Amino Acids Specifications / Monographs		
L-Valine	page	1/2
Issued Date: Dec, 4, 2015		

L-Valine¹

C₅H₁₁NO₂: 117.15

L-Valine, when dried, contains not less than 98.5 percent and not more than 101.0 percent of L-Valine ($C_5H_{11}NO_2$).

Description

White crystals or crystalline powder; tastes at first slightly sweet and then bitter.

Freely soluble in formic acid, soluble in water, practically insoluble in ethanol (95).

Dissolves in dilute hydrochloric acid.

Identification

Compare the infrared absorption spectrum of the sample with that of the standard by potassium bromide disc method.

Specifications

Item	Limit	Test
Specific rotation $[\alpha]_D^{20}$	+27.6 to +28.7°	AJI TEST 1 [Dried sample, C=8, 6mol/L HCI] ²
State of solution	Clear and colorless	AJI TEST 2
(Transmittance)	Not less than 98.0%	[0.5g in 20mL of H ₂ O, spectrophotometer, 430nm, 10mm cell thickness]
Chloride (Cl)	Not more than	AJI TEST 3
	0.020%	[0.5g, A-1, ref: 0.28mL of 0.01mol/L HCl]
Ammonium (NH ₄)	Not more than 0.02%	AJI TEST 4
		[D-1]
Sulfate (SO ₄)	Not more than 0.020%	AJITEST 5
		[0.85g, (1), ref: 0.35mL of 0.005mol/L H ₂ SO ₄]
Iron (Fe)	Not more than 10ppm	AJI TEST 6
		[0.75g, B-1, ref:0.75mL of Iron Std. (0.01mg/mL)]
Heavy metals (Pb)	Not more than 10ppm	AJI TEST 7
		[1.0g, (1), ref:1.0mL of Pb Std. (0.01mg/mL)]
Arsenic (As ₂ O ₃)	Not more than 1ppm	AJI TEST 8
		[2.0g, (1), ref: 2.0mL of As ₂ O ₃ Std.]
Related substances	1)Conforms ³	AJITEST 9
		[Test sample: 50µg, B-6-a, control; L-Val 0.25µg]
	2)Ile	AJI TEST 26 ⁴
	Not more than 0.40%	
	Any unspecified impurity	
	Not more than 0.20%	
	Total impurities	
	Not more than 1.00%	
Loss on drying	Not more than 0.20%	AJI TEST 11
		[1g, at 105°C for 3 hours]

AJINOMOTO CO,. INC.

Amino Acids Specifications / Monographs		
L-Valine	page	2/2
Issued Date: Dec, 4, 2015		

L-Valine

Specifications (cont'd)

Item	Limit	Test
Residue on ignition	Not more than 0.10%	AJI TEST 13
(Sulfated)		[1g, at 550°C to 650°C for 3 hours]
Assay	98.5 to 101.0%	AJI TEST 14
		[Dried sample, 120mg, (1), 3mL of formic acid, 50mL of acetic
		acid (100), 0.1mol/L HClO ₄ 1mL=11.72mg C ₅ H ₁₁ NO ₂]
pН	5.5 to 6.5	AJI TEST 33
		[0.5g in 20mL of H ₂ O]

The test for Endotoxin when the material will be used for manufacturing parenteral products is as follows:

Item	Limit	Test
Endotoxin	Less than 6.0EU/g	AJI TEST 34
		[C=1, kinetic-turbidimetric technique]

 $^{^{1}\,\,}$ This product, in terms of actual quality, conforms to USP, EP, and JP.

End of document

² Temperature coefficient of $[\alpha]_D^t$: -0.03°

Any secondary spot in the chromatogram obtained from the Test Solution is less intense than the principal spot in the chromatogram obtained from the Standard Solution: the number of those spots is not more than four and not more than 2.0% of the total impurities found.

⁴ Disregard limit: 0.05%