**IUPAC** 



# International Union of Pure and Applied Chemistry

#### JCAMP-DX for EMR (IUPAC RECOMMENDATIONS 2005)

Journal:	Pure and Applied Chemistry
Manuscript ID:	PAC-REC-05-09-13
Manuscript Type:	Recommendation
Date Submitted by the Author:	11-Sep-2005
Complete List of Authors:	Cammack, Richard; King's College, Pharmaceutical Sciences Research Division Fann, Yang; NINDS/NIH, Intramural IT Program Lancashire, Robert; University of the West Indies, Department of Chemistry Maher, John; University of Bristol, Department of Chemistry McIntyre, Peter; University of Glamorgan, School of Applied Sciences Morse, Reef; Illinois State University, Department of Chemistry
Keywords:	recommendations, Analytical Chemistry, EMR, EPR, ESR, spectroscopy, data standard, JCAMP-DX

powered by ScholarOne Manuscript Central<sup>™</sup>

1

IUPAC

# INTERNATIONAL UNION OF PURE AND APPLIED CHEMISTRY



COMMITTEE ON PRINTED AND ELECTRONIC PUBLICATIONS SUBCOMMITTEE ON ELECTRONIC DATA STANDARDS (JCAMP-DX)

# **JCAMP-DX** for EMR

# (IUPAC RECOMMENDATIONS 2005)

Prepared for publication by Richard Cammack<sup>1</sup>, Yang Fann<sup>2</sup>, Robert J. Lancashire<sup>3</sup><sup>‡</sup>, John P. Maher<sup>4</sup>, Peter S. McIntyre<sup>5</sup>, and Reef Morse<sup>6</sup>

<sup>1</sup> Pharmaceutical Sciences Research Division, King's College, Franklin-Wilkins Building, 150 Stamford St, London SE1 9NH, U.K.

<sup>2</sup> Intramural IT Program, NINDS/NIH, 10 Center Drive, MSC 1430, Bethesda, MD 20892-1430. U.S.A.

<sup>3</sup> Department of Chemistry, University of the West Indies, Mona, Kingston 7, JMAAW15, Jamaica

<sup>4</sup> School of Chemistry, University of Bristol, Bristol BS8 1TS, United Kingdom

<sup>5</sup> School of Applied Sciences, University of Glamorgan, Pontypridd, Mid-Glamorgan, CF37 1DL, United Kingdom

<sup>6</sup> Department of Chemistry, Illinois State University, Normal, IL 61790-4160, U.S.A.

*‡* Corresponding author

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

# **JCAMP-DX** for EMR

Richard Cammack<sup>1</sup>, Yang Fann<sup>2</sup>, Robert J. Lancashire<sup>3</sup>\*, John P. Maher<sup>4</sup>, Peter S. McIntyre<sup>5</sup>, and Reef Morse<sup>6</sup>

<sup>1</sup> Pharmaceutical Sciences Research Division, King's College London, Franklin-Wilkins Building, 150 Stamford St, London SE1 9NH, U.K.

<sup>2</sup> Intramural IT Program, NINDS/NIH, 10 Center Drive, MSC 1430, Bethesda, MD 20892-1430. U.S.A.

<sup>3</sup> Department of Chemistry, University of the West Indies, Mona, Kingston 7, JMAAW15, Jamaica

<sup>4</sup> School of Chemistry, University of Bristol, Bristol BS8 1TS, United Kingdom

<sup>5</sup> School of Applied Sciences, University of Glamorgan, Pontypridd, Mid-Glamorgan, CF37 1DL, United Kingdom

<sup>6</sup> Department of Chemistry, Illinois State University, Normal, IL 61790-4160, U.S.A.

\*Author to whom comments should be sent.

(IUPAC Recommendations 2005)

Abstract. In this document, we define a data exchange format initially formulated from discussions of an INTERNATIONAL UNION OF PURE AND APPLIED CHEMISTRY (IUPAC) limited term task group at the 35<sup>th</sup> Royal Society of Chemistry-ESR conference in Aberdeen 2002. The definition of this format is based on the IUPAC Joint Committee on Atomic and Molecular Physical Data Exchange (JCAMP-DX) protocols, which were developed for the exchange of infrared spectra [1] and extended to chemical structures [2], nuclear magnetic resonance data [3,4], mass spectra [5] and ion mobility spectra [6]. This standard of the JCAMP-DX was further extended to cover year 2000 compatible date strings and good laboratory practice [7] and the next release will cover the information needed for storing n-D data sets [8]. The aim of this paper is to adapt JCAMP-DX to the special requirements for EMR, electron magnetic resonance.

#### **1. INTRODUCTION**

JCAMP-DX is an evolving, open-ended, machine-independent, self-documenting file format for exchanging and archiving data from computerized laboratory instruments like spectrometers, diffractometers, and others, whose output is commonly represented as spectral (profile) plots, contours, or peak tables. The first JCAMP-DX protocol was designed to meet the need for exchanging infrared spectra between similar instruments of different manufacturers. The present document is the result of ongoing efforts by users and manufacturers to extend JCAMP-DX to other types of instrumental data.

A major objective of JCAMP-DX is to enable routine capture of data at the source to make it available for exchange, archiving, and entry into databases. All data are represented as labelled

4

5

7

9

10 11 12

13

14

15 16 17

18

19

21 22 23

24

27

29 30

31

32

37

40

41 42

43 44 45

47 48

51

57

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

fields of variable length using printable ASCII characters. A JCAMP-DX file is a text file which can be viewed, corrected, and annotated with ASCII text editors.

The JCAMP-DX protocol is non-proprietary. These specifications are copyrighted by the International Union of Pure and Applied Chemistry (IUPAC) solely for the purpose of linking them with the name JCAMP-DX. The right to copy these specifications for scientific purposes is hereby granted.

Use of the name JCAMP-DX to describe data files implies that they conform to the format and style described in the relevant protocols and the information content defined for a particular DATATYPE.

Use of the name JCAMP-DX in the description of software capability implies the ability to generate and read JCAMP-DX files as defined in the relevant published protocols for a particular DATATYPE (Section 4).

#### 2. SIGNIFICANCE AND USE

This version of JCAMP-DX provides for a description of the file structure to be used to accommodate a very wide range of EMR applications.

Inasmuch as it is very desirable for instrument data systems to be able to read and write files in a standard format directly, instrument vendors are encouraged to develop JCAMP-DX software for the instruments which they currently support. It is feasible for vendors to do so for a tightly defined CORE of information.

#### **3. BASIC STRUCTURE OF JCAMP-DX FILES.**

JCAMP-DX is a FILE specification. The basic element of JCAMP-DX files is the LABELLED-DATA-RECORD (LDR). LABELLED-DATA-RECORDS are combined into BLOCKS. A FILE may contain more than one BLOCK.

A simple FILE would look like

Data Block

##TITLE= Headers Data ##END=

The BLOCK structure was first described for IR COMPOUND files in Section 11 of Reference 1.

Link Block ##TITLE=

PROVISIONAL RECOMMENDATIONS **Do Not Cite – Do Not Implement - For Comment Only** 

9/11/2005 1:43 PM

#### IUPAC

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

Data Block 1	<i>Define total number of BLOCKS</i> ##TITLE=
Data Block 2	##END= ##TITLE=
Data Block 3	##END= ##TITLE=
File End	##END= ##END=

The NTUPLE structure was first described for NMR COMPLEX DATA STORAGE files in Section 7 of Reference 3.

NTUPLE File Start ##TITLE= Headers ##NTUPLES= ##PAGE= Data ##PAGE= Data ##END NTUPLES= File End ##END=

This is what we will normally mean by the word FILE.

The logical division of a JCAMP-DX file into CORE and NOTES subsets is a measure to separate the protocol for representing tabular data (CORE) which must be parsed by computer from that which is mainly for reference by humans.

#### 4 CORE.

1 2

19

21 22

23 24

29

31

32

37

40

41 42 43

46 47

48

51

57

The CORE portion of a JCAMP-DX BLOCK or FILE is the irreducible minimum JCAMP-DX file. The CORE provides a focus for instrument and software vendors to convert between inmemory forms and JCAMP-DX. The CORE contains one or more data arrays, corresponding attributes for defining units, scale factors, sizes, etc., of arrays for the data, plus key application dependent attributes and identification. The main objective of the CORE is to focus on the information which must be transferred to a foreign system in order for it to process and/or plot and label the data as if it had been generated internally.

**IUPAC** 

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

The CORE is divided into two subsets: CORE HEADER and CORE DATA.

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

#### 4.1 CORE HEADER

The CORE HEADER provides for overall identification of a JCAMP-DX file. Each record is prefixed with a ## sign and represented as ##HEADER= in the BLOCK or FILE. For example, ##TITLE=My First EMR Spectrum All the core headers are required elements. The CORE HEADER consists of: TITLE, JCAMP-DX, DATATYPE, DATA CLASS, ORIGIN, OWNER

In general information in the Header is either TEXT (free format) or STRING (reserved or predefined keywords) and information in the Data is either AFFN (ASCII FREE FORMAT NUMERIC) or ASDF (ASCII SQUEEZED DIFFERENCE FORM).

4.1.1 ##TITLE= (TEXT)

A description of the data file.

4.1.2 JCAMP-DX=(STRING) \$\$ for example JCAMP-DX \_writer.exe version 0.99

The VERSION NUMBER of JCAMP-DX, followed by a comment on the version of the software writing the file. *(Required)* 

4.1.3 ##DATA TYPE= (STRING).

Electron Magnetic Resonance spectroscopy now includes a wide variety of techniques all of which involve measurements which exploit the magnetic properties or ascertain the environment of the unpaired electron. The description of many of these as either ESR, electron spin resonance, or as EPR, electron paramagnetic resonance is commonplace in the literature. However the great diversity of techniques now applied to the measurement of paramagnetism, usually (but not always) in the presence of an external magnetic field, needs a generic descriptive term. In particular the emergence of imaging technology, of techniques such as ENDOR which sits between NMR and EPR/ESR, and of newer optical detection techniques, leads us to suggest an overall classification as 'EMR'. The use of the *historical* terms ESR or EPR thus falls under this overall classification, a situation which we do not wish to change. *(The existing IUPAC recommendation is to use the term EPR rather than ESR)*[9] The task group has therefore decided to adopt the generic title for the DATA TYPE as EMR, and to define all the METHODs in terms of either a MEASUREMENT or a SIMULATION.

#### e.g. ##DATA TYPE= EMR MEASUREMENT e.g. ##DATA TYPE= EMR SIMULATION (*Required*)

# 4.1.4.##DATA CLASS= (STRING).

This LDR contains the name of the type of tabular data, i.e., XYDATA, XYPOINTS, PEAK TABLE, PEAK ASSIGNMENTS, BLOCKS, or NTUPLES. Coates [10] first pointed out that the data for a number of different applications (IR and Raman spectra, GC retention times and NMR Chemical Shift) are so similar that JCAMP-DX files for all these types can be plotted or processed (smoothed, peak-picked, deconvoluted, etc.) by existing software for one of the techniques. This can clearly be extended to many other types to XY data which are represented in the XYDATA form, and probably also as XYPOINTS.

##DATA CLASS= is intended to provide an early indication to software for the purpose of determining whether or not it can process a given file. This seems important because it allows crossover of data processing routines between different techniques (Required)

#### 4.1.5 ##ORIGIN=(TEXT)

*The origin of the data, i.e. the name of the organisation. (Required)* 

# 4.1.6 ##OWNER=(TEXT)

The owner or author of the data. This can be "public domain" or the name of the person or organisation and may include a copyright note. *(Required)* 

# 4.2 CORE VARIABLE HEADER INFORMATION

The JCAMP-DX standard is easy to understand and expand. Many LDRs are already defined in previous JCAMP-DX protocols, and they should be used for electron magnetic resonance. However, in the case of the equipment parameters, the particular requirements of this technique call for some special LDRs. In the following part, definitions are given for LDRs that will allow a precise description of the equipment parameters for all EMR experiments (Note the syntax of ##.LABEL=.)

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

#### 4.2.1 ##.DETECTION MODE= CW or PULSE

*(required)* 

#### 4.2.1.1 CW

1 2

4

6

7

9 10 11

12

13

14 15 16

17

18 19

21

22

23

24

27

29

31

33

37

40

41

42 43

46 47

48

51

57

Continuous wave, the frequency (or field) dependence of the sample is measured in response to a periodic perturbation of microwave radiation to the sample.

#### 4.2.1.2 PULSE

One or more pulses of (e.g. microwave and/or RF) radiation are used to perturb the sample, and the recovery from this perturbation is measured as a function of a swept variable.

CW and pulse measurements are related via the Fourier Transform [11, 12, 13]. (*Required*)

#### 4.2.2 ##.METHOD= (STRING)

There are a wide variety of experimental methods used in EMR. The acronyms used are those described in the references used for the DETECTION MODE. The protocol description in this document concerns the following:

Abbreviations associated with other forms of EMR measurements not mentioned above, and which may need inclusion in the METHOD LDR in the future. Some of these will require extra EMR specific LDRs, but some may use the present set. Since many of these use pulse excitation, they will require methods to define pulse sequences.

ADMR – absorption detected magnetic resonance CIDEP – chemically induced dynamic electron polarization COSY - 2D correlation spectroscopy CYCLOPS – cyclically ordered phase sequence DECENT – decoupled ESEEM correlated to nuclear transition frequencies DEER – double electron electron resonance DEFENCE – deadtime free ESEEM by nuclear coherence-transfer echoes DONUT – double nuclear coherence transfer ELDOR – Electron-electron double resonance EPR – electron paramagnetic resonance (encompassed by SPECTRUM) ESR – electron spin resonance (encompassed by SPECTRUM) ESR-STM ESR scanning tunnelling microscopy ESE – electron spin envelope field sweep spectroscopy ESEEM – electron spin echo envelope modulation EXSCY (EXSY) – two dimensional exchange spectroscopy FDMR – fluorescence detected magnetic resonance FMR – very high field EPR FORTE – forbidden-transition labelled EPR HETEROCOSY – nD correlation spectroscopy

# PROVISIONAL RECOMMENDATIONS *Do Not Cite – Do Not Implement - For Comment Only*

8/26 P.O. 13757, Research Triangle Park, NC (919) 485-8700

\$ASQ585743 File000000.rtf

#### **IUPAC**

3	HFEPR – high field EPR
4	HYEND – hyperfine correlated ENDOR
6	HYSCORE - hyperfine sublevel correlation spectroscopy
7	LEFE – linear electric field effect
8	LODESR – longitudinal-detected ESR
9	I OD-PEPR - longitudinal-detected pulsed EPR
11	LOD-FEEFM - longitudinal-detected FEEFM
12	LOD-ESELVI - longitudinal-detected ESELVI
13	MARY magnetic field dependence of (affect on) resetion yield
14	MAR I – Inagliette field dependence of (effect off) feaction yield
15	MAS-EPR – magic angle spinning EPR
17	MODR – microwave optical double resonance
18	MQ-EPR – multiple quantum EPR
19	MQ-ELDOR – multiple quantum ELDOR
20	MQ-ENDOR – multiple quantum ENDOR
22	NZ-ESEEM - nuclear-Zeeman-Resolved ESEEM
23	PCDMR – photoconductive detected magnetic resonance
24	PEANUT – phase inverted echo-amplitude detected nutation
25	PEDRI – proton-electron double resonance imaging
20 27	PYESR – product yield detected ESR
28	RAS-EPR – right-angle spinning EPR
29	RYDMR – reaction yield detected magnetic resonance
30	SECSY – spin-echo correlated spectroscopy
31	SEDOR – spin-echo double resonance
33	SEESR – simultaneous electrochemistry ESR
34	SIFTER – single frequency technique for refocusing
35	SMART – single pulse matched resonance transfer
36 37	TREPR – time-resolved EPR
38	TREDMR – time resolved fluorescence detected magnetic resonance
39	
40	The list is intended to be as complete as negsible to allow research
41	The list is intended to be as complete as possible to anow researche

The list is intended to be as complete as possible to allow researchers to classify their particular form of spectrum. However, a number of the abbreviations are already encompassed by the current METHODs.

One possible route for defining these abbreviations might be by using a JCAMP-DX private LDR. Users can devise their own private labels to cover items they wish to include in the data file for completeness, e.g. ##\$RYDMR. Note however, that many software packages do not parse private labels and they may not be rewritten if saved from within an application. If used, the abbreviations need to have references.

4.2.2.1 DYNAMIC

Any of a number of electron magnetic resonance experiments in which a non-Boltzmann spin state distribution is produced by a combination of radical pairs or directly from the triplet state.

#### 4.2.2.2 ELDOR

Electron-Electron DOuble Resonance is a technique in which one frequency is used to pump an electronic transition at one frequency while observing at another frequency. It is often used to determine distances between centers containing unpaired electrons in different environments.

# 4.2.2.3 ENDOR

Electron-Nuclear Double Resonance has been described as EMR detected NMR and makes use of electron-nuclear hyperfine couplings between unpaired electrons and neighbouring nuclei.

# 4.2.2.4 ESEEM

Electron Spin Echo Envelope Modulation is a pulsed technique used for observing solid phase samples. [14]

# 4.2.2.5 ODMR

Any EMR experiment in which spin transitions are detected by optical means.

# 4.2.2.6 GONIOMETER

Any EMR experiment in which the sample angle, relative to the main magnetic field, is one of the axes.

# <u>4.2.2.7 HYSCORE</u>

The Hyperfine Sublevel Correlation Spectroscopy sequence was introduced in 1986 as a twodimensional four pulse experiment [15].

#### 4.2.2.8 KINETIC

Any EMR experiment in which time is one of the axes.

# 4.2.2.9 SATURATION

Any EMR experiment in which microwave power is one of the axes.

<u>4.2.2.10 SPECTRUM</u>

PROVISIONAL RECOMMENDATIONS **Do Not Cite – Do Not Implement - For Comment Only** 

9/11/2005 1:43 PM

\$ASQ585743\_File000000.rtf 10/26 P.O. 13757, Research Triangle Park, NC (919) 485-8700

2 3

4 5 6

7

9 10 11

12 13

14 15 16

17 18

19

21 22

23

24 25

27

29

30 31

32

37

40

41

42 43

45

47 48

51

55

57

59

#### **IUPAC**

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

Any EMR experiment in which the magnetic field is one of the axes.

#### 4.2.2.11 FID

Any EMR experiment in which the free induction decay is detected.

#### <u>4.2.2.12 TRIPLE</u>

Any EMR experiment, which involves the use two RF fields for modulating nuclear spins.

# 4.2.2.13 IMAGING

Any EMR experiment in which distance is one or more of the experimental axes.

# 4.2.2.14 SPECTRAL SPATIAL

Any EMR experiment in which distance and magnetic field are the experimental axes.

(METHOD is *Required*)

#### 4.2.3 ##.DETECTION METHOD= (TEXT)

Used where a resonator is not involved in the measurement and can name e.g. fluorescence, phosphorescence, photoconductivity, reaction yield. An alternative where appropriate is to give a wavelength for the detector, this may be applicable for HFEPR. (*Required*)

#### 4.2.4 ##.MICROWAVE FREQUENCY1= (AFFN)

Typical EMR experiments were conducted at a fixed microwave frequency ranged from few MHz to hundreds of GHz. To accommodate EMR experiments that utilized more than one source of microwave radiation, the number 1 and 2 are appended to the LDRs. The microwave frequency is expressed in Hz.

(Required)

#### 4.2.5 ##.MICROWAVE POWER1= (AFFN)

The power of microwave source in Watts. (Required)

4.2.6 ##.MICROWAVE PHASE1= (AFFN)

PROVISIONAL RECOMMENDATIONS Do Not Cite – Do Not Implement - For Comment Only 9/11/2005 1.43 PM \$ASQ585743 File000000.rtf

The phase of the microwave source in degrees. *(Required)* 

4.2.7 ##.MICROWAVE FREQUENCY2= (AFFN)

The frequency for the second microwave source in Hz. (*Required for ELDOR*)

4.2.8 ##.MICROWAVE POWER2= (AFFN)

The power for the second microwave source in Watts. (*Required for ELDOR*)

4.2.9 ##.MICROWAVE PHASE2= (AFFN)

The phase for the second microwave source in degrees. (*Required for ELDOR*)

4.2.10 ##.RECEIVER GAIN= (AFFN)

A scalar factor (gain) that is applied to the detection signal. (*Required*)

4.2.11 ##.MODULATION UNIT= (TEXT)

In a typical CW EMR experiment, the signal can be modulated by either field or frequency to achieve the enhancement of the signal resolution. The source of the modulation must be specified. (*Required for CW*)

4.2.12 ##.MODULATION AMPLITUDE= (AFFN)

The amplitude of the corresponding modulation unit. For example, the amplitude of field modulation is kept small compared to the linewidth in order to avoid over saturation of the signal. (*Required for CW*)

4.2.13 ##.MODULATION FREQUENCY= (AFFN)

The frequency of modulation unit. The typical used modulation frequency in an EMR experiment is 100 KHz. (*Required for CW*)

4.2.14 ##.RECEIVER HARMONIC= (AFFN)

The typical CW experiment is recorded in  $1^{st}$  harmonic (i.e. ##. RECEIVER HARMONIC=1). However, it can also be recorded as  $2^{nd}$  or  $3^{rd}$  Harmonics. (*Required for CW*)

4 5 6

7

9 10 11

12 13

14

15 16 17

18 19

21

22 23

24

29

31

37

40 41

42 43

47 48

51

57

#### IUPAC

# 4.2.15 ##.DETECTION PHASE= (AFFN)

The common used technique for EMR experiments is the phase-sensitive detection and is often calibrated to 0 degree relative to the reference arm. Therefore, the phase of the detector must be reported especially when the spectrum is not recorded in phase. (*Required for CW*)

4.2.16 ##.SCAN TIME= (AFFN)

The time (in seconds) required to complete one scan of the spectrum. (*Required*)

4.2.17 ##.NUMBER OF SCANS= (AFFN)

The total number of scans for a spectrum. (1 if a single scan) (*Required*)

4.2.18 ##.GONIOMETER ANGLE= (AFFN)

Assumes a single circle goniometer. The crystal orientation will also need to be recorded as will the crystal structure properties of the host lattice, and assuming the measurements are on a magnetically dilute sample chemical details of the host will be needed. *(Required if .METHOD=GONIOMETER)* 

4.2.19 ##.STATIC FIELD= (AFFN)

The static magnetic field in Tesla. This is commonly used when carrying out double resonance or spin echo experiments. (*Required for ENDOR*)

4.2.20 ##.SCANNED RF POWER= (AFFN)

In a double resonance experiment, say ENDOR, a constant power of RF source that was used to scan for nuclear transitions. (*Required for ENDOR*)

4.2.21 ##.PUMPED RF FREQUENCY 1= (AFFN)

In a double resonance experiment, say ENDOR, this is the frequency of the RF source which was used to scan for nuclear transitions (units are in Hz). (*Required for TRIPLE*)

4.2.22 ##.PUMPED RF POWER 1= (AFFN)

In a double resonance experiment, say ENDOR, this is the power of the RF source which was used to scan for nuclear transitions (units are in Watts). (*Required for TRIPLE*)

# 4.2.23 ##.GRADIENT THETA= (AFFN)

For EMR imaging experiments, this is the intensity of the gradient magnetic field along the theta angle (using spherical coordinates - units are in Tesla). (*Required for IMAGING*)

# 4.2.24 ##.GRADIENT PHI= (AFFN)

For EMR imaging experiments, this is the intensity of the gradient magnetic field along the phi angle (using spherical coordinates - units are in Tesla). (*Required for IMAGING*)

# 4.2.25 ##.GRADIENT STRENGTH IN THETA/PHI DIRECTION= (AFFN)

For EMR imaging experiments, this is the intensity of the total angular gradient magnetic field (using spherical coordinates - units are in Tesla) (*Required for IMAGING*)

# 4.2.26 ##.GRADIENT STRENGTH X= (AFFN)

For EMR imaging experiments, this is the intensity of the gradient magnetic field along the X axis (units are in Tesla). (*Required for IMAGING*)

# 4.2.27 ##.GRADIENT STRENGTH Y= (AFFN)

For EMR imaging experiments, this is the intensity of the gradient magnetic field along the Y axis (units are in Tesla). (*Required for IMAGING*)

# 4.2.28 ##.GRADIENT STRENGTH Z= (AFFN)

For EMR imaging experiments, this is the intensity of the gradient magnetic field along the Z axis (units are in Tesla). (*Required for IMAGING*)

4.2.29 ##.SIMULATION SOURCE= (TEXT)

The sources or tools used to simulate the spectrum (e.g. QPOW, Bruker, WinEPR, etc.). (*Required for EMR SIMULATION*)

4.2.30 ##.SIMULATION PARAMETERS= (TEXT)

The detailed descriptions of the EMR parameters that are needed to reproduce the simulated spectra such as microwave frequency, hyperfine coupling constants, rotation angles, static magnetic fields, scan range, linewidth, etc. *(Required for EMR SIMULATION)* 

# 4.3 CORE DATA

4

5 6

7

#### IUPAC

#### 4.3.1 ##XUNITS= (STRING) and ##YUNITS= (STRING)

*Here, the units of the axes can be given as one of the following keywords: For ##XUNITS=: DEGREES, HERTZ, KELVIN, SECONDS, TESLA, WATTS. For ##YUNITS=: POWER, INTENSITY and ARBITRARY UNITS. (Required)* 

4.3.2 ##FIRSTX= (AFFN) and ##LASTX= (AFFN)

*First and last actual abscissa values of ##XYDATA=. First tabulated abscissa times ##XFACTOR= should equal ##FIRSTX=. (Required for ##DATA CLASS=XYDATA)* 

4.3.3 ##FIRSTY= (AFFN)

*Here, the actual ordinate value corresponding to* ##FIRSTX= *is meant.* ##FIRSTY= *should be equal* ##YFACTOR= times the first Y-value in ##XYDATA=. (Required for ##DATA CLASS=XYDATA)

#### 4.3.4 ##XFACTOR= (AFFN) and ##YFACTOR= (AFFN)

The values of a spectrum may be converted to integer to save space and allow the DIFDUP format (see [1] Section 5). It is important to select a convenient scaling to keep the file within reasonable limits, but to store all significant digits. In such a case, the ##XFACTOR= and ##YFACTOR= LDR contain a floating-point number to be multiplied by the values in ##XYDATA= to arrive at the original data point value.

In most cases,  $\pm 32767$  is sufficient; therefore this is the recommended ordinate scaling. If larger scaling is necessary, it is required to give the actual unscaled maximum and minimum of the ordinates in the records ##MAXY= and ##MINY=.

For example, if a Y-value with 9 significant figures (e.g., 0.002457194) needs to be converted to an integer value for ASDF coding then:

a) divide by the maximum of the absolute Y-value (say, 0.346299765),

b) multiply by the largest integer value (MAXINT) necessary to place all significant figures left of the decimal point, and

c) convert to integers.

(Required for ASDF data types. Should be to 1.0 for AFFN)

4.3.5 ##NPOINTS= (AFFN)

The number of points in the data table is required for all data classes: XYDATA, XYPOINTS, PEAK TABLE, and PEAK ASSIGNMENTS. (Required, handled by VAR DIM in NTUPLE files)

# 4.4 CORE DATA TABLE

Data must be stored in one of the following data formats (Sections 4.4.1–4.4.4). Only one of these formats is allowed per DATA block, and the selected data table is given in the header, e.g., ##DATA CLASS=XYDATA.

#### 4.4.1 ##XYDATA= (AFFN or ASDF).

This LDR contains a table of spectral data with abscissa values at equal intervals specified by parameters defined in Section 4.3. The label is followed by a variable list, (X++(Y..Y)) where "..." indicates indefinite repeat of Y-values until the end of line and ++ indicates that X is incremented by (LASTX-FIRSTX) / (NPOINTS-1) between two Y-values. For discrete points, the AFFN form is allowed where each Y value is written out in full. This form creates large files, but is easily human-readable. The ASDF forms defined in JCAMP-DX 4.24 (Reference 1) produce much smaller files but at the cost of reduced human readability. In modern computers systems file size is no longer the restriction it used to be and the task group felt that users may be more comfortable with AFFN which would also allow the storage of numbers in scientific notation.

#### 4.4.2 ##XYPOINTS= (AFFN)

This LDR contains a table of spectral data with unequal abscissa increments. The label is followed by a variable list, (XY..XY). X and Y are separated by commas, data pairs are separated by semicolons or blanks. This LDR should not be used for peak tables. For example, X1, Y1; X2,Y2, ..., Xi, Yi.

#### 4.4.3 ##PEAK TABLE= (AFFN)

It is recommended to store peak information using ##PEAK ASSIGNMENTS=, however, for backward compatibility this definition is included. This data table contains a table of peaks where the peak data starts on the following line. The label is followed by the variable list (XY) or (XYW) for peak position, intensity, and width, where known, on the same line. The function used to calculate the peak width should be defined by a \$\$ comment in the line below the label. The peak groups are separated by a semicolon or space, components of a group are separated by commas.

#### 4.4.4 ##PEAK ASSIGNMENTS=(STRING)

Variable list: (XA), (XYA), or (XYWA)

After this LDR, a list of peaks and their assignments for each component is given in the following form:

(X1, [Y1, [W1]], <A1>)

• • • •

(Xi, [Yi, [Wi]], <Ai>)

X and Y indicate the location and height of each peak in units given by ##XUNITS= and ##YUNITS=. W stands for width in ##XUNITS=, and A represents a string describing the assignment enclosed in angle brackets.

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

The parentheses provide a start and end flag of each assignment. Square brackets indicate optional information. It is important for the technical readability to have the same format for the whole peak assignment table and describe it after the ##PEAK ASSIGNMENTS= LDR with (XA), (XYA), or (XYWA). This LDR should be followed by a comment, which gives the method of finding the peak.

#### **5. NOTES**

The notes portion of a JCAMP-DX file or block complements the core. Notes describe an experiment in greater detail than does ##TITLE=, including descriptions of equipment, method of observation, and data processing, as appropriate. Notes may contain information, which is not found in the native file in which data is originally collected by an instrument. Notes are placed before the core data section to permit them to be viewed without listing the whole file. The contents of the notes depend on the user as well as the technique or application. Notes will vary for different samples, sites, data systems, and applications employed in the experiment.

#### **5.1 GLOBAL NOTES**

These have been already defined in JCAMP-DX and are common to all spectroscopy types. The file headers, spectral and sample parameters are often the same for different analytical techniques. This allows us to implement many of the standard LDRs from the existing JCAMP-DX protocols. The list given below is only a selection from those allowed. A complete list can be found in the references [1-6].

At least one of the optional LDRs described in Sections 5.1.3-5.1.8 should be included in each JCAMP-DX file. This is important for later archiving, as these fields will yield more detailed information on the content of the data stored than a simple ##TITLE= field.

#### 5.1.1 ##LONG DATE= (STRING)

Date of measurement is required by many agencies and recommended in the year 2000 form: YYYY/MM/DD [HH:MM:SS[.SSS] [ $\pm$ XXXX]]. YYYY is the long format of the year, MM the number of the month, DD the number of the day, HH the hour, MM the minutes, SS.SSS the seconds and fractions of a second of the measurement,  $\pm$ XXXX is the difference to the UTC (e.g., +0100 is one hour difference to UTC). (Optional)

#### 5.1.2 ##SOURCE REFERENCE= (TEXT)

*Here, an identification of the original spectrum file in native format or library name and serial number is possible, for example. (Optional)* 

5.1.3 ##SAMPLE DESCRIPTION= (TEXT)

If the sample is not a pure compound, this field should contain its description, i.e., composition, origin, appearance, results of interpretation, etc. If the sample is a known compound, the following LDRs specify structure and properties, as appropriate. (Optional)

#### 5.1.4 ##IUPAC NAME=(STRING)

The use of IUPAC names for the sample has been recommended previously by the Commission on Molecular Structure and Spectroscopy of the Physical Chemistry Division in 1991. (Optional)

#### 5.1.5 ##CAS NAME= (STRING)

Sample name according to Chemical Abstracts naming conventions as described in the CAS Index Guide is required here. Examples can be found in Chemical Abstracts indices or the Merck Index. Greek letters are spelled out, and standard ASCII capitals are used for small capitals, Sub- and superscripts are indicated by prefixes / and  $\land$ . Example: alpha-D-glucopyranose, 1-(dihydrogen phosphate). (Ontional)

(Optional)

#### 5.1.6 ##NAMES= (STRING)

Here, the common, trade, or other names are allowed. Multiple names are placed on separate lines.

(Optional)

#### 5.1.7 ##MOLFORM= (STRING)

Another possibility of describing the sample is to write down the molecular formula. Elemental symbols are arranged with carbon first, followed by hydrogen, and then remaining element symbols in alphabetic order.

The first letter of each elemental symbol is capitalized. The second letter, if present, is lower case. One-letter symbols must be separated from the next symbol by a blank or digit. Sub- and superscripts are indicated by the prefixes: / and /, respectively and are terminated by the next nondigit. The slash may be omitted for subscripts. For readability, each atomic symbol may be separated from its predecessor by a space. For substances that are represented by dot disconnected formulas (hydrates, etc.), each fragment is represented in the above order, and the dot is represented by \*. Isotopic mass is specified by a leading superscript. D and T may be used for deuterium and tritium.

Examples: C2H4O2 or C2 H4 O2 (ethanoic acid) H2 /\17O (water, mass 17 oxygen) (Optional)

#### IUPAC

#### 5.1.8 ##CONCENTRATIONS= (STRING)

The list of the known components and their concentrations has the following form, where N stands for the name and C for the concentration of each component in units given with U in the form:

##CONCENTRATIONS= (NCU) (N1, C1, U1)

... (Ni, Ci, Ui)

The group for each component is enclosed in parentheses. Each group starts a new line and may continue on following lines.

(Optional)

# 5.1.9 ##SPECTROMETER/DATA SYSTEM= (TEXT)

This LDR contains manufacturers' name, model of spectrometer, software system, and release number, as appropriate in the form used by the manufacturer. (Optional)

# 5.1.10 ##DATA PROCESSING= (TEXT)

Here, all mathematical procedures used before storing the data in the JCAMP-DX file are described. This LDR is also important in peak assignments. (Optional)

5.1.11 ##XLABEL= (TEXT) and ##YLABEL= (TEXT) These LDRs give the possibility of labeling the axes. (Optional)

5.1.12 ##MAXX= (AFFN), ##MINX= (AFFN), ##MAXY= (AFFN), and ##MINY= (AFFN) These LDRs give the largest and smallest actual X- and Y-values in a spectrum. ##MAXY= and ##MINY= are required if the Y-range exceeds recommended ordinate scaling (Section 4.3.4). (Optional)

# **5.2 DATA-TYPE-SPECIFIC NOTES**

Data-type-specific labels are RESERVED labels that are defined by qualified user groups for a particular data-type. A data-type-specific label is distinguished by a data-type-specific label name which starts with a period (e.g., ##.MICROWAVE FREQUENCY=). Choice of period as distinguishing character is by analogy with the convention for data-structure names in Pascal and C. Effectively, the full label name is the concatenation of the data-type name and the label-name, with a period in between, i.e., ##EMR MEASURMENT• MICROWAVE FREQUENCY=.

#### 5.2.1 ##.RESONATOR= (TEXT)

In a typical EMR spectrometer the resonator (a.k.a. cavity) is used to hold the sample, however a resonant cavity is not required to achieve resonance (!). This LDR enables the cavity to be described, e.g.  $TE_{102}$  rectangular,  $TE_{011}$  cylindrical, helical, loop-gap, Fabry-Perot

#### 5.2.2 ##.TIME CONSTANT= (AFFN)

This is most appropriate with legacy measurements where a signal was measured and recorded on a chart recorder. With digital recording, especially where an integrating ADC is used, the interpretation of a time constant is problematic. The time constant is an output filter that utilizes resistive-capacitive filter to filter the output of the phase-sensitive detector to improve the signal to noise. The time constant is expressed in seconds.

5.2.3 ##.PUMPED RF FREQUENCY 2= (AFFN)

In a triple resonance experiment, this is the frequency of the RF source which was used for the third source (units are in Hz).

5.2.4 ##.PUMPED RF POWER 2= (AFFN)

In a triple resonance experiment, this is the power of the RF source which was used for the third source (units are in Watts).

5.2.5 ##DATA PROCESSING= (TEXT)

The descriptions of methods or algorithms used to process the data (e.g. FT).

5.2.6 ##.CALIBRATION STANDARD= (TEXT)

The chemical standard used internally or externally to calibrate the g value of an EMR signal. (e.g. DPPH or  $MgO_2$ ).

5.2.7 ##.X\_OFFSET= (AFFN)

The numerical offset of the global X axis.

Page

1

#### IUPAC

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

# 6. SUMMARY

The following table lists the LDRs discussed with their basic parameters shown.

#### Table 1. Relevant terms for EMR

Parameter	Allowable words and	Comments
	UNITS	
##TITLE=	TEXT	REQUIRED
##JCAMP-DX=	5.01 STRING	REQUIRED
##DATA TYPE=	EMR MEASUREMENT,	REQUIRED
	EMR SIMULATION	
##DATA CLASS=	XYDATA, XYPOINTS,	REQUIRED
	PEAK TABLE or	
	ASSIGNMENTS	
##ORIGIN=	TEXT	REQUIRED
##OWNER=	TEXT	REQUIRED
##LONGDATE=	YYYY/MM/DD	OPTIONAL
	[HH:MM:SSSS[.SSSS][±UU	
	ŪŪ]]	
##SPECTROMETER/DATA	TEXT	OPTIONAL
SYSTEM=		
##.DETECTION MODE=	CW, PULSE	REQUIRED
##.METHOD=	DYNAMIC, ELDOR,	REQUIRED
	ENDOR,ESEEM, ODMR,	
	GONIOMETER,	
	HYSCORE, KINETIC,	
	SATURATION,	
	SPECTRUM, FID, TRIPLE,	
	IMAGING, SPECTRAL	
	SPATIAL, STRING	4

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

##.RESONATOR=	TEXT	OPTIONAL
##. DETECTION METHOD=	TEXT	REOUIRED when not
		RESONATOR
##.MICROWAVE	HERTZ	REQUIRED
FREQUENCY1=		
##.MICROWAVE POWER1=	WATTS	REQUIRED
##.MICROWAVE PHASE1=	DEGREES	REQUIRED
##.MICROWAVE	HERTZ	REQUIRED for ELDOR
FREQUENCY2=		
##.MICROWAVE POWER2=	WATTS	REQUIRED for ELDOR
##.MICROWAVE PHASE2=	DEGREES	REQUIRED for ELDOR
##.RECEIVER GAIN=	DIMENSIONLESS	REQUIRED
	NUMBER	
##.MODULATION UNIT=	TESLA, HERTZ, LUMENS,	REQUIRED for CW
	STRING	
##.MODULATION AMPLITUDE=	NUMBER, IN UNITS OF	REQUIRED for CW
	MODULATION UNIT	
##.MODULATION	HERTZ	REQUIRED for CW
FREQUENCY=		
##.RECEIVER HARMONIC=	POSITIVE INTEGER,	REQUIRED for CW
	usually 1	
##.DETECTION PHASE=	DEGREES	REQUIRED for CW
##.TIME CONSTANT=	SECONDS	OPTIONAL
##.SCAN TIME=	SECONDS	REQUIRED
##.NUMBER OF SCANS=	INTEGER	REQUIRED
##.GONIOMETER ANGLE=	DEGREES	REQD if .METHOD=
		GONIOMETER
##.STATIC FIELD=	TESLA	<b>REQUIRED</b> for ENDOR
##.SCANNED RF POWER=	WATTS	<b>REQUIRED</b> for ENDOR
##.PUMPED RF FREQUENCY 1=	HERTZ	<b>REQUIRED</b> for TRIPLE
##.PUMPED RF POWER 1=	WATTS	<b>REQUIRED</b> for TRIPLE
##.PUMPED RF FREQUENCY 2=	HERTZ	OPTIONAL for TRIPLE
##.PUMPED RF POWER 2=	WATTS	OPTIONAL for TRIPLE
##DATA PROCESSING=	TEXT	OPTIONAL
##.CALIBRATION STANDARD=	TEXT	OPTIONAL
##.X_OFFSET=	NUMBER	
##.GRADIENT THETA=	TEXT, DEGREES	REQUIRED for IMAGING
##.GRADIENT PHI=	TEXT, DEGREES	<b>REQUIRED</b> for IMAGING
##.GRADIENT STRENGTH IN	TEXT, T/M	<b>REQUIRED</b> for IMAGING

# PROVISIONAL RECOMMENDATIONS *Do Not Cite – Do Not Implement - For Comment Only*

9/11/2005 1:43 PM

ASQ585743\_File000000.rtf 22/26 P.O. 13757, Research Triangle Park, NC (919) 485-8700

#### IUPAC

Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

THETA/PHI DIRECTION=		
##.GRADIENT STRENGTH X=	TEXT, T/M	<b>REQUIRED</b> for IMAGING
##.GRADIENT STRENGTH Y=	TEXT, T/M	<b>REQUIRED for IMAGING</b>
##.GRADIENT STRENGTH Z=	TEXT, T/M	REQUIRED for IMAGING
##.SIMULATION SOURCE=	TEXT	REQUIRED FOR EMR
		SIMULATION
##.SIMULATION	TEXT	REQUIRED FOR EMR
PARAMETERS=		SIMULATION
##SAMPLE DESCRIPTION=	TEXT	OPTIONAL
##CAS NAME=	TEXT	OPTIONAL
##STATE=	Solid, liquid, gas, etc. TEXT	OPTIONAL
##CONCENTRATION=	TEXT	OPTIONAL
##XUNITS=	DEGREES, HERTZ,	to be used as XLABEL if
	KELVIN, SECONDS,	##XLABEL is undefined.
	TESLA, WATTS, STRING	REQUIRED
##YUNITS=	PREDEFINED, can be	REQUIRED
	ARBITRARY UNITS,	
	STRING	
##XLABEL=	FIELD, TIME, ANGLE,	OPTIONAL
	TEMPERATURE, POWER,	
	TEXT	
##XFACTOR=	NUMBER	= 1 if no compression
##YFACTOR=	NUMBER	= 1 if no compression
##FIRSTX=	NUMBER	REQUIRED for XYDATA
##LASTX=	NUMBER	REQUIRED for XYDATA
##NPOINTS=	POSITIVE INTEGER	REQUIRED
##FIRSTY=	NUMBER	REQUIRED
##\$NOTES	TEXT	from legacy data
##END=		REQUIRED
		31

#### IUPAC

Example file	
##TITLE= VO(acac)2 in toluene @ 10-3M	
##ICAMP-DX= 5.00 $\$ converted ESP300E text file	
##DATA TYPE= FMR MEASUREMENT	
##DATA CLASS= XYDATA	
##ORIGIN= School of Chemistry University of Bristol	
##OWNER= public domain	
##LONGDATE= $2001/02/19$	
##SPECTROMETER/DATA SYSTEM= Bruker ESP30	00E
##INSTRUMENTAL PARAMETERS= X Band	
## DETECTION MODE= CW	\$\$ required 4.2.1
## METHOD= SPECTRUM	\$\$ required 4.2.2
## DETECTION METHOD= RESONATOR	\$\$ required 4.2.3
## MICROWAVE FREQUENCY1= $9.782513E+09$	\$\$ required 4.2.4 Hz (9.782513 GHz)
## MICROWAVE POWER1= 020	\$\$ required 4.2.5 Watts (20mW)
## MICROWAVE PHASE1= 0	\$\$ required 4.2.6
## RECEIVER GAIN= $2.0E+04$	\$\$ required 4.2.10
## MODULATION UNIT= TESLA \$\$ 1	required 4 2 11
##.MODULATION AMPLITUDE= 1.011E-03	\$\$ required 4.2.12 TESLA (1.011G)
##.MODULATION FREOUENCY= 100000	\$\$ required Hz (100kHz)
##.RECEIVER HARMONIC= 1	\$\$ required 4.2.14
##.DETECTION PHASE= 0	\$\$ required 4.2.15
##.SCAN TIME= 300	\$\$ required 4.2.16 (seconds)
##.NUMBER OF SCANS= 1	• • \$\$ required 4.2.17
##XUNITS= TESLA	\$ 1 gauss = 0.1 mT
##YUNITS= ARBITRARY UNITS	
##XFACTOR= .0001240537	
##YFACTOR= 1.245460	
##FIRSTX= 2.964866	
##LASTX= 4.064866	
##MAXY= 40810.00	
##MINY= -40552.00	
##DELTAX= .0005373718	
##NPOINTS= 2048	
##FIRSTY= -14.00000	
##XYDATA=(X++(YY))	
23900a1k9kmP7M2n5o7N1l3K1o5M9k1QK4K1p1O3j	3K4Nk0J6j4p8Q8L2r3k5L9OR6Lk9p9K
52/b3AbU03	
32/b/1/	
##END=	

\$ASQ585743\_File000000.rtf 24/26 P.O. 13757, Research Triangle Park, NC (919) 485-8700

 Draft for Publication in: Pure and Applied Chemistry - International Union of Pure and Applied Chemistry Subcommittee on Electronic Data Standards (JCAMP-DX) of the Committee on Printed and Electronic Publications

#### 7. ACKNOWLEDGEMENTS

The task group wishes to acknowledge financial support from IUPAC and to acknowledge technical contributions and feedback from instrument vendors and users.

#### 8. REFERENCES

1. R. S. McDonald and P. A. Wilks Jr. "JCAMP-DX: A Standard Form for Exchange of Infrared Spectra in Computer Readable Form", Appl. Spectrosc. 42, 151–162 (1988) and "JCAMP-DX. A standard format for the exchange of infrared spectra in computer readable format (IUPAC Recommendations 1991)", Pure Appl. Chem. 63, 1781–1792 (1991).

2. J. Gasteiger, B. M. P. Hendriks, P. Hoever, C. Jochum, H. Somberg. "JCAMP-CS: A Standard Exchange Format for Chemical Structure Information in Computer-Readable Form", Appl. Spectrosc. 45, 4–11 (1991)

3. A. N. Davies and P. Lampen. "JCAMP-DX for NMR", Appl. Spectrosc. 47, 1093–1099 (1993).

4. A.N. Davies, J. Lambert, R.J. Lancashire, P. Lampen with W. Conover, M. Frey, M. Grzonka, E. Williams, and D. Meinhart, "Guidelines for the Representation of Pulse Sequences for Solution-State Nuclear Magnetic Resonance Spectrometry (IUPAC Recommendations 2001)" Pure Appl. Chem., 73, 1749-1764, (2001).

5. P. Lampen, H. Hillig, A. N. Davies, M. Linscheid. "JCAMP-DX for Mass Spectrometry", Appl. Spectrosc. 48, 1545-1552 (1994).

6. J.I. Baumbach, P. Lampen, A.N. Davies. "IUPAC/JCAMP-DX. An International Standard for the Exchange of Ion Mobility Spectrometry Data", Pure & Appl. Chem., Vol. 73, No. 11, pp. 1765-1782, 2001.

7. P. Lampen, J. Lambert, R. J. Lancashire, R. S. McDonald, P. S. McIntyre, D. N. Rutledge, T. Fröhlich, A. N. Davies. "An Extension to the JCAMP-DX Standard File Format, JCAMP-DX V.5.01 (IUPAC Recommendations 1999)", Pure Appl. Chem. 71, 1549–1556 (1999).

8. for a draft, see http://www.jcamp.org

9. H. Kon, Recommendations for EPR/ESR nomenclature and conventions for presenting experimental data in publications (IUPAC recommendations 1989) *Pure Appl. Chem* **61**, (12), 2195-2200 (1989).

10. J.P. Coates, "Computers in Spectroscopy Part 3", Spectroscopy, 2, 14-18, 120-22, (1987).

11. "Electron Paramagnetic Resonance", J.A.Weil, J.R.Bolton, J.E.Wertz, Wiley-Interscience, 1994. ISBN 0 471 57234 9

12: "Principles of pulse electron paramagnetic resonance", A.Schweiger and G.Jeschke, Oxford UP., 2001. ISBN 0 19 850634 1

13. "Modern pulsed and continuous-wave electron spin resonance" edit by Larry Kevan and Michael Bowman, John Wiley & Sons, New York, ISBN 0-471-50274-X (1990).

14. L.G. Rowan, E.L. Hahn and W. B. Mims, "Electron spin echo envelope modulation", Phys. Rev. A, 137, 61 (1965).

Lehr, and t investigatio. 15. P. Hoefer, A. Grupp, H. Nebenfuehr, and M. Mehring, "Hyperfine sublevel correlation (HYSCORE) spectroscopy: a 2D ESR investigation of the squaric acid radical", Chem. Phys. Lett. 132, 279 (1986).