCCGTCACAAGCTT

Protein Expression

Medium: Insect Xpress

Antibiotics: Ampicillin

Procedure: 3L Sf9 cells at a density of 2×10^6 /ml were infected with recombinant DDR1 baculovirus (virus stock P2; 3ml of virus stock per 1000 ml cell culture). Cells were shaken at

110rpm at 27°C in an Infors shaker with a radius of 25 mm. 72 hours after infection the cultures were harvested by centrifugation at 900 g for 20 mins. Cell pellets were resuspended in 50 mM HEPES pH 7.5, 500 mM NaCl, 5 mM Imidazole, 5 % glycerol plus Merck Set III protease inhibitor and stored at -20°C.

Protein Purification

Procedure: Stored cell pellets were thawed and sonicated on ice for 5 mins (5 min cycle of 5 seconds on/10 seconds off). 1ml of 5% PEI (0.125 %) was added to the lysate. Lysate was centrifuged for 60 min at 4 degrees in a JA 25.50 rotor at 60,000 g (22,000 rpm). The supernatant was loaded onto Ni-IMAC resin (equilibrated in 50 mM HEPES pH 7.5, 500 mM NaCl, 5 mM Imidazole, 5 % glycerol) and rotated gently at 4°C for 1 hour. This was applied to a gravity column and washed and eluted with 50 mM HEPES pH 7.5, 500 mM NaCl, 5% glycerol, and 30-250mM imidazole. The final fraction contained 1M imidazole. Fractions containing protein (as seen by SDS-PAGE) were treated with TEV protease overnight at 4°C prior to gel filtration on an S200 gel filtration column using 50 mM HEPES pH 7.5, 300 mM NaCl and 0.5 mM TCEP as the running buffer. Additional purification was carried out by loading fractions from gel filtration onto a Ni-IMAC gravity column equilibrated in 50 mM HEPES pH 7.5, 300 mM NaCl and 0.5 mM TCEP and collecting the flow through, wash and first elution (30 mM imidazole). The flow through was concentrated to 8.4 mg/ml using a centrifugal concentrator.

Columns: Column 1: Ni-IMAC, then TEV; Column 2: S200; Column 3: ni rebind; Column 4: None; Column 5: None

Concentration: 8.41 mg/ml

Mass-spec Verification: 38147 Da, uncleaved – expected mass out of insect cells

Compound Exact Mass: 524.6

Structure Determination

Crystallization: *Crystallization Condition:* 20% PEG3350 10% ethylene glycol 0.1M bis-tris-propane pH6.5 0.2M sodium acetate; *Protein Concentration:* 9.08mg/ml; *Crystallization*

Ligands: *Compound 1:* PK010102a; *Compound 2:* PK010102a; *Crystallization Comments:* : In order to enhance the compound solubility in aqueous solution, 100 μ l of protein stock at 8.4 mg/ml was diluted in 2 ml of gel filtration buffer. 2 μ l of compound stock (to reach 1 mM proportion with the protein) was added to the dilution and mixed with a P1000 pipette to allow as much compound as possible to dissolve. The mixture was concentrated again to 100 μ l using a 10kDa cut-off Amicon centrifugal device, transferred to a 1.5 ml tube and spun down for 10 mins at 4 degrees in a microfuge at maximum speed to precipitate the remaining insoluble compound.

Data Collection: Data was collected at Diamond beamline i04-1, wavelength 0.92818;

Data Processing: Data was processed to a resolution of ~2.6 Å using Phaser, Aimless, XDS and buster.