

logo

NTDB id 1169 A1552VC RS11075 WP 000648511.1	MAKSAARSVWNGMAKAKKMKYFWEKDSAGKMKGEYSALTEALARKRKQGLPKKRRKSKSLFETSSKKKYTK	68
NTDB id 1402 DSB67 RS12670 WP 010643257.1	MKATQTLPLKNYRWKGINNSGKK.VSGQMLAISEIEVRDKLKDQHIQIKKLLKKGVSVLLAR.LTHRVRKSK	67
NTDB id 1112 NGFG RS09215 WP 003689811.1	MKTIIAPQLKNYRWKGINNSGKK.TSGNVLAMTEIEVRERLDAQHIKIKKLLKKGSIISFITK.ISHRVRKSK	70
NTDB id 1113 AAA85695.1 219..1451()	MAK.....NGGFSLFAKKEKRFIFEGRHSASDKLVNCEVSATFEEEARKKLAKRGIRPLQITRVK.....TSSKRKITQE	70
NTDB id 126128 AMD23 RS16860 WP 040107659.1	MAK.....NGGFSLFAKKEKRFIFEGRHSASDKLVNCEVSATFEEEARKKLAKRGIRPLQITRVK.....TSSKRKITQE	78
NTDB id 1252 GC085 RS07730 WP 011213805.1	MSAARSVVKTAKTAASAEQMSPFVWEGTDRKGVK.MKGEQTARNANLLRAELRRQGITPLVVVKPKPLFGA.AGKKVSAK	67
NTDB id 1198 PSJM300 03950 AFN76868.1	MDKNSPPLLTFFHYQGINKAGQK.MEGDIQARSLAIADLRKQGIIVTNKVVKKKRPLFDR.KNKKITQA	66
NTDB id 1016 ACIAD RS01680 WP 004920476.1	MAQKAIKNSVFTWEGLDKRGAK.IKGELSGVSPALVKAQLRKQGINPQKVRKKSISLFS..AGKKIKPM	69
NTDB id 1059 ABD1 RS01610 WP 000279215.1	MAAKKTQVMPVFAYEGVDRKGIK.LKGELPARNMALAKVTLRKQGITIRTIREKRKNIFEGLLKKVKPL	69
consensus	MAVKKAQMMPTFAYEGVDRKGVK.IKGELPAKNMALAKVTLRKQGVTVRNIIEKRKNIFEGLLKKKVTTL	69

logo

NTDB id 1169 A1552VC RS11075 WP 000648511.1	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	148
NTDB id 1402 DSB67 RS12670 WP 010643257.1	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	147
NTDB id 1112 NGFG RS09215 WP 003689811.1	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	150
NTDB id 1113 AAA85695.1 219..1451()	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	150
NTDB id 126128 AMD23 RS16860 WP 040107659.1	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	158
NTDB id 1252 GC085 RS07730 WP 011213805.1	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	147
NTDB id 1198 PSJM300 03950 AFN76868.1	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	146
NTDB id 1016 ACIAD RS01680 WP 004920476.1	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	149
NTDB id 1059 ABD1 RS01610 WP 000279215.1	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	149
consensus	DITITFTRQLATMIMKAGVPLVQAFEIVAGHNPIMRYMIMKVEGSSLSALRKHPYFDLYCNLVAGEESGVL	149

logo

NTDB id 1169 A1552VC RS11075 WP 000648511.1	PEVFERLATYREKSEQLRRAKVIKALIYPSMVVLVALGVSYLMLTMVPIPEFESMFKGFGAELPWFTQVQLKLSHWVQAVSL	228
NTDB id 1402 DSB67 RS12670 WP 010643257.1	SQVFERLATYREKSEQLRRAKVIKALIYPSMVVLVALGVSYLMLTRVPIPEFEKMFMTGFGADLPAFTQMVLNLSAWTQNWGP	227
NTDB id 1112 NGFG RS09215 WP 003689811.1	ESLDDKLAIYKEKTQAIRKKVKTALTYPVSVIAVAGLVFVMMIFVLPFAKFEVYANMGAELPLTQTVMMDSDFFVSYGW	230
NTDB id 1113 AAA85695.1 219..1451()	ESLDDKLAIYKEKTQAIRKKVKTALTYPVSVIAVAGLVFVMMIFVLPFAKFEVYANMGAELPALQTVMMDSDFFVSYGW	230
NTDB id 126128 AMD23 RS16860 WP 040107659.1	ETVLDITVATYKENLEALKGKIKKALFYPMVMAMVAMLVSAILLVWVVPQFEDVFKGFGAELPAFTQMIVNLSRFMVSWWW	238
NTDB id 1252 GC085 RS07730 WP 011213805.1	DIMLDKVATYKEKIEIKKKIKKALTYPVSVIAVAGLVFVMMIFVLPFAKFEVYANMGAELPALQTVMMDSDFFVSYGW	227
NTDB id 1198 PSJM300 03950 AFN76868.1	ETLLDRVATYKEKTEALKAKIKKAMNYPVIAVVLVAVIVTALLIKVVPQFQDVAFNFGAELPAFTLMVIGLSEALQAWWY	226
NTDB id 1016 ACIAD RS01680 WP 004920476.1	EIMLDRVAVYKEKSEVLKQKIKKAMKYPATVIVVAIVVTIILMVKVVPVFKDLFTSFAGADLPAFTQLVNMNSNMWDYWF	229
NTDB id 1059 ABD1 RS01610 WP 000279215.1	ETMLDRVAIYKEKSELLKQKIKKAMKYPATVIVVAIVVTIILMVKVVPVFKDLFTSFAGADLPAFTQMVMNMNSNMWDYWF	229
consensus	PEVFERLATYREKSEQLRRAKVIKALIYPSMVVLVALGVSYLMLTMVPIPEFESMFKGFGAELPWFTQVQLKLSHWVQAVSL	229

logo

NTDB id 1169 A1552VC RS11075 WP 000648511.1	WAFITIAAAIFGLKALRKNSFQIRLKTSLRLKFPITGNVLAKASIAKFSRTLATSFAAGIPILASLKTAKTSGNVHFE	308
NTDB id 1402 DSB67 RS12670 WP 010643257.1	FTGLSFVSLIISGKIMSQRSDSFRLLAARSSLRVPILGPVLSKAAIAKFSRTLATSFAAGIPILATLKTAKTSGNMHYQ	307
NTDB id 1112 NGFG RS09215 WP 003689811.1	MVLIALGFAIYGFLLKARSIKIQRMDAILLRMPFIFGDIVRKGTIARWGRRTATLFAAGVPLVDVLDSTAGAAGNLIYE	310
NTDB id 1113 AAA85695.1 219..1451()	MVLIALGFAIYGFLLKARSIKIQRMDAILLRMPFIFGDIVRKGTIARWGRRTATLFAAGVPLVDVLDSTAGAAGNLIYE	310
NTDB id 126128 AMD23 RS16860 WP 040107659.1	LMLFALVGAAGISFYAYKRSPAMQHGMDRLVLKVPITIGIMNHSSIAARFARTTAVTFKAGVPLVLEALGIVAGATGNSVYE	318
NTDB id 1252 GC085 RS07730 WP 011213805.1	IIFGALGGVVSFFYAKNHSLEFAQTIIDRVMLKLPVIGPILEKAAIARFARTLSITFAAGLPLVEALKSVAGATGNTIYA	307
NTDB id 1198 PSJM300 03950 AFN76868.1	VVLGAFVGAAYAFKQAHTRSEKFRNWFDRLMLKVPVVGDIYKSSVARFARTLSTTFAAGVPLVDALDSVAGATGNVVF	306
NTDB id 1016 ACIAD RS01680 WP 004920476.1	ILILVIGAAIAGFLEARKRSKRFRDQLDRLVLRPLVIGDLVYKAIARIYSRTLATTFAAGVPLIDALASTAGATNNTVYE	309
NTDB id 1059 ABD1 RS01610 WP 000279215.1	IMIIAIGAVIAAFLEAKRSKRFRDGLDKLALKLPFIFGDLVYKAIARIYSRTLATTFAAGVPLIDALESTAGATNNVIYE	309
consensus	WAFITIAAAIFGLKALRKNSFQIRLKTSLRLKFPITGNVLAKASIAKFSRTLATSFAAGIPILASLKTAKTSGNVHFE	309

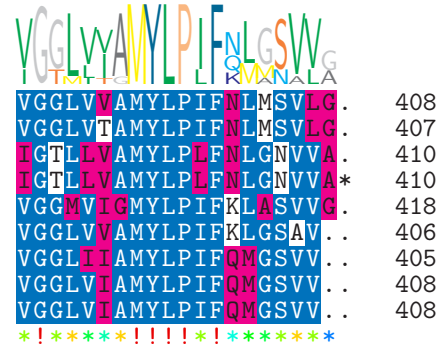
```

NTDB id 1169 A1552VC RS11075 WP 000648511.1
NTDB id 1402 DSB67 RS12670 WP 010643257.1
NTDB id 1112 NGFG RS09215 WP 003689811.1
NTDB id 1113 AA885695.1 219..1451( )
NTDB id 126128 AMD23 RS16860 WP 040107659.1
NTDB id 1252 GC085 RS07730 WP 011213805.1
NTDB id 1198 PSJM300 03950 AFN76868.1
NTDB id 1016 ACIAD RS01680 WP 004920476.1
NTDB id 1059 ABD1 RS01610 WP 000279215.1
consensus

```



logo



-  non conserved
-  similar
-  $\geq 50\%$ conserved