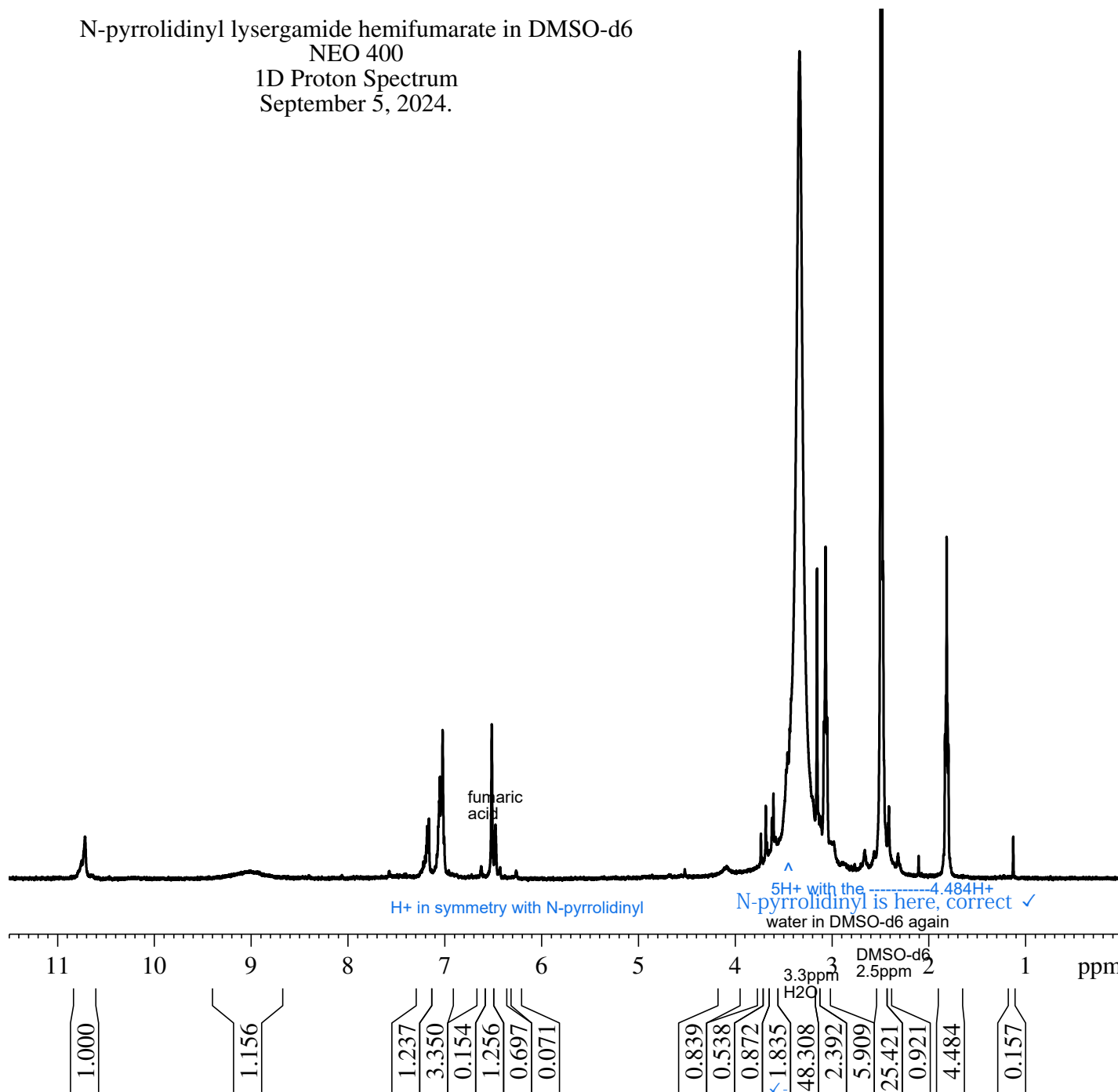


N-pyrrolidinyl lysergamide hemifumarate in DMSO-d6
 NEO 400
 1D Proton Spectrum
 September 5, 2024.

Current Data Parameters
 NAME Syntharise090524
 EXPNO 410
 PROCNO 1

F2 - Acquisition Parameters
 Date_ 20240905
 Time 11.57 h
 INSTRUM Avance Neo 400
 PROBHD Z163739_0039 (
 PULPROG zg30
 TD 65536
 SOLVENT DMSO
 NS 16
 DS 0
 SWH 8196.722 Hz
 FIDRES 0.250144 Hz
 AQ 3.9976959 sec
 RG 101
 DW 61.000 usec
 DE 13.54 usec
 TE 296.6 K
 D1 0.10000000 sec
 TD0 1
 SFO1 400.3424721 MHz
 NUC1 1H
 P0 3.33 usec
 P1 10.00 usec
 PLW1 14.20300007 W

F2 - Processing parameters
 SI 65536
 SF 400.3400073 MHz
 WDW EM
 SSB 0
 LB 0.30 Hz
 GB 0
 PC 1.00



H+ in symmetry with N-pyrrolidinyl

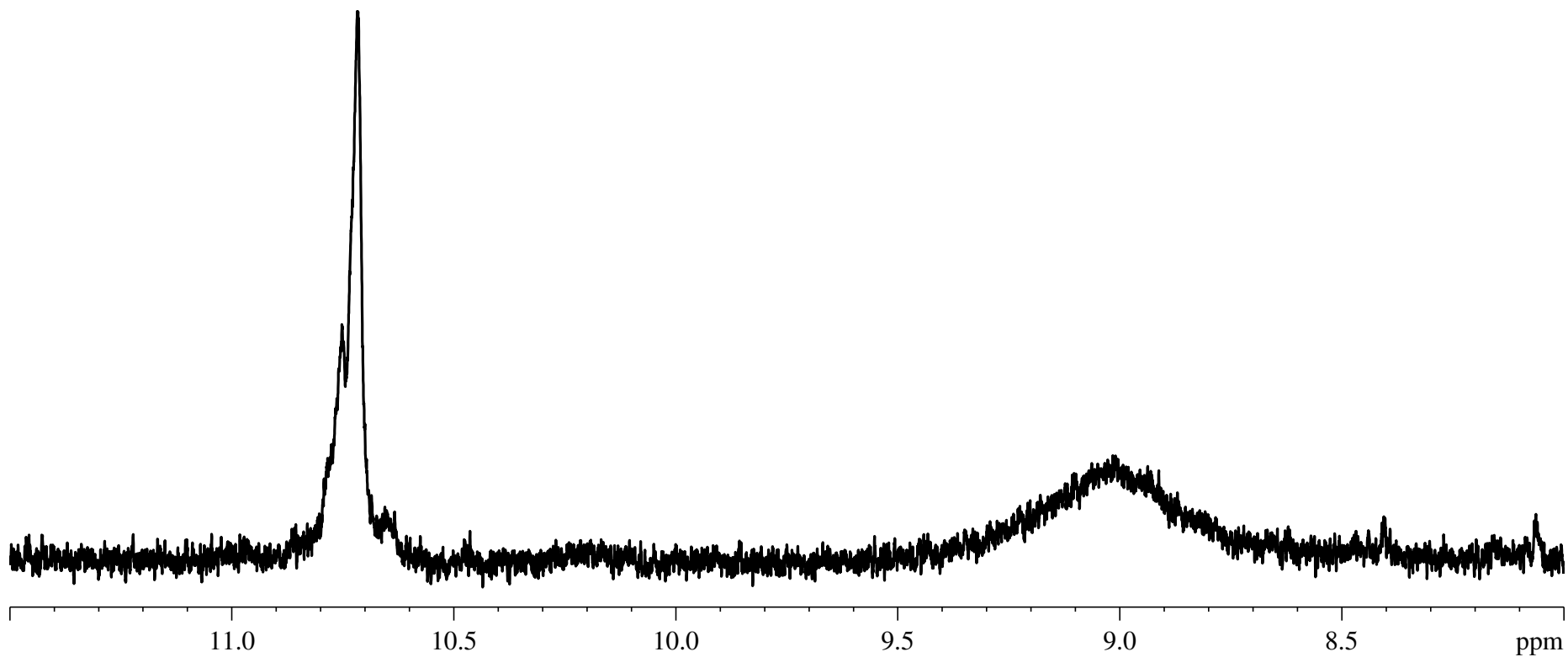
5H+ with the N-pyrrolidinyl is here, correct ✓
 water in DMSO-d6 again

symmetry DMSO-d6 mixes with the 3H+ too for the N-methyl on structure

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— 10.717

— 9.023

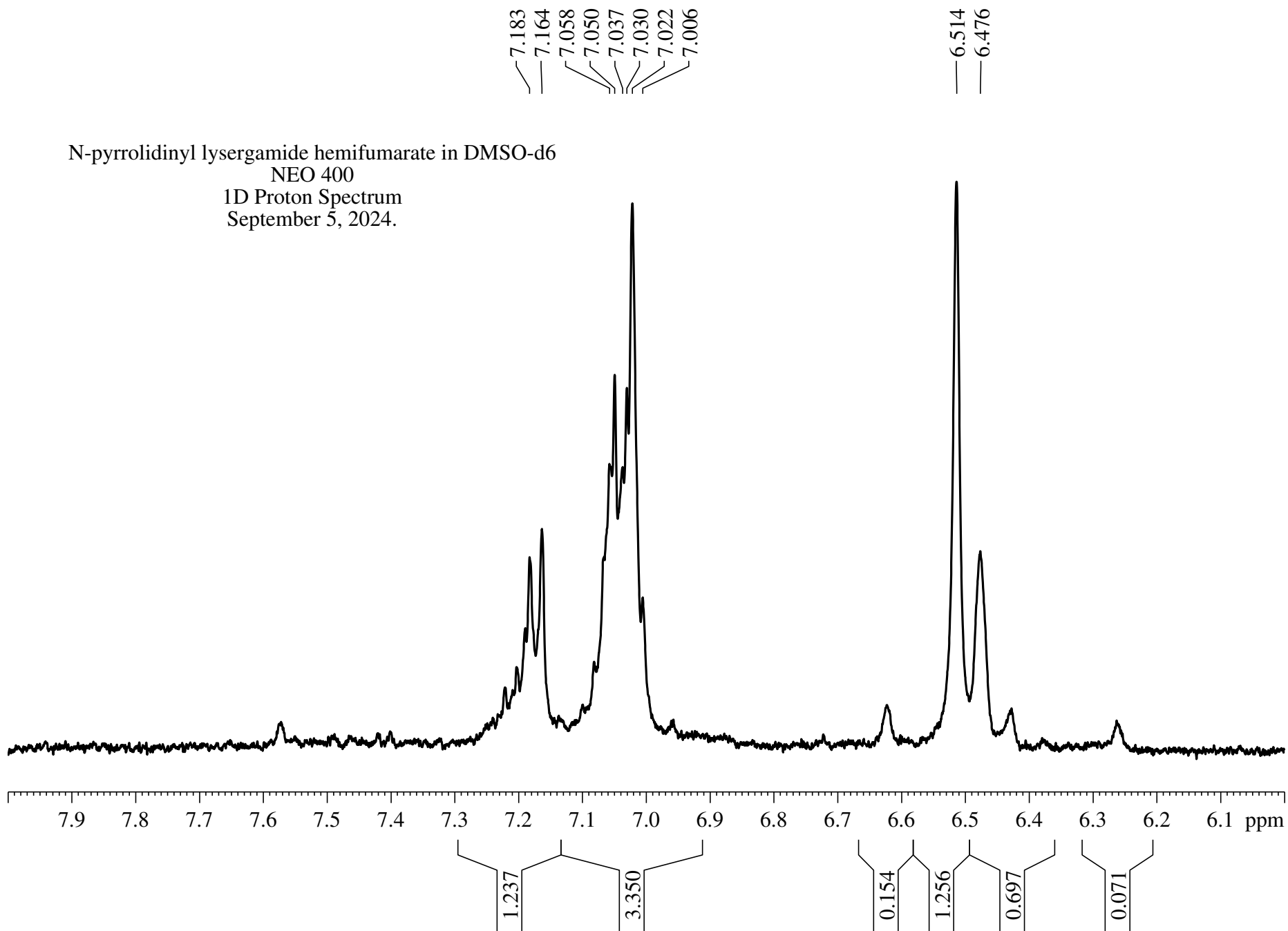


1.000

1.156

ppm

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