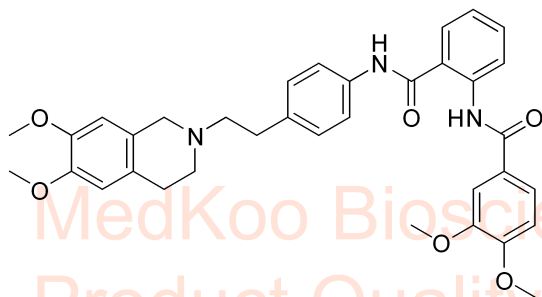


# WIKI4 HNMR Analysis, solvent: DMSO-d6



8.574  
8.570  
8.563  
8.559  
8.513  
8.511  
8.495  
8.492  
8.486  
8.484  
8.465  
8.463  
7.903  
7.884  
7.882  
7.864  
7.368  
7.362  
7.351  
7.346  
7.310  
7.306  
7.298  
7.294  
7.065  
7.059  
7.048  
7.042  
4.194  
4.177  
4.160  
3.825  
3.308  
3.280  
2.516  
2.511  
2.507  
2.502  
2.498  
2.141  
2.123  
2.106  
1.240



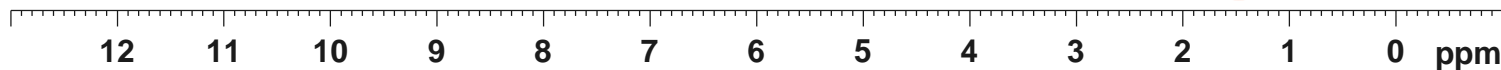
MedKoo Cat#: 406349  
Name: WIKI4  
CAS#: 838818-26-1  
Lot#: B7B09C21  
Chemical Formula: C<sub>29</sub>H<sub>23</sub>N<sub>5</sub>O<sub>3</sub>S  
Exact Mass: 521.15216  
Molecular Weight: 521.59

Current Data Parameters  
NAME WIKI4  
EXPNO 10  
PROCNO 1

F2 - Acquisition Parameters  
INSTRUM spect  
PROBHD 5 mm PABBO BB-  
PULPROG zg30  
TD 65536  
SOLVENT DMSO  
NS 16  
DS 2  
SWH 8012.820 Hz  
FIDRES 0.122266 Hz  
AQ 4.0894465 sec  
RG 256  
DW 62.400 usec  
DE 6.50 usec  
TE 1526.7 K  
D1 1.00000000 sec  
TD0 1

==== CHANNEL f1 =====  
SFO1 400.1324710 MHz  
NUC1 1H  
P1 13.00 usec  
PLW1 19.00000000 W

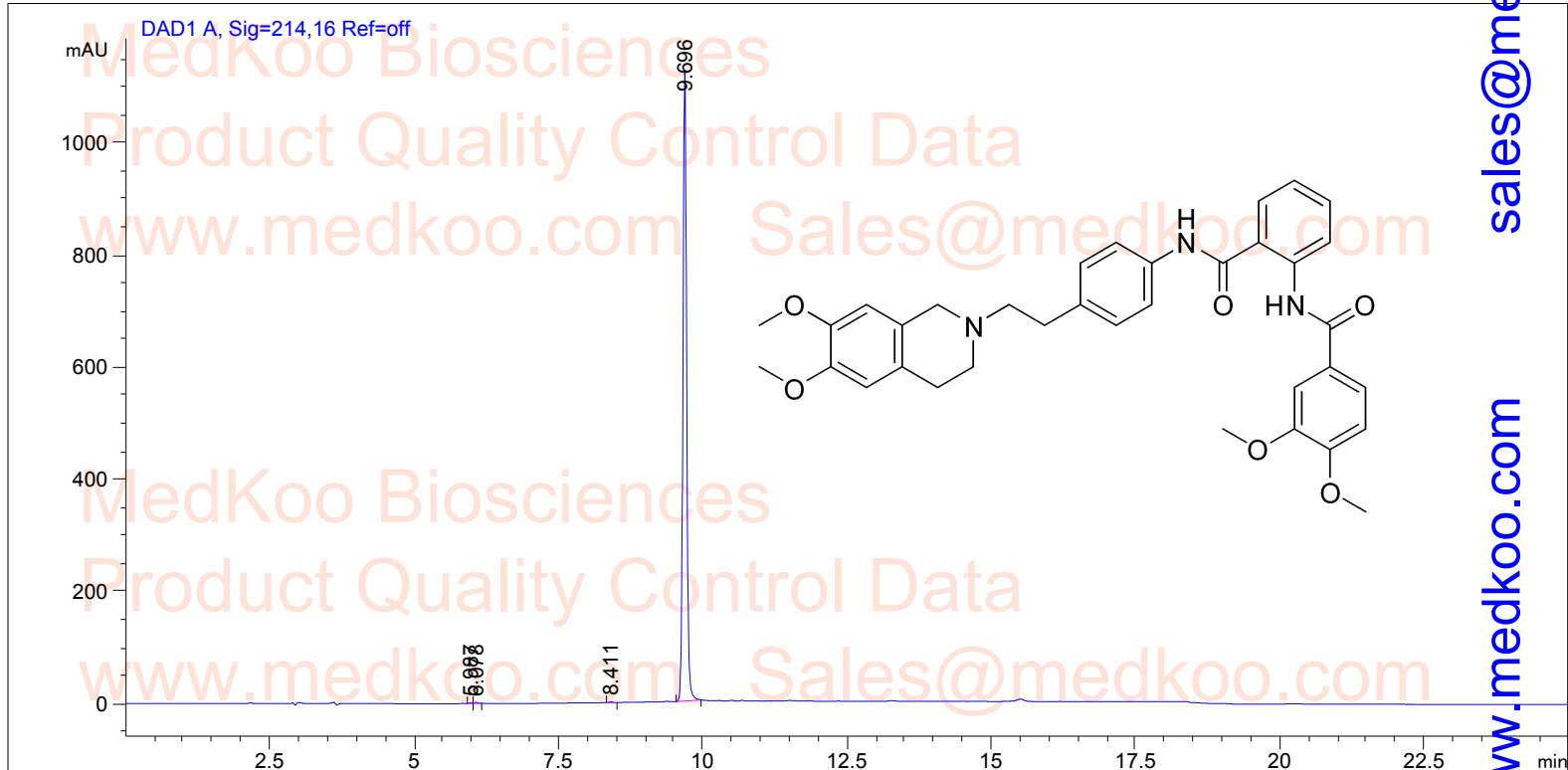
F2 - Processing parameters  
SI 65536  
SF 400.1300000 MHz  
WDW no  
SSB 0  
LB 0 Hz  
GB 0  
PC 1.00



# WIKI4 HPLC Analysis

User : System  
 Instrument : Instrument 1  
 Acq. Method : C:\CHEM32\1\DATA\Normal.M  
 Analysis Method : C:\CHEM32\1\METHODS\Normal.M

Seq. Line : 3  
 Location : Vial 29  
 Inj : 1  
 Inj Volume : 10.0 µl



## Area Percent Report

Sorted By : Signal  
 Multiplier : 1.0000  
 Dilution : 1.0000

MedKoo Cat#: 406349

Name: WIKI4

CAS#: 838818-26-1

Lot#: B7B09C21

Chemical Formula: C<sub>29</sub>H<sub>23</sub>N<sub>5</sub>O<sub>3</sub>S

Exact Mass: 521.15216

Molecular Weight: 521.59

Use Multiplier & Dilution Factor with ISTDs

Signal 1: DAD1 A, Sig=214,16 Ref=off

Peak #	Ret. Time [min]	Type	Width [min]	Area [mAU*s]	Height [mAU]	Area %
1	5.987	BV	0.0457	5.37040	1.86152	0.1058
2	6.078	VB	0.0533	7.10145	2.00657	0.1398
3	8.411	BB	0.0587	6.79239	1.85439	0.1338
4	9.696	BB	0.0705	5059.04346	1125.25195	99.6207

Total : 5078.30769 1130.97443

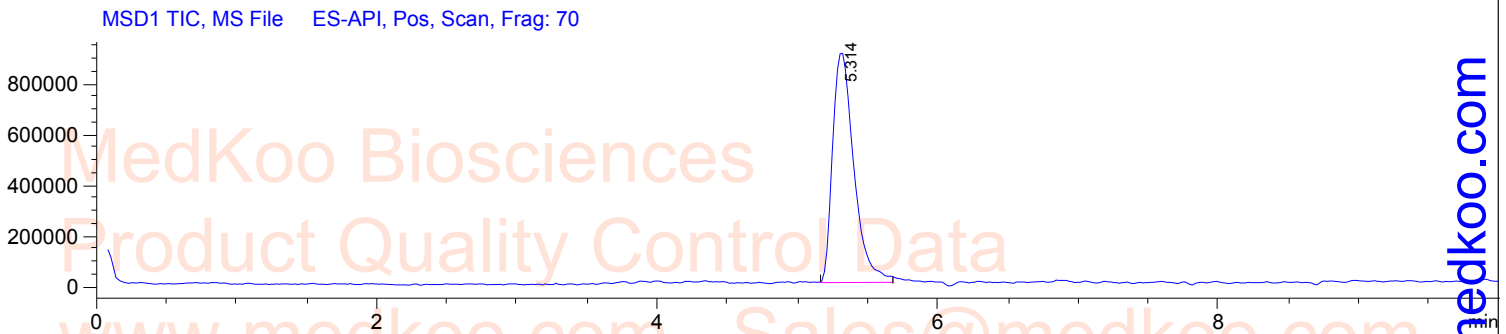
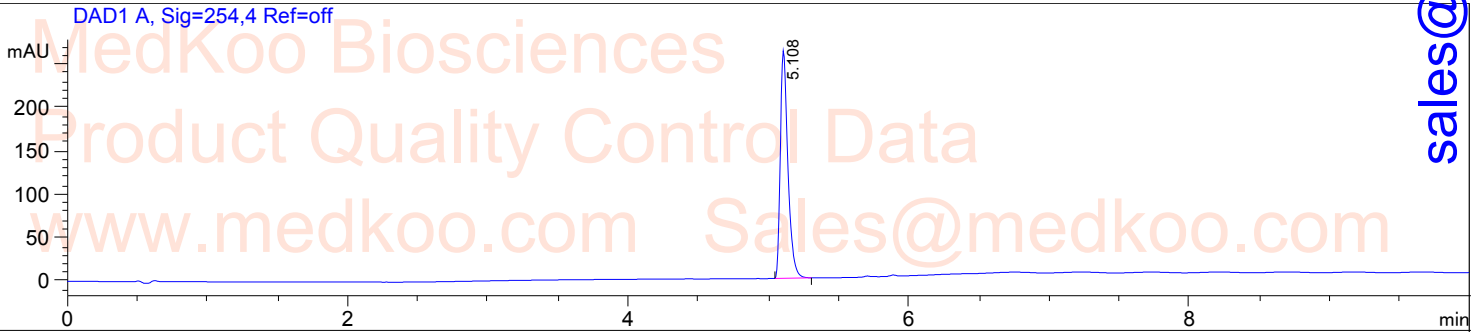
sales@medkoo.com

www.medkoo.com

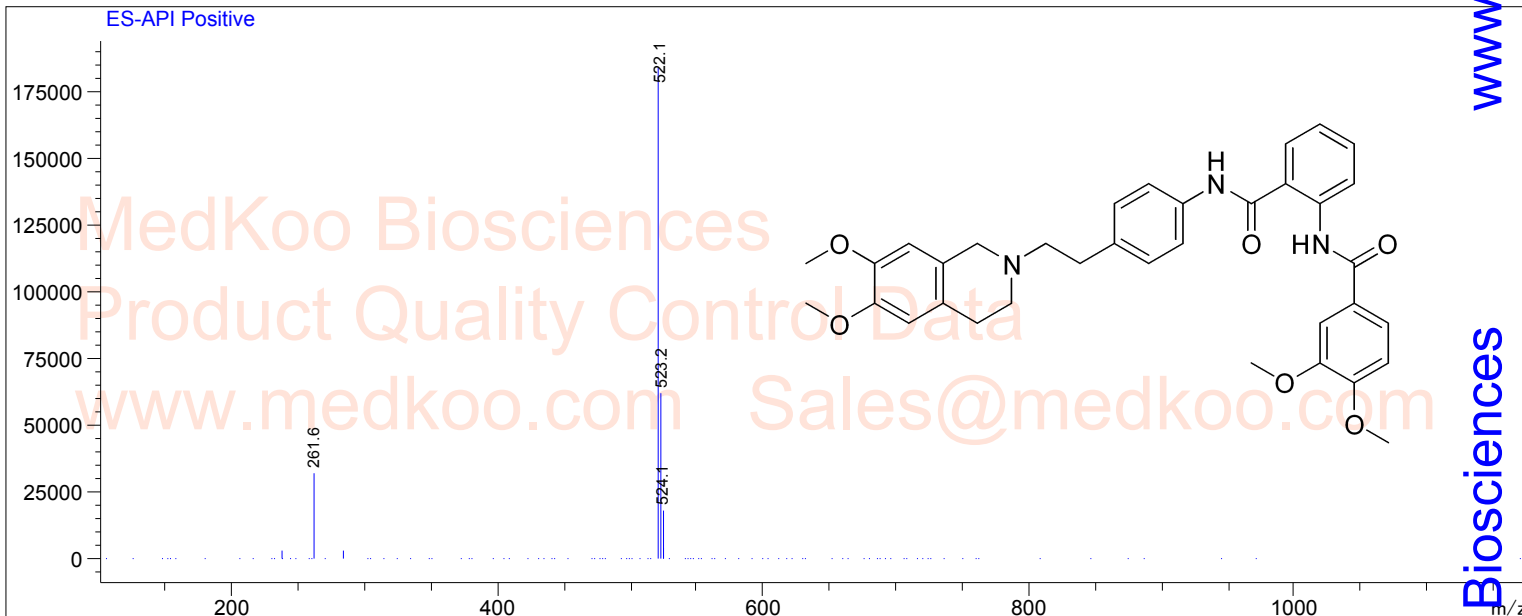
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# WIKI4 LC/MS Analysis

Sample Name : WIKI4 Tgt Mass (EZX) :  
Acq. Operator : [BSB4] Location : Vial 24  
Spec. Reported : MS Integration Inj : 1  
Inj Volume : 1 ul  
Acq. Method : C:\Chem32\1\METHODS\TEST KR POS.M  
Analysis Method : C:\Chem32\1\METHODS\TEST KR POS.M :



Ret. Time: 5.31 <<<< POSITIVE SPECTRA >>>>



MedKoo Cat#: 406349  
Name: WIKI4  
CAS#: 838818-26-1  
Lot#: B7B09C21  
Chemical Formula: C<sub>29</sub>H<sub>23</sub>N<sub>5</sub>O<sub>3</sub>S  
Exact Mass: 521.15216  
Molecular Weight: 521.59

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