



Sample ID SD260512-044 (138897)	Matrix Edible
Tested for blend.	
Received May 12, 2026	Reported May 29, 2026
Analyses executed KTM	

Laboratory note: COA Update: Photo updated per client request.

KTM - Kratom

Analyzed May 12, 2026 | Instrument HPLC VWD | Method SOP-KTM
The expanded Uncertainty of the Kratom analysis is approximately ±7.81% at the 95% Confidence Level

Analyte	LOD ppm	LOQ ppm	Result %	Result mg/g
7-hydroxy Mitragynine (7HMG)	0.008	0.025	ND	ND
MGM-15 (MGM)	0.186	0.562	ND	ND
Mitragynine (MITG)	0.018	0.054	ND	ND
Speciogynine (SPEG)	0.007	0.02	ND	ND
Speciociliatine (SPCL)	0.004	0.011	ND	ND



UI Unidentified
ND Not Detected
N/A Not Applicable
NT Not Reported
LOD Limit of Detection
LOQ Limit of Quantification
<LOQ Detected
>ULOL Above upper limit of linearity
CFU/g Colony Forming Units per 1 gram
TNTC Too Numerous to Count



DEA license: RP0611043
ISO/IEC 17025:2017 Acc. 85368



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Authorized Signature

Brandon Starr

Brandon Starr, Quality Assurance Manager
Fri, 29 May 2026 17:18:48 -0700

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Sample Indoblend 20ct Gravity - Watermelon



Sample ID SD260512-046 (138899)	Matrix Edible
Tested for blend.	
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KTM - Kratom

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MGM-15 (MGM)	0.186	0.562	ND	ND
Mitragynine (MITG)	0.018	0.054	ND	ND
Speciogynine (SPEG)	0.007	0.02	ND	ND
Speciocilatin (SPCL)	0.004	0.011	ND	ND



UI Unidentified
ND Not Detected
N/A Not Applicable
NT Not Reported
LOD Limit of Detection
LOQ Limit of Quantification
<LOQ Detected
>ULOL Above upper limit of linearity
CFU/g Colony Forming Units per 1 gram
TNTC Too Numerous to Count



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Sample Indoblend 20ct Gravity - Blue Razz



Sample ID SD260512-043 (138896)	Matrix Edible
Tested for blend.	
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KTM - Kratom

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MGM-15 (MGM)	0.186	0.562	ND	ND
Mitragynine (MITG)	0.018	0.054	ND	ND
Speciogynine (SPEG)	0.007	0.02	ND	ND
Speciocillatine (SPCL)	0.004	0.011	ND	ND



UI Unidentified
ND Not Detected
N/A Not Applicable
NT Not Reported
LOD Limit of Detection
LOQ Limit of Quantification
<LOQ Detected
>ULOL Above upper limit of linearity
CFU/g Colony Forming Units per 1 gram
TNTC Too Numerous to Count



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Sample Indoblend 20ct Gravity - Strawberry



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Tested for blend.	
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KTM - Kratom

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Speciociliatine (SPCL)	0.004	0.011	ND	ND



UI Unidentified
ND Not Detected
N/A Not Applicable
NT Not Reported
LOD Limit of Detection
LOQ Limit of Quantification
<LOQ Detected
>ULOL Above upper limit of linearity
CFU/g Colony Forming Units per 1 gram
TNTC Too Numerous to Count



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