

**Customer Information**

Rocktomic Labs LLC
Private Label Supplements & Dropshipping
1615 Lakes Pkwy, Suite C, Lawrenceville,
GA 30043
www.rocktomic.com

CERTIFICATE OF ANALYSIS

Product	Xenoestrogens Support	Lot Number	250103-RT2551
Item Number	RT2551	Country of Origin	USA

TEST	SPECIFICATION	RESULTS	METHOD
Identity			
Capsule Appearance	Clear hard-shell capsule, size 0, filled with powder	Pass	DOC-1779 (WI-QC-017)
Powder Color	Greenish, To Match Standard	Pass	DOC-1779 (WI-QC-017)
Powder Odor	Characteristic, To Match Standard	Pass	DOC-1779 (WI-QC-017)
Fingerprint Identity	Must Conform to Standard	Pass	UHPLC-DAD

Purity			
Total Aerobic Microbial Count	$\leq 1 \times 10^7$ cfu/g	Pass	USP <2021> or equivalent
Total Combined Yeast and Mold Count	$\leq 1 \times 10^5$ cfu/g	Pass	USP <2021> or equivalent
Enterobacterial Count (Bile-Tolerant Gram Neg. Bacteria)	$\leq 1 \times 10^4$ cfu/g	Pass	02-209-01
<i>Staphylococcus aureus</i>	None Detected in 10 g	Pass	USP <2022> or equivalent
<i>Escherichia coli</i>	None Detected in 10 g	Pass	USP <2022> or equivalent
<i>Salmonella spp.</i>	None Detected in 10 g	Pass	USP <2022> or equivalent
Lead (Pb)	Conforms to USP <232> Elemental Impurities Limits (Heavy Metals) $\leq 5 \mu\text{g}$ per 2 capsules	Pass	ICP-MS
Arsenic (As)	Conforms to USP <232> Elemental Impurities Limits (Heavy Metals) $\leq 15 \mu\text{g}$ per 2 capsules	Pass	ICP-MS
Cadmium (Cd)	Conforms to USP <232> Elemental Impurities Limits (Heavy Metals) $\leq 5 \mu\text{g}$ per 2 capsules	Pass	ICP-MS
Mercury (Hg)	Conforms to USP <232> Elemental Impurities Limits (Heavy Metals) $\leq 2 \mu\text{g}$ per 2 capsules	Pass	ICP-MS



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Strength			
Iodine	240-683 µg / 2 capsules	461 µg / 2 capsules	ICP-MS

Customer Approval	
Signature/Date	Final Disposition
	PASS

This product has been reviewed and approved by the Rocktomic Labs. The results reported in this certificate of analysis (COA) are truthful and representative of the lot. The finished product and all raw materials used to manufacture the batch have been found to meet required specification limits in accordance with 21 CFR 111 (cGMP) regulations and the product is released.