B-01

Revision 1 Correction 1

PHARMACOPOEIAL DISCUSSION GROUP

CORRECTION

CODE: B-01 NAME: Amino Acid Determination

(Correction of the sign-off document Revision 1 signed on 26 October 2016)

Item to be corrected:

"Methodologies of Amino Acid Analysis General Principles"

The last sentence in "METHOD 8- Precolumn NBD-F Derivatization General Principle" was changed as below;

"Profile analysis was <u>can be</u> achieved for about 1.5 mg μg of protein hydrolysates in the final precolumn labeling reaction mixture for HPLC."

Harmonised provisions:

Provision	EP	JP	USP
Introduction	+	+	+
Apparatus	+	+	+
General Precautions	+	+	+
Reference Standard Material	+	+	+
Calibration of Instrumentation	+	+	+
Repeatability	+	+	+
Sample Preparation	+	+	+
Internal Standards	+	+	+
Protein Hydrolysis	+	+	+
Methodologies of Amino Acid Analysis General Principles	+	+	+
Data Calculation and Analysis	+	+	+

+ will adopt and implement; - will not stipulate

Non-harmonised provisions:

None

B-01

Local requirements

EP	JP	USP
None	None	Footnote : "Suitable standards may be obtained from NIST (Gaithersburg, MD), Beckman Instruments (Fullerton, CA), Sigma Chemical (St. Louis, MO), Pierce (Rockford, IL), or Agilent (Palo Alto, CA)."

European Pharmacopoeia

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AMINO ACID ANALYSIS

2 3 Amino acid analysis refers to the methodology used to determine the 4 amino acid composition or content of proteins, peptides, and other pharmaceutical 5 preparations. Proteins and peptides are macromolecules consisting of covalently 6 bonded amino acid residues organized as a linear polymer. The sequence of the 7 amino acids in a protein or peptide determines the properties of the molecule. 8 Proteins are considered large molecules that commonly exist as folded structures 9 with a specific conformation, while peptides are smaller and may consist of only a 10 few amino acids. Amino acid analysis can be used to quantify protein and 11 peptides, to determine the identity of proteins or peptides based on their amino 12 acid composition, to support protein and peptide structure analysis, to evaluate 13 fragmentation strategies for peptide mapping, and to detect atypical amino acids 14 that might be present in a protein or peptide. It is necessary to hydrolyze a 15 protein/peptide to its individual amino acid constituents before amino acid 16 analysis. Following protein/peptide hydrolysis, the amino acid analysis procedure 17 can be the same as that practiced for free amino acids in other pharmaceutical 18 preparations. The amino acid constituents of the test sample are typically 19 derivatized for analysis. 20 **Apparatus** 21 22 23 Methods used for amino acid analysis are usually based on a 24 chromatographic separation of the amino acids present in the test sample. Current 25 techniques take advantage of the automated chromatographic instrumentation 26 designed for analytical methodologies. An amino acid analysis instrument will 27 typically be a low-pressure or high-pressure liquid chromatograph capable of 28 generating mobile phase gradients that separate the amino acid analytes on a 29 chromatographic column. The instrument must have postcolumn derivatization 30 capability, unless the sample is analyzed using precolumn derivatization. The 31 detector is usually an ultraviolet-visible or fluorescence detector depending on the 32 derivatization method used. A recording device (e.g., integrator) is used for

33 transforming the analog signal from the detector and for quantitation. It is 34 preferred that instrumentation be dedicated particularly for amino acid analysis. 35 **General Precautions** 36 37 38 Background contamination is always a concern for the analyst in 39 performing amino acid analysis. High purity reagents are necessary (e.g., low 40 purity hydrochloric acid can contribute to glycine contamination). Analytical 41 reagents are changed routinely every few weeks using only high-pressure liquid 42 chromatography (HPLC) grade solvents. Potential microbial contamination and 43 foreign material that might be present in the solvents are reduced by filtering 44 solvents before use, keeping solvent reservoirs covered, and not placing amino 45 acid analysis instrumentation in direct sunlight. 46 Laboratory practices can determine the quality of the amino acid analysis. 47 Place the instrumentation in a low traffic area of the laboratory. Keep the 48 laboratory clean. Clean and calibrate pipets according to a maintenance schedule. 49 Keep pipet tips in a covered box; the analysts may not handle pipet tips with their 50 hands. The analysts may wear powder-free latex or equivalent gloves. Limit the 51 number of times a test sample vial is opened and closed because dust can 52 contribute to elevated levels of glycine, serine, and alanine. 53 A well-maintained instrument is necessary for acceptable amino acid 54 analysis results. If the instrument is used on a routine basis, it is to be checked 55 daily for leaks, detector and lamp stability, and the ability of the column to 56 maintain resolution of the individual amino acids. Clean or replace all instrument 57 filters and other maintenance items on a routine schedule. 58 **Reference Standard Material** 59 60 61 Acceptable amino acid standards are commercially available for amino 62 acid analysis and typically consist of an aqueous mixture of amino acids. When determining amino acid composition, protein or peptide standards are analyzed 63 64 with the test material as a control to demonstrate the integrity of the entire

procedure. Highly purified bovine serum albumin has been used as a proteinstandard for this purpose.

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Calibration of Instrumentation

70 Calibration of amino acid analysis instrumentation typically involves 71 analyzing the amino acid standard, which consists of a mixture of amino acids at a 72 number of concentrations, to determine the response factor and range of analysis 73 for each amino acid. The concentration of each amino acid in the standard is 74 known. In the calibration procedure, the analyst dilutes the amino acid standard 75 to several different analyte levels within the expected linear range of the amino 76 acid analysis technique. Then, replicates at each of the different analyte levels 77 can be analyzed. Peak areas obtained for each amino acid are plotted versus the 78 known concentration for each of the amino acids in the standard dilution. These 79 results will allow the analyst to determine the range of amino acid concentrations 80 where the peak area of a given amino acid is an approximately linear function of 81 the amino acid concentration. It is important that the analyst prepare the samples for amino acid analysis so that they are within the analytical limits (e.g., linear 82 83 working range) of the technique employed in order to obtain accurate and 84 repeatable results.

85 Four to six amino acid standard levels are analyzed to determine a 86 response factor for each amino acid. The response factor is calculated as the 87 average peak area or peak height per nmol of amino acid present in the standard. 88 A calibration file consisting of the response factor for each amino acid is prepared 89 and used to calculate the concentration of each amino acid present in the test 90 sample. This calculation involves dividing the peak area corresponding to a given 91 amino acid by the response factor for that amino acid to give the nmol of the 92 amino acid. For routine analysis, a single-point calibration may be sufficient; 93 however, the calibration file is updated frequently and tested by the analysis of 94 analytical controls to ensure its integrity.

95

96 97	Repeatability
97 98	Consistent high quality amino acid analysis results from an analytical
99	laboratory require attention to the repeatability of the assay. During analysis of
100	the chromatographic separation of the amino acids or their derivatives, numerous
101	peaks can be observed on the chromatogram that correspond to the amino acids.
102	The large number of peaks makes it necessary to have an amino acid analysis
103	system that can repeatedly identify the peaks based on retention time and integrate
104	the peak areas for quantitation. A typical repeatability evaluation involves
105	preparing a standard amino acid solution and analyzing many replicates (i.e., six
106	analyses or more) of the same standard solution. The relative standard deviation
107	(RSD) is determined for the retention time and integrated peak area of each amino
108	acid. An evaluation of the repeatability is expanded to include multiple assays
109	conducted over several days by different analysts. Multiple assays include the
110	preparation of standard dilutions from starting materials to determine the variation
111	due to sample handling. Often the amino acid composition of a standard protein
112	(e.g., bovine serum albumin) is analyzed as part of the repeatability evaluation.
113	By evaluating the replicate variation (i.e., RSD), the laboratory can establish
114	analytical limits to ensure that the analyses from the laboratory are under control.
115	It is desirable to establish the lowest practical variation limits to ensure the best
116	results. Areas to focus on to lower the variability of the amino acid analysis
117	include sample preparation, high background spectral interference due to quality
118	of reagents and/or laboratory practices, instrument performance and maintenance,
119	data analysis and interpretation, and analyst performance and habits. All
120	parameters involved are fully investigated in the scope of the validation work.
121 122	Sample Preparation
123	
124	Accurate results from amino acid analysis require purified protein and
125	peptide samples. Buffer components (e.g., salts, urea, detergents) can interfere
126	with the amino acid analysis and are removed from the sample before analysis.
127	Methods that utilize postcolumn derivatization of the amino acids are generally
128	not affected by buffer components to the extent seen with precolumn

129	derivatization methods. It is desirable to limit the number of sample
130	manipulations to reduce potential background contamination, to improve analyte
131	recovery, and to reduce labor. Common techniques used to remove buffer
132	components from protein samples include the following methods: (1) injecting
133	the protein sample onto a reversed-phase HPLC system, removing the protein
134	with a volatile solvent containing a sufficient organic component, and drying the
135	sample in a vacuum centrifuge; (2) dialysis against a volatile buffer or water; (3)
136	centrifugal ultrafiltration for buffer replacement with a volatile buffer or water;
137	(4) precipitating the protein from the buffer using an organic solvent (e.g.,
138	acetone); and (5) gel filtration.
139	
140	Internal Standards
141 142	It is recommended that an internal standard be used to monitor physical
143	and chemical losses and variations during amino acid analysis. An accurately
144	known amount of internal standard can be added to a protein solution prior to
145	hydrolysis. The recovery of the internal standard gives the general recovery of
146	the amino acids of the protein solution. Free amino acids, however, do not
147	behave in the same way as protein-bound amino acids during hydrolysis because
148	their rates of release or destruction are variable. Therefore, the use of an internal
149	standard to correct for losses during hydrolysis may give unreliable results. It will
150	be necessary to take this point under consideration when interpreting the results.
151	Internal standards can also be added to the mixture of amino acids after hydrolysis
152	to correct for differences in sample application and changes in reagent stability
153	and flow rates. Ideally, an internal standard is an unnaturally occurring primary
154	amino acid that is commercially available and inexpensive. It should also be
155	stable during hydrolysis, its response factor should be linear with concentration,
156	and it needs to elute with a unique retention time without overlapping other amino
157	acids. Commonly used amino acid standards include norleucine, nitrotyrosine,
158	and α -aminobutyric acid.
159	

160	Protein Hydrolysis
161	
162	Hydrolysis of protein and peptide samples is necessary for amino acid
163	analysis of these molecules. The glassware used for hydrolysis must be very
164	clean to avoid erroneous results. Glove powders and fingerprints on hydrolysis
165	tubes may cause contamination. To clean glass hydrolysis tubes, boil tubes for 1
166	hour in 1 N hydrochloric acid or soak tubes in concentrated nitric acid or in a
167	mixture of concentrated hydrochloric acid and concentrated nitric acid (1:1).
168	Clean hydrolysis tubes are rinsed with high-purity water followed by a rinse with
169	HPLC grade methanol, dried overnight in an oven, and stored covered until use.
170	Alternatively, pyrolysis of clean glassware at 500°C for 4 hours may also be used
171	to eliminate contamination from hydrolysis tubes. Adequate disposable
172	laboratory material can also be used.
173	Acid hydrolysis is the most common method for hydrolyzing a protein
174	sample before amino acid analysis. The acid hydrolysis technique can contribute
175	to the variation of the analysis due to complete or partial destruction of several
176	amino acids. Tryptophan is destroyed; serine and threonine are partially
177	destroyed; methionine might undergo oxidation; and cysteine is typically
178	recovered as cystine (but cystine recovery is usually poor because of partial
179	destruction or reduction to cysteine). Application of adequate vacuum (\leq less
180	than 200 μ m of mercury or 26.7 Pa) or introduction of an inert gas (argon) in the
181	headspace of the reaction vessel can reduce the level of oxidative destruction. In
182	peptide bonds involving isoleucine and valine the amido bonds of Ile-Ile, Val-Val,
183	Ile-Val, and Val-Ile are partially cleaved; and asparagine and glutamine are
184	deamidated, resulting in aspartic acid and glutamic acid, respectively. The loss of
185	tryptophan, asparagine, and glutamine during an acid hydrolysis limits
186	quantitation to 17 amino acids. Some of the hydrolysis techniques described are
187	used to address these concerns. Some of the hydrolysis techniques described (i.e.,
188	Methods 4-11) may cause modifications to other amino acids. Therefore, the
189	benefits of using a given hydrolysis technique are weighed against the concerns
190	with the technique and are tested adequately before employing a method other
191	than acid hydrolysis.

192 A time-course study (i.e., amino acid analysis at acid hydrolysis times of 193 24, 48, and 72 hours) is often employed to analyze the starting concentration of 194 amino acids that are partially destroyed or slow to cleave. By plotting the 195 observed concentration of labile amino acids (i.e., serine and threonine) versus 196 hydrolysis time, the line can be extrapolated to the origin to determine the starting 197 concentration of these amino acids. Time-course hydrolysis studies are also used 198 with amino acids that are slow to cleave (e.g., isoleucine and valine). During the 199 hydrolysis time course, the analyst will observe a plateau in these residues. The 200 level of this plateau is taken as the residue concentration. If the hydrolysis time is 201 too long, the residue concentration of the sample will begin to decrease, indicating 202 destruction by the hydrolysis conditions.

203 An acceptable alternative to the time-course study is to subject an amino 204 acid calibration standard to the same hydrolysis conditions as the test sample. 205 The amino acid in free form may not completely represent the rate of destruction 206 of labile amino acids within a peptide or protein during the hydrolysis. This is 207 especially true for peptide bonds that are slow to cleave (e.g., Ile-Val bonds). 208 However, this technique will allow the analyst to account for some residue 209 destruction. Microwave acid hydrolysis has been used and is rapid but requires special equipment as well as special precautions. The optimal conditions for 210 211 microwave hydrolysis must be investigated for each individual protein/peptide 212 sample. The microwave hydrolysis technique typically requires only a few 213 minutes, but even a deviation of one minute may give inadequate results (e.g., 214 incomplete hydrolysis or destruction of labile amino acids). Complete 215 proteolysis, using a mixture of proteases, has been used but can be complicated, 216 requires the proper controls, and is typically more applicable to peptides than 217 proteins. 218 NOTE—During initial analyses of an unknown protein, experiments with 219 various hydrolysis time and temperature conditions are conducted to determine 220 the optimal conditions.

Method 1

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Acid hydrolysis using hydrochloric acid containing phenol is the most

225 common procedure used for protein/peptide hydrolysis preceding amino acid

analysis. The addition of phenol to the reaction prevents the halogenation oftyrosine.

Hydrolysis Solution: 6 N hydrochloric acid containing 0.1% to 1.0% ofphenol.

230 Procedure—

231 Liquid Phase Hydrolvsis—Place the protein or peptide sample in a 232 hydrolysis tube, and dry. [NOTE—The sample is dried so that water in the sample 233 will not dilute the acid used for the hydrolysis.] Add 200 µL of Hydrolysis 234 Solution per 500 µg of lyophilized protein. Freeze the sample tube in a dry ice-235 acetone bath, and flame seal in vacuum. Samples are typically hydrolyzed at 236 110°C for 24 hours in vacuum or inert atmosphere to prevent oxidation. Longer 237 hydrolysis times (e.g., 48 and 72 hours) are investigated if there is a concern that 238 the protein is not completely hydrolyzed.

239 *Vapor Phase Hydrolysis*—This is one of the most common acid hydrolysis 240 procedures, and it is preferred for microanalysis when only small amounts of the 241 sample are available. Contamination of the sample from the acid reagent is also 242 minimized by using vapor phase hydrolysis. Place vials containing the dried 243 samples in a vessel that contains an appropriate amount of *Hydrolysis Solution*. 244 The *Hydrolysis Solution* does not come in contact with the test sample. Apply an 245 inert atmosphere or vacuum (\leq less than 200 µm of mercury or 26.7 Pa) to the 246 headspace of the vessel, and heat to about 110°C for a 24-hour hydrolysis time. 247 Acid vapor hydrolyzes the dried sample. Any condensation of the acid in the 248 sample vials is minimized. After hydrolysis, dry the test sample in vacuum to 249 remove any residual acid.

250 251

Method 2

- 252 Tryptophan oxidation during hydrolysis is decreased by using
- 253 mercaptoethanesulfonic acid (MESA) as the reducing acid.
- 254 **Hydrolysis Solution:** 2.5 M MESA solution.

255	Vapor Phase Hydrolysis—About 1 to 100 µg of the protein/peptide
256	under test is dried in a hydrolysis tube. The hydrolysis tube is placed in a larger
257	tube with about 200 μ L of the <i>Hydrolysis Solution</i> . The larger tube is sealed in
258	vacuum (about 50 µm of mercury or 6.7 Pa) to vaporize the Hydrolysis Solution.
259	The hydrolysis tube is heated to 170°C to 185°C for about 12.5 minutes. After
260	hydrolysis, the hydrolysis tube is dried in vacuum for 15 minutes to remove the
261	residual acid.
262	
263	Method 3
264 265	Tryptophan oxidation during hydrolysis is prevented by using thioglycolic
266	acid (TGA) as the reducing acid.
267	Hydrolysis Solution—A solution containing 7 M hydrochloric acid, 10%
268	of trifluoroacetic acid, 20% of thioglycolic acid, and 1% of phenol.
269	Vapor Phase Hydrolysis—About 10 to 50 µg of the protein/peptide
270	under test is dried in a sample tube. The sample tube is placed in a larger tube
271	with about 200 μ L of the <i>Hydrolysis Solution</i> . The larger tube is sealed in
272	vacuum (about 50 μ m of mercury or 6.7 Pa) to vaporize the TGA. The sample
273	tube is heated to 166°C for about 15 to 30 minutes. After hydrolysis, the sample
274	tube is dried in vacuum for 5 minutes to remove the residual acid. Recovery of
275	tryptophan by this method may be dependent on the amount of sample present.
276 277	Method 4
278 279	Cysteine-cystine and methionine oxidation is performed with performic
280	acid before the protein hydrolysis.
281	Oxidation Solution —The performic acid is prepared fresh by mixing
282	formic acid and 30 percent hydrogen peroxide (9:1), and incubated at room
283	temperature for 1 hour.
284	Procedure —The protein/peptide sample is dissolved in 20 μ L of formic
285	acid, and heated at 50°C for 5 minutes; then 100 μ L of the Oxidation Solution is
286	added. The oxidation is allowed to proceed for 10 to 30 minutes. In this reaction,
287	cysteine is converted to cysteic acid and methionine is converted to methionine

288	sulfone. The excess reagent is removed from the sample in a vacuum centrifuge.
289	This technique may cause modifications to tyrosine residues in the presence of
290	halides. The oxidized protein can then be acid hydrolyzed using Method 1 or
291	Method 2.
292 293	Method 5
294 295	Cysteine-cystine oxidation is accomplished during the liquid phase
296	hydrolysis with sodium azide.
297	Hydrolysis Solution: 6 N hydrochloric acid containing 0.2% of phenol,
298	to which is added sodium azide to obtain a final concentration of 0.2% (w/v). The
299	added phenol prevents halogenation of tyrosine.
300	Liquid Phase Hydrolysis—The protein/peptide hydrolysis is conducted
301	at about 110°C for 24 hours. During the hydrolysis, the cysteine-cystine present
302	in the sample is converted to cysteic acid by the sodium azide present in the
303	Hydrolysis Solution. This technique allows better tyrosine recovery than Method
304	4, but it is not quantitative for methionine. Methionine is converted to a mixture
305	of the parent methionine and its two oxidative products, methionine sulfoxide and
306	methionine sulfone.
307 308	Method 6
309 310	Cysteine-cystine oxidation is accomplished with dimethyl sulfoxide
311	(DMSO).
312	Hydrolysis Solution: 6 N hydrochloric acid containing 0.1% to 1.0% of
313	phenol, to which DMSO is added to obtain a final concentration of 2% (v/v).
314	Vapor Phase Hydrolysis—The protein/peptide hydrolysis is conducted at
315	about 110°C for 24 hours. During the hydrolysis, the cysteine-cystine present in
316	the sample is converted to cysteic acid by the DMSO present in the Hydrolysis
317	Solution. As an approach to limit variability and compensate for partial
318	destruction, it is recommended to evaluate the cysteic acid recovery from
319	oxidative hydrolyses of standard proteins containing 1 to 8 mol of cysteine per
320	mol protein. The response factors from protein/peptide hydrolysates are typically

321	about 30% lower than those for nonhydrolyzed cysteic acid standards. Because
322	histidine, methionine, tyrosine, and tryptophan are also modified, a complete
323	compositional analysis is not obtained with this technique.
324 325	Method 7
326 327	Cysteine-cystine reduction and alkylation is accomplished by a vapor
328	phase pyridylethylation reaction.
329	Reducing Solution —Transfer 83.3 μ L of pyridine, 16.7 μ L of 4-
330	vinylpyridine, 16.7 μ L of tributylphosphine, and 83.3 μ L of water to a suitable
331	container, and mix.
332	Procedure —Add the protein/peptide (between 1 and 100 μ g) to a
333	hydrolysis tube, and place in a larger tube. Transfer the Reducing Solution to the
334	large tube, seal in vacuum (about 50 μ m of mercury or 6.7 Pa), and incubate at
335	about 100°C for 5 minutes. Then remove the inner hydrolysis tube, and dry it in a
336	vacuum desiccator for 15 minutes to remove residual reagents. The
337	pyridylethylated protein/peptide can then be acid hydrolyzed using previously
338	described procedures. The pyridylethylation reaction is performed
339	simultaneously with a protein standard sample containing 1 to 8 mol of cysteine
340	per mol protein to improve accuracy in the pyridylethyl-cysteine recovery.
341	Longer incubation times for the pyridylethylation reaction can cause
342	modifications to the α -amino terminal group and the ϵ -amino group of lysine in
343	the protein.
344	Method 8
345 346	Cysteine-cystine reduction and alkylation is accomplished by a liquid
347	phase pyridylethylation reaction.
348	Stock Solutions—Prepare and filter three solutions: 1 M Tris
349	hydrochloride (pH 8.5) containing 4 mM edetate disodium (Stock Solution A), 8
350	M guanidine hydrochloride (Stock Solution B), and 10% of 2-mercaptoethanol in
351	water (<i>Stock Solution C</i>).

352	Reducing Solution —Prepare a mixture of <i>Stock Solution B</i> and <i>Stock</i>
353	Solution A (3:1) to obtain a buffered solution of 6 M guanidine hydrochloride in
354	0.25 M Tris hydrochloride.
355	Procedure —Dissolve about 10 μ g of the test sample in 50 μ L of the
356	Reducing Solution, and add about 2.5 μ L of Stock Solution C. Store under
357	nitrogen or argon for 2 hours at room temperature in the dark. To achieve the
358	pyridylethylation reaction, add about 2 μ L of 4-vinylpyridine to the protein
359	solution, and incubate for an additional 2 hours at room temperature in the dark.
360	The protein/peptide is desalted by collecting the protein/peptide fraction from a
361	reversed-phase HPLC separation. The collected sample can be dried in a vacuum
362	centrifuge before acid hydrolysis.
363 364	Method 9
365	
366	Cysteine-cystine reduction and alkylation is accomplished by a liquid
367	phase carboxymethylation reaction.
368	Stock Solutions—Prepare as directed for <i>Method 8</i> .
369	Carboxymethylation Solution —Prepare a solution containing 100 mg of
370	iodoacetamide per mL of alcohol.
371	Buffer Solution —Use the <i>Reducing Solution</i> , prepared as directed for
372	Method 8.
373	Procedure —Dissolve the test sample in 50 μ L of the <i>Buffer Solution</i> , and
374	add about 2.5 μ L of <i>Stock Solution C</i> . Store under nitrogen or argon for 2 hours
375	at room temperature in the dark. Add the <i>Carboxymethylation Solution</i> in a ratio
376	1.5 fold per total theoretical content of thiols, and incubate for an additional 30
377	minutes at room temperature in the dark. [NOTE—If the thiol content of the
378	protein is unknown, then add 5 μ L of 100 mM iodoacetamide for every 20 nmol
379	of protein present.] The reaction is stopped by adding excess of 2-
380	mercaptoethanol. The protein/peptide is desalted by collecting the protein/peptide
381	fraction from a reversed-phase HPLC separation. The collected sample can be
382	dried in a vacuum centrifuge before acid hydrolysis. The S-carboxyamidomethyl-

383	cysteine formed will be converted to S-carboxymethylcysteine during acid
384	hydrolysis.
385	Method 10
386 387	Cysteine-cystine is reacted with dithiodiglycolic acid or dithiodipropionic
388	acid to produce a mixed disulfide. [NOTE—The choice of dithiodiglycolic acid or
389	dithiodipropionic acid depends on the required resolution of the amino acid
390	analysis method.]
391	Reducing Solution—A solution containing 10 mg of dithiodiglycolic acid
392	(or dithiodipropionic acid) per mL of 0.2 M sodium hydroxide.
393	Procedure —Transfer about 20 μ g of the test sample to a hydrolysis tube,
394	and add 5 μL of the Reducing Solution. Add 10 μL of isopropyl alcohol, and then
395	remove all of the sample liquid by vacuum centrifugation. The sample is then
396	hydrolyzed using Method 1. This method has the advantage that other amino acid
397	residues are not derivatized by side reactions, and the sample does not need to be
398	desalted prior to hydrolysis.
399 400	
401 402	METHOD 11 Asparagine and glutamine are converted to aspartic acid and glutamic
403	acid, respectively, during acid hydrolysis. Asparagine and aspartic acid residues
404	are added and represented by Asx, while glutamine and glutamic acid residues are
405	added and represented by Glx. Proteins/peptides can be reacted with bis(1,1-
406	trifluoroacetoxy)iodobenzene (BTI) to convert the asparagine and glutamine
407	residues to diaminopropionic acid and diaminobutyric acid residues, respectively,
408	upon acid hydrolysis. These conversions allow the analyst to determine the
409	asparagine and glutamine content of a protein/peptide in the presence of aspartic
410	acid and glutamic acid residues.
411	Reducing Solutions —Prepare and filter three solutions: a solution of 10
412	mM trifluoroacetic acid (Solution A), a solution of 5 M guanidine hydrochloride
413	and 10 mM trifluoroacetic acid (Solution B), and a freshly prepared solution of
414	dimethylformamide containing 36 mg of BTI per mL (Solution C).

415 **Procedure**—In a clean hydrolysis tube, transfer about 200 µg of the test 416 sample, and add 2 mL of *Solution A* or *Solution B* and 2 mL of *Solution C*. Seal 417 the hydrolysis tube in vacuum. Heat the sample at 60°C for 4 hours in the dark. 418 The sample is then dialyzed with water to remove the excess reagents. Extract the 419 dialyzed sample three times with equal volumes of n-butyl acetate, and then 420 lyophilize. The protein can then be acid hydrolyzed using previously described 421 procedures. The α,β -diaminopropionic and α,γ -diaminobutyric acid residues do 422 not typically resolve from the lysine residues upon ion-exchange chromatography 423 based on amino acid analysis. Therefore, when using ion-exchange as the mode 424 of amino acid separation, the asparagine and glutamine contents are the 425 quantitative difference in the aspartic acid and glutamic acid content assayed with 426 underivatized and BTI-derivatized acid hydrolysis. [NOTE—The threonine, 427 methionine, cysteine, tyrosine, and histidine assayed content can be altered by 428 BTI derivatization; a hydrolysis without BTI will have to be performed if the 429 analyst is interested in the composition of these other amino acid residues of the 430 protein/peptide.]

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Methodologies of Amino Acid Analysis General Principles

Many amino acid analysis techniques exist, and the choice of any one 434 435 technique often depends on the sensitivity required from the assay. In general, 436 about one-half of the amino acid analysis techniques employed rely on the 437 separation of the free amino acids by ion-exchange chromatography followed by 438 postcolumn derivatization (e.g., with ninhydrin or *o*-phthalaldehyde). Postcolumn 439 detection techniques can be used with samples that contain small amounts of 440 buffer components, such as salts and urea, and generally require between 5 and 10 441 ug of protein sample per analysis. The remaining amino acid techniques typically 442 involve precolumn derivatization of the free amino acids (e.g., phenyl 443 isothiocyanate; 6-amino-quinolyl-N-hydroxysuccinimidyl carbanate or o-444 phthalaldehyde; (dimethylamino) azobenzenesulfonyl chloride; 9-445 fluorenylmethylchloroformate; and, 7-fluoro-4-nitrobenzo-2-oxa-1,3-diazole) 446 followed by reversed-phase HPLC. Precolumn derivatization techniques are very

447 sensitive and usually require between 0.5 and 1.0 μ g of protein sample per 448 analysis but may be influenced by buffer salts in the samples. Precolumn 449 derivatization techniques may also result in multiple derivatives of a given amino 450 acid, which complicates the result interpretation. Postcolumn derivatization 451 techniques are generally influenced less by performance variation of the assay 452 than precolumn derivatization techniques. 453 The following *Methods* may be used for quantitative amino acid analysis. 454 Instruments and reagents for these procedures are available commercially. 455 Furthermore, many modifications of these methodologies exist with different 456 reagent preparations, reaction procedures, chromatographic systems, etc. Specific 457 parameters may vary according to the exact equipment and procedure used. 458 Many laboratories will utilize more than one amino acid analysis technique to 459 exploit the advantages offered by each. In each of these Methods, the analog 460 signal is visualized by means of a data acquisition system, and the peak areas are 461 integrated for quantification purposes. 462 463 METHOD 1 – Postcolumn Ninhydrin Detection General Principle 464 465 Ion-exchange chromatography with postcolumn ninhydrin detection is one 466 of the most common methods employed for quantitative amino acid analysis. As 467 a rule, a Li-based cation-exchange system is employed for the analysis of the 468 more complex physiological samples, and the faster Na-based cation-exchange 469 system is used for the more simplistic amino acid mixtures obtained with protein 470 hydrolysates (typically containing 17 amino acid components). Separation of the 471 amino acids on an ion-exchange column is accomplished through a combination 472 of changes in pH and cation strength. A temperature gradient is often employed 473 to enhance separation. 474 When the amino acid reacts with ninhydrin, the reactant has characteristic 475 purple or yellow color. Amino acids, except imino acid, give a purple color, and 476 show the maximum absorption at 570 nm. The imino acids such as proline give a 477 yellow color, and show the maximum absorption at 440 nm. The postcolumn 478 reaction between ninhydrin and amino acid eluted from column is monitored at

479 440 and 570 nm, and the chromatogram obtained is used for the determination of480 amino acid composition.

481 Detection limit is considered to be 10 pmol for most of the amino acid
482 derivatives, but 50 pmol for proline. Response linearity is obtained in the range
483 of 20 to 500 pmol with correlation coefficients exceeding 0.999. To obtain good
484 composition data, samples larger than 1 µg before hydrolysis are best suited for
485 this amino acid analysis of protein/peptide.

486

487 METHOD 2 – Postcolumn OPA Fluorometric Detection General Principle
 488

o-Phthalaldehyde (OPA) reacts with primary amines in the presence of
thiol compound, to form highly fluorescent isoindole products. This reaction is
utilized for the postcolumn derivatization in analysis of amino acids by ion-

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492 exchange chromatography. The rule of the separation is the same as *Method 1*.

493 Instruments and reagents for this form of amino acid analysis are available

494 commercially. Many modifications of this methodology exist.

Although OPA does not react with secondary amines (imino acids such as proline) to form fluorescent substances, the oxidation with sodium hypochlorite allows secondary amines to react with OPA. The procedure employs a strongly acidic cation-exchange column for separation of free amino acids followed by postcolumn oxidation with sodium hypochlorite and postcolumn derivatization using OPA and thiol compound such as *N*-acetyl-L-cysteine and 2-

mercaptoethanol. The derivatization of primary amino acids are not noticeablyaffected by the continuous supply of sodium hypochlorite.

503 Separation of the amino acids on an ion-exchange column is accomplished 504 through a combination of changes in pH and cation strength. After postcolumn 505 derivatization of eluted amino acids with OPA, the reactant passes through the 506 fluorometric detector. Fluorescence intensity of OPA-derivatized amino acids are 507 monitored with an excitation wavelength of 348 nm and an emission wavelength 508 of 450 nm.

509 Detection limit is considered to be a few tens of picomole level for most of 510 the amino acid derivatives. Response linearity is obtained in the range of a few

511	picomole level to a few tens of nanomole level. To obtain good compositional
512	data, the starting with greater than 500 ng of sample before hydrolysis is best
512	
	suited for the amino acid analysis of protein/peptide.
514 515	METHOD 3 – Precolumn PITC Derivatization General Principle
516 517	Phenylisothiocyanate (PITC) reacts with amino acids to form
518	phenylthiocarbamyl (PTC) derivatives which can be detected with high sensitivity
519	at 245 nm. Therefore, precolumn derivatization of amino acids with PITC
520	followed by a reversed-phase HPLC separation with UV detection is used to
521	analyze the amino acid composition.
522	After the reagent is removed under vacuum, the derivatized amino acids
523	can be stored dry and frozen for several weeks with no significant degradation. If
524	the solution for injection is kept cold, no noticeable loss in chromatographic
525	response occurs after three days.
526	Separation of the PTC-amino acids on a reversed-phase HPLC with ODS
527	column is accomplished through a combination of changes in concentrations of
528	acetonitrile and buffer ionic strength. PTC-amino acids eluted from column are
529	monitored at 254 nm.
530	Detection limit is considered to be 1 pmol for most of the amino acid
531	derivatives. Response linearity is obtained in the range of 20 to 500 pmol with
532	correlation coefficients exceeding 0.999. To obtain good compositional data,
533	samples larger than 500 ng of protein/peptide before hydrolysis is best suited for
534	this amino acid analysis of proteins/peptides.
535 536	METHOD 4 – Precolumn AQC Derivitization General Principle
537 538	Precolumn derivatization of amino acids with 6-aminoquinolyl-N-
539	hydroxysuccinimidyl carbamate (AQC) followed by reversed-phase HPLC
540	separation with fluorometric detection is used.
541	6-Aminoquinolyl-N-hydroxysuccinimidyl carbamate (AQC) reacts with
542	amino acids to form stable, fluorescent unsymmetric urea derivatives (AQC-
543	amino acids) which are readily amenable to analysis by reversed-phase HPLC.

544	Therefore, precolumn derivatization of amino acids with AQC followed by
545	reversed-phase HPLC separation is used to analyze the amino acid composition.
546	Separation of the AQC-amino acids on ODS column is accomplished
547	through a combination of changes in concentrations of acetonitrile and salt.
548	Selective fluorescence detection of the derivatives with excitation wavelength at
549	250 nm and emission wavelength at 395 nm allows for the direct injection of the
550	reaction mixture with no significant interference from the only major fluorescent
551	reagent by-product, 6-aminoquinoline. Excess reagent is rapidly hydrolyzed ($t_{1/2}$
552	<15 seconds) to yield 6-aminoquinoline, N-hydroxysuccinimide and carbon
553	dioxide, and after 1 minute no further derivatization can take place.
554	Peak areas for AQC-amino acids are essentially unchanged for at least 1
555	week at room temperature, and the derivatives have more than sufficient stability
556	to allow for overnight automated chromatographic analysis.
557	Detection limit is considered to be ranging from <i>ca</i> . 40 to 320 fmol for
558	each amino acid, except for Cys. Detection limit for Cys is approximately 800
559	fmol. Response linearity is obtained in the range of 2.5 to 200 μ M with
560	correlation coefficients exceeding 0.999. Good compositional data could be
561	obtained from the analysis of derivatized protein hydrolysates containing as little
562	as 30 ng of protein/peptide.
563	
564	METHOD 5 – Precolumn OPA Derivatization General Principle
565 566	Precolumn derivatization of amino acids with <i>o</i> -phthalaldehyde (OPA)
567	followed by reversed-phase HPLC separation with fluorometric detection is used.
568	This technique does not detect amino acids that exist as secondary amines (e.g.,
569	proline).
570	<i>o</i> -Phthalaldehyde (OPA) in conjunction with a thiol reagent reacts with
571	primary amine groups to form highly fluorescent isoindole products. 2-
572	Mercaptoethanol of 3-mercaptopropionic acid can be used as the thiol. OPA itself
573	does not fluoresce and consequently produces no interfering peaks. In addition,
574	its solubility and stability in aqueous solution, along with the rapid kinetics for the
575	reaction, make it amenable to automated derivatization and analysis using an

576	autosampler to mix the sample with the reagent. However, lack of reactivity with
577	secondary amino acids has been predominant drawback. This method does not
578	detect amino acids that exist as secondary amines (e.g., proline). To compensate
579	for this drawback, this technique may be combined with another technique
580	described in Method 7 or Method 8.
581	Precolumn derivatization of amino acids with OPA is followed by a
582	reversed-phase HPLC separation. Because of the instability of the OPA-amino
583	acid derivative, HPLC separation and analysis are performed immediately
584	following derivatization. The liquid chromatograph is equipped with a
585	fluorometric detector for the detection of derivatized amino acids. Fluorescence
586	intensity of OPA-derivatized amino acids is monitored with an excitation
587	wavelength of 348 nm and an emission wavelength of 450 nm.
588	Detection limits as low as 50 fmol via fluorescence have been reported,
589	although the practical limit of analysis remains at 1 pmol.
590	
591	METHOD 6 – Precolumn DABS-Cl Derivatization General Principle
592 593	Precolumn derivatization of amino acids with
594	(dimethylamino)azobenzenesulfonyl chloride (DABS-Cl) followed by reversed-
595	phase HPLC separation with visible light detection is used.
595 596	(Dimethylamino)azobenzenesulfonyl chloride (DABS-Cl) is a
597	chromophoric reagent employed for the labeling of amino acids. Amino acids
598	labeled with DABS-Cl (DABS-amino acids) are highly stable and show the
599	maximum absorption at 436 nm.
600	DABS-amino acids, all 19 naturally occurring amino acids derivatives,
601	can be separated on an ODS column of a reversed-phase HPLC by employing
602	gradient systems consisting of acetonitrile and aqueous buffer mixture. Separated
603	DABS-amino acids eluted from column are detected at 436 nm in the visible
604	region.
605	This Method can analyze the imino acids such as proline together with the
606	amino acids at the same degree of sensitivity, DABS-Cl derivatization method
607	permits the simultaneous quantification of tryptophan residues by previous

608	hydrolysis of the protein/peptide with sulfonic acids such as
609	mercaptoethanesulfonic acid, p-toluenesulfonic acid or methanesulfonic acid
610	described under Method 2 in "Protein Hydrolysis". The other acid-labile residues,
611	asparagine and glutamine, can also be analysed by previous conversion into
612	diaminopropionic acid and diaminobutyric acid, respectively, by treatment of
613	protein/peptide with BTI described under Method 11 in "Protein Hydrolysis".
614	The non-proteinogenic amino acid, norleucine cannot be used as internal
615	standard in this method, as this compound is eluted in a chromatographic region
616	crowded with peaks of primary amino acids. Nitrotyrosine can be used as an
617	internal standard, because it is eluted in a clean region.
618	Detection limit of DABS-amino acid is about 1 pmol. As little as 2 to 5
619	pmol of an individual DABS-amino acid can be quantitatively analysed with
620	reliability, and only 10 to 30 ng of the dabsylated protein hydrolysate is required
621	for each analysis.
622 623	METHOD 7 – Precolumn FMOC-Cl Derivatization General Principle
624	•
625	Precolumn derivatization of amino acids with 9-fluorenylmethyl
626	chloroformate (FMOC-Cl) followed by reversed-phase HPLC separation with
627	fluorometric detection is used.
628	9-Fluorenylmethyl chloroformate (FMOC-Cl) reacts with both primary
629	and secondary amino acids to form highly fluorescent products. The reaction of
630	FMOC-Cl with amino acid proceeds under mild conditions in aqueous solution
631	and is completed in 30 seconds. The derivatives are stable, only the histidine
632	derivative showing any breakdown. Although FMOC-Cl is fluorescent itself, the
633	reagent excess and fluorescent side-products can be eliminated without loss of
634	FMOC-amino acids.
635	FMOC-amino acids are separated by a reversed-phase HPLC using ODS
636	column. The separation is carried out by gradient elution varied linearly from a
637	mixture of acetonitrile methanol and acetic acid buffer (10:40:50) to a mixture of
638	acetonitrile and acetic acid buffer (50:50), and 20 amino acid derivatives are
639	separated in 20 minutes. Each derivative eluted from column is monitored by a

640	fluorometric detector set at an excitation wavelength of 260 nm and an emission
641	wavelength of 313 nm.
642	The detection limit is in the low fmol range. A linearity range of 0.1 to 50
643	μ M is obtained for most of the amino acids.
644 645 646	METHOD 8 – Precolumn NBD-F Derivatization General Principle
647	Precolumn derivatization of amino acids with 7-fluoro-4-nitrobenzo-2-
648	oxa-1.3-diazole (NBD-F) followed by reversed-phase HPLC separation with
649	fluorometric detection is used.
650	7-fluoro-4-nitrobenzo-2-oxa-1.3-diazole (NBD-F) reacts with both
651	primary and secondary amino acids to form highly fluorescent products. Amino
652	acids are derivatized with NBD-F by heating to 60°C for 5 minutes.
653	NBD-amino acid derivatives are separated on an ODS column of a
654	reversed-phase HPLC by employing gradient elution system consisting of
655	acetonitrile and aqueous buffer mixture, and 17 amino acid derivatives are
656	separated in 35 minutes. ε-Aminocaproic acid can be used as an internal standard,
657	because it is eluted in a clean chromatographic region. Each derivative eluted
658	from column is monitored by a fluorometric detector set at an excitation
659	wavelength of 480 nm and an emission wavelength of 530 nm.
660	The sensitivity of this method is almost the same as for precolumn OPA
661	derivatization method (Method 5), excluding proline to which OPA is not
662	reactive, and might be advantageous for NBD-F against OPA. The detection limit
663	for each amino acid is about 10 fmol. Profile analysis can be achieved for about
664	1.5 μ g of protein hydrolysates in the final precolumn labeling reaction mixture for
665	HPLC.
666	
667	Data Calculation and Analysis
668 669	When determining the amino acid content of a protein/peptide
670	hydrolysate, it should be noted that the acid hydrolysis step destroys tryptophan
671	and cysteine. Serine and threonine are partially destroyed by acid hydrolysis,
672	while isoleucine and valine residues may be only partially cleaved. Methionine

673 can undergo oxidation during acid hydrolysis, and some amino acids (e.g., glycine 674 and serine) are common contaminants. Application of adequate vacuum (\leq not 675 less than 200 µm of mercury or 26.7 Pa) or introduction of inert gas (argon) in the 676 headspace of the reaction vessel during vapor phase hydrolysis can reduce the 677 level of oxidative destruction. Therefore, the quantitative results obtained for cysteine, tryptophan, threonine, isoleucine, valine, methionine, glycine, and serine 678 679 from a protein/peptide hydrolysate may be variable and may warrant further 680 investigation and consideration. 681 682 **CALCULATIONS** 683 684 Amino Acid Mole Percent—This is the number of specific amino acid 685 residues per 100 residues in a protein. This result may be useful for evaluating amino acid analysis data when the molecular weight of the protein under 686 investigation is unknown. This information can be used to corroborate the 687 688 identity of a protein/peptide and has other applications. Carefully identify and 689 integrate the peaks obtained as directed for each *Procedure*. Calculate the mole 690 percent for each amino acid present in the test sample by the formula: 691 $100r_{U}/r$, 692 in which r_U is the peak response, in nmol, of the amino acid under test; and r is 693 the sum of peak responses, in nmol, for all amino acids present in the test sample. 694 Comparison of the mole percent of the amino acids under test to data from known 695 proteins can help establish or corroborate the identity of the sample protein. 696 Unknown Protein Samples—This data analysis technique can be used to 697 estimate the protein concentration of an unknown protein sample using the amino 698 acid analysis data. Calculate the mass, in µg, of each recovered amino acid by the 699 formula: 700 $mM_W/1000$, 701 in which *m* is the recovered quantity, in nmol, of the amino acid under test; and 702 M_W is the average molecular weight for that amino acid, corrected for the weight 703 of the water molecule that was eliminated during peptide bond formation. The

sum of the masses of the recovered amino acids will give an estimate of the total

mass of the protein analyzed after appropriate correction for partially and

706 completely destroyed amino acids. If the molecular weight of the unknown

707 protein is available (i.e., by SDS-PAGE analysis or mass spectroscopy), the

amino acid composition of the unknown protein can be predicted. Calculate the

number of residues of each amino acid by the formula:

710

$m/(1000M/M_{WT}),$

711 in which *m* is the recovered quantity, in nmol, of the amino acid under test; *M* is

the total mass, in μ g, of the protein; and M_{WT} is the molecular weight of the unknown protein.

714 Known Protein Samples—This data analysis technique can be used to 715 investigate the amino acid composition and protein concentration of a protein 716 sample of known molecular weight and amino acid composition using the amino 717 acid analysis data. When the composition of the protein being analyzed is known, 718 one can exploit the fact that some amino acids are recovered well, while other 719 amino acid recoveries may be compromised because of complete or partial 720 destruction (e.g., tryptophan, cysteine, threonine, serine, methionine), incomplete 721 bond cleavage (i.e., for isoleucine and valine) and free amino acid contamination 722 (i.e., by glycine and serine).

723 Because those amino acids that are recovered best represent the protein, 724 these amino acids are chosen to quantify the amount of protein. Well-recovered 725 amino acids are, typically, aspartate-asparagine, glutamate-glutamine, alanine, 726 leucine, phenylalanine, lysine, and arginine. This list can be modified based on experience with one's own analysis system. Divide the quantity, in nmol, of each 727 728 of the well-recovered amino acids by the expected number of residues for that 729 amino acid to obtain the protein content based on each well-recovered amino acid. 730 Average the protein content results calculated. The protein content determined 731 for each of the well-recovered amino acids should be evenly distributed about the 732 mean. Discard protein content values for those amino acids that have an 733 unacceptable deviation from the mean. Typically \geq greater than 5% variation 734 from the mean is considered unacceptable. Recalculate the mean protein content 735 from the remaining values to obtain the protein content of the sample. Divide the

736	content of each amino acid by the calculated mean protein content to determine
737	the amino acid composition of the sample by analysis.
738	Calculate the relative compositional error, in percentage, by the formula:
739	$100m/m_{s}$,
740	in which m is the experimentally determined quantity, in nmol per amino acid
741	residue, of the amino acid under test; and m_S is the known residue value for that
742	amino acid. The average relative compositional error is the average of the
743	absolute values of the relative compositional errors of the individual amino acids,
744	typically excluding tryptophan and cysteine from this calculation. The average
745	relative compositional error can provide important information on the stability of
746	analysis run over time. The agreement in the amino acid composition between the
747	protein sample and the known composition can be used to corroborate the identity
748	and purity of the protein in the sample.
749	