

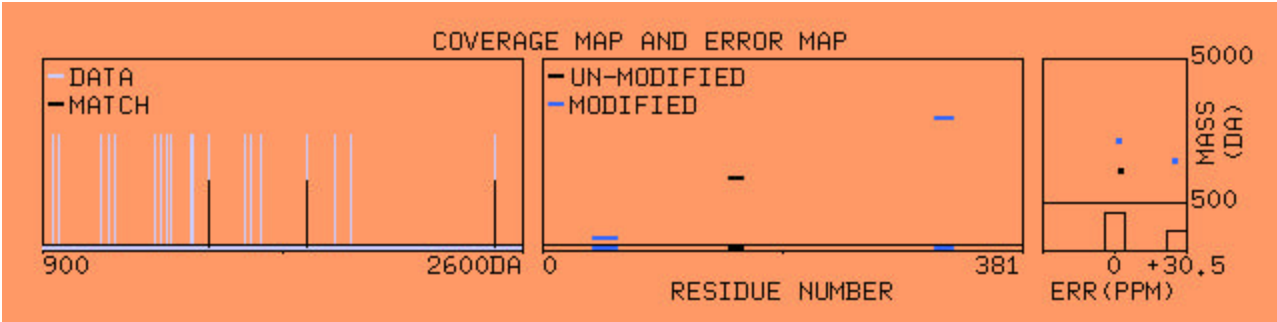
ProFound - Search Result Details


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

Details for rank 1 candidate in search 20060404123243-1E9C-192168001107

1. FTTSGH1393 1252948 1254090 [+1 L=1143 r=-1.227] (FTT1234 1252948 1254090 +)
=====>ref|YP_222152.1| choloylglycine hydrolase family protein [Brucella abortus biovar 1str. 9-941] Score = 166 bits (420), Expect = 1e-39=====>ref|YP_170192.1| choloylglycine hydrolase family protein [Francisella tularensissubsp. tularensis Schu 4] Score = 765 bits (1976), Expect = 0.0

Sample ID : Schu4 235 [Pass:1]
Measured peptides : 19
Matched peptides : 3
Min. sequence coverage: 13%



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
1492.771	M	1492.767	3	149	160	1	QALQQYKVWSDK
1837.949	M	1837.902	25	312	327	0	LGNQTMFDHTAFTVIK
							(1)+O@M;
2500.228	M	2500.224	2	41	60	1	TMEWGFNWSWQLIYIPKGTK
							(1)+O@M;

Unmatched Monoisotopic Masses:

935.478 962.444 1109.620 1140.661 1158.685 1302.768 1323.717 1340.757 1357.777
1424.789 1434.802 1621.954 1638.911 1674.971 1940.935 1994.033

Search again using unmatched masses:

Search again

in

All taxa