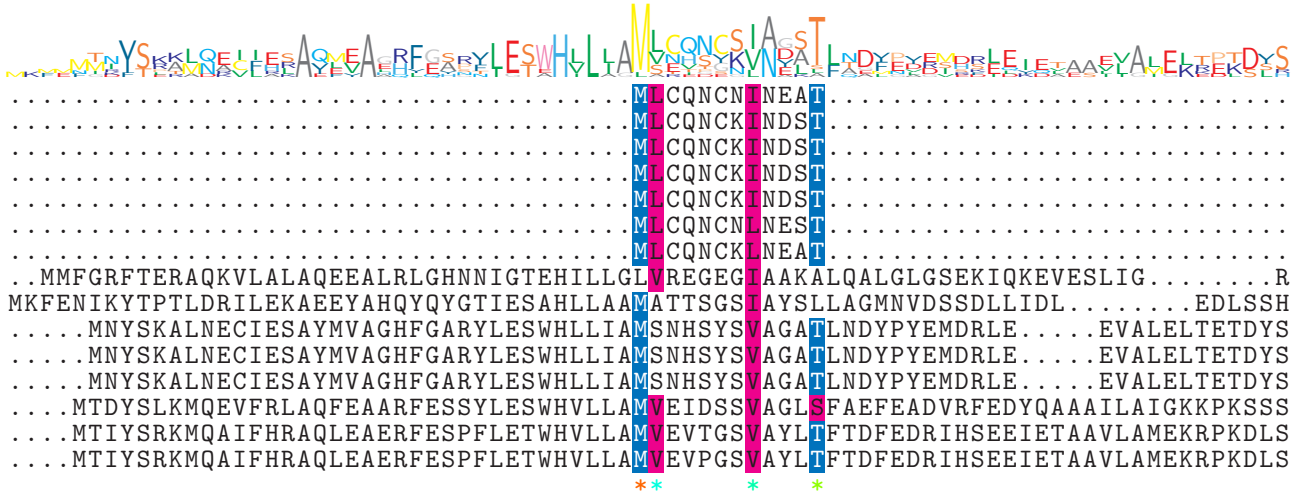


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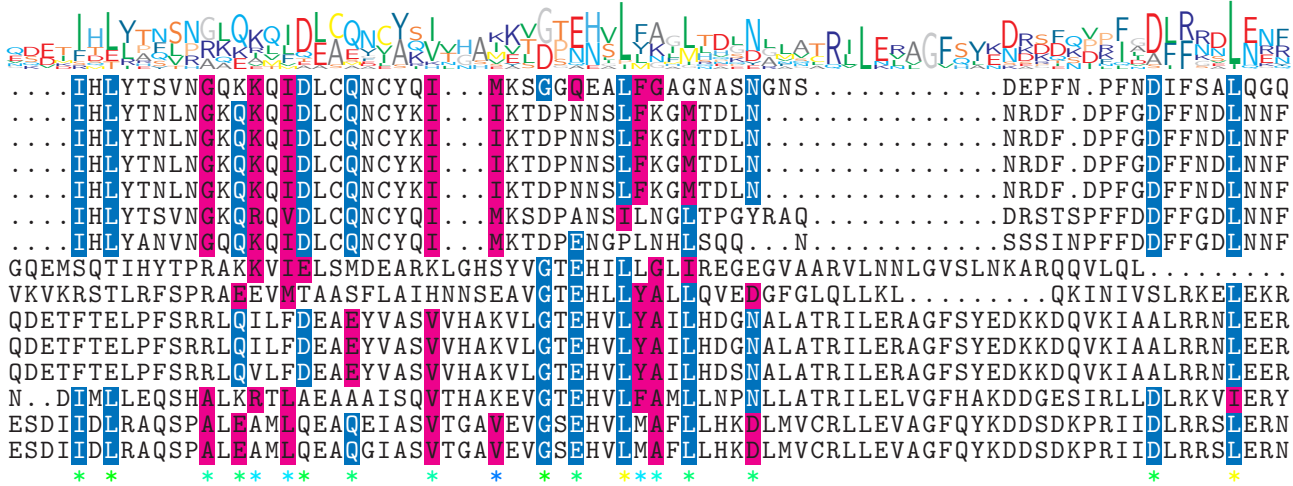
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NTDB	id	1017338	ABHI20	RS06385	WP	010922489.1
NTDB	id	378	SMU	RS02690	WP	002263569.1
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NTDB	id	329	STU	RS10020	WP	011225298.1
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consensus						

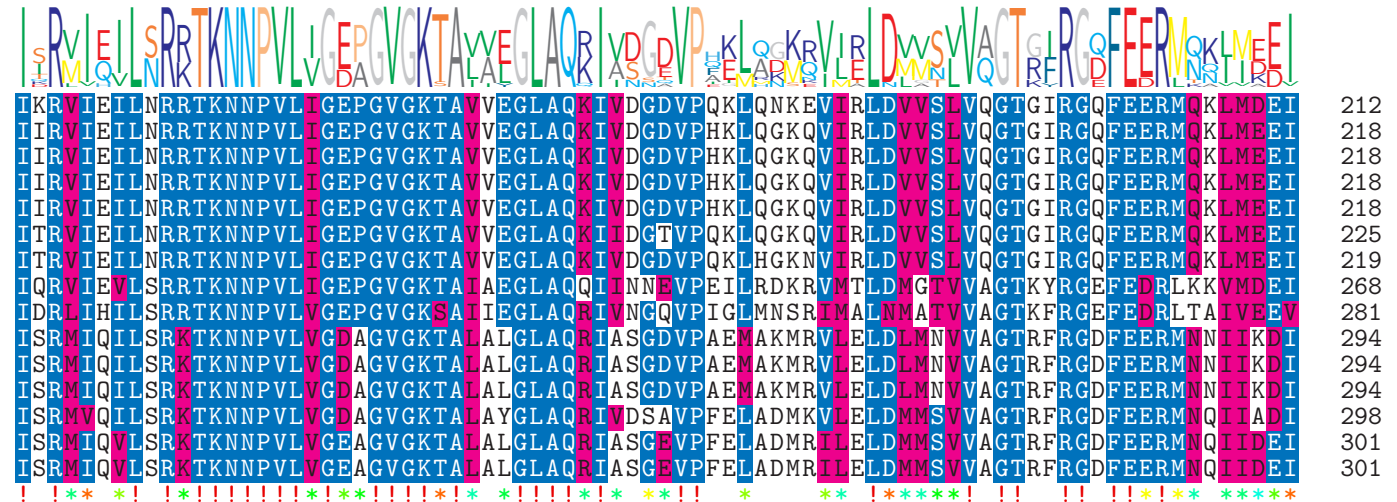
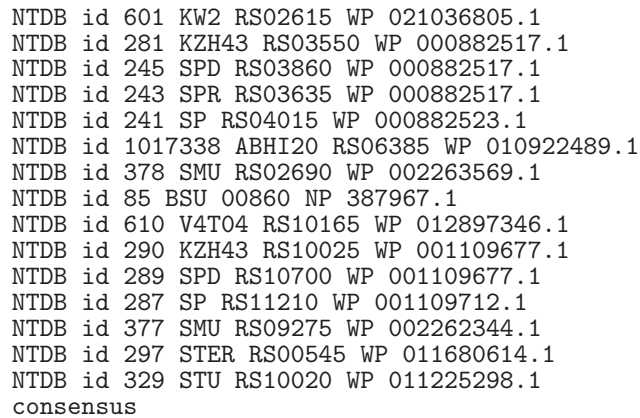


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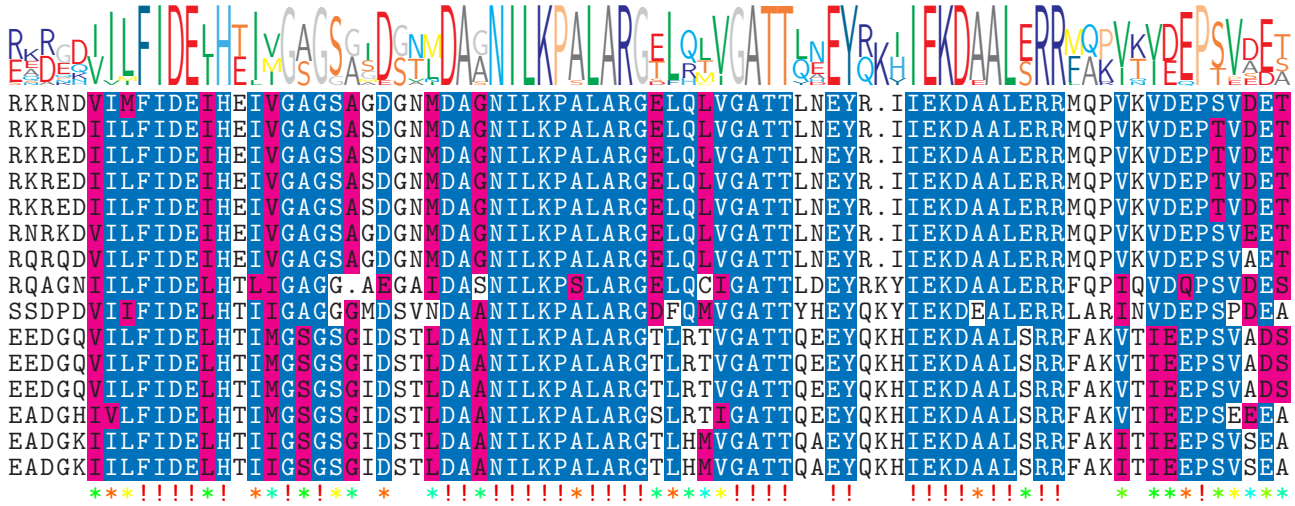
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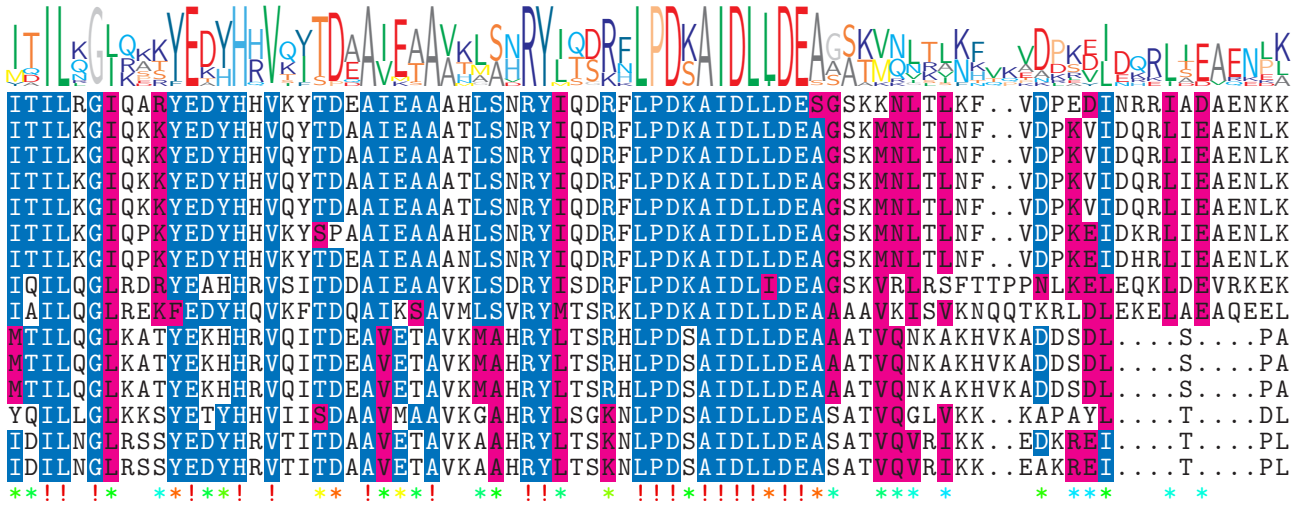
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NTDB	id	601	KW2	RS02615	WP	021036805.1
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NTDB id 329 STU RS10020 WP 011225298.1	GAPPGYVGYDEGGELTEKVRNK	677
consensus	!*!!!!*!*!*!*!!!!*!! !!!**!*!*!*!!!!!!* * !*!*?!! *!! !! ! !* **!!!!!! !	

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NTDB id 378 SMU RS02690 WP 002263569.1	EANVGFGASREG....RTNSVLGELGNFFSPEFMNRFDGIEFQALS	683
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NTDB id 610 V4T04 RS10165 WP 012897346.1	DKTVGFGAKNITADYSAMKSRILEELKRHYRPEFLNRIDENIVFHSLESQETIEQIVKIMSKSLIKRLAEQDIHVKLTPSA	756
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consensus	* *!!!!** ** *! * !!!!! ! * ! !* * * *! *!* * ** * * * * *	

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X non conserved
X similar
X ≥ 50% conserved