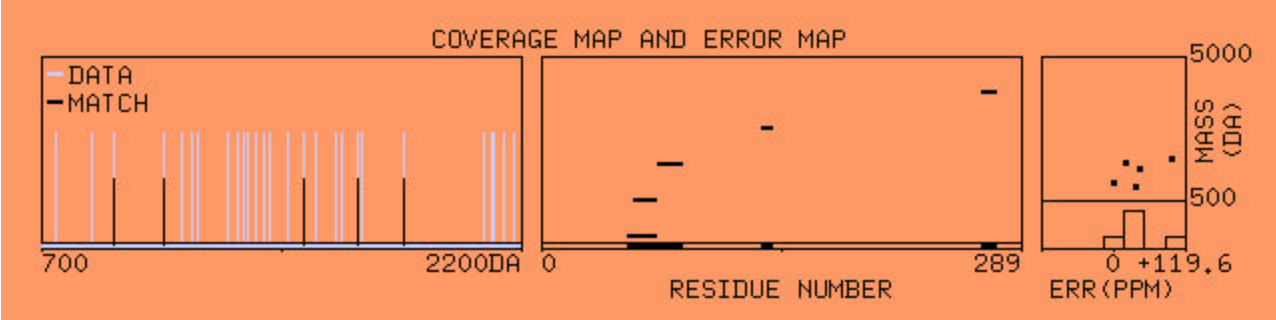



ProFound - Search Result Details



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Details for rank 1 candidate in search 20060330151530-16EC-192168001107

1. FTTSGH0341 329238 330104 [+3 L= 867 r=-1.179] (FTT0314 329238 330104 +)  
====>ref|YP\_123996.1| Elongation factor Ts (EF-Ts) [Legionella pneumophila str. Paris]emb|CAH12830.1| Elongation factor Ts (EF-Ts) [Legionella pneumophila str. Paris] Score = 342 bits (876), Expect = 1e-92====>ref|YP\_169363.1| protein chain elongation factor EF-Ts [Francisella tularensissubsp. tularensis Schu 4] Score = 543 bits (1398), Expect = e-153  
Sample ID : Schu4 171 [Pass:0]  
Measured peptides : 28  
Matched peptides : 5  
Min. sequence coverage: 18%



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
927.550	M	927.513	40	133	140	0	IGENIQVR
1086.592	M	1086.591	1	266	275	1	LDVGEGIEKK
1518.835	M	1518.767	45	56	70	0	VAAEGVIEVYAADGR
1689.932	M	1689.893	23	71	85	0	AILLEINSETDFVAR
1833.120	M	1832.937	100	53	70	1	ASRVAAEGVIEVYAADGR

Unmatched Monoisotopic Masses:

747.628 856.541 1139.957 1171.072 1191.321 1287.969 1316.284 1337.711 1344.839  
1373.808 1395.698 1417.761 1469.929 1561.531 1621.974 1638.913 1702.161 2083.119  
2112.036 2113.111 2149.288 2178.057 2179.710

Search again using unmatched masses:

Search again

 in 

All taxa