



DN LABORATORY

A Govt. Approves Testing Laboratory Under "the Drugs & Cosmetics Act 1940 And Rules " There Under
Approved By: FDA Haryana Ayush Haryana: An ISO 9001 : 2015 & GLP Certified Lab.
 Address: Ind. Area, Phase-1, Phanchkula, Haryana.
 Mob: +91 9538239428, email id: dnlabpk@gmail.com, manisha256@gmail.com

Certificate of Analysis

Form-47, 160 (A) Report of test or Analysis by Approved Institution

Sample Name	Resveratrol Fisetin Capsules			Received Date	02-01-2026
Customer Information	Reverse Clinics by Threads Physio Orthosport LLP, F-10/4, DLF Phase 1, Sector-27, Gurugram, Haryana 122001			Ref. No.	NIL
Supplied By	NS			Report No.	DNL/01/26-27/15B/C
Batch Size	NS	LOT No.	CW-090	Sample Qty.	60 Capsules
Mfg. Date	01-2026	Exp. Date	02-01-2028	Fssai. Lic. No.	20825005004121
Date of Analysis	03-01-2026	Date of Completion	09-01-2026	Protocol ID	IHS

RESULT OF ANALYSIS

Description	Brown Powder filled in transparent hard gelatin capsule		
Average Fill weight	: 550 mg		
Disintegration Time	: 10-11 min		(NMT – 30 min)
Assay	: Each Serving Capsule (on an average fill) contains: -		
Composition	Claim	Observed	Method
Resveratrol	240 mg	240.10 mg	HPLC
Fisetin	100 mg	99.95 mg	HPLC
Coenzyme Q10	100 mg	100.15 mg	HPLC
NMN	60 mg	59.85 mcg	HPLC
Quercetin	30 mg	30.20 mg	HPLC
Ashwagandha	15 mg	15.10 mg	HPLC
Black Pepper	5 mg	4.95 mg	HPLC
Identification	Conform	Complies	IR Spectrum
Characterstics	Brown Powder	Complies	Organoleptic
Odor & Taste	Characterstics	Conform	Organoleptic
Solubility	Soluble in Water	Complies	Turbidity Meter
Assay	NLT 98%	99.35%	HPLC
Refractive Index @ 25 C	1.05-1.25 at 20°C	1.14	Refractometer
Specific Gravity	1.670-1.695 at 20°C	1.688	Densimeter
Particle Size	95% pass 80 mesh	Conforms	CP 2015
Loss on drying	NMT 1%	0.45%	USP<731>
Residue on Ignition	NMT 0.5%	0.21%	USP<281>
Residue on Solvents	NMT 100 ppm	Conforms	NLS-QCS-1007
Residual Pesticides	NMT 10 ppm	3 ppm	Eu. Ph.<2.8.13>
pH 5 % in water	3-6 at 25°C	4.2	USP<791>
Ash Content	NMT 1%	0.29 %	USP<561>
Moisture	NMT 5 %	2.35 %	AOAC
Acid Insoluble Ash	NMT 0.1%	0.03%	Muffle Furnace
Bulk Density	45-60g/100mL	53g/100mL	Bulk Density Apparatus
Tapped Density	70-90 g/100mL	83g/100mL	Tapped Density Apparatus
Heavy Metals:			
Total Heavy Metals	10 ppm Max	Conforms	USP<231> ICP-MS
Lead	NMT - 2 ppm	0.53 ppm	USP <231> ICP-MS
Arsenic	NMT - 1 ppm	0.39 ppm	USP <231> ICP-MS
Cadmium	NMT - 1 ppm	0.23 ppm	USP <231> ICP-MS
Mercury	NMT - 0.1 ppm	0.02 ppm	USP <231> ICP-MS
Microbial Examination:			
Total Plate Count	NMT 1000 cfu/g	100 cfu/g	USP<61>
Yeast & Mould	NMT 100 cfu/g	15 cfu/g	USP<61>
E. Coli	Absent/g	Absent	USP<61>
Salmonella	Absent/g	Absent	USP<61>
Staphylococcus Aureus	Absent/g	Absent	USP<61>
Pseudomonas Aeruginosa	Absent/g	Absent	USP<61>

09-01-2026

Date of Completion


 Signature of Person-in-charge of testing



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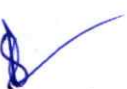
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OPINION : In the opinion of the undersigned the above sample is of standard quality as defined in the Act and Rules made there under for the reasons given below

The sample confirms to IP BP USP BIS ISO AYUR test specification with respect to above test only.

09-01-2026
Date of Completion


Signature of Person-in-charge of testing

Note :

1. This report is not to be reproduced wholly or in part and cannot be used as evidence in the court of law and should not be used in any advertising media without our special permission in writing.
2. Samples (s) not drawn by us, unless otherwise stated.
3. Total liability of our analytical division is limited to the invoiced amount.
4. Sample will be destroyed after one month from date of issue of test certificate unless otherwise specified.
5. Results given in reports are released to sample tested.