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Reaction of Chlorosulfonyl Isocyanate (CSI) with Fluorosubstituted Alkenes: Evidence of a Concerted Pathway for Reaction of CSI with Fluorosubstituted Alkenes (PREPRINT)

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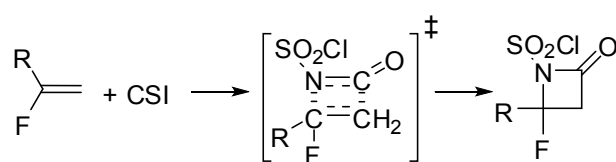
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Abstract: Concerted reactions are indicated for the electrophilic addition of chlorosulfonyl isocyanate with monofluoroalkenes. A vinyl fluorine atom on an alkene raises the energy of a step-wise transition state more than the energy of the competing concerted pathway. This energy shift induces CSI to react with monofluoroalkenes by a one-step process. The low reactivity of CSI with monofluoroalkenes, stereospecific reactions, the absence of 2:1 uracil products with neat fluoroalkenes and quantum chemical calculations support a concerted pathway.

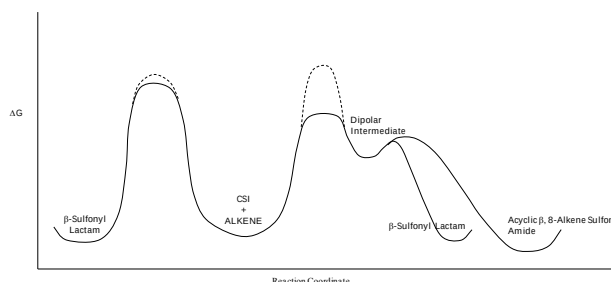
Introduction: Chlorosulfonyl isocyanate (CSI) is the most reactive and versatile isocyanate.¹ CSI reacts with alkenes to give chlorosulfonyl *beta*-lactams that are readily reduced to *beta*-lactams.^{2,3} This reaction sequence provides a synthetic route to *beta*-lactam antibiotics.⁴ Fluorine in *beta*-lactam antibiotics have the fluorine atom attached to the periphery of the compound while the *beta*-lactam ring, the location which interacts with Penicillin binding proteins and *beta*-lactamases, remains unchanged. We demonstrate here a method to synthesize this new class of compounds with the fluorine located on the *beta*-lactam ring.

Reactions of CSI with hydrocarbon alkenes are reported to proceed through an open-ion dipolar intermediate.^{1,3,5} Moriconi suggests that some 1,2-disubstituted olefins retain stereochemistry through fast collapse of the dipolar intermediate.^{3,5} *Ab initio* calculations show that [2 + 2]

cycloadditions between alkenes and isocyanates can react via a concerted transition state with zwitterionic character.⁶ These calculations also found that electron-donating groups on the alkene, or electron-withdrawing groups on the isocyanate, lower the activation energy and induce the nature of the reaction to become more synchronous.⁶ Calculations also support a concerted process for the cycloaddition of isocyanates with aldehydes.⁷ Quantum chemical calculations and photoelectron spectral data show that substituting a hydrogen with a fluorine atom on the pi-bond of an alkene does not significantly alter the molecular energy of the pi-bond,⁸ and therefore, the HOMO and LUMO orbital energies for a concerted pathway should not be altered either. On the other hand, the energy for a dipolar stepwise pathway is raised significantly by the vinyl fluorine atom through its strong inductive effect.⁹ This perturbation of the Free Energy profile is described in Figure 1 where the fluorine atom raises the transition state energy significantly for the step-wise process, but it only increases the energy of the concerted pathway by a modest amount. In Figure 1 the solid line represents the energy profile for hydrocarbon alkenes while the dashed line describes the pathway for monofluoroalkenes. Therefore, alkenes with a vinyl fluorine atom may allow a concerted process to compete with or completely dominate the step-wise pathway. Both concerted and step-wise pathways might be realized for reactions of CSI with appropriately substituted fluoroalkenes. The product stereochemistry and perhaps even the regiochemistry might be influenced by changing from an open-ion dipolar intermediate compared to a one-step concerted pathway.

Figure 1

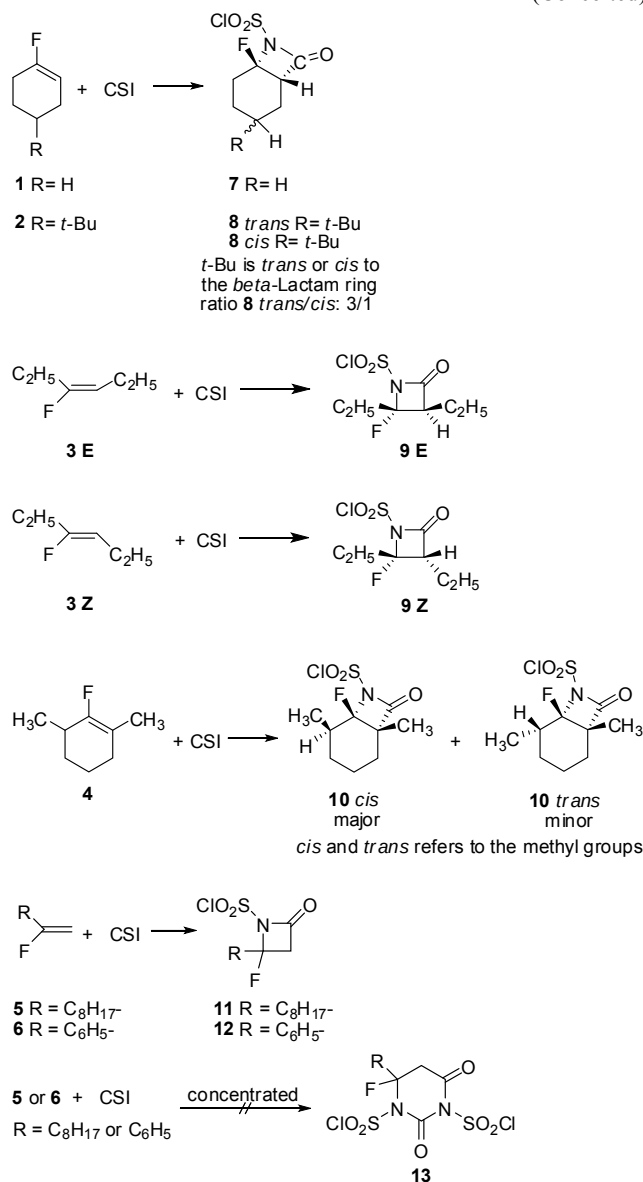
Free Energy Diagram for Reaction of Chlorosulfonyl Isocyanate (CSI) with Hydrocarbon Alkenes and Fluorocarbon Alkenes



Results and Discussion: CSI is a sluggish electrophile and it reacts poorly in solution with alkenes that contain an electron-withdrawing vinyl fluorine¹⁰. We found that neat reactions of CSI with these less reactive fluoroalkenes proceed smoothly and in good yield. Neat reactions of CSI with these monofluoroalkenes allow for the synthesis of *beta*-fluorolactams under “Green Chemistry” conditions. Thus, dialkylsubstituted monofluoroalkenes like the 1-fluorocyclohexenes (1), (2), 3-fluorohex-3-enes 3 (E) and 3 (Z), and the trialkylsubstituted fluorocyclohexene (4) react with CSI to give the chlorosulfonyl *beta*-fluorolactams (7), 8 *cis/trans*, 9(E), 9(Z), and 10 *cis/trans*, respectively (Scheme 1). A stereospecific reaction of CSI with 3 (E) and 3 (Z) is consistent with a concerted process for this series of fluoroalkenes. Product regiochemistry was confirmed by the carbonyl ¹³C NMR three bond coupling with fluorine (*J*_{C-F} = 3-

6 Hz). The nitrogen of the *beta*-lactams is bonded to the carbon with the fluorine since the developing positive charge in the concerted transition state prefers to be on the carbon stabilized by back-bond resonance from fluorine.

Scheme 1
(Concerted)



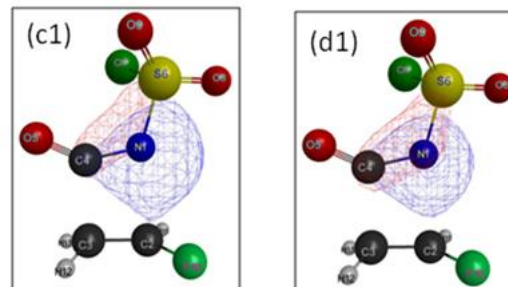
The regiochemistry of the *beta*-sulfonyl fluorolactam products did not change when a third alkyl group was incorporated in fluoroalkene **4** as indicated by the three bond fluorine to carbonyl coupling of 3 Hz in the *beta*-lactams (**10** *cis/trans*). Assignment of the carbons from **10** *cis* and **10** *trans* were apparent from the magnitude of the carbon-fluorine coupling and from DEPT and HSQC experiments. The *cis/trans* stereochemistry of **10** was assigned using a 1-dimension ROESY experiment. Irradiating the upfield methyl adjacent to the carbonyl of the major isomer enhanced the methyl on the methine carbon. Irradiating the upfield methyl of the minor isomer enhanced the methine hydrogen on the

minor isomer. Irradiating the methine hydrogen's of each isomer separately confirmed the experiments irradiating the methyl groups above.

Products from 2-fluorodec-1-ene (**5**) and 2-fluoro-2-phenylethene (**6**) decomposed at elevated temperatures. The *beta*-sulfonyl fluorolactams (**11** and **12**) were formed with **5** or **6** and CSI in methylene chloride at room temperature (Scheme 1). At high concentrations of **5** or **6**, approaching the reaction conditions used for fluoroalkenes **1**, **2**, **3** (**E**), **3** (**Z**) and **4**, uracil products **13** were not formed. At these high concentrations we would expect capture of a dipolar intermediate by a second molecule of CSI to give uracil products like those reported for the reaction of CSI with hydrocarbon alkenes that can support stable dipolar intermediates.^{1a,2a,11} Thus we suggest that fluoroalkenes **1** through **6** react by a concerted pathway.

Quantum chemical calculations at the MP2/6-311G(d,p) level of theory^{12a-c,13} also support our claim of a one-step process for reaction of CSI with fluoroalkenes as described in Figure 1. Transition states for the concerted pathway and a portion of the stepwise pathway were calculated for reaction of CSI with vinyl fluoride (Supporting Information). Intrinsic reaction coordinate calculations were performed to trace the minimum energy paths connecting the transition states to the corresponding local minima; i.e., reactants and products. The step-wise transition state, which is 60.9 kcal/mol above separated CSI + fluoroethene reactants, was found to be 26.6 kcal/mol higher in energy than the concerted transition state (34.3 kcal/mol above reactants.) The concerted transition state is not orthogonal as reported for ketene cycloadditions where the orbitals mix by a $[\pi^2(s) + \pi^2(a)]$ process¹⁴ A six electron process, involving the lone pair on nitrogen represented as $\omega^2 [\pi^2(s) + \pi^2(s) + \omega^2(s)]$, would allow for a concerted cyclization where the alkene carbon atoms and the O=C=N- moiety of CSI are in the same plane. Calculated localized molecular orbitals of the cyclic 2+2 transition state for the cycloaddition of CSI to vinyl fluoride show significant mixing between the C-N pi bond in CSI and the nitrogen lone pair electrons (Figure 2).

Figure 2



(c1) are the electrons of the C-N pi bond. (d1) the lone pair electrons of the nitrogen atom.

Our data support a concerted reaction of CSI with these less reactive fluoroalkenes because:

- (1) Reactions with **3** (**E**) and **3** (**Z**) are stereospecific.
- (2) Neat reactions of CSI with **1**, **2**, **3** (**E**), **3** (**Z**), **4**, **5**, and **6** do not give uracil products.

(3) A concerted pathway is supported by quantum chemical calculations.

We are investigating the parameters that seem to influence a change of mechanism for reactions of fluoroalkenes with CSI.

Experimental Section: Diethylaminosulfur trifluoride was added to cyclohexanones in methylene chloride to give mixtures of 1,1-difluorocyclohexanes and 1-fluorocyclohexenes. After water work-up, the methylene chloride was removed by distillation and the mixture was distilled through a vigreux column to give enriched 1-fluorocyclohexenes **1**, **2** and **4** containing various amounts of 1,1-difluorocyclohexanes. Acyclic fluoroalkenes **3E**¹⁵, **3Z**¹⁵ and **5**¹⁵, **6**¹⁶ were prepared as described in the literature. The products were isolated by chromatography (column or preparative thin layer), or in one case by crystallization. The following procedure is representative.

To 156 mg (1.00 mmol) 4-*tert*-butyl-1-fluorocyclohexene (**4**) in a small round bottom flask was added 155 mg, 96 microliter (1.10 mmol) chlorosulfonyl isocyanate (CSI). The stirred mixture was heated to 65-70° C for one hour and then cooled. Methylene chloride (2-3 mL) was added, followed by dropwise addition of ice water. The organic layer was separated and the aqueous layer extracted with methylene chloride. The combined organic extractions were washed with 2 % aqueous sodium bicarbonate, dried over anhyd. magnesium sulfate and concentrated. ¹⁹F NMR analysis on the crude mixture showed **8** *cis/trans* to be formed in a ratio of 1.0/3.0, respectively. Column chromatography (10 g silica gel) of the crude mixture with hexanes/chloroform gave a 194 mg, 65%, of pure **8** *cis*/**8** *trans* in a ratio of 1.0/2.6 respectively. Reactions of fluoroalkenes **1**, **2**, **3E**, **3Z** with CSI were done similarly. Spectral and exact mass data are listed in the **Supporting Information** section.

CSI (1.10 mmol) was added to fluoroalkenes **5** or **6** (1.00 mmol) in 0.2 to 4 mL methylene chloride at 0° C. The mixture was allowed to warm to room temperature and then stirred for four hours. Work-up was accomplished as described above for the reactions with **5** and **6**. Product **11** was obtained 90% pure (¹⁹F NMR) by preparative TLC while product **12** was isolated by crystallization from ether. Crystals from **12** decomposed in several minutes at room temperature, but were sufficiently stable in solution to obtain spectral data. Wet crystals of **12** were kept cold during transportation for X-Ray analysis at low temperature.

Acknowledgment: Support for this work was provided by the National Science Foundation (NSF-RUI Grant No. CHE-0640547), and Research Associates of PLNU (alumni support group). We would also like to acknowledge our use of the 400 MHz NMR at the University of San Diego obtained by support from the National Science Foundation (NSF MRI Grant No. CHE-0417731). We thank Dr. Richard Kondratt at the Mass Spectrometry Center, the University of California, Riverside for the Exact Mass data. We also want to thank Dr. Leroy Lafferty at San Diego State University for conducting the 1-dimensional ROESY experiments at 600 MHz.

Supporting Information Available: Spectral data to characterize the products, X-Ray data for **12** and quantum chemical data are available on line at <http://pubs.acs.org>.

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Supporting Information

Reaction of Chlorosulfonyl Isocyanate (CSI) with Fluorosubstituted Alkenes: Evidence for a Concerted Pathway with CSI and Fluorosubstituted Alkenes

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Tabulated ¹H, ¹⁹F, and ¹³C NMR, Infrared, Exact Mass and Isolated Yield Data

7: Isolated (50%) by column chromatography on silica gel with hexanes/methylene chloride. ¹H NMR 400 MHz (CDCl₃) δ = 1.59-1.72 (m, 3H); 1.81-1.99 (m, 2H); 2.01-2.21 (m, 2H); 2.69-2.80 (m, 1H); 3.52-3.63 (m, 1H). ¹⁹F NMR 376 MHz (CDCl₃) δ = -112.8 (m). ¹³C NMR 100.6 MHz (CDCl₃) δ = 15.5 (d, *J* = 8 Hz); 16.0 (s); 18.4 (s); 24.7 (d, *J* = 25 Hz); 53.5 (d, *J* = 21 Hz); 102.9 (d, *J* = 248 Hz); 160.1 (d, *J* = 4 Hz). IR (KBr) neat 1832 cm⁻¹. Exact mass [MH]⁺ calcd. for C₇H₁₀N)₃FSCl 242.00539; found 242.00470.

8 *trans/cis*: Isolated (65%) as a 2.6/1.0 ratio *trans/cis* by column chromatography as described above. ¹H NMR 400 MHz (CDCl₃) δ = 0.89 (s, 9H); 1.20-2.30 (m, 6H); [*trans* 2.55-2.75 (m) and *cis* 2.78-2.90 (m), 1H]; [*cis* 3.50 (m) and *trans* 3.67 (dm, *J* = 13 Hz), 1H]. ¹⁹F NMR 376 MHz (CDCl₃) *trans* δ = -117.7 (m) and *cis* -114.1 (m), ratio 3/1, respectively on the crude reaction mixture. ¹³C NMR 100.6 MHz (CDCl₃) **8 *trans*** δ = 19.2 (d, *J* = 8 Hz); 21.6 (s); 26.4 (d, *J* = 26 Hz); 26.7 (s); 33.2 (s); 40.0 (s); 57.5 (d, *J* = 21 Hz); 105.2 (d, *J* = 248 Hz); 161.7 (d, *J* = 6 Hz). **8 *cis*** δ = 21.2 (d, *J* = 9 Hz); 22.9 (s); 26.9 (s); 29.4 (d, *J* = 26 Hz); 32.9 (s); 43.2 (s); 55.5 (d, *J* = 22 Hz); 105.1 (d, *J* = 246 Hz); 162.9 (d, *J* = 4 Hz). IR (KBr) neat mixture *trans* 1826 cm⁻¹ *cis* 1838 cm⁻¹. Exact mass [MH]⁺ calcd. for C₁₁H₁₈NO₃FSCl 298.067996; found 298.068000.

9E: Isolated (50%) by column chromatography as described above. ¹H NMR 400 MHz (CDCl₃) δ = 1.18 (t, *J* = 7.4 Hz, 6H); 1.65-1.98 (m, 2H); 2.03-2.23 (m, 1H); 2.54-2.68 (m, 1H); 3.42-3.52 (m, 1H). ¹⁹F NMR 376 MHz (CDCl₃) δ = -119.4 (ddd, *J* = 30.5, 13.7 and 9.2 Hz). ¹³C NMR 100.6 MHz (CDCl₃) δ = 7.6 (d, *J* = 4 Hz); 11.6 (s); 18.5 (d, *J* = 2 Hz); 24.7 (d, *J* = 28 Hz); 63.1 (d, *J* = 24 Hz); 108.2 (d, *J* = 247 Hz); 162.2 (d, *J* = 5 Hz). IR (KBr) neat 1830 cm⁻¹. Exact mass [MH]⁺ calcd. for C₇H₁₂NO₃FSCl 244.0210; found 244.0202.

9Z: Isolated (55%) by column chromatography as described above. ¹H NMR 400 MHz (CDCl₃) δ = 1.11 (t, *J* = 7.6 Hz, 3H); 1.14 (t, *J* = 7.4 Hz, 3H); 1.78-1.99 (m, 2H); 2.10-2.29 (m, 1H); 2.47-2.60 (m, 1H); 3.36-3.43 (m, 1H). ¹⁹F NMR 376 MHz (CDCl₃) δ = -137.3 (dt, *J* = 27.5 and 6.9 Hz). ¹³C NMR 100.6 MHz (CDCl₃) δ = 7.8 (d, *J* = 4 Hz); 11.7 (s); 17.7 (d, *J* = 5 Hz); 27.5 (d, *J* = 28 Hz); 60.2 (d, *J* = 22 Hz); 107.6 (d, *J* = 249 Hz); 162.4 (d, *J* = 1.5 Hz). IR (KBr) neat 1833 cm⁻¹. Exact mass, negative ion ESI [M⁺-H] calcd. for C₇H₁₀NO₃FSCl 242.0054; found 242.0051.

10 *cis/trans*: *cis* and *trans* refers to the two methyl groups on the cyclohexane ring. Isolated (48%) by column chromatography as described above. ^1H NMR 600 MHz (C_6D_6) δ = [*cis* 1.15 (dd, J = 7.0 and 1.8 Hz) and *trans* 1.26 (d, J = 7.0 Hz, 3H)]; [*trans* 1.30 (d, J = 2.9 Hz) and *cis* 1.33 (d, J = 2.9 Hz, 3H)]; *cis* and *trans* 1.43-1.62 (m, 2H); *cis* and *trans* 1.62-1.73 (m, 2H); *cis* and *trans* 1.80-1.96 (m, 2H); [*cis* 2.26 (m) and *trans* 2.78 (m), 1H]. ^{19}F NMR 376 MHz (CDCl_3) *trans* δ = -135.3 (s); *cis* -138.6 (brd. s), ratio of 1.0/1.1, respectively on the crude reaction mixture. ^{13}C NMR 150.8 MHz (C_6H_6) assignments supported by DEPT and HSQC experiments. **10** *cis* δ = 15.2 (CH_3 , d, J = 8.4 Hz); 16.1 (CH_3 , d, J = 7.9 Hz); 16.0 (CH_2 , s); 26.0 (CH_2 , d, J = 4.5 Hz); 28.7 (CH_2 , s); 31.5 (CH , d, J = 24.7 Hz); 59.9 (C adj. to the carbonyl, d, J = 20.2 Hz); 108.7 (d, J = 256.4 Hz); 166.3 (d, J = 2.8 Hz). **10** *trans*: δ = 14.4 (CH_3 , d, J = 9.0 Hz); 14.7 (CH_3 , d, J = 2.8 Hz); 17.1 (CH_2 , s); 25.6 (CH_2 , d, J = 7.3 Hz); 28.8 (CH_2 , s); 32.4 (CH , d, J = 24.1 Hz); 61.8 (C adj. to the carbonyl, d, J = 18.0 Hz); 111.2 (d, J = 256.4 Hz); 166.7 (d, J = 2.8 Hz). IR (KBr) neat mixture 1834 cm^{-1} . Exact mass, negative ion ESI [$\text{M}^+ - \text{H}$] calcd. for $\text{C}_9\text{H}_{12}\text{NO}_3\text{FSCl}$ 268.0210; found 268.0212.

11: Decomposition produced 8% side products during purification by preparative thin layer chromatography on silica gel with chloroform/methanol (95:5). Isolated in 33% yield. ^1H NMR 400 MHz (CDCl_3) δ = 0.89 (t, J = 7.0 Hz, 3H); 1.29 (m, 10H); 1.38-1.62 (m, 2H); 2.06-2.26 (m, 1H); 2.44-2.56 (m, 1H); 3.33-3.48 (m, 2H). ^{19}F NMR 376 MHz (CDCl_3) δ = -120.9 (m). The 8% impurity around -131 to -132 ppm is from decomposition during purification by TLC. ^{13}C NMR 100.6 MHz (CDCl_3). δ = 14.0 (s); 22.5 (s); 23.5 (s); 23.7 (s); 29.0 (s); 29.1 (d, J = 14.0 Hz); 31.7 (s); 48.9 (d, J = 25.1 Hz); 76.8 (d, J = 4.8 Hz); 105.7 (d, J = 246.1 Hz); 158.6 (d, J = 3.0 Hz). IR (KBr) neat 1831 cm^{-1} . Exact mass, negative ion ESI [$\text{M}^+ - \text{H}$] calcd. for $\text{C}_{11}\text{H}_{18}\text{NO}_3\text{FSCl}$ 298.0680; found 298.0716.

12: Yield (65%) by ^{19}F NMR with 4-fluoroanisole as internal standard. ^1H NMR 400 MHz (CDCl_3) δ = 3.62-3.85 (m, 2H); 7.51 (m, 3H); 7.61 (m, 2H). ^{19}F NMR 376 MHz (CDCl_3) δ = -129.0 (t, J = 10.5 Hz). ^{13}C NMR 100.6 MHz (CDCl_3). δ = 53.4 (d, J = 25 Hz); 103.7 (d, J = 246 Hz); 125.2 (d, J = 8 Hz); 129.2 (s); 130.9 (s); 132.0 (d, J = 29 Hz); 158.9 (d, J = 2 Hz). IR (KBr) neat 1834 cm^{-1} .

Automation directory: /home/organic/vnmrsys/data/studies/auto_2008.06.13_01
Sample id : /home/organic/vnmrsys/data/plnu/s_DMH29_C01
Sample : DMH29_C

Pulse Sequence: s2pu1

Solvent: d2o

Ambient temperature

Operator: plnu

File: DMH29_C_Carbon_01

Mercury-400BB "pandora.scst.sandiego.edu"

Relax. delay 5.000 sec

Pulse 45.0 degrees

Acq. time 1.300 sec

Width 24154.6 Hz

128 repetitions

OBSERVE C13, 100.6198270 MHz

DECOUPLE H1, 400.1601851 MHz

Power 41 dB

continuously on

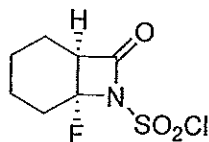
WALTZ-16 modulated

DATA PROCESSING

Line broadening 0.5 Hz

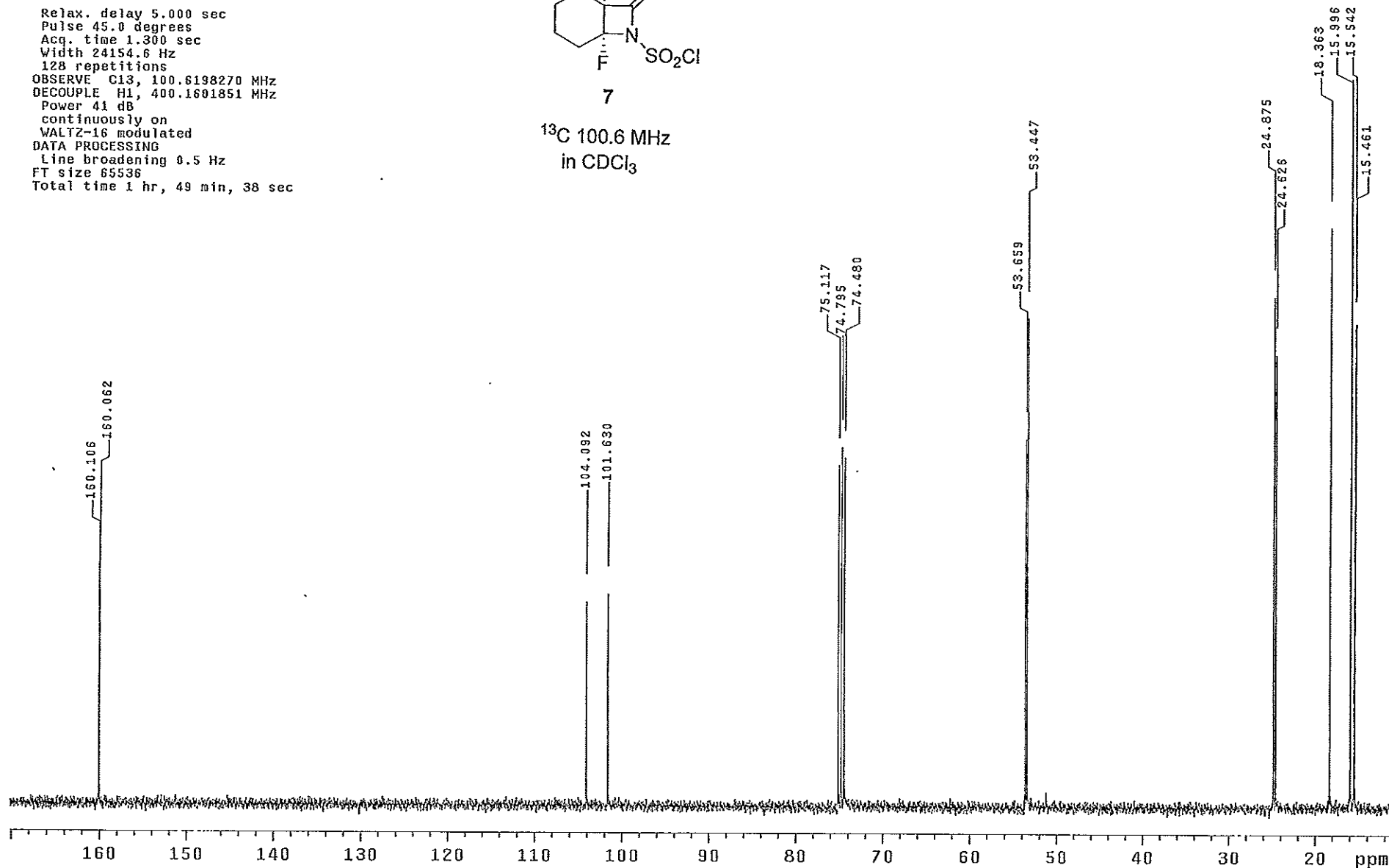
FT size 65536

Total time 1 hr, 49 min, 38 sec



7

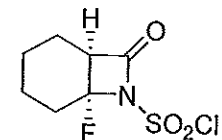
¹³C 100.6 MHz
in CDCl₃



Automation directory: /home/organic/vnmrsys/data/studies/auto_2008.06.13_01
Sample id : /home/organic/vnmrsys/data/plnu/s_DMH29_F01
Sample : DMH29_F

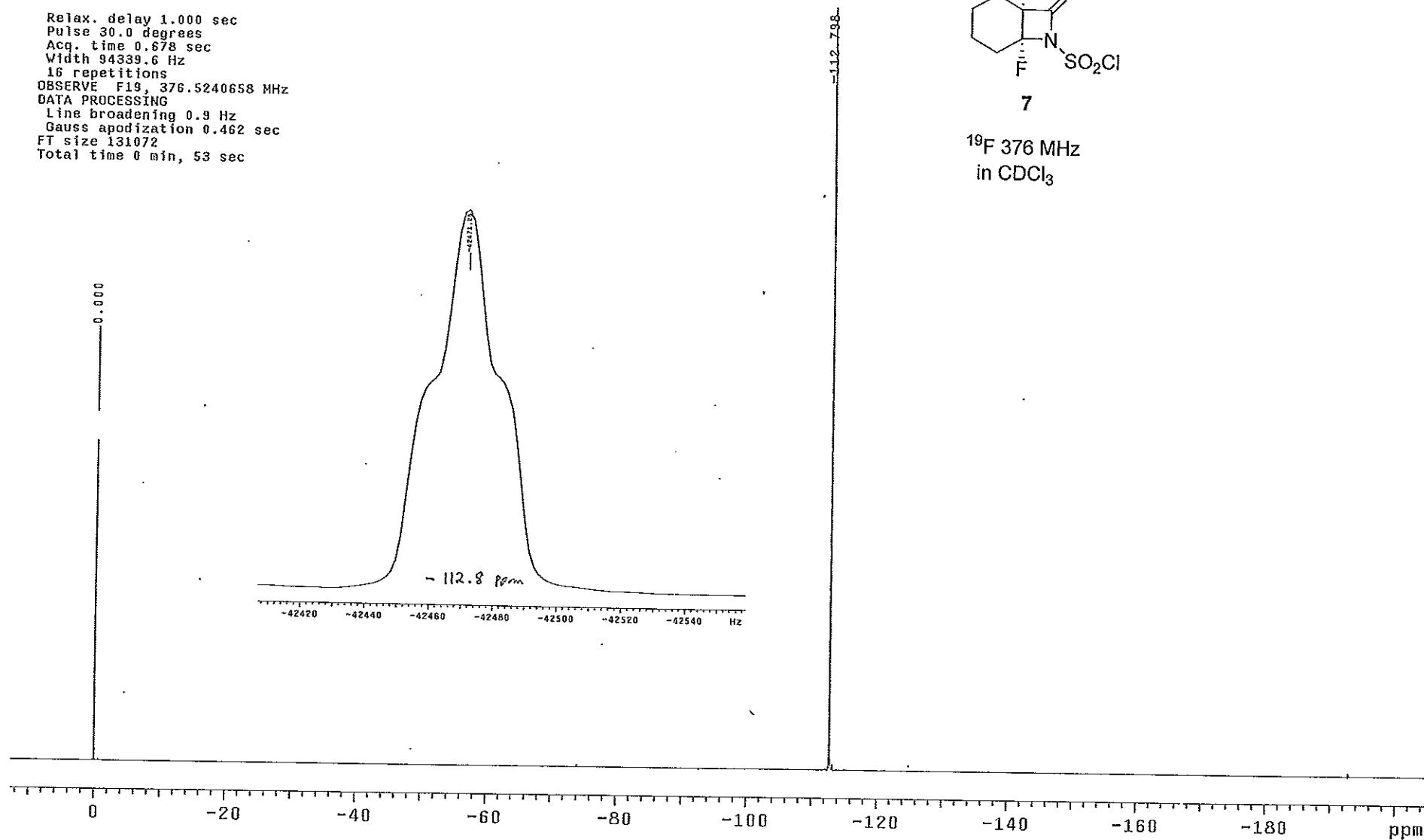
Pulse Sequence: s2pu1
Solvent: d2o
Ambient temperature
Operator: plnu
File: DMH29_F_Fluorine_01
Mercury-400BB "pandora.scst.sandiego.edu"

Relax. delay 1.000 sec
Pulse 30.0 degrees
Acq. time 0.678 sec
Width 94339.6 Hz
16 repetitions
OBSERVE F19, 376.5240658 MHz
DATA PROCESSING
Line broadening 0.9 Hz
Gauss apodization 0.462 sec
FT size 131072
Total time 0 min, 53 sec



7

^{19}F 376 MHz
in CDCl_3



S5

Automation directory: /home/organic/vnmrsys/data/studies/auto_2008.06.13_01
 Sample id : /home/organic/vnmrsys/data/plnu/s_DMH29_H01
 Sample : DMH29_H

Pulse Sequence: s2pul

Solvent: cdcl3

Ambient temperature

Operator: plnu

File: DMH29_H_Proton_01

Mercury-400BB "pandora.scst.sandiego.edu"

Relax. delay 2.000 sec

Pulse 45.0 degrees

Acq. time 1.998 sec

Width 6402.0 Hz

8 repetitions

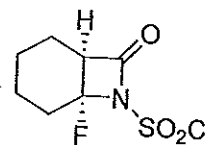
OBSERVE H1, 400.1571242 MHz

DATA PROCESSING

Line broadening 0.5 Hz

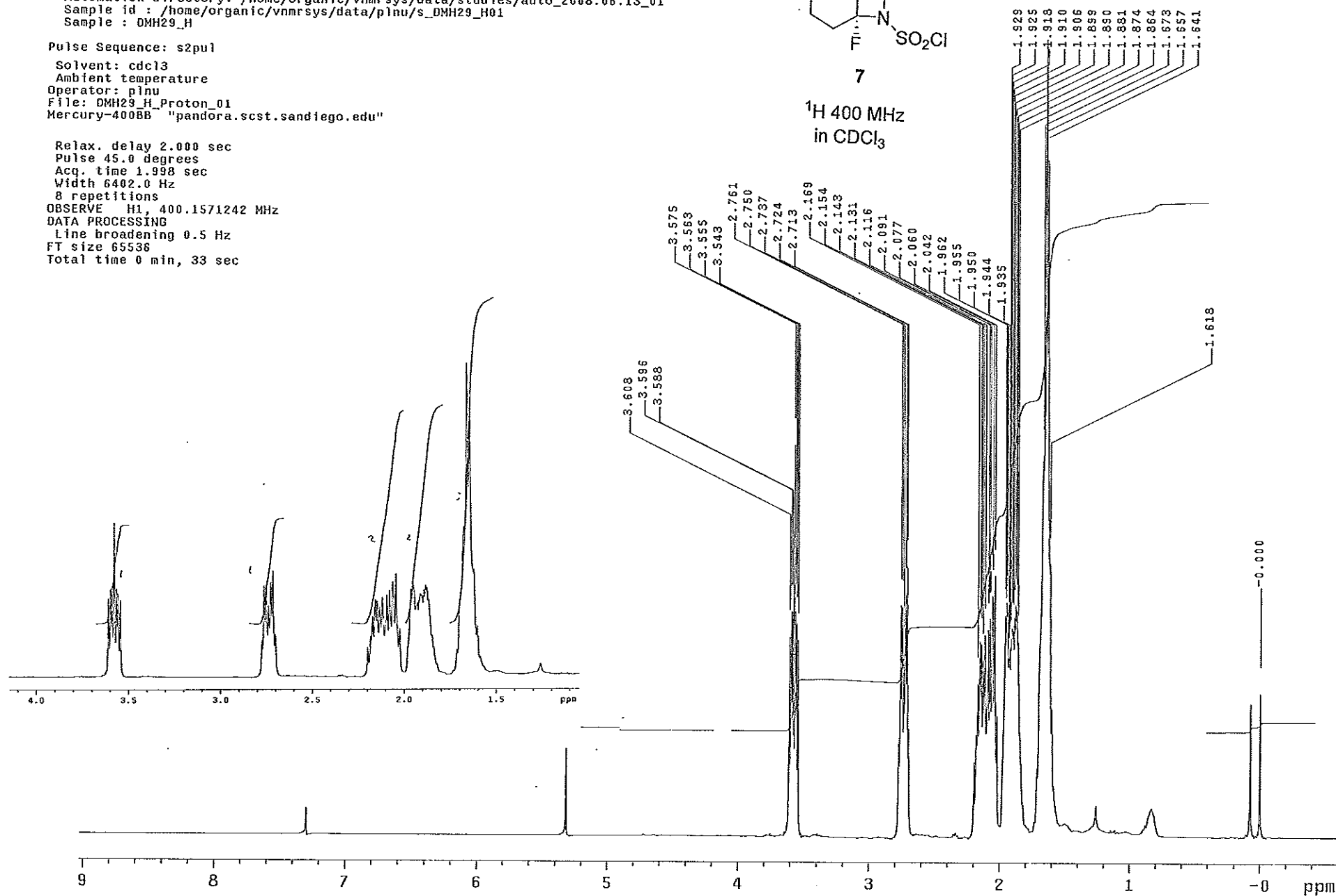
FT size 65536

Total time 0 min, 33 sec



7

¹H 400 MHz
in CDCl₃

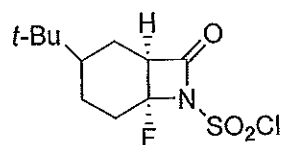


Automation directory: /home/organic/vnmrsys/data/studies/auto_2007.06.21
 Sample id : /home/organic/vnmrsys/data/plnu/s_KD33_H01
 Sample : KD33_H

Pulse Sequence: s2pul
 Solvent: cdc13
 Temp. 20.0 C / 293.1 K
 Operator: organic
 File: KD33_H_Proton_01
 Mercury-400BB "pandora"

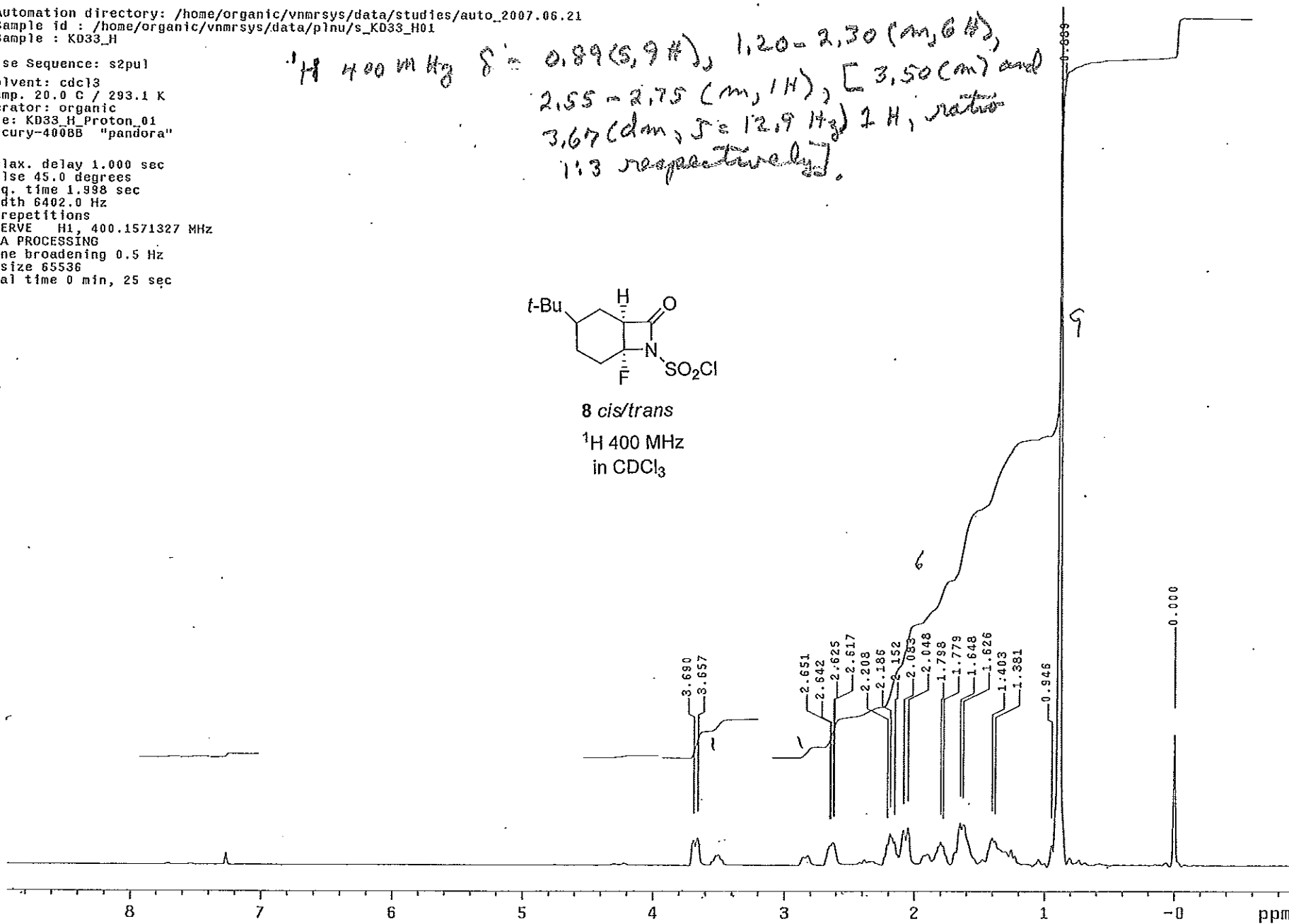
Relax. delay 1.000 sec
 Pulse 45.0 degrees
 Acq. time 1.998 sec
 Width 6402.0 Hz
 8 repetitions
 OBSERVE H1, 400.1571327 MHz
 DATA PROCESSING
 Line broadening 0.5 Hz
 FT size 65536
 Total time 0 min, 25 sec

^1H 400 MHz δ = 0.89 (s, 9H), 1.20-2.30 (m, 6H),
 2.55-2.75 (m, 1H), [3.50 (m) and
 3.67 (dm, J = 12.9 Hz) 2H, ratio
 1:3 respectively].



8 cis/trans

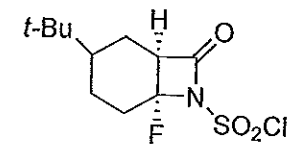
^1H 400 MHz
 in CDCl_3



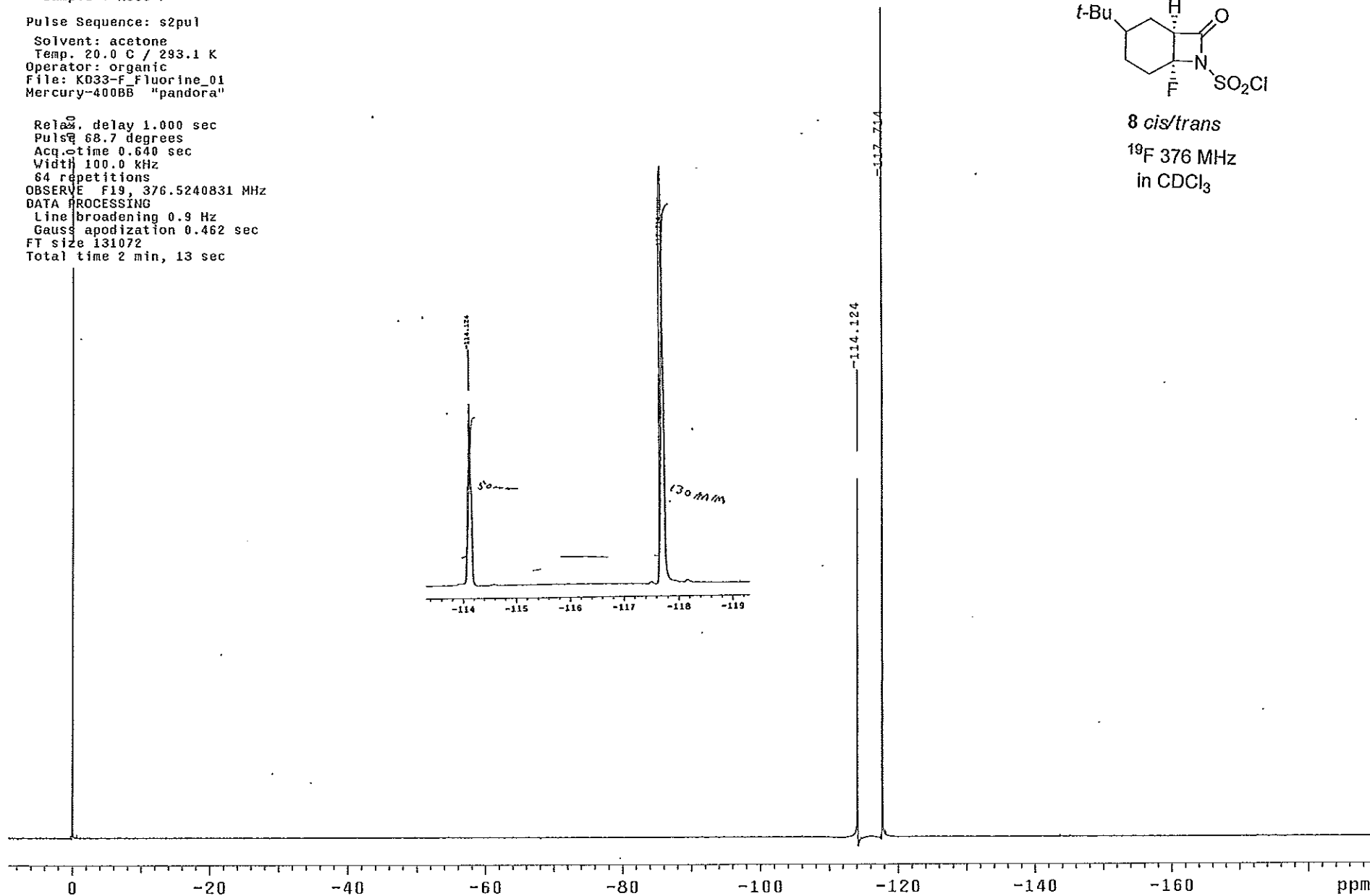
Automation directory: /home/organic/vnmrsys/data/studies/auto_2007.06.21
Sample id : /home/organic/vnmrsys/data/plnu/s_K033-F01
Sample : K033-F

Pulse Sequence: s2pul
Solvent: acetone
Temp. 20.0 C / 293.1 K
Operator: organic
File: K033-F_Fluorine_01
Mercury-400BB "pandora"

Relax. delay 1.000 sec
Pulse 68.7 degrees
Acq. time 0.640 sec
Width 100.0 kHz
64 repetitions
OBSERVE F19, 376.5240831 MHz
DATA PROCESSING
Line broadening 0.9 Hz
Gauss apodization 0.462 sec
FT size 131072
Total time 2 min, 13 sec



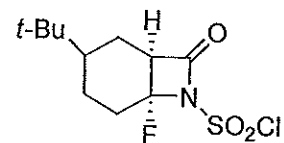
8 cis/trans
¹⁹F 376 MHz
in CDCl₃



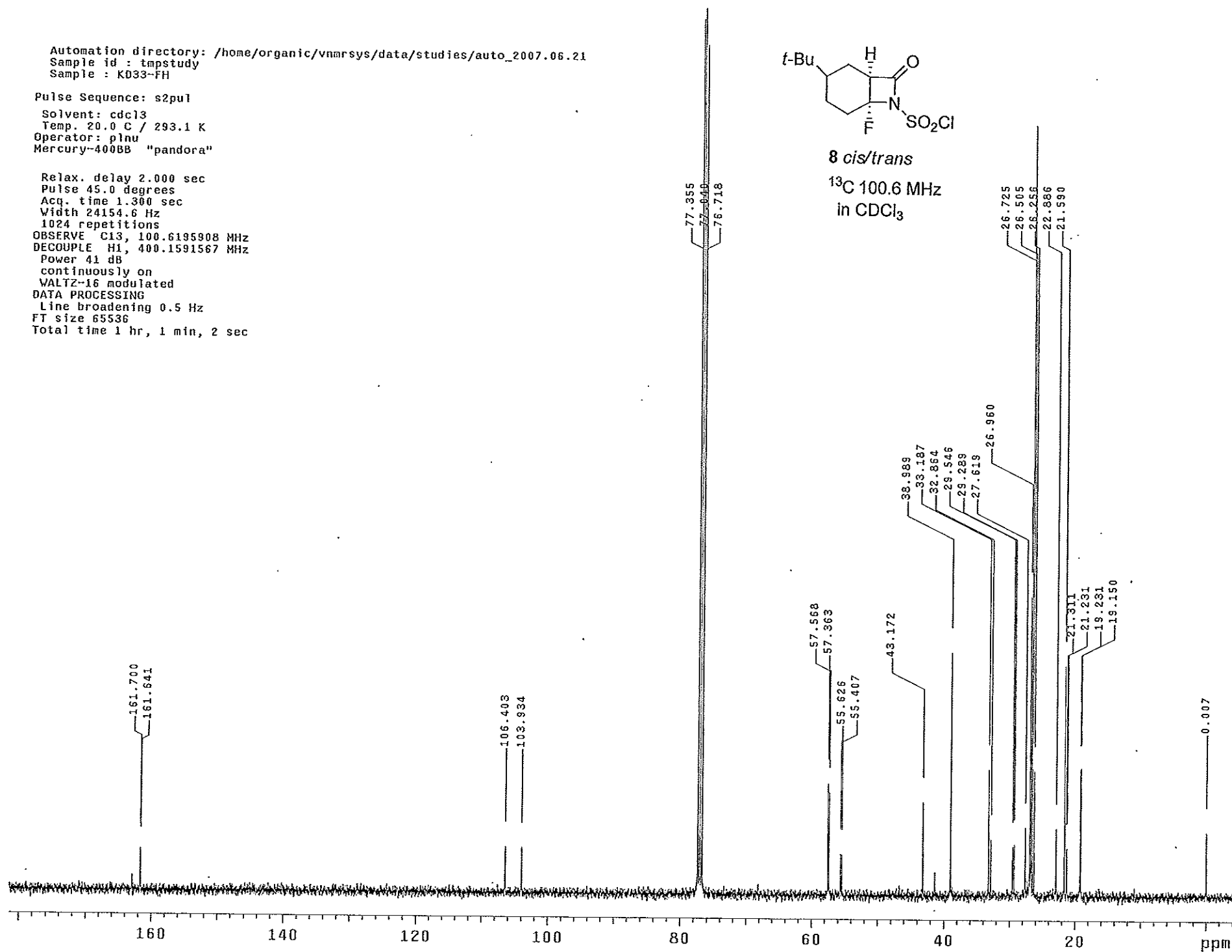
Automation directory: /home/organic/vnmrsys/data/studies/auto_2007.06.21
Sample id : tmpstudy
Sample : KD33-FH

Pulse Sequence: s2pul
Solvent: cdcl3
Temp. 20.0 C / 293.1 K
Operator: plnu
Mercury-400BB "pandora"

Relax. delay 2.000 sec
Pulse 45.0 degrees
Acq. time 1.300 sec
Width 24154.6 Hz
1024 repetitions
OBSERVE C13, 100.6195908 MHz
DECOUPLE H1, 400.1591567 MHz
Power 41 dB
continuously on
WALTZ-16 modulated
DATA PROCESSING
Line broadening 0.5 Hz
FT size 65536
Total time 1 hr, 1 min, 2 sec



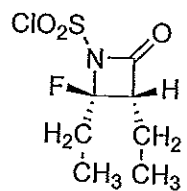
8 cis/trans
¹³C 100.6 MHz
in CDCl₃



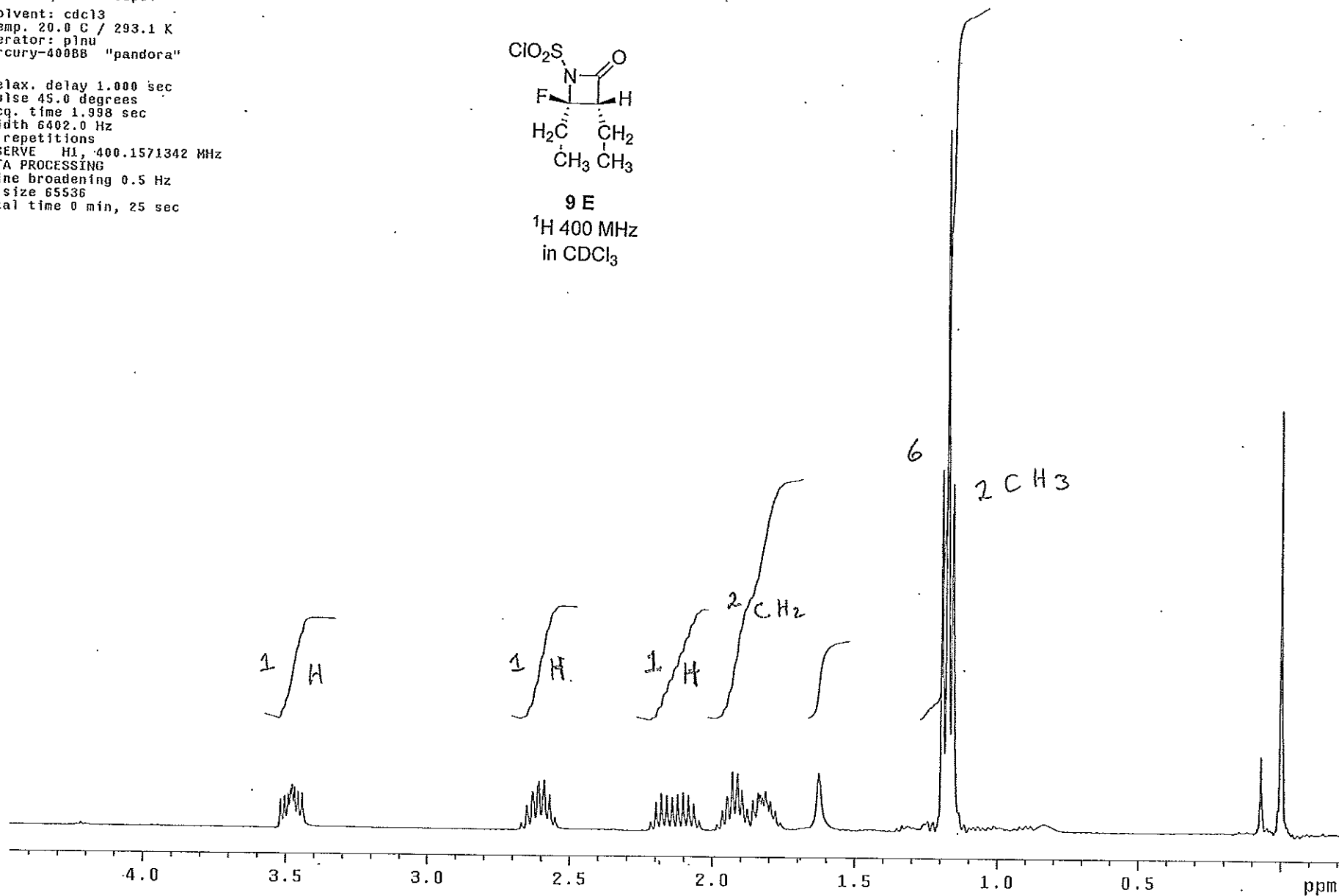
Automation directory: /home/organic/vnmrsys/data/studies/auto_2007.07.12
Sample id : tmpstudy
Sample : KB-45_H

Pulse Sequence: s2pul
Solvent: cdcl3
Temp. 20.0 C / 293.1 K
Operator: plnu
Mercury-400BB "pandora"

Relax. delay 1.000 sec
Pulse 45.0 degrees
Acq. time 1.998 sec
Width 6402.0 Hz
8 repetitions
OBSERVE H1, 400.1571342 MHz
DATA PROCESSING
Line broadening 0.5 Hz
FT size 65536
Total time 0 min, 25 sec



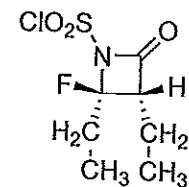
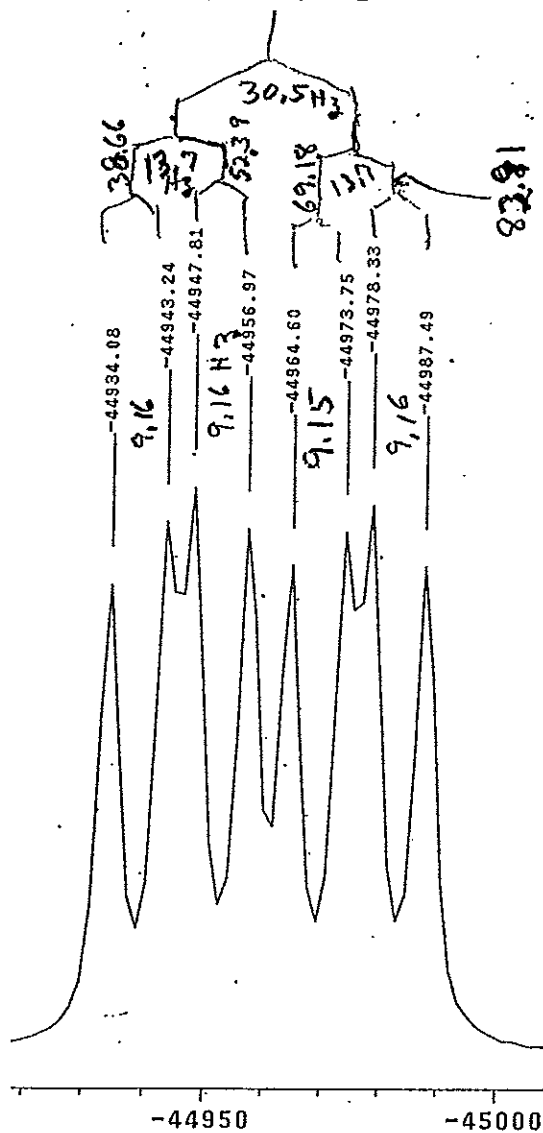
9 E
 ^1H 400 MHz
in CDCl_3



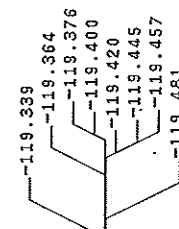
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 Sample id : tmpstudy
 Sample : KD-45_F

Pulse Sequence: s2pu1
 Solvent: cdcl3
 Temp. 20.0 C / 293.1 K
 Operator: plnu
 Mercury-400BB "pandora"

Relax delay 1.000 sec
 Pulse 68.7 degrees
 Acq. time 0.640 sec
 Width 100.0 kHz
 16 repetitions
 OBSERVE F19, 376.5240808 MHz
 DATA PROCESSING
 Line broadening 0.9 Hz
 Gauss apodization 0.462 sec
 FT size 131072
 Total time 0 min, 53 sec



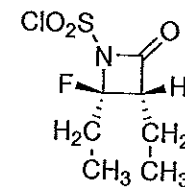
9 E
¹⁹F 376 MHz
 in CDCl₃



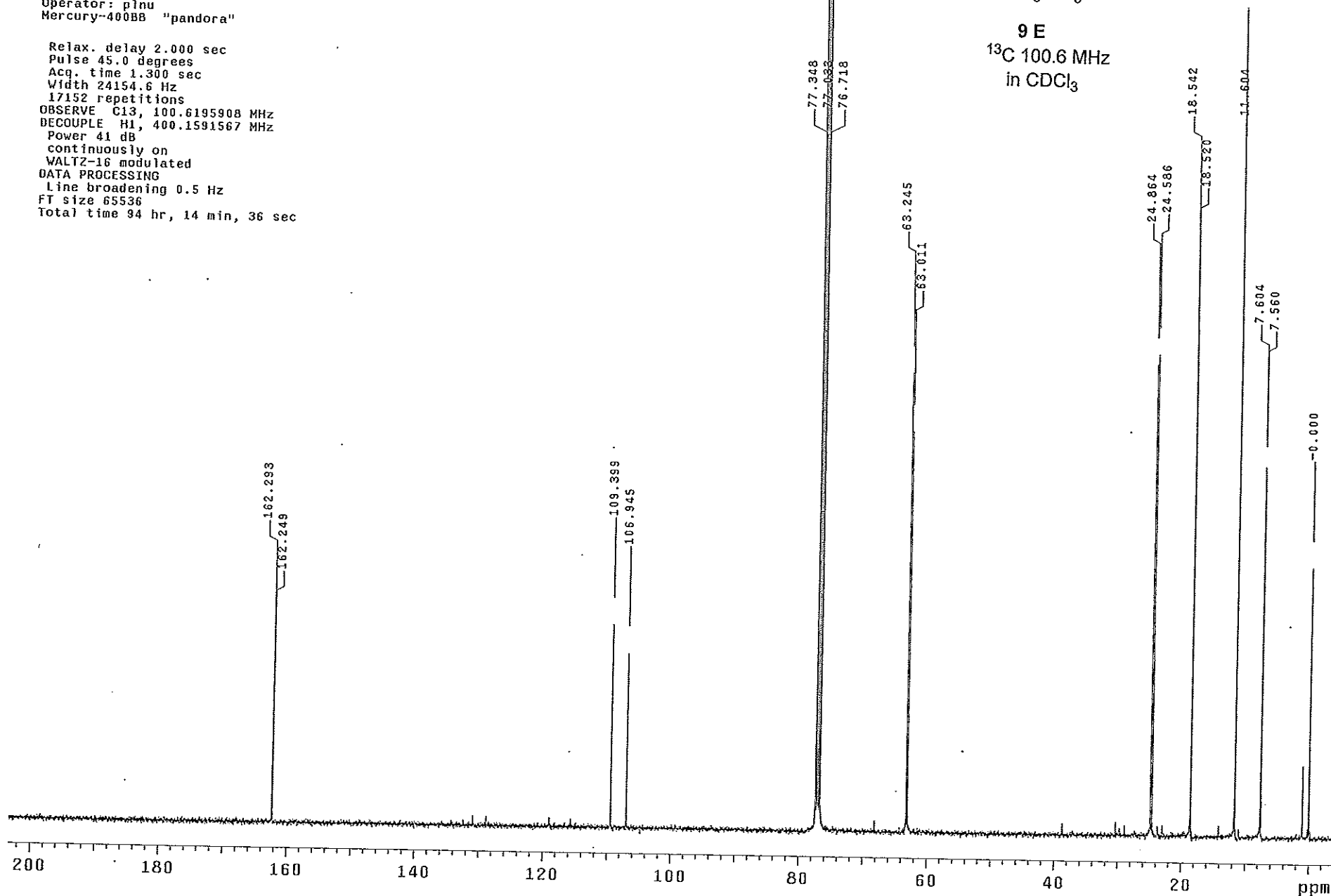
Automation directory: /home/organic/vnmrsys/data/studies/auto_2007.07.12
Sample id : tmpstudy
Sample : K0-45_C

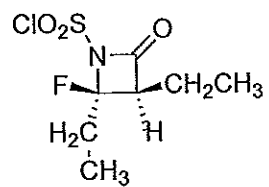
Pulse Sequence: s2pu1
Solvent: cdcl3
Temp. 20.0 C / 293.1 K
Operator: plnu
Mercury-400BB "pandora"

Relax. delay 2.000 sec
Pulse 45.0 degrees
Acq. time 1.300 sec
Width 24154.6 Hz
17152 repetitions
OBSERVE C13, 100.6195908 MHz
DECOUPLE H1, 400.1591567 MHz
Power 41 dB
continuously on
WALTZ-16 modulated
DATA PROCESSING
Line broadening 0.5 Hz
FT size 65536
Total time 94 hr, 14 min, 36 sec



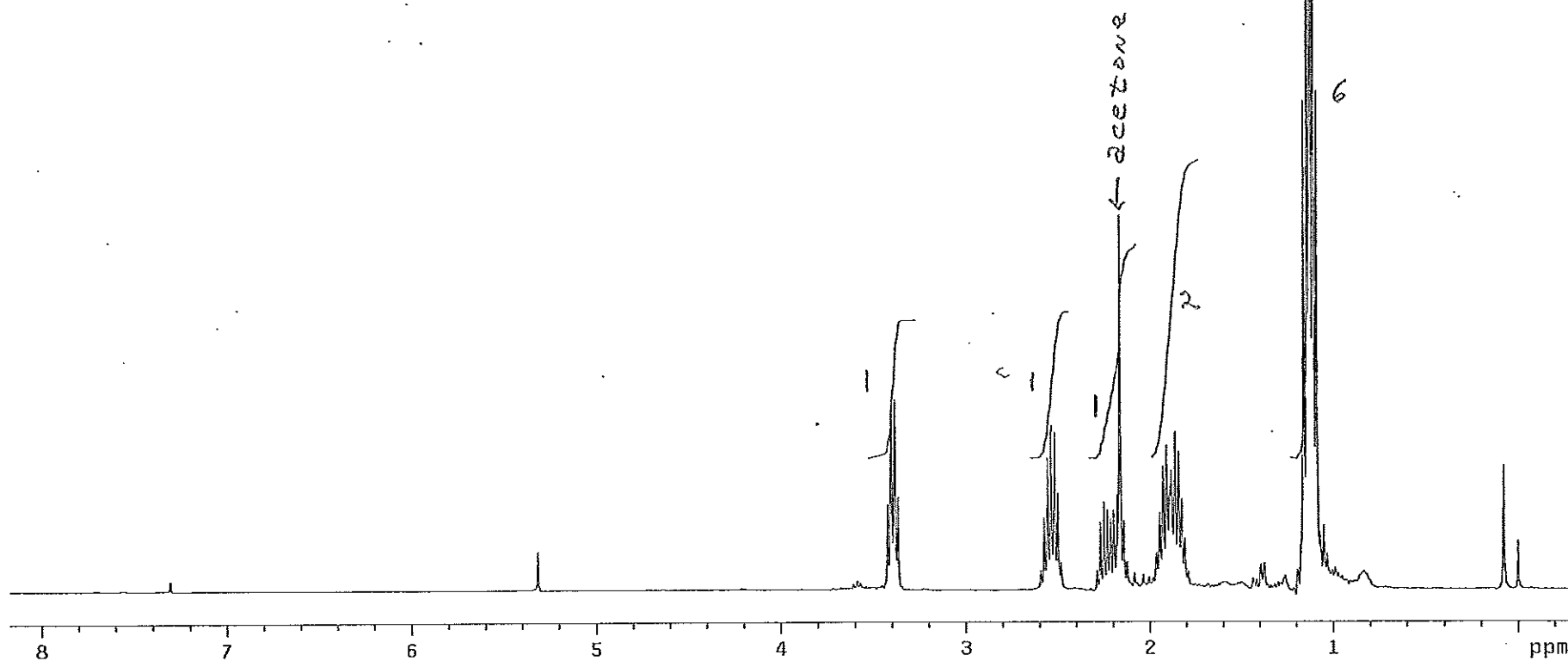
9 E
¹³C 100.6 MHz
in CDCl₃

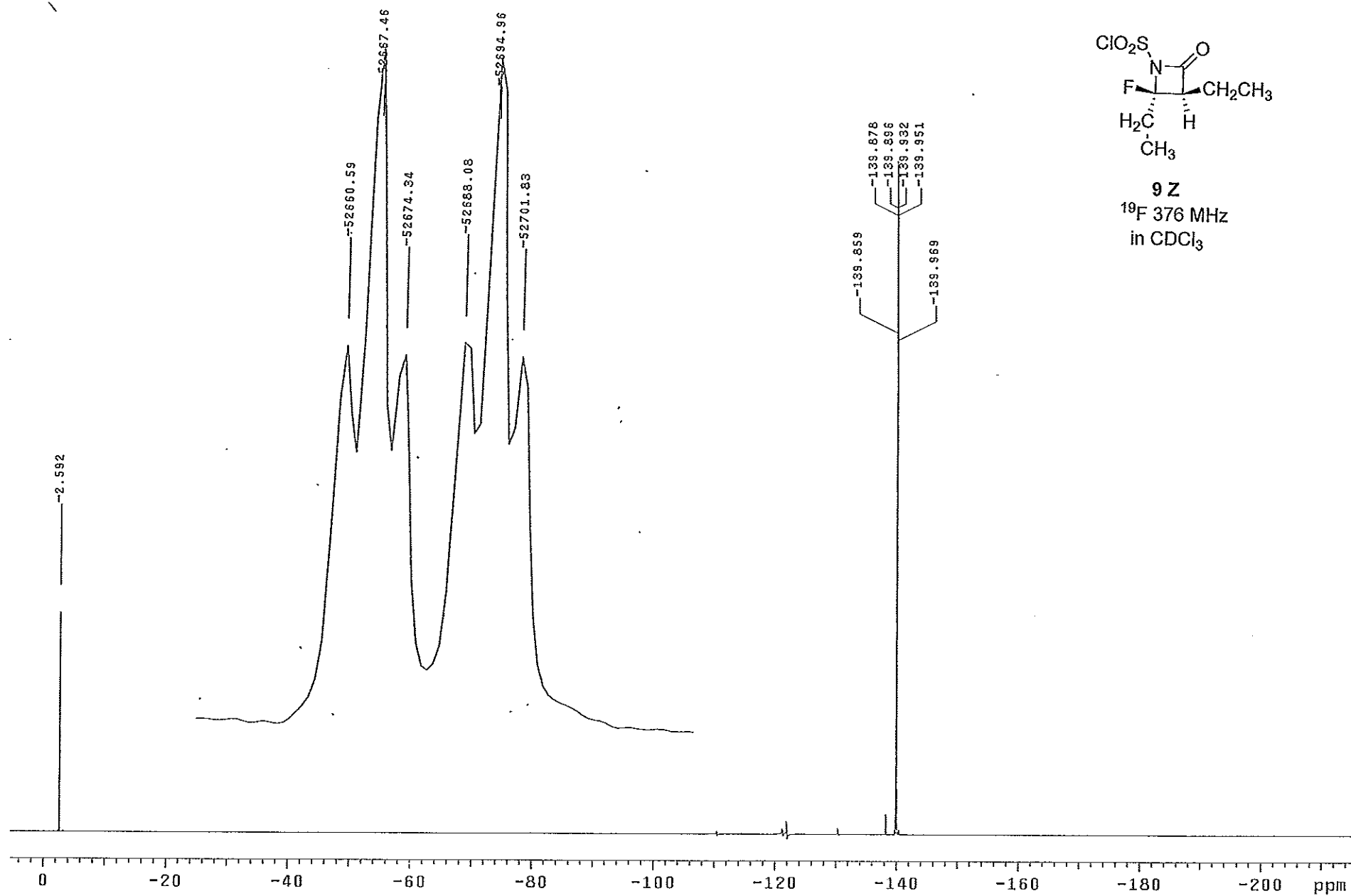




9Z

¹H 400 MHz
in CDCl₃





Automation directory: /home/organic/vnmrsys/data/studies/auto_2008.06.05
Sample id : /home/organic/vnmrsys/data/plnu/s_DMH27_C01
Sample : DMH27_C

Pulse Sequence: s2pul

Solvent: d2o

Ambient temperature

Operator: plnu

File: DMH27_C_Carbon_01

Mercury-400BB "pandora.scst.sandiego.edu"

Relax. delay 5.000 sec

Pulse 45.0 degrees

Acq. time 1.300 sec

Width 24154.6 Hz

64 repetitions

OBSERVE C13, 100.6195812 MHz

DECOUPLE H1, 400.1601851 MHz

Power 41 dB

continuously on

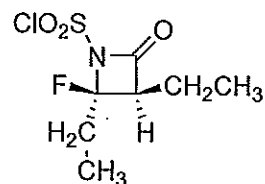
WALTZ-16 modulated

DATA PROCESSING

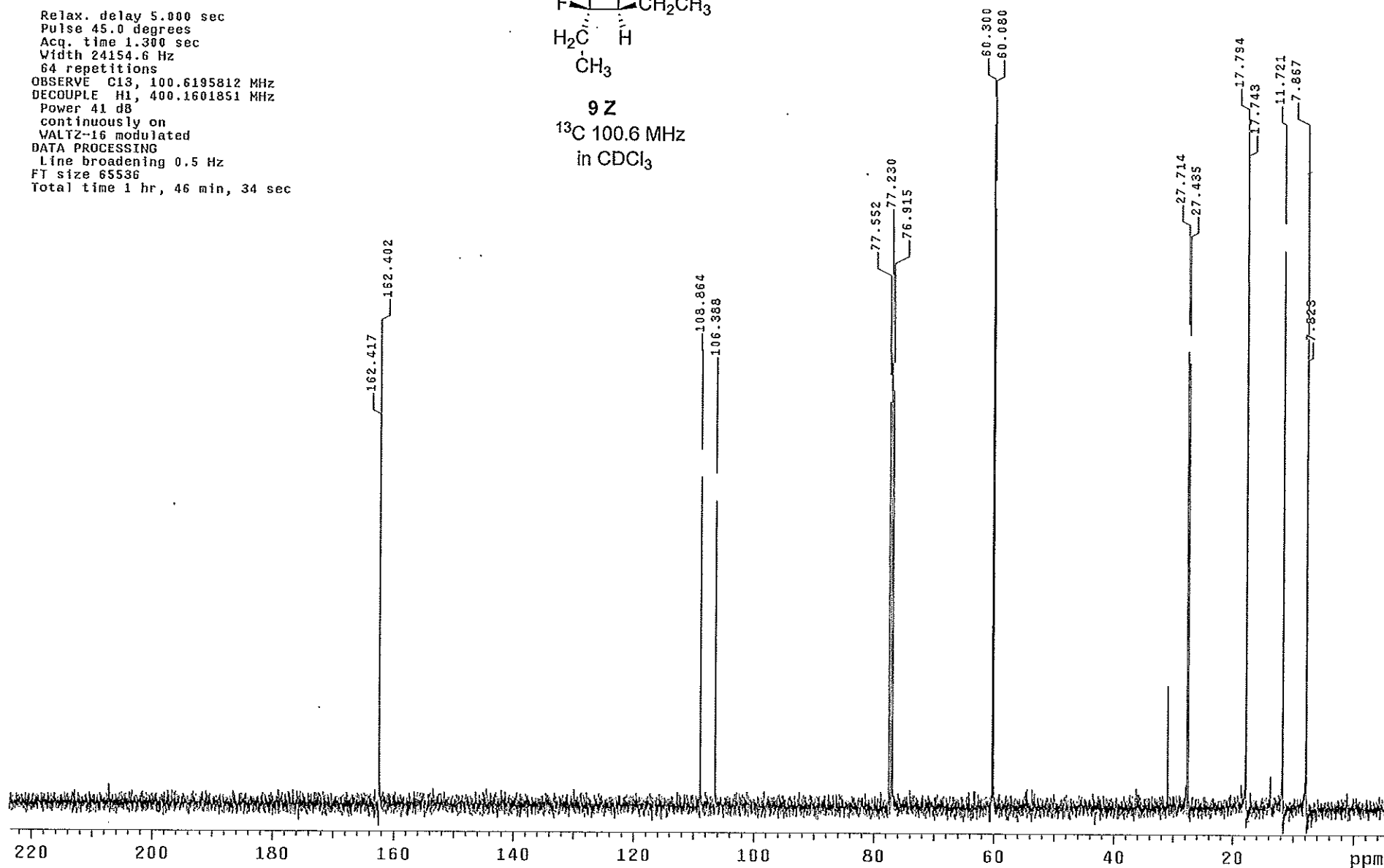
Line broadening 0.5 Hz

FT size 65536

Total time 1 hr, 46 min, 34 sec



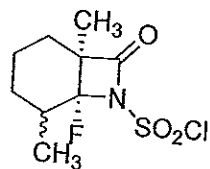
9 Z
¹³C 100.6 MHz
in CDCl₃



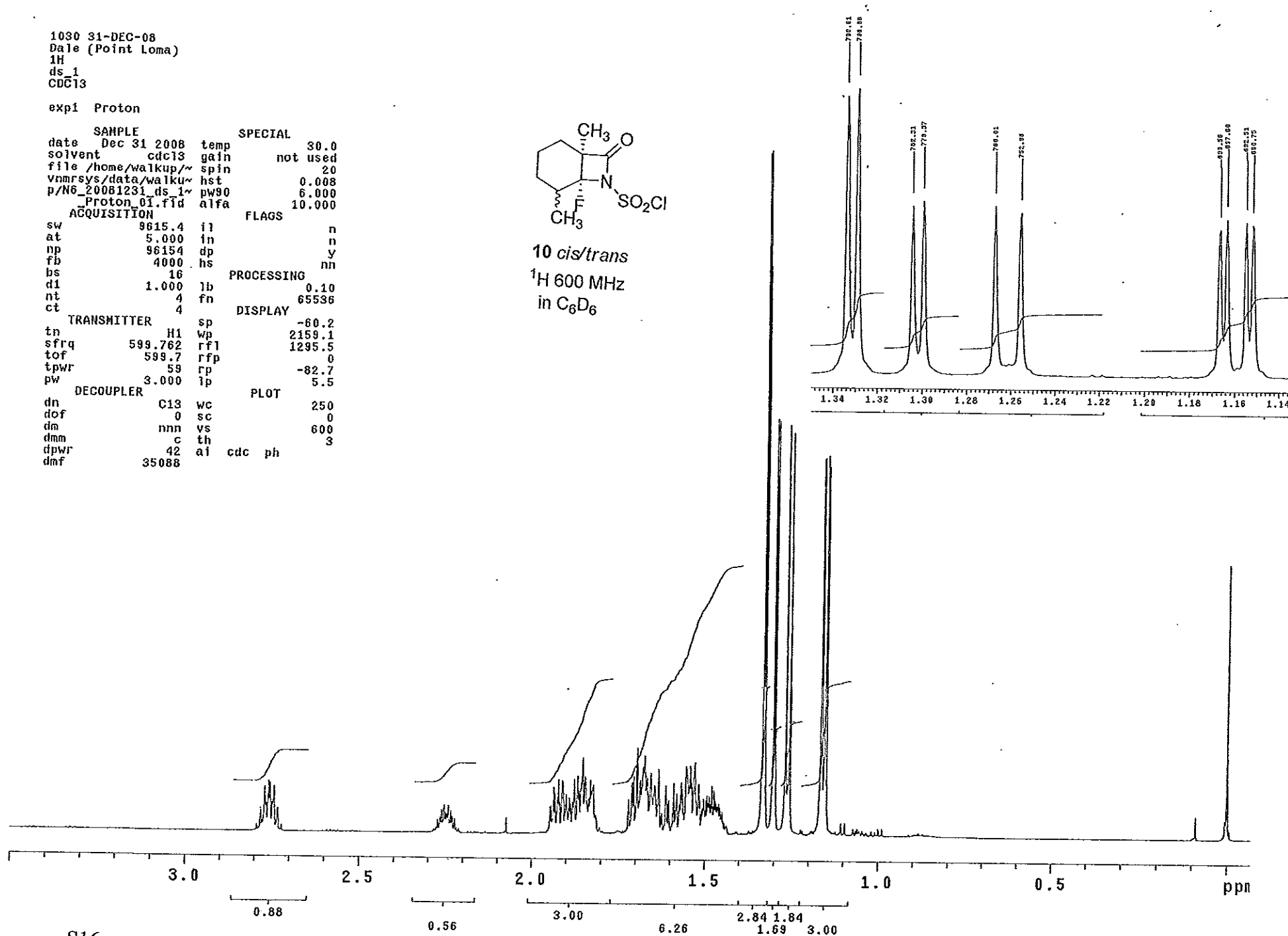
1030 31-DEC-08
 Dale (Point Loma)
 1H
 ds_1
 CDC13

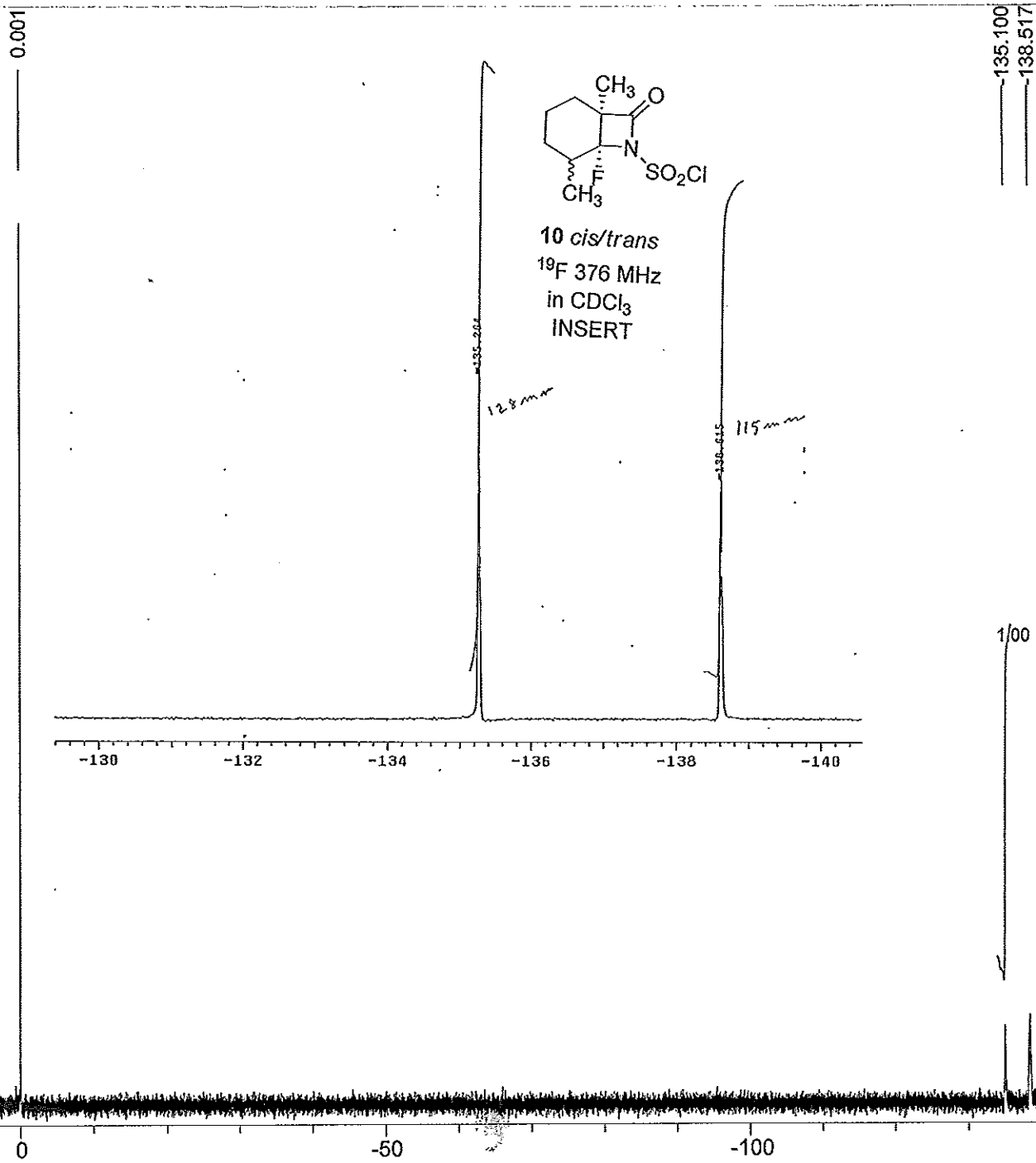
exp1 Proton

SAMPLE		SPECIAL	
date	Dec 31 2008	temp	30.0
solvent	cdc13	gain	not used
file	/home/walkup/~	spin	20
nmr	sys/data/walku~	hst	0.008
p/N6_20081231	ds_1	pw90	6.000
Proton_01.fid	alfa	10.000	
ACQUISITION		FLAGS	
sw	9615.4	il	n
at	5.000	in	n
np	96154	dp	y
fb	4000	hs	nn
bs	16	PROCESSING	
d1	1.000	lb	0.10
nt	4	fn	65536
ct	4	DISPLAY	
TRANSMITTER		sp	-60.2
tn	H1	wp	2159.1
sfrq	599.762	rfl	1295.5
tof	599.7	rff	0
tpwr	59	rp	-82.7
pw	3.000	lp	5.5
DECOUPLER		PLOT	
dn	C13	wc	250
dof	0	sc	0
dm	nnn	vs	600
dmm	c	th	3
dpwr	42	cdc	ph
dmf	35088		



10 cis/trans
¹H 600 MHz
 in C₆D₆





S17

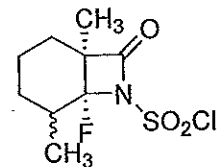
USER: Undefined -- DATE: 12/09/08 (11:47)

FI: 56.465	SW1: 59880	OF1: -12998.3	PTS1d: 262144
EX: zg.ppg	PW: 15.0 us	PD: 1.0 sec	NA: 8
		LB: 0.0	Nuts - temp

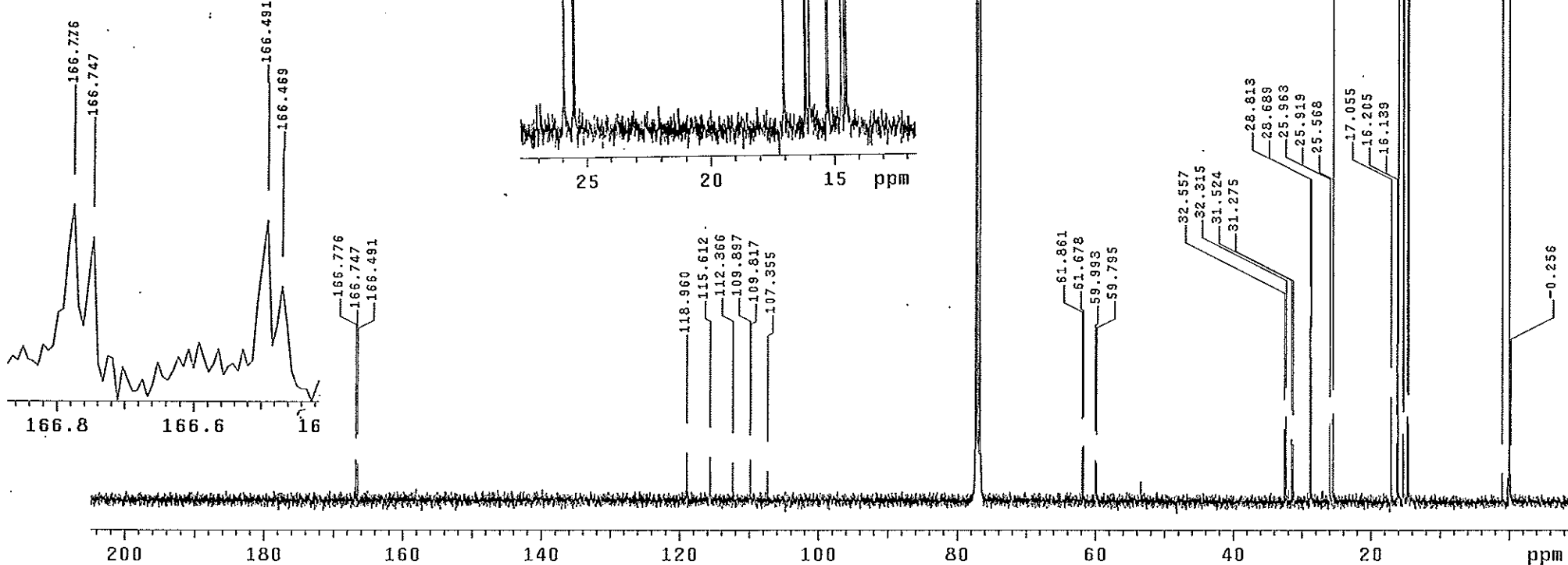
Automation directory: /home/organic/vnmrsys/data/studies/auto_2008.07.15
 Sample id : /home/organic/vnmrsys/data/plnu/s_KD27_C01
 Sample : KD27_C

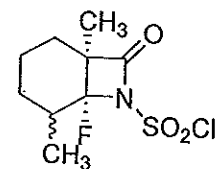
Pulse Sequence: s2pu1
 Solvent: cdc13
 Ambient temperature
 Operator: plnu
 File: KD27_C_Carbon_01
 Mercury-400BB "pandora.scst.sandiego.edu"

Relax. delay 10.000 sec
 Pulse 45.0 degrees
 Acq. time 1.300 sec
 Width 24154.6 Hz
 3824 repetitions
 OBSERVE C13, 100.6195908 MHz
 DECOUPLE H1, 400.1591567 MHz
 Power 41 dB
 continuously on
 WALTZ-16 modulated
 DATA PROCESSING
 Line broadening 0.5 Hz
 FT size 65536
 Total time 16 hr, 4 min, 46 sec



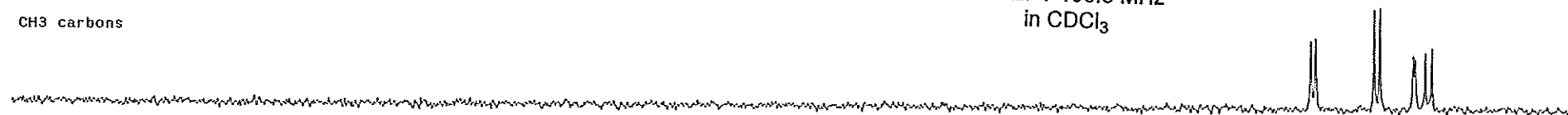
10 cis/trans
¹³C 100.6 MHz
 in CDCl₃



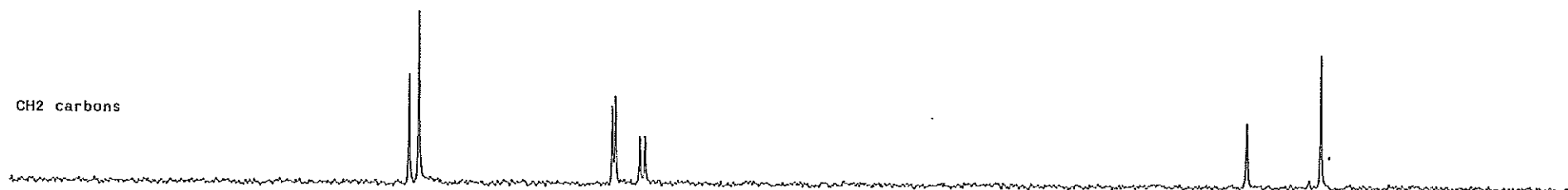


10 *cis/trans*
DEPT 150.8 MHz
in CDCl₃

CH₃ carbons



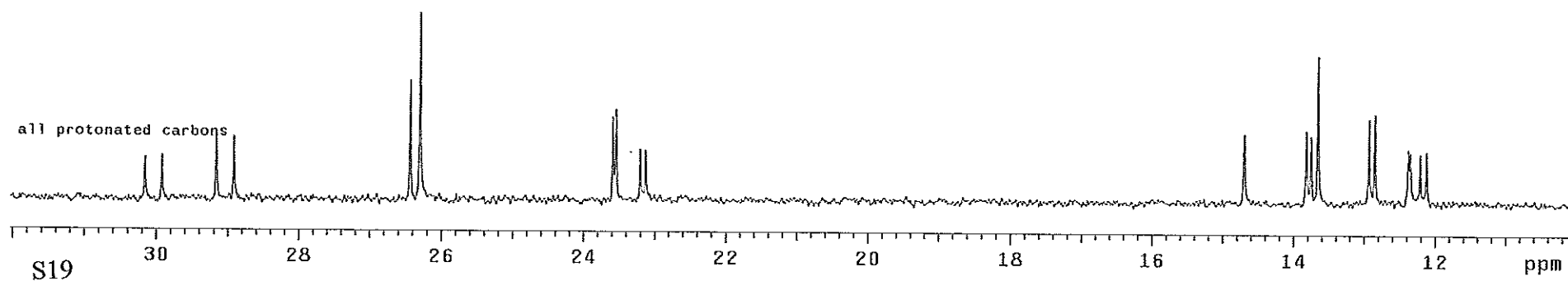
CH₂ carbons



CH carbons

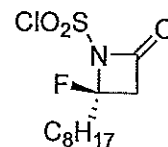


all protonated carbons

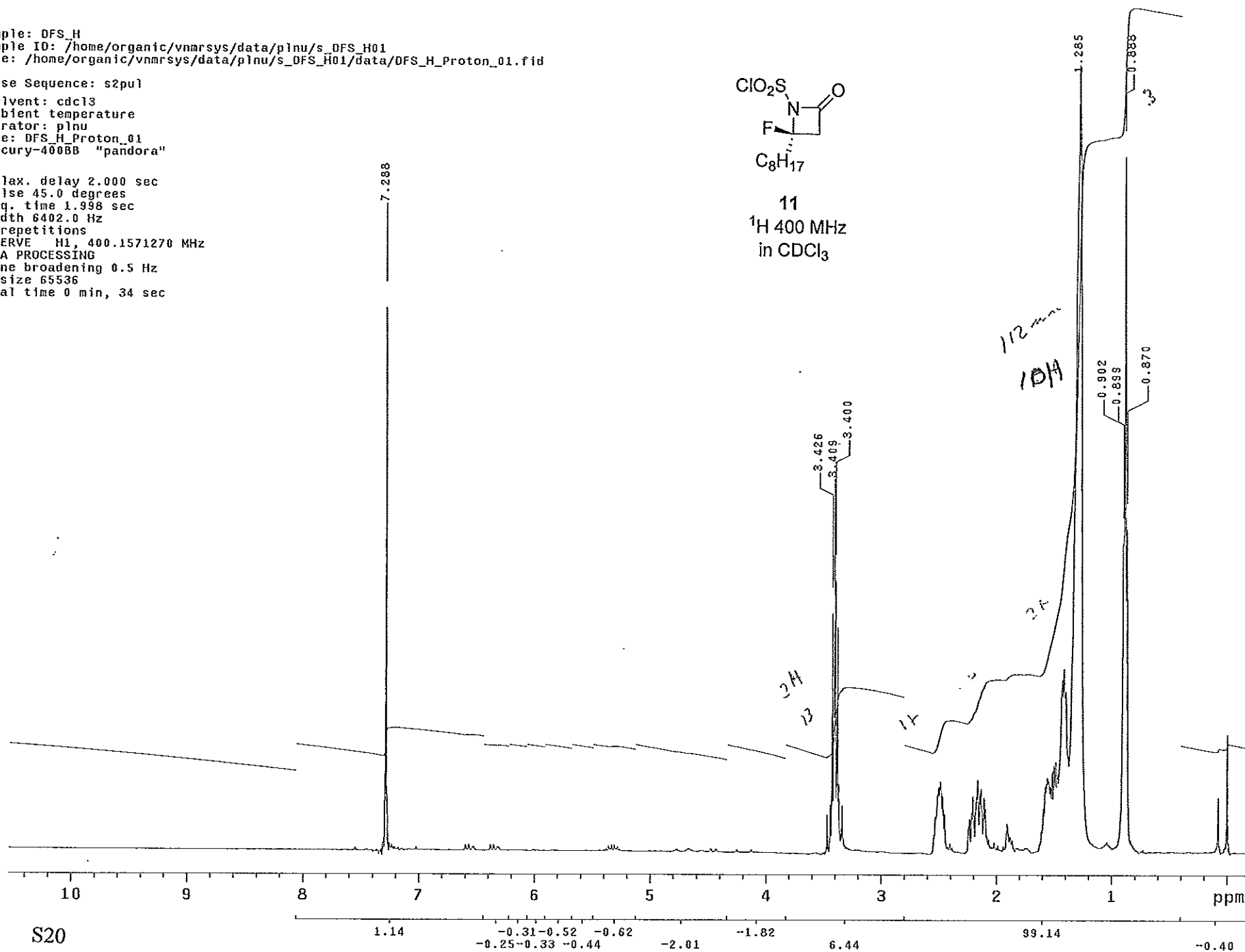



```
Pulse Sequence: s2pu1
Solvent: cdcl3
Ambient temperature
Operator: plnu
File: DFS_H_Proton_01
Mercury-400BB "pandora"
```

```
Relax. delay 2.000 sec
Pulse 45.0 degrees
Acq. time 1.998 sec
Width 6402.0 Hz
8 repetitions
OBSERVE H1, 400.1571270 MHz
DATA PROCESSING
Line broadening 0.5 Hz
FT size 65536
Total time 0 min, 34 sec
```



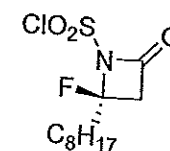
11
¹H 400 MHz
 in CDCl₃



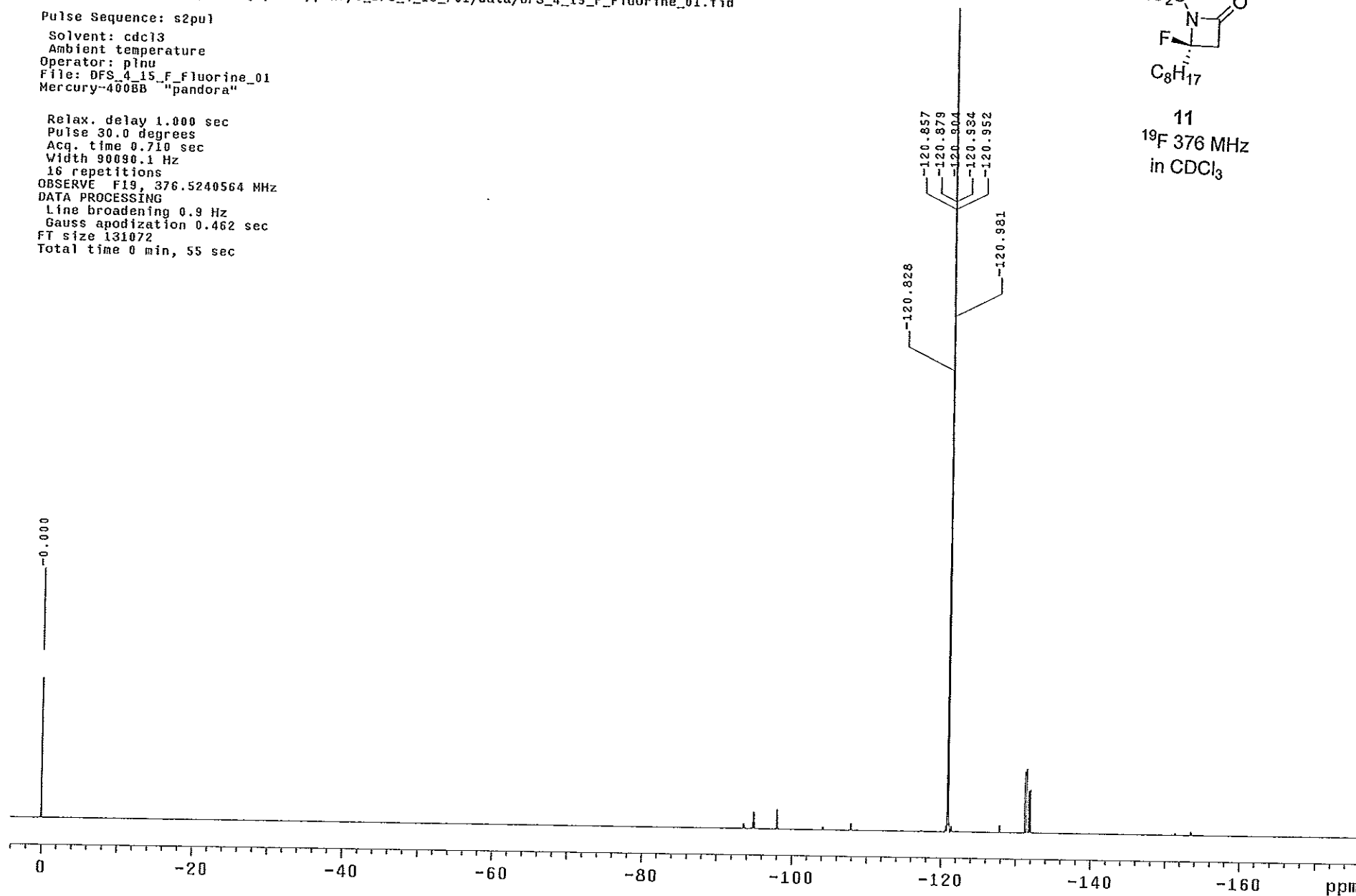
Sample: DFS_4_15_F
Sample ID: /home/organic/vnmrsys/data/plnu/s_DFS_4_15_F01
File: /home/organic/vnmrsys/data/plnu/s_DFS_4_15_F01/data/DFS_4_15_F_Fluorine_01.fid

Pulse Sequence: s2pu1
Solvent: cdc13
Ambient temperature
Operator: plnu
File: DFS_4_15_F_Fluorine_01
Mercury-400BB "pandora"

Relax. delay 1.000 sec
Pulse 30.0 degrees
Acq. time 0.710 sec
Width 90090.1 Hz
16 repetitions
OBSERVE F19, 376.5240564 MHz
DATA PROCESSING
Line broadening 0.9 Hz
Gauss apodization 0.462 sec
FT size 131072
Total time 0 min, 55 sec



11
 ^{19}F 376 MHz
in CDCl_3

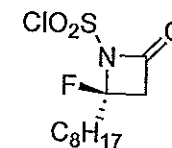


S21

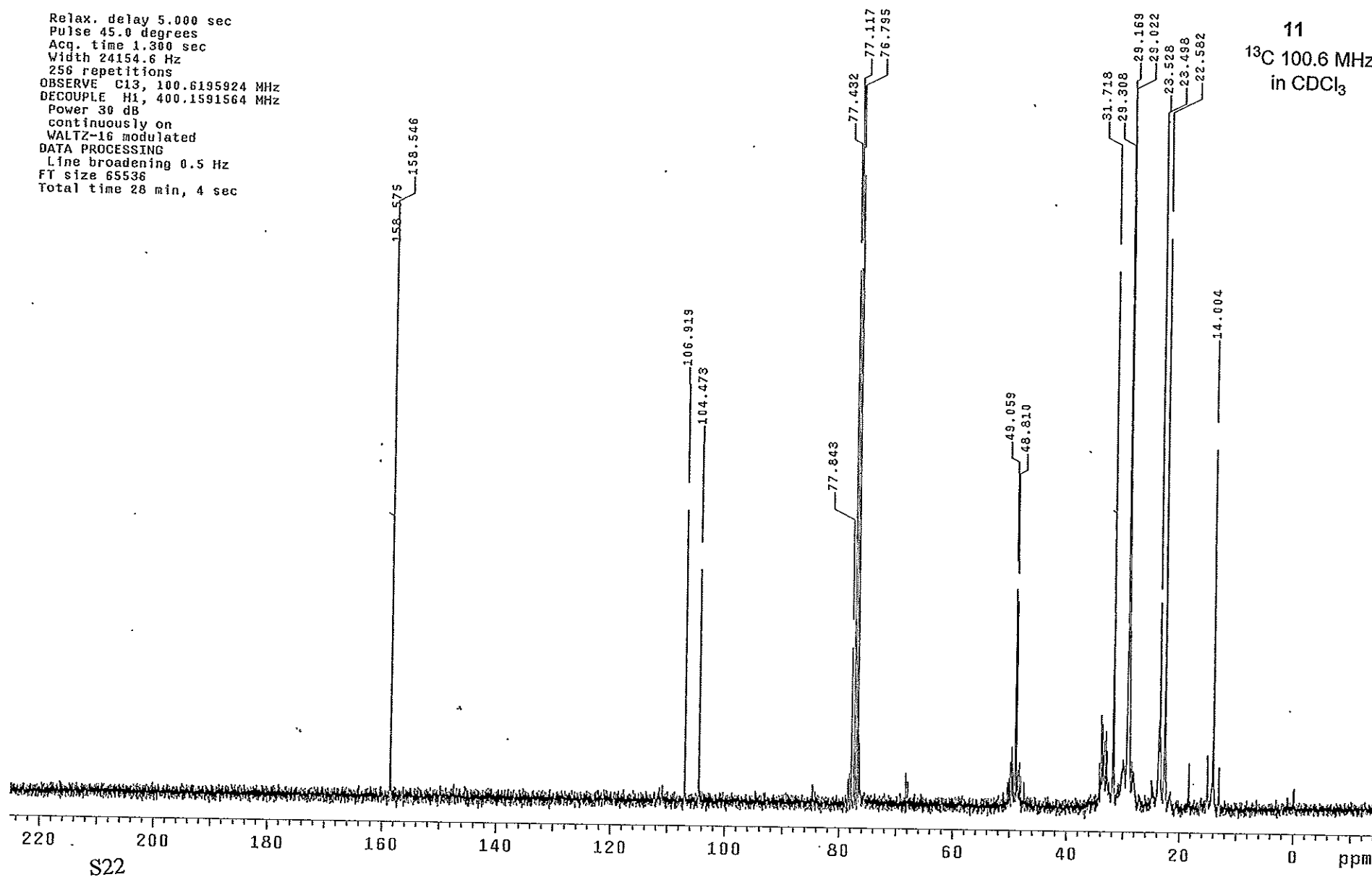
Sample: DFS_4_15_C
 Sample ID: /home/organic/vnmrsys/data/plnu/s_DFS_4_15_C01
 File: /home/organic/vnmrsys/data/plnu/s_DFS_4_15_C01/data/DFS_4_15_C_Carbon_01.fid

Pulse Sequence: s2pu1
 Solvent: cdc13
 Ambient temperature
 Operator: plnu
 File: DFS_4_15_C_Carbon_01
 Mercury-400BB "pandora"

Relax. delay 5.000 sec
 Pulse 45.0 degrees
 Acq. time 1.300 sec
 Width 24154.6 Hz
 256 repetitions
 OBSERVE C13, 100.6195924 MHz
 DECOUPLE H1, 400.1591564 MHz
 Power 30 dB
 continuously on
 WALTZ-16 modulated
 DATA PROCESSING
 Line broadening 0.5 Hz
 FT size 65536
 Total time 28 min, 4 sec



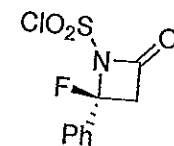
11
¹³C 100.6 MHz
 in CDCl₃



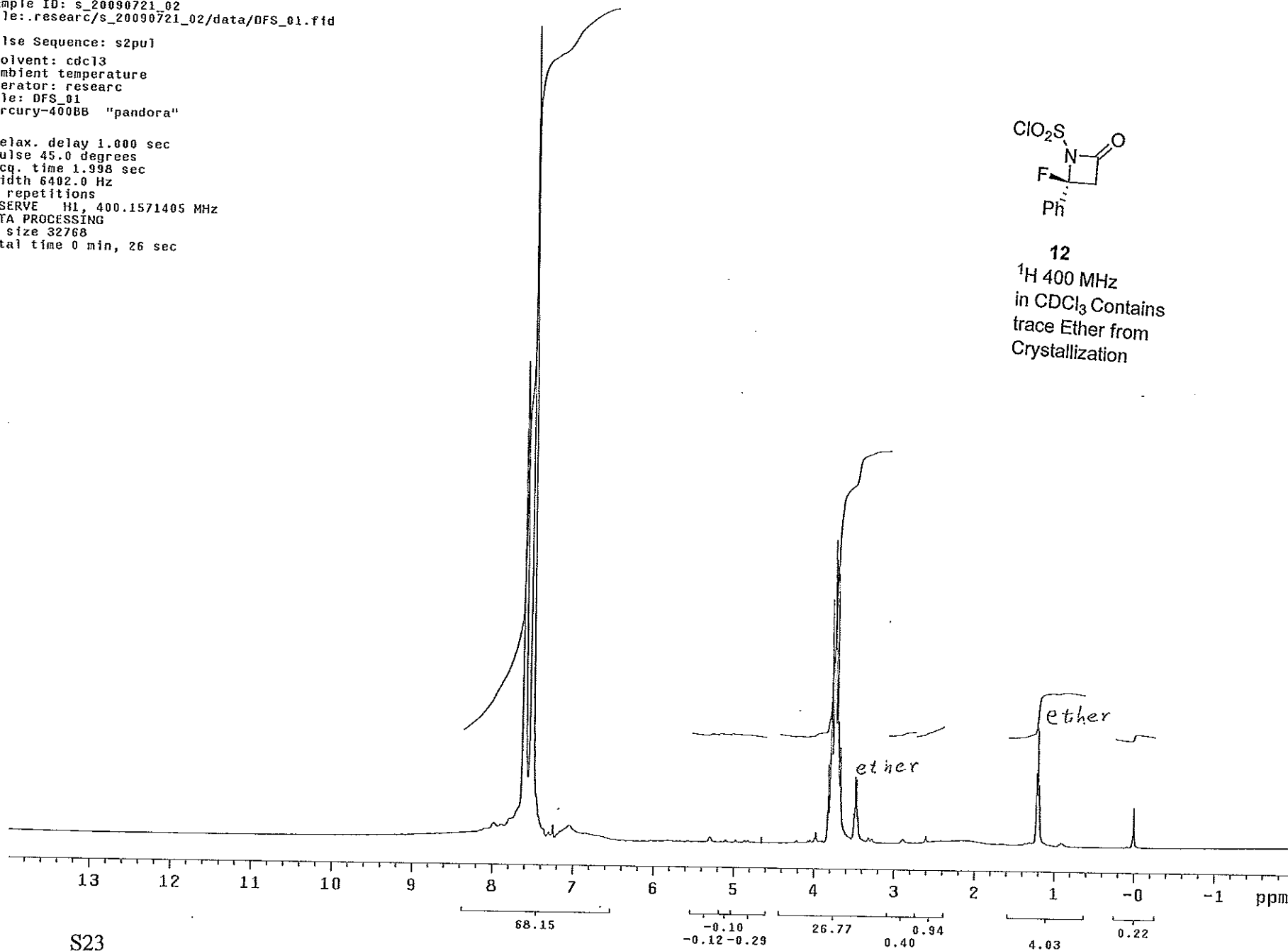
Sample: DFS
Sample ID: s_20090721_02
File: .research/s_20090721_02/data/DFS_01.fid

Pulse Sequence: s2pu1
Solvent: cdcl3
Ambient temperature
Operator: research
File: DFS_01
Mercury-400BB "pandora"

Relax. delay 1.000 sec
Pulse 45.0 degrees
Acq. time 1.998 sec
Width 6402.0 Hz
8 repetitions
OBSERVE H1, 400.1571405 MHz
DATA PROCESSING
FT size 32768
Total time 0 min, 26 sec

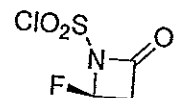
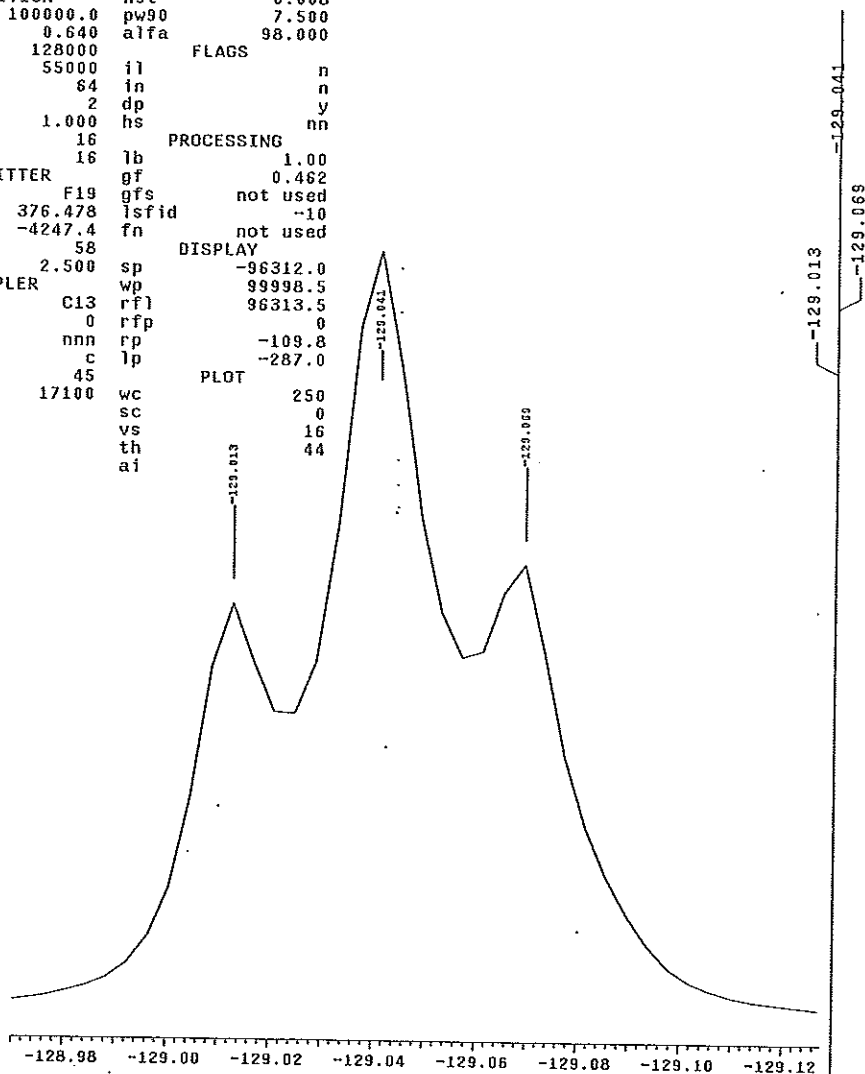


12
¹H 400 MHz
in CDCl₃ Contains
trace Ether from
Crystallization



expl Fluorine

SAMPLE		SPECIAL	
date	Jul 30 2009	temp	not used
solvent	cdcl3	gain	not used
file	exp	spin	not used
ACQUISITION		hst	0.008
sw	100000.0	pw90	7.500
at	0.640	alfa	98.000
np	128000	FLAGS	
fb	55000	il	n
bs	64	in	n
ss	2	dp	y
d1	1.000	hs	nn
nt	16	PROCESSING	
ct	16	lb	1.00
tn	F19	gf	0.462
sfrq	376.478	gfs	not used
tof	-4247.4	lsfid	-10
tpwr	58	fn	not used
pw	2.500	DISPLAY	
dn	C13	sp	-96312.0
dot	0	wp	99998.5
dm	nnn	rfl	96313.5
dmm	c	rflp	0
dpwr	45	rp	-109.8
dmt	17100	lp	-287.0
		PLOT	
		wc	250
		sc	0
		vs	16
		th	44
		ai	

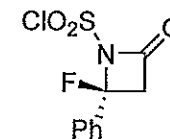


12
 ^{19}F 376 MHz
in CDCl_3

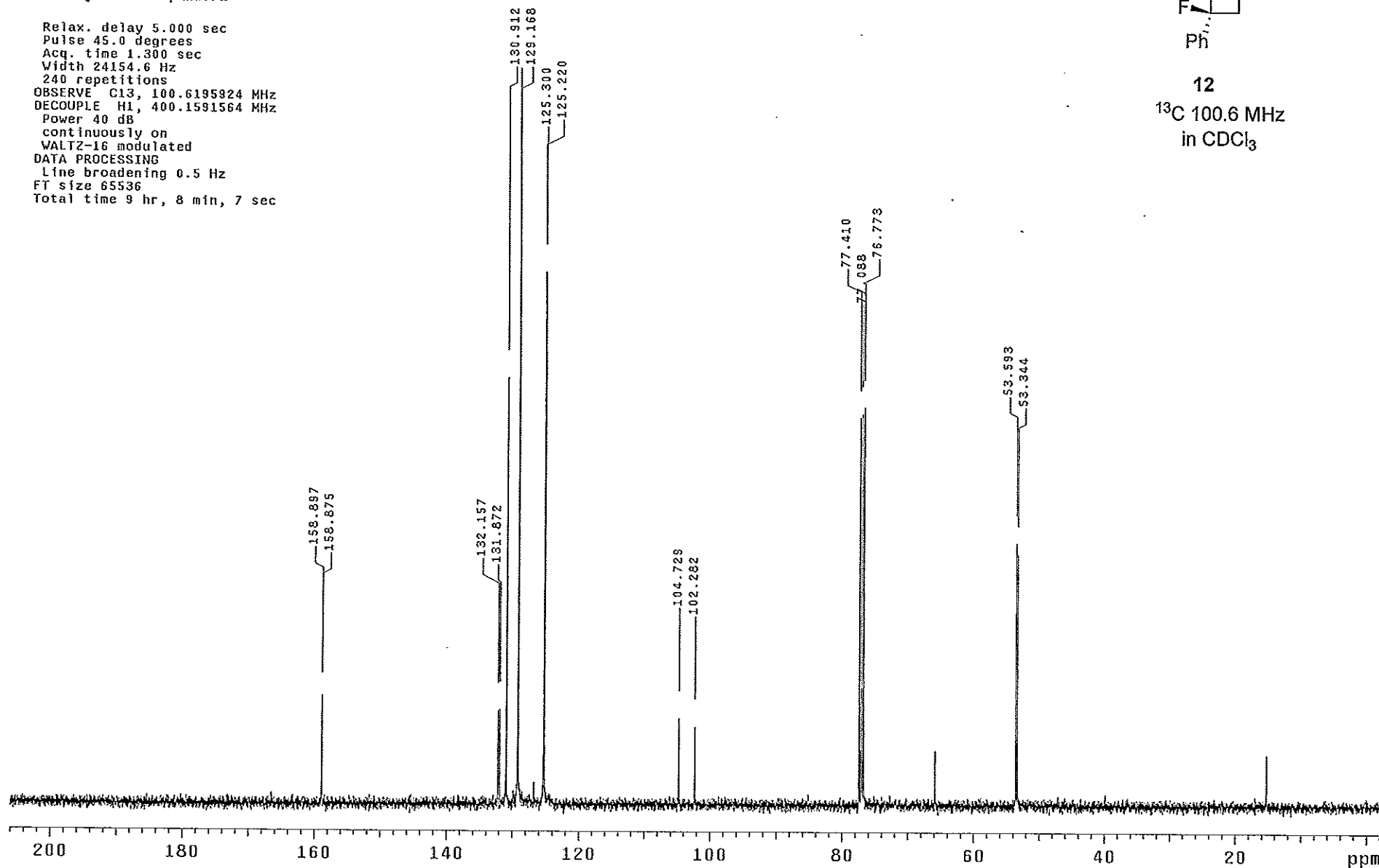
Sample: DFS
Sample ID: s_20090721_01
File: research/s_20090721_01/data/DFS_01.fid

Pulse Sequence: s2pu1
Solvent: cdcl3
Ambient temperature
Operator: research
File: DFS_01
Mercury-400BB "pandora"

Relax. delay 5.000 sec
Pulse 45.0 degrees
Acq. time 1.300 sec
Width 24154.6 Hz
240 repetitions
OBSERVE C13, 100.6195924 MHz
DECOUPLE H1, 400.1591564 MHz
Power 40 dB
continuously on
WALTZ-16 modulated
DATA PROCESSING
Line broadening 0.5 Hz
FT size 65536
Total time 9 hr, 8 min, 7 sec



12
¹³C 100.6 MHz
in CDCl₃



S25

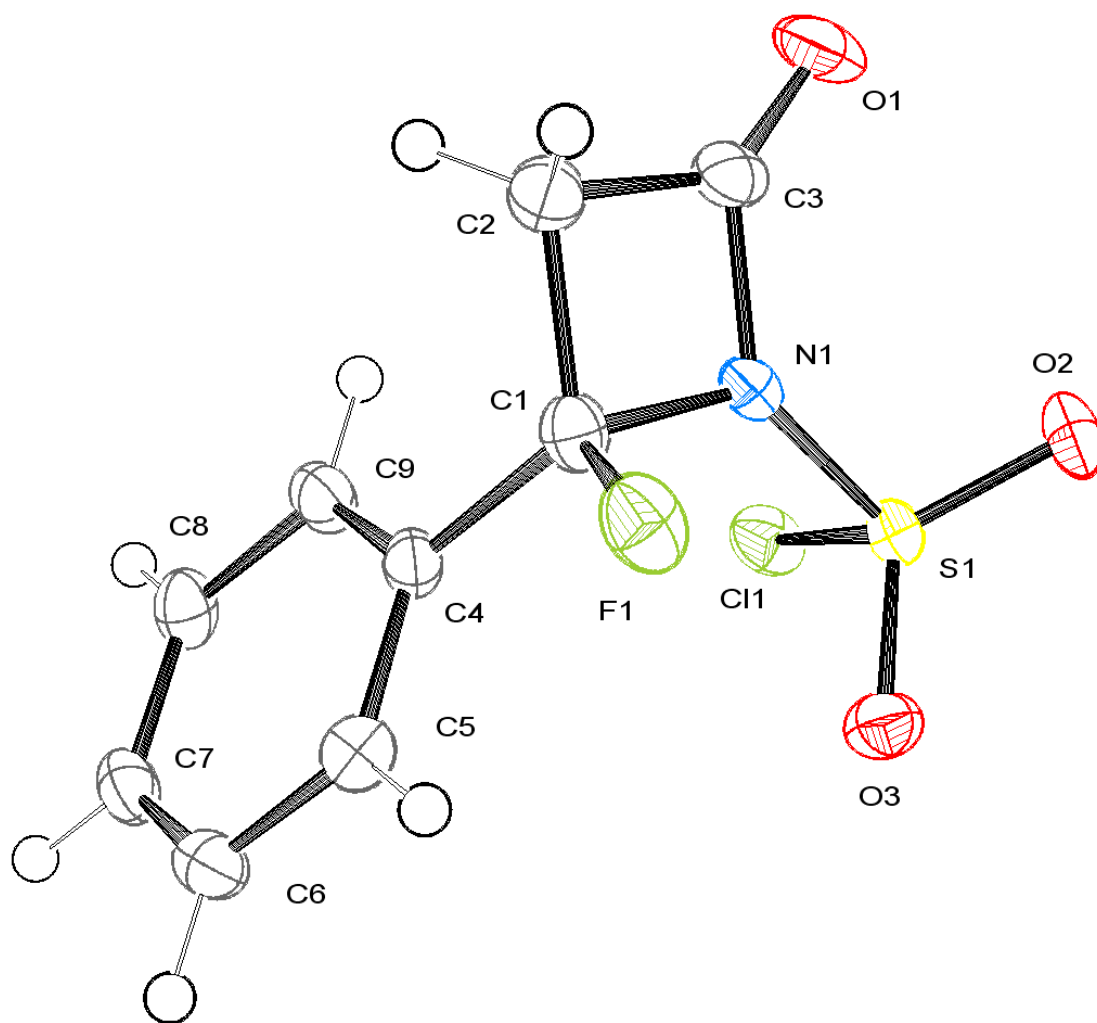


Table 1. Crystal data and structure refinement for plnu05.

Identification code	plnu05	
Empirical formula	C ₉ H ₇ Cl F N O ₃ S	
Formula weight	263.67	
Temperature	120(2) K	
Wavelength	1.54178 Å	
Crystal system	Monoclinic	
Space group	P2(1)/n	
Unit cell dimensions	a = 13.4185(5) Å	α = 90°.
	b = 5.6716(3) Å	β = 98.008(3)°.
	c = 13.7167(6) Å	γ = 90°.
Volume	1033.72(8) Å ³	
Z	4	
Density (calculated)	1.694 Mg/m ³	
Absorption coefficient	5.265 mm ⁻¹	
F(000)	536	
Crystal size	0.25 x 0.17 x 0.11 mm ³	
Crystal color, habit	Colorless Rod	
Theta range for data collection	4.97 to 65.54°.	
Index ranges	-14 ≤ h ≤ 15, -6 ≤ k ≤ 6, -15 ≤ l ≤ 16	
Reflections collected	5633	
Independent reflections	1710 [R(int) = 0.0248]	
Completeness to theta = 65.00°	96.9 %	
Absorption correction	Multi-scan	
Max. and min. transmission	0.5951 and 0.3528	
Refinement method	Full-matrix least-squares on F ²	
Data / restraints / parameters	1710 / 0 / 146	
Goodness-of-fit on F ²	1.042	
Final R indices [I > 2σ(I)]	R1 = 0.0287, wR2 = 0.0731	
R indices (all data)	R1 = 0.0319, wR2 = 0.0748	
Extinction coefficient	0.0010(3)	
Largest diff. peak and hole	0.261 and -0.312 e.Å ⁻³	

Table 2. Atomic coordinates ($\times 10^4$) and equivalent isotropic displacement parameters ($\text{\AA}^2 \times 10^3$) for plnu05. $U(\text{eq})$ is defined as one third of the trace of the orthogonalized U^{ij} tensor.

	x	y	z	$U(\text{eq})$
C(1)	4343(1)	3211(3)	6656(1)	22(1)
C(2)	3613(1)	5327(4)	6491(2)	27(1)
C(3)	3488(1)	5188(4)	7570(2)	24(1)
C(4)	5422(1)	3414(3)	6474(1)	19(1)
C(5)	5857(1)	1624(3)	5987(1)	22(1)
C(6)	6844(1)	1838(3)	5806(1)	24(1)
C(7)	7399(1)	3811(4)	6112(1)	24(1)
C(8)	6970(2)	5586(4)	6610(2)	25(1)
C(9)	5983(1)	5391(4)	6790(1)	24(1)
Cl(1)	5793(1)	3521(1)	9275(1)	27(1)
F(1)	3920(1)	1210(2)	6204(1)	30(1)
N(1)	4163(1)	3233(3)	7699(1)	21(1)
O(1)	3060(1)	6206(3)	8141(1)	33(1)
O(2)	3915(1)	1555(2)	9306(1)	28(1)
O(3)	5020(1)	-445(2)	8254(1)	26(1)
S(1)	4620(1)	1614(1)	8632(1)	20(1)

Table 3. Bond lengths [Å] and angles [°] for plnu05.

C(1)-F(1)	1.378(2)	C(3)-C(2)-H(2B)	113.9
C(1)-N(1)	1.483(2)	C(1)-C(2)-H(2B)	113.9
C(1)-C(4)	1.508(3)	H(2A)-C(2)-H(2B)	111.1
C(1)-C(2)	1.547(3)	O(1)-C(3)-N(1)	130.84(19)
C(2)-C(3)	1.514(3)	O(1)-C(3)-C(2)	138.70(19)
C(2)-H(2A)	0.9900	N(1)-C(3)-C(2)	90.43(14)
C(2)-H(2B)	0.9900	C(9)-C(4)-C(5)	119.77(17)
C(3)-O(1)	1.185(2)	C(9)-C(4)-C(1)	120.10(16)
C(3)-N(1)	1.427(2)	C(5)-C(4)-C(1)	120.13(17)
C(4)-C(9)	1.386(3)	C(4)-C(5)-C(6)	119.89(18)
C(4)-C(5)	1.387(3)	C(4)-C(5)-H(5)	120.1
C(5)-C(6)	1.388(3)	C(6)-C(5)-H(5)	120.1
C(5)-H(5)	0.9500	C(7)-C(6)-C(5)	120.39(18)
C(6)-C(7)	1.377(3)	C(7)-C(6)-H(6)	119.8
C(6)-H(6)	0.9500	C(5)-C(6)-H(6)	119.8
C(7)-C(8)	1.386(3)	C(6)-C(7)-C(8)	119.77(17)
C(7)-H(7)	0.9500	C(6)-C(7)-H(7)	120.1
C(8)-C(9)	1.386(3)	C(8)-C(7)-H(7)	120.1
C(8)-H(8)	0.9500	C(9)-C(8)-C(7)	120.22(19)
C(9)-H(9)	0.9500	C(9)-C(8)-H(8)	119.9
Cl(1)-S(1)	2.0087(6)	C(7)-C(8)-H(8)	119.9
N(1)-S(1)	1.6247(16)	C(8)-C(9)-C(4)	119.96(18)
O(2)-S(1)	1.4128(14)	C(8)-C(9)-H(9)	120.0
O(3)-S(1)	1.4137(14)	C(4)-C(9)-H(9)	120.0
		C(3)-N(1)-C(1)	94.16(14)
F(1)-C(1)-N(1)	109.23(15)	C(3)-N(1)-S(1)	134.47(13)
F(1)-C(1)-C(4)	109.44(15)	C(1)-N(1)-S(1)	131.35(13)
N(1)-C(1)-C(4)	116.66(16)	O(2)-S(1)-O(3)	122.92(9)
F(1)-C(1)-C(2)	111.11(16)	O(2)-S(1)-N(1)	108.39(8)
N(1)-C(1)-C(2)	87.13(14)	O(3)-S(1)-N(1)	107.40(8)
C(4)-C(1)-C(2)	121.36(16)	O(2)-S(1)-Cl(1)	106.32(6)
C(3)-C(2)-C(1)	88.28(15)	O(3)-S(1)-Cl(1)	106.94(6)
C(3)-C(2)-H(2A)	113.9	N(1)-S(1)-Cl(1)	103.18(6)
C(1)-C(2)-H(2A)	113.9		

Table 4. Anisotropic displacement parameters ($\text{\AA}^2 \times 10^3$) for plnu05. The anisotropic displacement factor exponent takes the form: $-2\pi^2 [h^2 a^{*2} U^{11} + \dots + 2 h k a^* b^* U^{12}]$

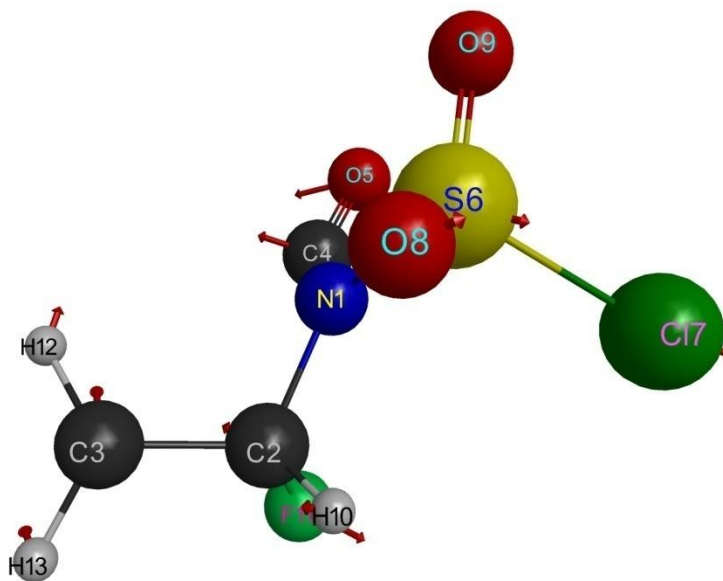
	U ¹¹	U ²²	U ³³	U ²³	U ¹³	U ¹²
C(1)	21(1)	23(1)	22(1)	-1(1)	4(1)	-4(1)
C(2)	20(1)	30(1)	31(1)	7(1)	3(1)	2(1)
C(3)	15(1)	22(1)	35(1)	0(1)	4(1)	0(1)
C(4)	19(1)	22(1)	18(1)	3(1)	5(1)	0(1)
C(5)	22(1)	22(1)	21(1)	0(1)	0(1)	-2(1)
C(6)	22(1)	27(1)	24(1)	-1(1)	5(1)	4(1)
C(7)	18(1)	32(1)	22(1)	4(1)	6(1)	0(1)
C(8)	25(1)	25(1)	26(1)	-1(1)	7(1)	-8(1)
C(9)	26(1)	23(1)	25(1)	-3(1)	10(1)	-1(1)
Cl(1)	18(1)	30(1)	34(1)	-5(1)	2(1)	-3(1)
F(1)	24(1)	32(1)	33(1)	-8(1)	7(1)	-9(1)
N(1)	18(1)	24(1)	22(1)	1(1)	8(1)	3(1)
O(1)	23(1)	34(1)	44(1)	-6(1)	11(1)	8(1)
O(2)	22(1)	38(1)	25(1)	2(1)	10(1)	-4(1)
O(3)	26(1)	20(1)	32(1)	0(1)	4(1)	2(1)
S(1)	16(1)	21(1)	22(1)	1(1)	6(1)	-1(1)

Table 5. Hydrogen coordinates ($\times 10^4$) and isotropic displacement parameters ($\text{\AA}^2 \times 10^{-3}$) for plnu05.

	x	y	z	U(eq)
H(2A)	3932	6804	6305	32
H(2B)	2989	4990	6037	32
H(5)	5478	255	5778	27
H(6)	7140	617	5468	29
H(7)	8074	3955	5982	29
H(8)	7354	6941	6828	30
H(9)	5690	6612	7131	29

Quantum Chemical Data for CSI and Vinyl Fluoride

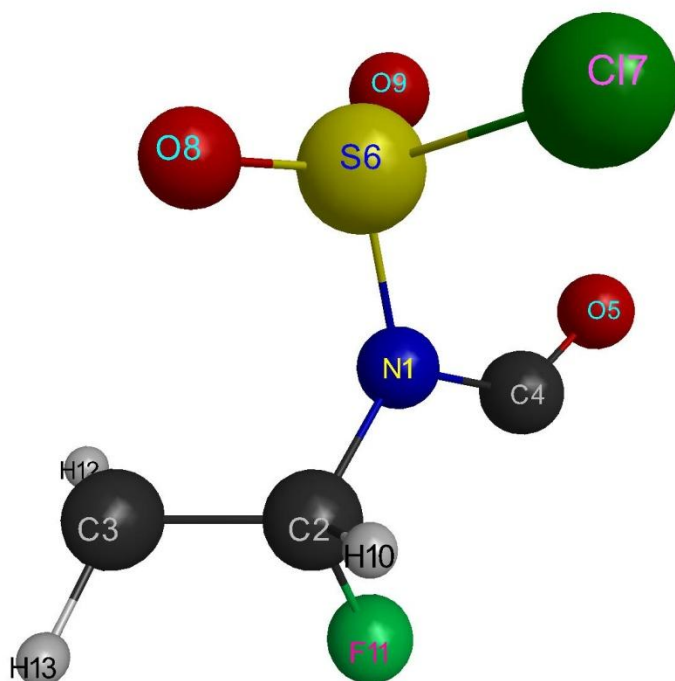
Structure 1. Stepwise transition state geometry and normal mode corresponding to the single imaginary frequency of $119.2i \text{ cm}^{-1}$.



Cartesian coordinates (in angstroms)

N	0.665493	0.418156	-0.179637
C	1.506227	0.451207	1.051268
C	2.816076	-0.226468	0.891209
C	0.878744	1.350863	-1.139308
O	0.301163	1.700881	-2.119813
S	-0.653670	-0.675017	-0.220016
CL	-2.049923	0.370244	0.897448
O	-0.248118	-1.814620	0.556094
O	-1.125028	-0.729734	-1.571876
H	0.910104	-0.008886	1.843423
F	1.634490	1.793043	1.371460
H	3.222614	-0.339300	-0.111212
H	3.495338	-0.174672	1.737348

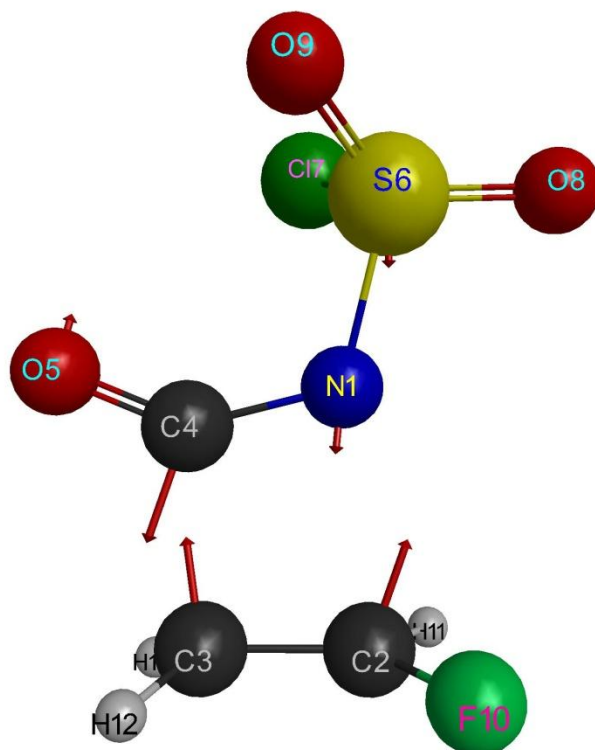
Structure 2. Local reactive intermediate in stepwise mechanism



Cartesian coordinates (in angstroms)

N	0.643432	0.535411	-0.128092
C	1.528550	0.597228	1.110207
C	2.360275	-0.631964	1.415735
C	0.472412	1.663876	-0.893380
O	-0.311075	1.901562	-1.765771
S	-0.525950	-0.674258	-0.221505
CL	-2.146779	0.312321	0.624224
O	-0.119179	-1.688164	0.708468
O	-0.838309	-0.902352	-1.599683
H	0.881375	0.873960	1.947915
F	2.357640	1.655669	0.844106
H	2.822503	-1.098021	0.544829
H	3.026637	-0.442723	2.257462

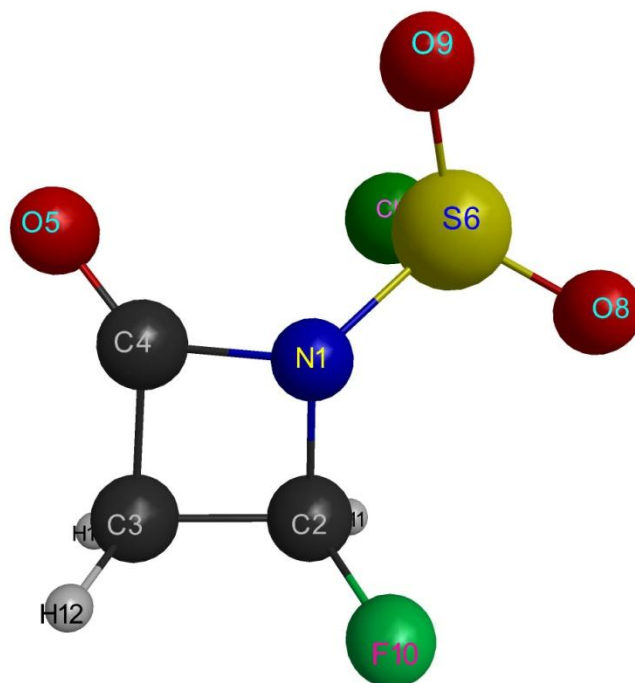
Structure 3. Concerted transition state geometry and normal mode corresponding to the single imaginary frequency of $599.6i\text{ cm}^{-1}$.



Cartesian coordinates (in angstroms)

N	0.457119	-0.170217	0.154261
C	2.473411	-0.391100	0.971303
C	2.745437	0.713232	0.177182
C	1.045789	0.835911	-0.493058
O	0.846808	1.714573	-1.258577
S	-1.132957	-0.515751	-0.228693
CL	-2.121174	1.133346	0.627833
O	-1.477952	-1.660921	0.568354
O	-1.383246	-0.451139	-1.645827
F	2.772563	-1.587991	0.563583
H	2.093155	-0.356071	1.985310
H	3.361003	0.549561	-0.701659
H	2.847900	1.663904	0.690610

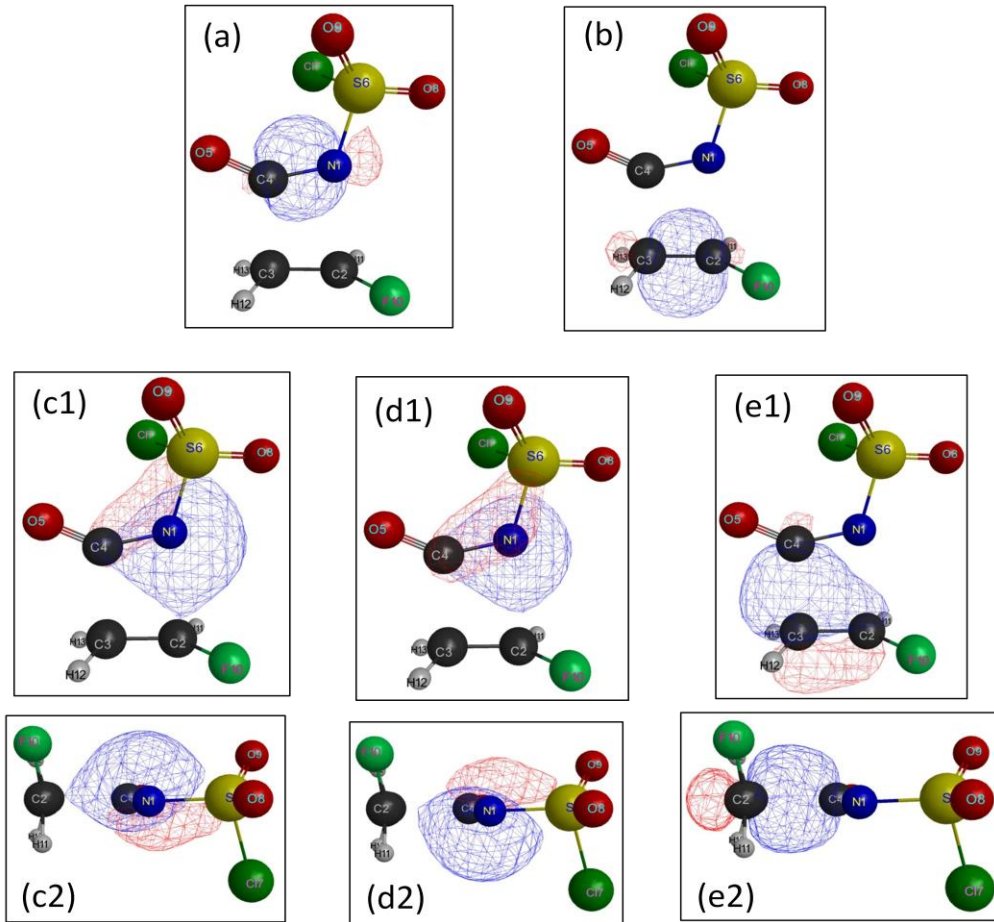
Structure 4. Reaction product local minimum



Cartesian coordinates (in angstroms)

N	0.618167	-0.137219	-0.240307
C	1.569329	-0.303473	0.858611
C	2.443394	0.792026	0.219812
C	1.293181	1.024826	-0.763196
O	0.976592	1.825037	-1.586651
S	-0.986867	-0.565398	-0.260045
Cl	-1.794172	0.986050	0.886519
O	-1.109205	-1.760749	0.524399
O	-1.475615	-0.399241	-1.595299
F	2.130019	-1.539889	0.892142
H	1.144927	-0.082138	1.840211
H	3.339505	0.382031	-0.248867
H	2.681197	1.654629	0.843558

Localized molecular orbitals of the cyclic 2+2 transition state, 5a-e2.



5a-e. RHF/6-311G(d,p) energy localized molecular orbitals at the concerted transition state.

(a) C-N sigma bond; (b) C-C sigma bond, (c1,c2) two views of the C-N pi bond; (d1,d2) two views of the N atom lone pair; (e1,e2) two views of the vinyl pi bond.

Stationary point	E(MP2/6-311G(d,p) ^a	Zero-point energy ^b	Relative energy ^c
CSI + CH ₂ =CHF	-1352.507608	0.067482	0.0
Stepwise reactive intermediate	-1352.422724	0.073312	56.9
Stepwise TS	-1352.412566	0.069565	60.9
Concerted TS	-1352.461899	0.069305	29.8
Product	-1352.540651	0.073210	-17.1

^a In hartrees.

^b In hartrees, scaled by 0.9748

^c In kcal/mol, including scaled zero point energies.