

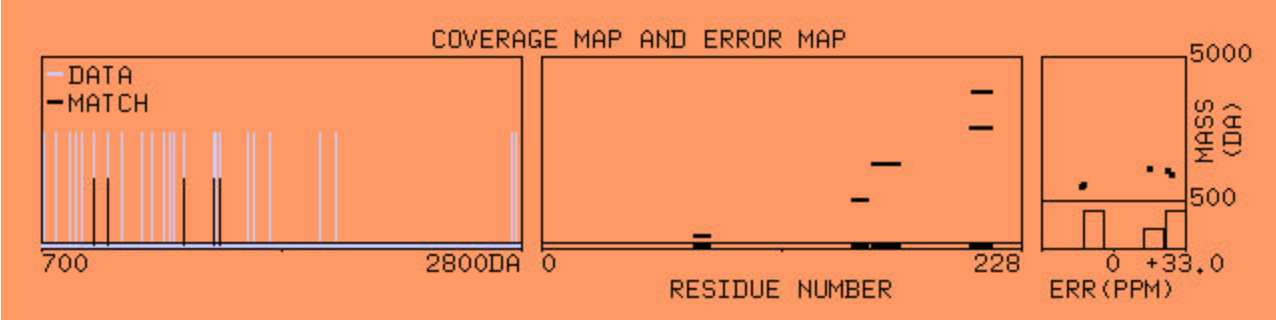
ProFound - Search Result Details


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

Details for rank 1 candidate in search 20060404162256-1BE4-192168001107

1. FTTSGH1760 1619463 1618780 [-1 L= 684 r=-1.196] (FTT1557 1619463 1618780 -)  
====>ref|YP\_123579.1| hypothetical protein lpp1255 [Legionella pneumophila str. Paris]  
ref|YP\_126605.1| hypothetical protein lpl1254 [Legionella pneumophila str. Lens] Score = 208 bits (530), Expect = 1e-52====>ref|YP\_170481.1| Two-component response regulator [Francisella tularensis subsp.tularensis Schu 4] Score = 439 bits (1130), Expect = e-122

Sample ID : Schu4 260 [Pass:0]  
Measured peptides : 28  
Matched peptides : 5  
Min. sequence coverage: 19%



Note: click on the  symbol to change column format.

Measured	Avg/	Computed	Error	 Residues	Missed		
Mass(M)	Mono	Mass	 (ppm)	Start	To	Cut	Peptide sequence
926.530	M	926.543	-14	148	156	0	DGVIIIPVSK
994.640	M	994.653	-13	73	81	0	VPIILLTAR
1327.797	M	1327.761	27	205	215	0	INIPNFIQTIR
1455.892	M	1455.855	25	204	215	1	KINIPNFIQTIR
1479.901	M	1479.877	17	158	171	0	ELALLAILVQNAGR

Unmatched Monoisotopic Masses:

714.100 768.610 826.470 828.476 850.511 882.514 1051.607 1144.599 1182.589  
1241.697 1263.728 1279.733 1462.827 1464.750 1604.872 1630.781 1703.486 1917.042  
1994.050 2762.478 2763.307 2774.392 2780.337

Search again using unmatched masses:

Search again

 in 

All taxa