REPORT NUMBER







REPORT OF ANALYSIS For: (35416) OPEN FARM INC OF-Turkey Bone Broth OF440092122- Turkey

	Level Found		Reporting			Analyst-	Verified-
Analysis	As	Received	Units	Limit	Method	Date	Date
Sample ID: OF440092122	Lab Number: 13907510	Date Sample	d: 2022-0 9	9-23			
Listeria	I	negative	org/125g	1	RapidChek/AOAC RI 020401	kkb0-2022/10/02	jzh4-2022/10/02
E. coli (generic)		n.d.	cfu/g	10	AOAC 2018.13	Kat6-2022/10/01	snl7-2022/10/01
Salmonella	I	negative	org/375g	1	RapidChek/AOAC RI 030301; AFNOR SDI 34/01-04/10	kkb0-2022/10/02	jzh4-2022/10/02

All results are reported on an AS RECEIVED basis., n.d. = not detected, cfu = colony forming unit

For questions please contact:

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The result(s) issued on this report only reflect the analysis of the sample(s) submitted.

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Detailed Method Description(s)

E. coli and Total Coliform using 3M Petrifilm

Sample analysis follows MWL MI 292 which is based on AOAC 2018.13. A representative sample is obtained and added to phosphate buffer. Aliquots of the sample are withdrawn and placed on Petrifilm plates. The plates are incubated for 18 to 24 hours. After incubation, the plates are counted to determine the number of generic E. coli and total coliforms present. The color of the colony and the presence of gas differentiate a generic coliform from E. coli. The levels are reported as colony forming units (cfu).

Salmonella - Lateral Flow

Samples are analyzed following MWL MI 195 which is based on the RapidChek Select Salmonella User Guide. A representative sample is obtained and combined with a selective media and allowed to incubate. After incubation, a test strip is used for Salmonella determination. Results are reported as negative or presumptive positive.

Listeria Lateral Flow

Samples are analyzed following MWL MI 194 which is based on the RapidChek Listeria User Guide. A representative sample is obtained and combined with a selective growth media. It is incubated for 40-48 hours. After incubation, an aliquot is heated for 10 minutes, and a test strip for Listeria detection is used. Results are reported as negative or presumptive positive. This procedure does not speciate Listeria.

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