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## Research Article

# MALDI-TOF/TOF *de novo* sequence analysis of 2-D PAGE-separated proteins from *Halorhodospira halophila*, a bacterium with unsequenced genome

Because protein identifications rely on matches with sequence databases, high-throughput proteomics is currently largely restricted to those species for which comprehensive sequence databases are available. The identification of proteins derived from organisms with unsequenced genomes mainly depends on homology searching. Here, we report the use of a simplified, gel-based, chemical derivatization strategy for *de novo* sequence analysis using a MALDI-TOF/TOF mass spectrometer. This approach allows the determination of *de novo* peptide sequences of up to 20 amino acid residues in length. The protocol was applied on a proteomic study of 2-D PAGE-separated proteins from *Halorhodospira halophila*, an extremophilic eubacterium with yet unsequenced genome. Using three different homology-based search algorithms, we were able to identify more than 30 proteins from this organism using subpicomole quantities of protein.

**Keywords:** Chemically assisted fragmentation / Guanidination / Homology search / Sulfonation  
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## 1 Introduction

Proteomics is playing a pivotal role in the postgenome era in helping to define the functional role of genes. MS, hyphenated with a range of electrophoretic and multi-dimensional chromatographic separation techniques, has emerged as a key platform technology in proteomics for the rapid and high-throughput identification, characterization, and quantitation of proteins [1]. Typically, proteins are digested using trypsin and the resultant peptides are then subjected to MS analysis. The tryptic peptides provide a characteristic PMF which can be used to identify proteins. Although this approach is useful to identify pro-

teins in simple mixtures, peptide sequence information obtained by MS/MS is required to identify individual proteins in more complex samples [2]. Sophisticated algorithms (e.g., SEQUEST and MASCOT) have been developed to aid in this process, starting from peptide MS/MS data whereby peptides are identified by correlating the uninterpreted MS/MS spectra with simulated (predicted) product ion spectra derived from peptides of the same mass contained in the databases. While the above-mentioned algorithms for protein identification from peptide MS/MS data have enjoyed considerable success, their utility is directly related to the quality of the product ion spectra and depends on the availability of database information about the proteins under investigation. For proteins not contained within sequence databases, it is necessary to determine partial or complete amino acid sequences using either manual or automated *de novo* peptide sequence analysis methods.

Since manual *de novo* sequencing is a very time-consuming process, several software tools were developed that deduce an amino acid sequence from an MS/MS

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**Abbreviations:** MQ, Milli-Q; nrdb, nonredundant database; PYP, photoactive yellow protein

spectrum. Interpretation of MS/MS spectra relies on measuring the mass differences between adjacent fragment ion peaks of one of the major ion series, *i.e.*, b-series (ions containing N-terminus) or y-series (ions containing C-terminus), which are common in tryptic peptides. However, most of *de novo* sequencing software tools inevitably suffer from inherent limitations of MS/MS spectral analysis, including incomplete b- and y-ion series (gaps), the presence of other peptide-derived peaks such as a-ions, internal fragments and neutral losses of water or ammonia [3]. The recent introduction of MALDI-TOF/TOF MS technology offers the advantage of MALDI ionization with MS/MS in a TOF instrument [4]. *De novo* sequencing of underivatized peptides using MALDI-TOF/TOF MS has recently been demonstrated by Yergey *et al.* [5]. However, compared to the MS/MS spectra of doubly charged ESI-generated ions, MS/MS spectra of singly charged MALDI ions contain more ions from other fragment ion series. Therefore, a number of derivatization methods have been proposed to improve the fragmentation of singly charged ions.

One approach to facilitate the interpretation of MS/MS spectra is to enrich a series of fragments by attaching a strongly negatively or positively charged group to the N-terminus of peptides. Keough *et al.* [6] demonstrated that the N-terminus can be derivatized by acylation with 2-sulfobenzoic acid cyclic anhydride or chloro-sulfonylacetyl chloride. N-Terminal sulfonic acid derivatives were subsequently proposed for peptide sequencing by ESI-MS, MALDI-TOF MS [7], and MALDI-TOF/TOF MS [8]. The introduction of a sulfo group facilitates the MS/MS fragmentation of singly charged peptide ions by providing a second “mobile” proton which lowers amide bond strength and allows more facile unimolecular decay [9]. Since sulfonation reagents react with amino groups, this derivatization results in the modification of both the N-termini and the  $\epsilon$ -amines of lysine-containing peptides. Therefore, Keough *et al.* [10] expanded their approach and combined guanidination of lysine residues with the addition of a sulfonic acid group to the N-terminus. Following guanidination of lysine  $\epsilon$ -amines, introduction of sulfonic acid groups to tryptic peptides is possible solely at the N-terminus. We further improved this method by performing the Lys side-chains’ modification directly on gel-separated proteins prior to tryptic digestion. In this way, removal of the molar excess of guanidination reagents can simply be accomplished during the destaining step of the gel spots [11].

Today, most of the proteomic studies of extremophilic bacteria have been performed on members of the Archaea for which a sequenced genome is available.

Zhu *et al.* [12] applied the MudPIT approach to analyze the proteome of *Methanococcus jannaschii*, an autotrophic methanoarchaeon and the first member of the Archaea with a completely sequenced genome [12]. A shotgun approach was also used to identify the *Sulfolobus solfataricus* P2 proteome, a thermo-acidophilic crenarchaeon [13]. The cytosolic and membrane proteome of *Halobacterium salinarum* has been analyzed using PMF and LC-MS/MS techniques [14, 15]. *H. salinarum* is a member of the halophilic Archaea and an important model organism to study adaptations necessary for living in salty habitats. The genome sequence of *Halobacterium* species NRC-1 has completely been determined [16].

The extremely halophilic purple phototrophic bacterium *Halorhodospira* (formerly *Ectothiorhodospira*) *halophila* shows a photophobic response toward intense blue light. The wavelength dependence of this response corresponds with the absorption spectrum of the photoactive yellow protein (PYP), which suggests this protein to be the primary photoreceptor for this response [17]. Photoreceptors allow living organisms to make optimal use of the light conditions for growth and development and/or the protection from light damage. Various types of light-induced sensory responses have been characterized physiologically in detail. However, the molecular basis of this type of response is only slowly emerging. While the PYP protein is extremely well studied at the physical level, direct proof of a link between PYP and negative phototaxis is lacking. Moreover, in other species that produce PYP, a link with phototaxis has never been reported. A major limitation to the study of the physiological function of PYP is the fact that sequence information about the genome is not available (a DOE funded sequence program is currently running) and, more in particular, genetic techniques are poorly developed in this organism. There is limited information about the flanking regions of the PYP gene [18] in *H. halophila* and other species, but except for the presence of the biosynthetic genes for the production of the cofactor (*p*-coumaric acid) and for its covalent attachment to the protein, there are no generalizations that provide clues concerning the role of PYP.

Here, as a proof of principle, we applied our improved MS identification approach to identify a number of 2-D PAGE-separated proteins from *H. halophila*. (Partial) Sequences of tryptic peptides were submitted to homology search for identification of the corresponding protein. For this purpose, we applied three different homology-based search algorithms: FASTS, MS-BLAST, and MS-Homology [19–21].

## 2 Materials and methods

### 2.1 Materials

Urea, ammonium persulfate, CBB G-250, and agarose were obtained from Amersham Biosciences (Uppsala, Sweden). Iodoacetamide, CHAPS, DTT, and TEMED were from Fluka (Buchs, Switzerland). IPG strips, SDS, glycine, and ampholytes were purchased from BioRad (Hercules, CA, USA). The acrylamide/bisacrylamide solution was obtained from National Diagnostics (Atlanta, GE, USA), and the solvents for mass spectrometric sample preparation were from Biosolve (Valkenswaard, The Netherlands).

### 2.2 Bacterial growth and preparation of extracts

*H. halophila* SL-1 was grown anaerobically under tungsten illumination at 30°C in medium 253 described by DSMZ, and 2.5 mL of these cultures were used to inoculate 250 mL anaerobically prepared medium. Cultures were grown anaerobically under tungsten illumination or green and blue light conditions at 30°C and harvested at the late-exponential growth phase. After washing with dH<sub>2</sub>O, the cells were resuspended in 100 mM Tris-HCl, pH 8.0, supplemented with 50 µg DNase and 0.5 mM PMSF, and fractionated by sonication, followed by centrifugation to remove the cell debris (14 000 rpm, 30 min, 4°C).

### 2.3 2-DE and analysis

After determination of the protein concentration with the Protein Assay Kit (BioRad), approximately 250 µg of protein was mixed with IPG rehydration buffer (8 M urea, 2% w/v CHAPS, 0.3% DTT, final volume = 360 µL). The strips were allowed to rehydrate for 7 h and to focus (IEF) using a Multiphor II system (Amersham Biosciences) running the following program: 150 V (30'), 150 V (120'), 300 V (30'), 300 V (45'), 3500 V (90'), 3500 V (540'), 500 V (10'), and held at 500 V. The temperature was kept at 18°C. After completion of the IEF program, the IPG strips were equilibrated in a 50 mM Tris-HCl solution, pH 8.8, containing 6 M urea, 30% glycerol, 2% SDS, and 1% DTT, for 10 min, after which the solution was replaced with the same solution, except that DTT was exchanged by 5% iodoacetamide. The strips were then placed on the home-casted vertical SDS-PAGE gels and subjected to electrophoresis at 10 mA/gel for 15 min, followed by a ±5 h run at 20 mA/gel until the bromophenol blue front reached the bottom of the gel. Staining was performed using CBB G-250. The 2-D-gel images were digitized using a GS-710 densitometer (BioRad) and analyzed with the accompanying PDQuest 7.1 software (BioRad).

### 2.4 In-gel guanidination

Guanidination was performed by adding 5 µL of Milli-Q (MQ) water, 11 µL of 7 N ammonium hydroxide (Merck, Darmstadt, Germany), and 3 µL of a freshly prepared 7.5 M O-methylisourea hemisulfate solution (Acros, Geel, Belgium) to the excised spots. The samples were vortexed briefly and incubated at 65°C. After incubation for 2 h, the guanidinated samples were taken from the oven and the remainder of the solution was discarded. The gel pieces containing the guanidinated samples were desalted and destained in one step. Two washes using 150 µL of 200 mM ammonium bicarbonate in 50% ACN/MQ (30 min at 30°C) were performed and subsequently the gel pieces were dried in a SpeedVac (Thermo Savant, Holbrook, NY).

### 2.5 Trypsin digestion and sulfonation

A volume of 8 µL digestion buffer (50 mM ammonium bicarbonate, pH 7.8) containing 150 ng modified trypsin *per* microliter (Promega, Madison, WI) was added to the dried gel spots and the tubes were kept on ice for 45 min to allow the gel pieces to be completely soaked with the protease solution. Digestion was performed overnight at 37°C, the supernatants were recovered and the resulting peptides were extracted twice with 35 µL of 60% ACN/0.1% DIEA. The extracts were pooled and dried in the SpeedVac. The peptides were redissolved in 4 µL of 12.5 mM ammonium bicarbonate and 50% ACN/MQ, and 2 µL was mixed with 2 µL of the sulfonation solution. The sulfonation reagent was prepared by dissolving 2 mg 2-sulfobenzoic acid cyclic anhydride in 1 mL dry THF to attain a 0.01 mM solution. The tubes were briefly vortexed and reacted for 15 min at room temperature. Upon sulfonation, the samples were not desalted, a fraction of the samples was mixed with matrix solution and spotted on the MALDI plate.

### 2.6 MS and MS/MS

A 4700 Proteomics Analyzer (Applied Biosystems, Foster City, CA) with TOF/TOF optics was used for all MALDI-MS and MS/MS applications. Samples were prepared by mixing 0.7 µL of the sample with 0.7 µL matrix solution (7 mg/mL  $\alpha$ -cyano-4-hydroxycinnamic acid (CHCA) in 50% ACN containing 0.1% TFA) and spotted on a stainless steel 192-well target plate. They were allowed to air-dry at room temperature, and were then inserted in the mass spectrometer and subjected to mass analysis. The mass spectrometer was externally calibrated with a mixture of angiotensin I, Glu-fibrino-peptide B, ACTH (1–17), and ACTH (18–39). For MS/MS experiments, the instrument was externally calibrated with fragments of Glu-fibrino-peptide B.

All of the sulfonated peptides were subjected to MS/MS using a MALDI-TOF/TOF instrument. In an initial study, using this method, in which the fragmentation spectra resulting from high- and low-energy CID experiments were compared, the authors concluded that the difference in fragmentation and the effect on database search results were surprisingly small [22]. The major difference observed is the presence of high-energy fragment ions (w-ions) in the high-energy CID spectra of some peptides. When the CID mode (gas on, collision energy 0.5 to >3.5 keV) is applied, a larger number of low-molecular weight fragments (immonium ions, internal fragments) have been observed [23]. However, it has also been shown that the use of high-energy CID results in a loss of sequence information, as the y-ion abundance decreases at both higher gas pressure and higher collision energy [24]. Therefore, we performed all fragmentation experiments with the collision energy set at 1 keV and no gas in the collision chamber (low-energy CID).

## 2.7 Database searches

*De novo* determined peptide sequences were deduced manually and used for similarity searches using the FASTS, MS-BLAST, and the MS-Homology algorithm. On-line submissions were performed using MS-BLAST at the Heidelberg server (<http://dove.embl-heidelberg.de/Blast2/msblast.html>). Searches were performed against the nonredundant database (nrdb) using standard settings. The FASTS algorithm ([http://fasta.bioch.virginia.edu/fasta\\_www/cgi/](http://fasta.bioch.virginia.edu/fasta_www/cgi/)) was carried out using standard settings, and searches were performed against the NCBI/BLAST nrdb with BLOSUM 50 as search matrix. MS-Homology searches (Protein Prospector 4.0.5) were performed on the UCSF server against the NCBI nrdb using BLOSUM 50 as search matrix (<http://prospector.ucsf.edu/ucsfhtml4.0/mshomology.htm>).

The software used for similarity searches does not discriminate between the isobaric amino acids Ile and Leu. Therefore, all mass increments of 113 Da between consecutive y-ions were arbitrarily designated as Ile. The FASTS search results were considered significant if the E-value was below  $1.0e-04$ . The MS-BLAST search results were considered significant if the resulting scores were higher than the threshold score indicated in the software. In order for a particular protein in the database to generate a hit, MS-Homology must find homologous sequences for the minimum number of peptides required to match. The scoring method used is based on a mutation matrix such as the one used in the BLAST and FASTA programs. The final score is calculated by adding the scores for the individual peptide alignments together. If

there are several possible alignments of a given peptide, then the highest scoring alignment is used in the calculation. As the searches are based on similarity, proteins identified with lower scores must have the same generic function as the first hit. Proteins were considered positively identified only if all three search algorithms yielded the same homologous protein in the first hit. It has been demonstrated that indirect evidence can add to the significance of an identification [20]. Therefore, the identifications were further validated by using information such as the cleavage specificity of trypsin and sequence information resulting from known preferential fragmentation patterns of sulfonated peptides [8].

## 3 Results and discussion

The total protein extracts from *H. halophila* grown anaerobically under yellow or green/blue light were separated by 2-D PAGE. From these two gels, 100 spots were randomly selected and manually excized. The proteins were guanidinated in-gel and desalted/destained in one single step as described previously. Subsequently, the guanidinated proteins were enzymatically cleaved with trypsin and, after extraction, the peptides were sulfonated [11]. For 74 spots, we observed a good PMF of the sulfonated peptides, suitable for *de novo* MALDI MS/MS analysis. For the other 26 spots, we observed none or a very weak PMF, with signal intensities that were too low for MALDI MS/MS fragmentation. Previous experiments have indicated that sulfonic acid-derivatized peptides have poorer positive-ion sensitivity than the corresponding native peptides. The introduction of a negative charge usually leads to a decrease or even loss in signal intensity in positive mode [8]. Keough *et al.* [6] also noticed a decreased intensity of sulfonated peptides in the positive mode, compared to the negative ion mode and, apparently, some peptides show no signal above the noise level [6]. Recently, it was also demonstrated that the presence of the strong negative charge of the sulfonic group can create problems for sample desalting on RP media (low yield for less hydrophobic peptides) [25]. However, in our approach, guanidination is performed in-gel, and therefore, an additional desalting step to remove the excess of reagents can be omitted [11].

All fragmentation experiments were performed with the collision energy set at 1 keV and no gas in the collision chamber (low-energy CID). Typically, the most intense peaks in the PMF were selected for MS/MS analysis. In most spots, three to six peptide sequences, with a length varying between 5 and 20 amino acids, were obtained *de novo* (Tables 1a and b). In all fragment spectra, an initial loss of the sulfonic acid derivative was observed

( $\Delta m = 184$  Da). By simple manual calculation of the differences between the adjacent *y*-ion fragments, the amino acid sequence could readily be interpreted. As an example, the protein in spot 5 was identified as fructose-1,6-bisphosphate aldolase (Table 1a). Upon sulfonation, four peptides were subjected to MALDI MS/MS analysis. In all fragmentation spectra, except one, we observed a complete *y*-ion series that could easily be interpreted, facilitating *de novo* sequencing (Figs. 2a–d). MS/MS spectra of derivatized peptides having an internal homo-Arg (guanidinated lysine) also produced complete *y*-ion series (Fig. 3a). However, the *y*-ion series were often accompanied with (Yi–17)-ion series, due to the neutral loss of  $\text{NH}_3$ . Although this (Yi–17) series is seen in the fragment spectra up to the step where the basic residue is cleaved off, its presence does not interfere with the sequence interpretation (Figs. 2a–d). Please note that the loss of the sulfonic group (–184 Da) and the neutral loss of ammonia (–17 Da) is summed as an initial loss of –201 Da.

During PSD experiments with derivatized peptides enhanced fragmentation at Pro residues and a reduced abundance in fragmentation at the C-terminal side of Pro has been observed [15]. Here, and in a previous study,

TOF/TOF analysis of peptides containing an internal Pro indicated that *y*-ions resulting from cleavage on the N-terminal side of Pro are enhanced while *y*-ions resulting from cleavage on the C-terminal side of Pro are less abundant or almost completely depleted [8]. Therefore, peptides with a Pro residue near the N-terminus of a peptide can limit the amount of sequence information and cause gaps in the derived sequence (Figs. 2d, 3c). If the Pro residue occurs in the middle or near the C-terminus of the peptide this effect is less detrimental (Fig. 2c), and even for derivatized peptides containing internal homo-Arg and Pro residues we were able to derive uninterrupted peptide sequences of 20 amino acids (Fig. 4b and c). As proteomic strategies are becoming increasingly reliant on the use of automated database search algorithms, incorporation of “fragmentation rules”, such as the observed preferential cleavage at Xxx-Pro peptide bonds, into the database search algorithms will aid in the development of more effective tools for high-throughput protein identification. Furthermore, the occurrence of “nonsequence” specific ion fragments, such as the neutral loss of ammonia from peptides with internal Arg or homo-Arg, can be used to improve predictive models of peptide fragmentation for *de novo* sequence analysis.

**Table 1a.** Identified proteins from *Halorhodospira halophila* (yellow light)

Spot <sup>a)</sup>	Protein <sup>b)</sup>	Identification FASTS <sup>c)</sup>	Pept. <sup>d)</sup>	AA <sup>e)</sup>	<i>E</i> -score <sup>f)</sup>	MS-Blast score <sup>g)</sup>	MS-Hom. score <sup>h)</sup>
1, 2	71909086	Porin, Gram-negative type	5	50	3.50e-05	135	124
3	37527547	S-Adenosylmethionine synthetase	6	54	2.20e-17	202	216
4	39936346	Elongation factor Tu	6	73	2.90e-29	336	376
5	26991638	Fructose-1,6-biphosphate aldolase	4	32	6.60e-14	115	186
6	33634721	Phosphoribulokinase	4	33	5.40e-05	95	129
7	9392587	Sarcosine-dimethylglycine methyltransferase	4	31	0.00021	–	113
8	34497819	Acetoacetyl-CoA reductase	4	51	4.30e-10	149	159
9	74317158	Triosephosphate isomerase	3	29	7.80e-12	156	121
10	34498798	Adenylate kinase	3	31	1.10e-15	177	191
11	53804425	2,3,4,5-Tetrahydropyridine-2,6-dicarboxylate <i>N</i> -succinyltransferase	5	45	1.40e-08	167	144
12	47574096	Pentose-5-phosphate-3- epimerase	3	29	0.00046	105	98
13	67941974	Superoxide dismutase	8	67	5.70e-17	313	298
14	68304953	DsrC	1	13	1.60e-07	99	82
15	78700374	Nucleoside diphosphate kinase	3	49	4.00e-29	279	259
16	132132	Ribulose biphosphate carboxylase small chain	3	34	1.10e-06	107	113
17	53805138	Pterin-4- $\alpha$ -carbinolamine dehydratase	4	64	4.00e-13	188	132
18	69951812	Cold-shock protein, DNA-binding	2	21	2.70e-08	112	115

**Table 1b.** Identified proteins from *Halorhodospira halophila* (green/blue light)

Spot <sup>a)</sup>	Protein <sup>b)</sup>	Identification FASTS <sup>c)</sup>	Pept. <sup>d)</sup>	AA <sup>e)</sup>	E-score <sup>f)</sup>	MS-Blast score <sup>g)</sup>	MS-Hom. score <sup>h)</sup>
1	77166263	Chaperone protein dnaK (Hsp70)	4	35	1.60e-11	186	155
2	54294307	30S ribosomal protein S1	3	34	1.60e-13	152	181
3	53762519	Chaperonin GroEL (HSP60 family)	6	83	8.60e-26	383	299
4	71899446	ATP synthase F1, alpha subunit	2	35	1.00e-10	152	127
5	71550918	Ribulose-bisphosphate carboxylase	4	30	1.60e-05	99	160
6	37527547	S-Adenosylmethionine synthetase (methionine adenosyltransferase) (AdoMet synthetase)	6	56	1.00e-22	194	257
7	56461311	Fructose/tagatose bisphosphate aldolase	5	29	4.40e-11	109	156
8	33862805	Phosphoribulokinase	3	36	1.40e-16	158	191
9	33152351	NADH-dependent enoyl-ACP reductase	4	40	1.20e-12	161	190
10	2497482	Adenylate kinase (ATP-AMP transphosphorylase)	4	37	3.1e-09	174	195
11	52006362	Dissimilatory sulfite reductase	1	13	0.00097	73	70
12	77866837	Nucleoside diphosphate kinase	2	24	4.70e-10	101	132
13	1402737	Major cold shock protein	1	11	9.70e-05	77	71

a) Spot number according to the position on the 2-D PAGE (Fig. 1).

b) NCBI *Entrez* entries (<http://www.ncbi.nih.gov/Entrez/>).

c) Identification based on FASTS search result.

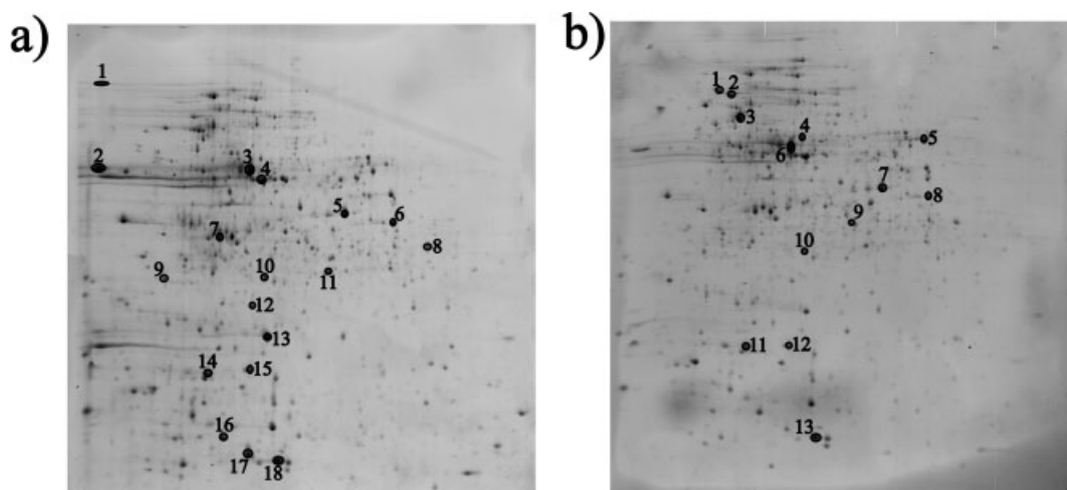
d) Number of peptide sequences used in the query.

e) Total number of amino acids used in the query.

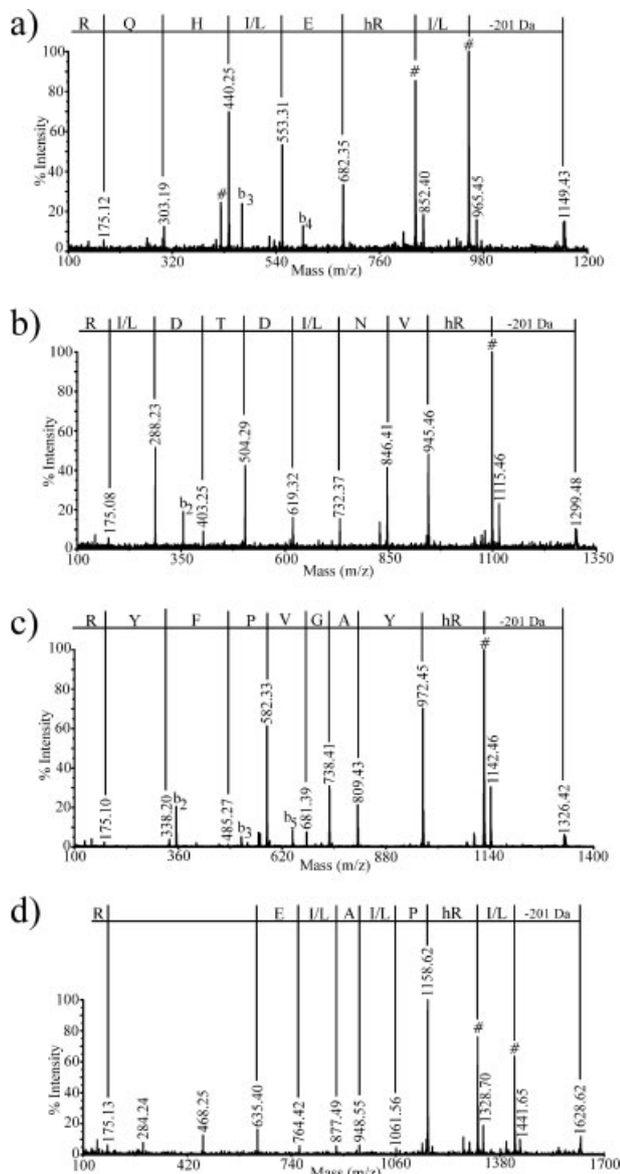
f) In FASTS, the E(N) value reports the number of times the score should be obtained by chance against a database of size N. For searches against the NCBI nonredundant protein database  $N \approx 2075116$ .

g) MS-BLAST score for searches against the nonredundant protein database at <http://dove.embl-heidelberg.de/Blast2/msblast.html>.

h) MS-Homology (Protein Prospector 4.0.5) score against the NCBI nonredundant protein database.

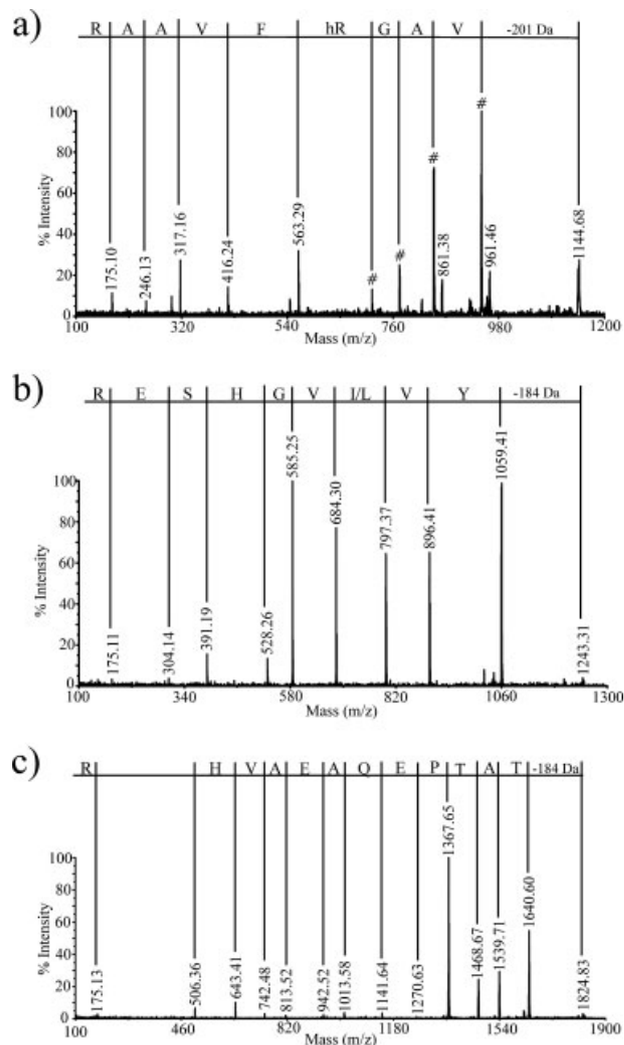


**Figure 1.** 2-D PAGE-separated proteins from *H. halophila* grown anaerobically under tungsten illumination (a) or green and blue light (b). Spots in which the protein was positively identified are numbered according to Table 1.



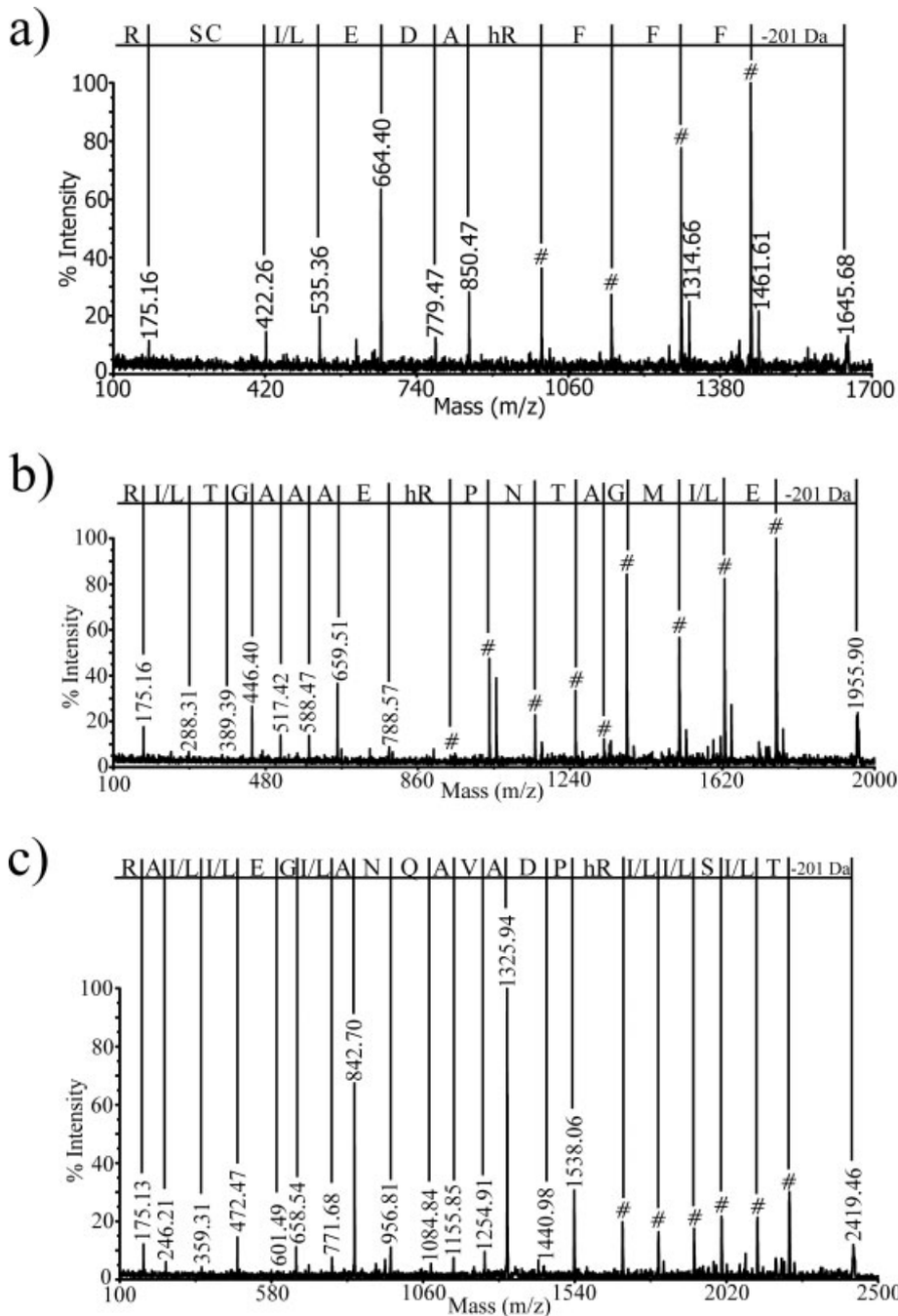
**Figure 2.** MALDI MS/MS spectra (positive ion mode) of the in-gel guanidinated and sulfonated tryptic peptide mixture of spot 5 (Fig. 1). Fragment spectra of peptide IKEIHQR (a), KVNIDDIR (b), KYAGVPFYR (c), and IKPIAIE (d). All labeled fragment ions are  $y$ -ions, ( $y$ -17)-ions resulting from the neutral loss of  $\text{NH}_3$  are indicated as #. Where appropriate, other fragment ions are indicated ( $b$ -ions). *De novo* derived sequence information is indicated in the one-letter code (homoarginine (hR)). Loss of the sulfonation label is indicated as  $-184$  Da or as a loss of  $-201$  Da including the neutral loss of ammonia ( $-17$  Da).

The *de novo* determined peptide sequences were used to identify the proteins by sequence similarity searching, as reviewed by Liska and Shevchenko [26]. Given the size and growth of the current databases, it is possible that many proteins already have homologs in a database.



**Figure 3.** MALDI MS/MS spectra (positive ion mode) of the in-gel guanidinated and sulfonated tryptic peptide mixture of spot 9 (Fig. 1). Fragment spectra of peptide VAGKFVAAR (a), YVIGHSER (b), and TATPEQAEAVH (c). Labeling is as in legend Fig. 2.

Database searching with MS-derived *de novo* peptide sequences allows the proteomic identification of proteins from organisms whose genomes have not been sequenced. However, MS and sequence similarity searches are difficult to combine. Conventional database search algorithms like BLAST or FASTA are optimized for accurate sequence queries that are longer than 35 amino acid residues. Usually, peptide sequences obtained by MS/MS do not exceed the length of a tryptic peptide, typically comprising 10–15 amino acids and, therefore, the statistical significance of retrieved hits is often ambiguous. Several database searching approaches have been reported that accommodate specific requirements of MS/MS sequencing. MS-driven BLAST (MS-



**Figure 4.** MALDI MS/MS spectra (positive ion mode) of the in-gel guanidinated and sulfonated tryptic peptide mixture of spot 15 (Fig. 1). Fragment spectra of peptide FFFKA-DEI(CS)R (a), EIMGATNP-KEAAAGTIR (b), and TISIIKP-DAVAQNAIGEIIAR (c). Labeling is as in legend Fig. 2.

BLAST) is a database search protocol for identifying unknown proteins by sequence similarity to homologous proteins available in a database. MS-BLAST utilizes redundant, degenerate, and partially inaccurate peptide sequence data obtained by *de novo* interpretation of MS/MS spectra. MS-BLAST does not allow gaps within individual peptides, while gaps between peptides are not penalized and can be of arbitrary length. Therefore, all peptide sequences obtained by the interpretation of acquired MS/MS are assembled into a single searching

string in arbitrary order [20, 27]. MS-Homology is a database searching tool from the UCSF Mass Spectrometry Facility (Protein Prospector 4.0.5) that performs homology-based searches [21]. The program allows the comparison of a number of *de novo* derived peptide sequences, followed by the maximum number of amino acid substitutions allowed for each sequence, against a selected database. Different peptides from the same unknown protein can be entered in the list. A database search will look for proteins containing peptides identical

or homologous to the listed sequences. The quality of the results will be dependent on the number of peptides sequenced and the accuracy of the sequence information entered, as well as on database completeness and species to species sequence variability for the peptides entered. It is also possible to enter a part of the sequence as a mass, along with a tolerance factor. FASTS is a recently reported sequence similarity search algorithm designed to use *de novo* sequence data from organisms lacking comprehensive sequence data. FASTS searches databases using peptide sequences of unknown order, evaluating all possible arrangements of the peptides. The algorithm uses the heuristic FASTA comparison strategy to accelerate the search, but also uses alignment probability, rather than a similarity score, as the criterion for alignment optimality. Because the true order of the query peptides used by FASTS is not known, FASTS only requires that the aligned peptides do not overlap [19].

The *de novo* derived sequence information from each spot was combined in one search query and analyzed using the three search algorithms. Only when the top results (first hits) from the three searches yielded the same protein, the identification was considered as positive. Most search queries included 30–70 amino acids, resulting from three to six peptide sequences (Supplementary Table 1). Using these queries, all three homology-based search algorithms yielded identifications with a score significantly better than the threshold score (Tables 1a and b). Three proteins were identified using only two *de novo* derived peptide sequences, and another three proteins were identified by using only one peptide sequence (>ten amino acid residues). In the latter cases, although all three algorithms yielded the same homologous protein, confirming a positive identification, the identification scores dropped significantly (Tables 1a and b). Surprisingly, we observed that the identification score obtained varied slightly according to the position of the peptide sequences in the query using MS-BLAST and, to a lesser extent, using the FASTS search algorithm. As it was formerly suggested that the peptide sequences can be used in an arbitrary way, this observation will be the subject of further investigation.

Using this approach, we were able to identify 31 proteins in the 74 spots, in which a PMF was observed, from both gels (42%) (Table 1a and b). In some of the fragmentation spectra none or insufficient sequence information was derived, most likely because of the weak intensity of the derivatized precursor. For other spots, sufficient *de novo* sequence information was obtained, but the homology search algorithms yielded a protein identification with a score below the threshold value, or no identification at all. According to simulation results, sequence-based meth-

ods, such as MS-BLAST and FASTS, are able to detect  $\pm 50\%$  of homologous sequences at the sequence identity level of  $\pm 50\%$ . The success of identification by sequence similarity searches will also depend on the number of recognized peptides from a digested protein. It has been calculated that, as more peptides are analyzed and matched, proteins of less similarity can be identified, the limit being around 50% identity. The simplest cases are those in which the proteins in question are highly conserved and can thus be identified *via* the sequences of their homologous proteins in other species. This strategy fails when the proteins are insufficiently similar. If the organism being studied is very distantly related to any organism with a sequenced genome, the likelihood of protein identification decreases [19, 28].

Some of the proteins we identified are definitively involved in the adaptation to halophilic life conditions. Sarcosine dimethylglycine *N*-methyltransferase (SDMT) of *Ectothiorhodospira halochloris* catalyzes the three-fold methylation of glycine to betaine, with *S*-adenosylmethionine (SAM) acting as the methyl group donor. Glycine betaine is accumulated in cells living in high salt concentrations in order to balance the osmotic pressure. SAM has an important role in DNA methylation and cell signaling. *S*-Adenosylmethionine synthetase catalyzes the formation of *S*-adenosylmethionine from *L*-methionine and ATP. However, a complete study of the photo-responses of *H. halophila*, and the role of PYP in this process, will require a differential display study of 2-D PAGE separated proteins from cells grown under different light conditions.

#### 4 Concluding remarks

The rapid and accurate identification of proteins is the primary goal of modern proteomics. MS/MS can generate some useful sequence information. However, manual interpretation of peptide spectra for *de novo* sequencing is often prohibitively challenging because of variation in favored ion fragmentation sites, the chemical nature of amino acid side chains and their relative order in a peptide backbone, and the presence of side-products such as neutral loss ions, contaminants, or noise peaks. Improvement of the fragmentation efficiency of peptides is of particular importance for MALDI-generated ions, because the predominant singly charged ions in MALDI generally fragment less good than doubly charged ions. The approach demonstrated here, consisting of *de novo* sequence analysis of derivatized peptides and homology-based identification, is a powerful technique for the identification of proteins with no genomic or other database information.

For 75% of the spots we observed a PMF upon in-gel guanidination and sulfonation of the extracted peptides. An apparently bad feature of the sulfonic acid derivatized peptides is their lower intensity in positive mode analysis, partly due to the suppression effect of the strong negative charge from the sulfonic group. The poorer positive ion-sensitivity is counterbalanced by a far more efficient fragmentation of sulfonated compared with non-sulfonated peptides. TOF/TOF analysis of underivatized peptides typically results in complex fragment spectra. After guanidination and sulfonation, a contiguous series of  $\gamma$ -ions was observed in almost all of the fragmentation spectra (Supplementary Table 1). The  $\gamma$ -ion series could easily be interpreted (manually or by using an algorithm) facilitating *de novo* sequencing.

The occurrence of “nonsequence” specific ion fragments, such as the neutral loss of ammonia, and preferential fragmentation pathways, such as the Xxx-Pro bond, can be used to improve predictive models of peptide fragmentation for *de novo* sequence analysis. The current understanding of the fragmentation mechanisms is still insufficient to ensure a high correlation between theoretically predicted MS/MS spectra and experimental results.

MS-Homology, MS-BLAST, and FASTS methods provide independent means of evaluating the statistical significance of hits and, therefore, it is not necessary to compare retrospectively the matched peptide sequences with actual tandem mass spectra to rule out false positive hits. As previously reported, the peptide sequences in the query were arbitrarily chosen [19, 28]. However, in this study we observed that the identification scores vary, according to the position of the sequences in the query, when applying the MS-BLAST and FASTS algorithm, a contradiction that will be the subject of a further study.

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