

The ABRF Edman Sequencing Research Group 2009 Study: Comparison of Edman and Mass Spectrometry Techniques for N-terminal Sequencing

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Introduction

For decades, high sensitivity Edman sequencing has been the method of choice for determining the N-terminal amino acid sequence of proteins¹. However, the major but by no means only limitation of this technique is its inability to obtain amino acid sequence from N-terminally blocked proteins. Mass spectrometric techniques for protein sequence analysis do not suffer from this limitation though unequivocal determination of protein N-termini on a routine basis has been elusive. The advantages of mass spectrometric techniques have in recent years driven investigators to look beyond Edman chemistry to find alternative technologies to obtain N-terminal sequence. Several mass spectrometry methodologies have been published, primarily for proteomics analyses, which may be quicker, less costly and more sensitive than Edman sequencing. Because such techniques involve a range of biochemical and instrumental methodologies having different advantages and limitations the ESRG has created a study to ascertain how reliably they can produce N-terminal amino acid sequence information and to compare those results to those obtained by automated Edman sequencing.

The ESRG 2009 study was designed to allow the participants freedom to use their analytical technique of choice to obtain as much N-terminal amino acid sequence information as possible from two test proteins in solution. Approximately one nanomole of each sample was provided so laboratories were able to attempt a variety of techniques with the goal of obtaining each protein's N-terminal sequence. Results of the analyses from participating laboratories are illustrated. Included is a comparison of methodology and instrumentation used by the laboratories to determine the N-terminal sequence of the protein.

Materials and Methods

TEST SAMPLE 1

E. coli BL21 was transformed with a pTrcHisTOPO plasmid containing a cDNA encoding the yeast alcohol dehydrogenase (ADH1) protein. Expression resulted in an ADH1 protein with a 35 amino acid leader sequence containing a hexahis-tag.

Alcohol Dehydrogenase 1 (Yeast) SP_P00330

GGSHHHHHHGMASMTGGQGRDLVDDDDKPTLMSIPETQKGVIFYESHGKLEYKDI PVFKFKANELLINVKYSVGHDTDLHAWHGDFWPLVFKLPLVGGHGGAGVGVGMGENVKGWKIGDYAGIKWLNGSCMAYCEBNGENSCPHADLSGYTHDGSFQYATADVAQAHI PQGDTLQAQVPI LCAGITVYFKALKSAMLMAGHWVAI SGAAGGSLGSLAVQYAKAMGVRVLGIDGGGKEELFRSIEGGVFPIDFTKEKDI VGVAVLKATDGGAHGVINVSVEAAIEASTRYVRANQTTVLYGMPAGAKCCSDVFNQVKSISVGSYVGNRDRTRALDFPARGLVKSPKIVGLSLPTEIYERKMEKQI VGRYVDVTSK

BOLD = 6-histidine-tagged beta galactosidase (cloning vector)

After transformation, *E. coli* was incubated in the presence of ampicillin (50µg/mL) and expression was induced with IPTG (0.5mM) in a New Brunswick Bioflow III fermentor with a 5L vessel. *E. coli* cells were harvested by centrifugation. Cells were re-suspended and lysed in a 25mM Tris-HCl pH 8.0, 0.3 M NaCl, 10mM Imidazole, 0.5% Triton-X 100. The lysate was loaded on a Ni Sepharose High Performance (GE Healthcare) resin column. The Ni column was washed with 25mM Tris-HCl pH 8.0, 0.3 M NaCl, 10mM Imidazole. After washing, the ADH1 protein was eluted with 25mM Tris-HCl pH 8.0, 0.3 M NaCl, 250mM Imidazole. Triton-X 100 was removed by washing 3X with Bio-Beads SM-2 Adsorbent (BioRad). ADH1 purity was assessed with MALDI-TOF, SDS-PAGE gel and Edman sequencing. The ADH1 was quantitated by amino acid analysis. One nmol of ADH1 was aliquoted in 1.5 ml low protein retention microcentrifuge tubes (Fisher) and dried for distribution to the study participants.

TEST SAMPLE 2

Glyceraldehyde-3-phosphate dehydrogenase, Rabbit Muscle, (G3P) was purchased from Sigma. (Cat. No. G5262)

Glyceraldehyde-3-phosphate dehydrogenase (Rabbit) SP_P46406

VKVGNGFRIGRLVTRAAFPNSKGVDDVAINDPFLDLHYVMYFQYDSHGKPFHTVKAENGKLVINGKATITFQERDPAINKWDGADAGEYVVESTGVPFTMEKAGAHKGGAKRVIISAPSDAPMFVGNVHKEKYNDSIKVSNASCTTCLAPLAKLIDHPFVLEGLMTVTHAIATATQKTVDFGSKLWRDGRGAQNIIIPASTGAAKAVGKVIPELNGKLGTMARFVPTFNVSVDLTCRLEKAAKDYDKKVKVQMSGEPGLKILGYTDEDVVSCDFNSATHSSTFDAGAGIALNDHFVFKLISWINDYFNGSNRVDLQVHAIASLKE

G3P was weighed and re-suspended in 0.1% TFA. One nmol was aliquoted in 1.5 mL low protein retention microcentrifuge tubes (Fisher) and dried for distribution to the study participants.

Table 1. Sequence Call by Participating Laboratory

Analysis Method	ESRG Lab #	N-terminal Sequence Data		Correct Protein ID	# Correct AA Calls
		Sequence	Protein ID		
Bottom-Up	001	GGSHHHHHHGMASMTGGQGRDLVDDDDKPTLMSIPETQKGVIFYESHGKLEYKDI PVFKFKANELLINVKYSVGHDTDLHAWHGDFWPLVFKLPLVGGHGGAGVGVGMGENVKGWKIGDYAGIKWLNGSCMAYCEBNGENSCPHADLSGYTHDGSFQYATADVAQAHI PQGDTLQAQVPI LCAGITVYFKALKSAMLMAGHWVAI SGAAGGSLGSLAVQYAKAMGVRVLGIDGGGKEELFRSIEGGVFPIDFTKEKDI VGVAVLKATDGGAHGVINVSVEAAIEASTRYVRANQTTVLYGMPAGAKCCSDVFNQVKSISVGSYVGNRDRTRALDFPARGLVKSPKIVGLSLPTEIYERKMEKQI VGRYVDVTSK	ADH1	Yes	12
	005			Yes	12
	011			Yes	7
	016			Yes	9
	018			Yes	11
	028			No	30
	029			No	34
	030			No	31
	034			No	42
	036			No	18
	037			No	29
	038			No	25
Edman	020			No	22
	022			No	0
	023			No	31
	026			No	30
	033			No	30
	035			No	34
	039			No	30
	040			No	21
	043			No	31
	044			No	21
	045			No	34
	046			No	34
Top-Down	001			Yes	44
	002			Yes	44
	003			Yes	19
	007			Yes	16
	010			No	18
	012			No	21
	013			Yes	0
	015			Yes	10
	016			Yes	18
	021			Yes	17
	022			Yes	31
	024			Yes	17
025			No	9	

Analysis Method	ESRG Lab #	N-terminal Sequence Data		Correct Protein ID	# Correct AA Calls
		Sequence	Protein ID		
Bottom-Up	001	VKVGNGFRIGRLVTRAAFPNSKGVDDVAINDPFLDLHYVMYFQYDSHGKPFHTVKAENGLKLVINGKATITFQERDPAINKWDGADAGEYVVESTGVPFTMEKAGAHKGGAKRVIISAPSDAPMFVGNVHKEKYNDSIKVSNASCTTCLAPLAKLIDHPFVLEGLMTVTHAIATATQKTVDFGSKLWRDGRGAQNIIIPASTGAAKAVGKVIPELNGKLGTMARFVPTFNVSVDLTCRLEKAAKDYDKKVKVQMSGEPGLKILGYTDEDVVSCDFNSATHSSTFDAGAGIALNDHFVFKLISWINDYFNGSNRVDLQVHAIASLKE	G3P	Yes	8
	005			Yes	9
	011			Yes	10
	016			Yes	8
	018			Yes	9
	028			Yes	20
	029			Yes	35
	030			Yes	49
	034			Yes	75
	036			Yes	30
	037			Yes	41
	038			Yes	33
Edman	020			Yes	26
	022			Yes	40
	023			Yes	26
	026			Yes	40
	033			Yes	26
	035			Yes	64
	039			Yes	24
	040			Yes	37
	043			Yes	18
	044			Yes	30
	045			Yes	30
	Top-Down	001			Yes
002				Yes	44
003				Yes	26
007				Yes	32
010				Yes	30
012				Yes	29
013				Yes	22
015				Yes	32
016				Yes	25
021				Yes	26
022				Yes	26
024				Yes	12
025			Yes	13	

Box color code: ■ Correct amino acid call based on sequence data alone ■ Protein ID from a data base search was used to identify the amino acid
■ Incorrect amino acid call
 Labs A, B, C and D data were contributed by ESRC committee members
 * ESRC 003, 003, 025 and 026 Proteins identified from Bottom-Up data (not shown)
 ** Many laboratories using the Top-Down technique also supplied C-terminal sequence data (data not shown)
 *** ESRC 021 evidence for the complete N-terminal sequence not present in the MS/MS spectra

Table 2. Instrumentation and Other Parameters used by Participating Laboratories

Analysis Method	ESRG Lab #	Instrumentation and Methods						
		Manufacturer	Instrument	Ionization	Reflection	Detection System	Reagent Solvent	
Bottom-Up	001	Waters/Micromass	QTOF Premier	ESI	reflector	Trap & Q-TOF	0.1% TFA	
	002	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	003	Waters/Micromass	QTOF Premier	ESI	reflector	Trap & Q-TOF	0.1% TFA	
	005	ABI	494HT	ESI	reflector	TOF	0.1% TFA	
	009	ABI	494HT	ESI	reflector	TOF	0.1% TFA	
	014	ABI	494HT	ESI	reflector	TOF	0.1% TFA	
	016	ABI	494HT	ESI	reflector	TOF	0.1% TFA	
	018	Waters/Micromass	QTOF Premier	ESI	reflector	Trap & Q-TOF	0.1% TFA	
	021	Waters/Micromass	QTOF Premier	ESI	reflector	Trap & Q-TOF	0.1% TFA	
	028	Waters/Micromass	QTOF Premier	ESI	reflector	Trap & Q-TOF	0.1% TFA	
	Edman	005	ABI	494HT	ESI	reflector	TOF	0.1% TFA
		009	ABI	494HT	ESI	reflector	TOF	0.1% TFA
014		ABI	494HT	ESI	reflector	TOF	0.1% TFA	
016		ABI	494HT	ESI	reflector	TOF	0.1% TFA	
018		ABI	494HT	ESI	reflector	TOF	0.1% TFA	
020		ABI	494HT	ESI	reflector	TOF	0.1% TFA	
022		ABI	494HT	ESI	reflector	TOF	0.1% TFA	
023		ABI	494HT	ESI	reflector	TOF	0.1% TFA	
026		ABI	494HT	ESI	reflector	TOF	0.1% TFA	
033		ABI	494HT	ESI	reflector	TOF	0.1% TFA	
Top-Down		001	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA
		002	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA
	003	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	007	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	010	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	012	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	013	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	015	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	016	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	021	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	022	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	
	024	Braker	Ultrahigh	MALDI	reflector	TOF	0.1% TFA	

Fig. 1 Example Data Supplied for the Edman Degradation Chemistry Approach from ESRC-A – Test Sample 2

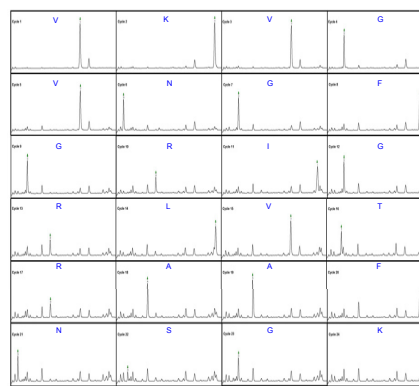


Fig. 2 Example Data Supplied for the Bottom-Up Approach from ESRC-016 – Test Sample 2

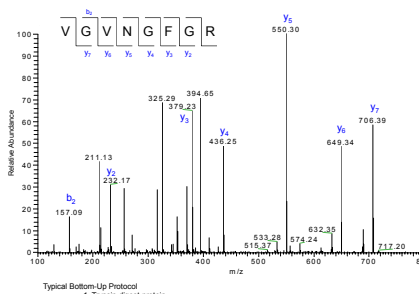


Fig. 3 Example ISD Data Supplied for the Top-Down Approach from ESRC-022 – Test Sample 1

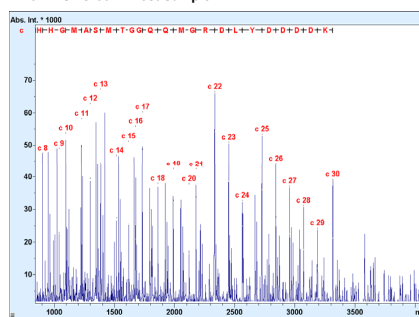


Table 3 Comparison of the Techniques used to Determine the N-terminal Sequence

Sequence Analysis Technique	Mechanism	Advantages	Disadvantages
Edman Degradation	Sequential chemical degradation of the protein by coupling of PITC to the N-terminus followed by cleavage with TFA and identification of the resulting phenylthiohydantoin-amino acid by liquid chromatography and UV detection.	Direct determination of the protein N-terminal sequence. All commonly occurring amino acids identifiable. Stable PTMs identified by retention time compared to the known modified PTH-AA standard.	N-terminal blockage (e.g. N-acetylation) prevents sequencing. Length of sequence read limited by problematic. Requires high-purity chemicals. High cost. Low throughput. 20 to 40 minutes per cycle for each residue. Commercial production of instrument ceased June 2008.
Top-Down MALDI In-Source Decay ^{1,2}	Matrix generated hydrogen radiol mediated fragmentation of intact protein in the ion source typically with higher laser power and chemical additives. Fragmentation occurs in the nanosecond time scale.	Rapid, high throughput, low cost. C-terminal sequence can also be determined. Stable PTMs identifiable. Isolation of terminal fragments for mass possible but instrument configuration dependent. Instruments commonly available.	ISD doesn't provide precursor ion selection. Sample must be purified. May not yield complete N-terminal sequence i.e. protein termini must be accessible to ionization/fragmentation process. Cannot distinguish isobaric amino acids. Background ion interference in the low mass region - the entire ion series representing the N-terminus may not be present.
Top-Down ESI Electron Transfer/ Dissociation ³	Electron transfer from Antiradicals anions to multiply protonated precursor cations resulting in radical new technique. Fragmentation occurs in the millisecond time scale.	Preserves PTM structures, rapid, high throughput. Precursor ion selection allows MS of known precursor ion mass. Extensive peptide backbone fragmentation.	ETD fragmentation requires multiply protonated precursors. For MS ⁿ of CAD generated low charge state precursors ETD efficiency decreases the entire ion series representing the N-terminus may not be present. Cannot distinguish isobaric amino acids. Relatively new technique. Fewer instruments in use. Requires high resolution mass analyzer to obtain charge state of protein precursor - higher cost of instrumentation. Only structurally accessible portions of the precursor are fragmented.
Bottom-Up ESI MS/MS MALDI TOF/MS MALDI PSD ⁴	Enzymatic or chemical cleavage of the protein followed by mass analysis of the peptide mixture and bioinformatic analysis to identify the N-terminus. Biochemical techniques to obtain the N-terminal peptide prior to mass analysis.	Generates smaller fragments that are amenable to analysis on a wide variety of instrumentation. High throughput, low cost depending on specific technique used.	Successful isolation of the N-terminal peptide is dependent, possibly requiring multiple biochemical approaches. Isobaric amino acids not determined. Absence of the complete ion series in the mass spectrum is common and prevents interpretation of the precursor acid sequence for the entire peptide.

Conclusions

- Edman sequencing remains a reliable means for determining the N-terminal sequence of an unblocked protein. The majority of labs that used this approach were able to easily sequence both test proteins. However, no one that used this technique was able to identify Test Sample 1 because of the long vector sequence.
- The Top-Down approach shows great promise for determining the N- (and C) terminal sequence of a protein in solution. The majority of labs using this approach correctly identified long strings of amino acids near the N- (and C) terminus. Low mass ions (< 700 m/z) were poorly resolved, making the first 5-7 amino acids difficult to call. Most labs easily identified both proteins.
- The Bottom-Up approach (without modification to the N-terminal amino acid) works well when the mass fingerprint can be matched to a protein in a data base. Those labs using the bottom-up approach found the N-terminal fragment of Test Sample 2 and were able to MS/MS sequence this fragment.
- Labs using a mass spectrometry approach often depend on the protein ID from a data base search to fill in the following:
 - Amino acids with isobaric masses (Ile/Leu and Gln/Lys)
 - Regions with poor ion signal (peptide bonds associated with Pro)
 - Missing low mass ions (< 700 m/z for ISD)
- For those performing Edman chemistry, sequence length was typically shorter for Test Sample 1 (contained a 6 histidine-tag) than Test Sample 2. This matches observations from the ESRC 2008 study that found histidine-tags are more difficult to sequence than other proteins. The histidine-tag was not a problem for those using an MS approach.

References

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