

logo

NTDB id 600 KW2 RS01765 WP 011675413.1	MATKK.....KTNFDDITKKYGA	ERDKALADALALIEKD	FGKGSIMRLGEA	.ANQKVSVVSSGSLALD	I	ALGAGGYPKGR	74	
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NTDB id 228 SPD RS09265 WP 001085462.1	.MAKK.....PKKLEEISKKFGA	EREKALNDALKLIEKD	FGKGSIMRLGER	.AEQKVQVMSSGSLALD	I	ALGSGGYPKGR	73	
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consensus		***	***	*	***!	*!!*****	*	***!!*!!*!!*!!*!!*!!*!!*!!

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consensus

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I I E I Y G P E S S G K T T V A L H A V A Q A Q K E G G I A A F I D A E H A L D P A Y A A L G V N I D E L L S Q P D S G E Q G L E I A G K L I D S G A V D L	153
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I V E I Y G P E S S G K T T V A L H A V A E V Q K Q G G T A A Y I D A E H A L D P K Y A T A L G V N I D D L L S Q P D T G E Q G L E I A D A L V S S G A V D I	140
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consensus

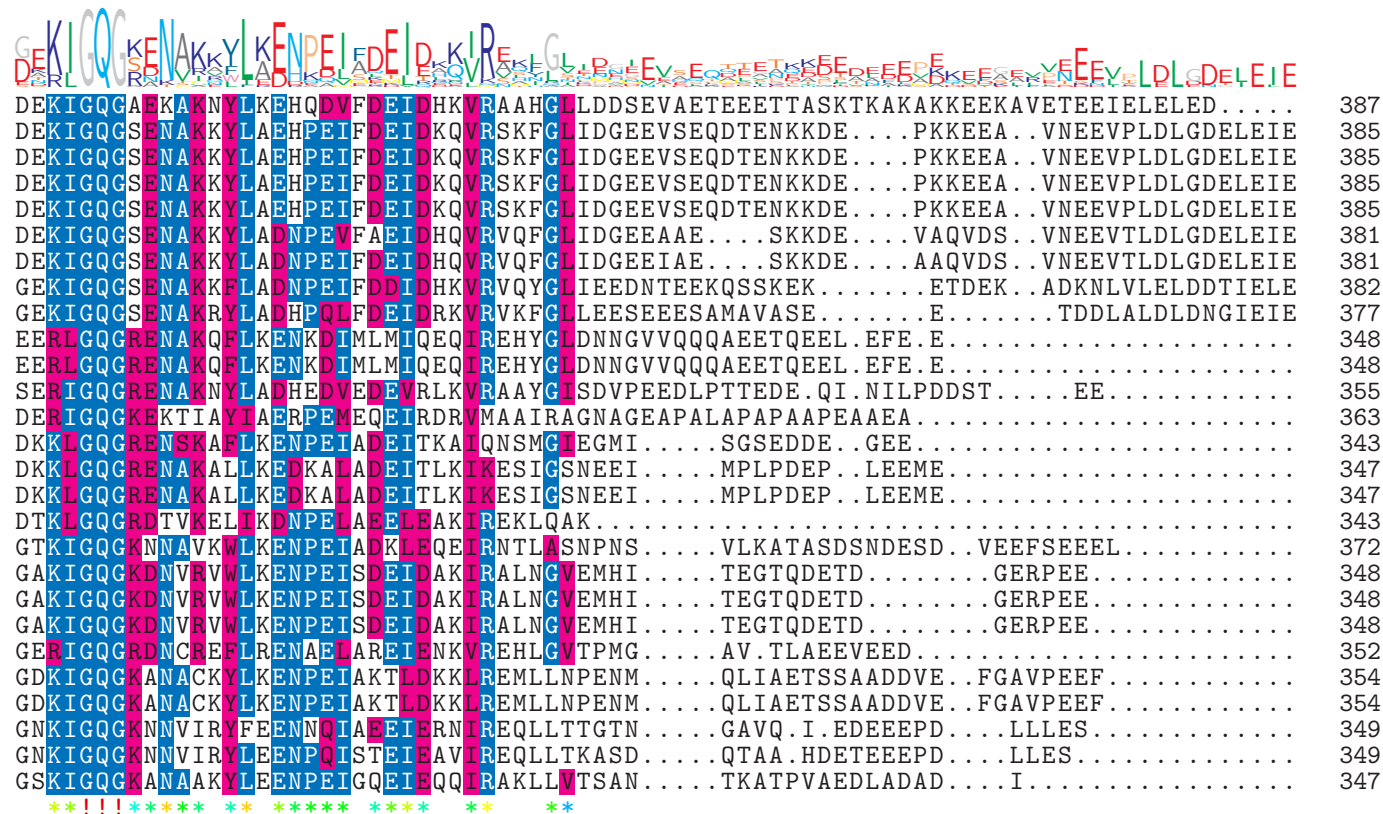
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consensus	*!!**!* *	

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consensus



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consensus			

- X non conserved
- X similar
- X ≥ 50% conserved