

CLIENT / BRAND Reserve Research	BATCH ID RR-20260817-HDF8	CLIENT BATCH / LOT BAC003-001	COMPOUND Bacteriostatic Water (benzyl alcohol 0.9% w/v)	SAMPLE (AS RUN) RR-20260817-HDF8-01	DATE RECEIVED 17 AUG 2026
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DATE ANALYZED
02 SEP 2026

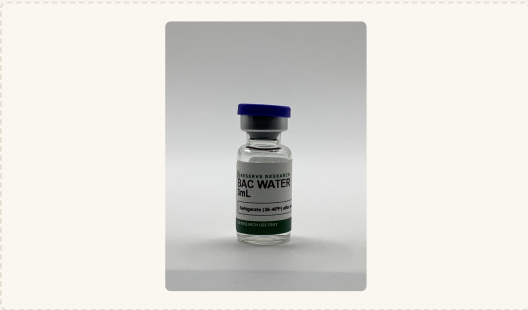
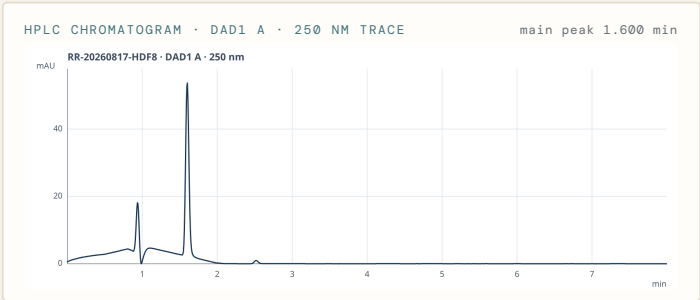
CHROMATOGRAPHIC PURITY 99.99% OF TOTAL PEAK AREA	BENZYL ALCOHOL CONTENT – MG/ML 9.13 LABEL CLAIM: 9 MG/ML (0.9% W/V) – 101% OF CLAIM ≈ 27.4 MG PER VIAL (CALCULATED FROM LABELED FILL VOLUME OF 3 ML – FILL VOLUME NOT INDEPENDENTLY VERIFIED)	RETENTION TIME 1.600 MINUTES · MAIN PEAK
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Overall: **Conforms** to specification — **Identity Conforms** (RT + UV vs. reference); sample meets the acceptance criteria below. SPEC SET IS CLIENT / METHOD-DEFINED

SUMMARY OF RESULTS

TEST	SPECIFICATION	RESULT	STATUS
Appearance	Clear, colorless solution	Clear, colorless	Conforms
Purity (HPLC, area %)	≥ 95.0%	99.99%	Conforms
Benzyl alcohol content	0.9% w/v (9 mg/mL)	9.13 mg/mL (101% of claim)	Conforms

CHROMATOGRAM & SAMPLE REFERENCE



How to read this. Bacteriostatic water is sterile water preserved with 0.9% benzyl alcohol, which allows a vial to be entered multiple times. This certificate verifies by HPLC that benzyl alcohol is present at its labeled concentration and that no unexpected UV-active compounds are in the vial. Purity is the percentage of detected material that is benzyl alcohol. This certificate does not assess sterility, endotoxin, or microbial content.

AUTHORIZED SIGNATORY · KMD ANALYTICAL
Kevin Doud Kevin Doud
Laboratory Director

DATE OF ISSUE
02 SEP 2026

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