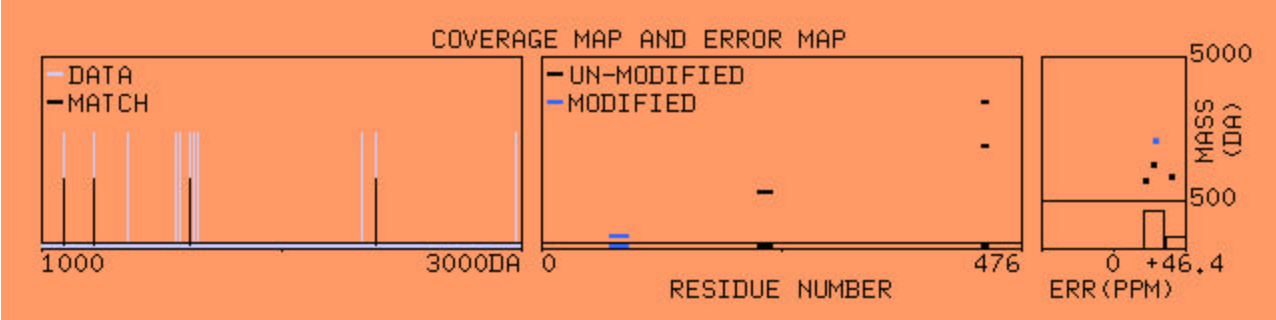



ProFound - Search Result Details



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Details for rank 1 candidate in search 20060330143627-184C-192168001107

1. FTTSGH1740 1603293 1601866 [-1 L=1428 r=-1.216] (FTT1539 1603293 1601866 -)
=====>ref|ZP_00368662.1| probable periplasmic protein Cj0092 [Campylobacter lari
RM2100]gb|EAL55107.1| probable periplasmic protein Cj0092 [Campylobacter lari RM2100]
Score = 152 bits (385), Expect = 2e-35=====>ref|YP_170467.1| hypothetical protein FTT1539c
[Francisella tularensis subsp.tularensis Schu 4] Score = 927 bits (2397), Expect = 0.0
Sample ID : Schu4 162 [Pass:0]
Measured peptides : 11
Matched peptides : 4
Min. sequence coverage: 9%



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
1092.474	M	1092.451	21	437	444	0	NFSYDEYR
1220.593	M	1220.546	39	437	445	1	NFSYDEYRK
1616.956	M	1616.913	27	216	231	0	SLTTGFGNLSGLLPIK
2395.162	M	2395.095	28	69	88	0	SAEEILNDVMENYIDQNNLR

(1)+O@M;

Unmatched Monoisotopic Masses:

1365.847 1564.784 1581.883 1639.914 1655.867 2341.658 2979.389

Search again using unmatched masses:

Search again in All taxa