MATLAB Import/Export Routines Version 1.2 User Manual CTF MEG[™] System





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MATLAB Import/Export Routines User Manual

Version 1.2

VSM MedTech Ltd.





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$\mathsf{CTF} \mid \mathbf{MEG}^{^{\scriptscriptstyle{\mathsf{M}}}}$

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Warnings and Cautions





WARNING

These programs must not be used for clinical applications. If CTF MEG data are processed by this tool, they should not be later employed for clinical and/or diagnostic purposes.

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Introduction



1. Introduction

The popularity of MATLAB for data analysis has resulted in the need for a package of MATLAB programs that will allow the reading of CTF datasets into the MATLAB environment, and the preparation of CTF datasets containing the results of MATLAB processing.

To meet this need, VSM has prepared MATLAB programs for manipulating CTF datasets:

- (1) **readCTFds**: Reads the files that define a CTF dataset, but not the sensor data.
- (2) **getCTFdata**: Reads the sensor data.
- (3) setCTFDataBalance: Balances the data.
- (4) getCTFBalanceCoefs: Reads the balance tables that are included in each dataset.
- (5) writeCTFds: Creates a new CTF-format dataset.
- (6) addCTFtrial: Adds trials to a dataset created by writeCTFds.

The following programs have been prepared for reading and writing MRIs and head-model files:

- (6) readCTFMRI: Reads MRIs in CTF format (version 4)
- (7) writeCTFMRI: Creates MRI files that are compatible with MRIViewer.
- (8) readCTFhdm: Reads multiple-local spheres head-model files.
- (9) writeCTFhdm: Creates head-model files that are compatible with CTF MEG[™] Analysis Software (DipoleFit and SAM beamformer programs).

This document describes the use of these programs. The formats of CTF files are described in document "CTF MEG™ File Formats", PN900-0088.

These programs have been verified to transfer CTF datasets to and from MATLAB correctly (MATLAB6.5 running under Windows XP), but within MATLAB anything can be done to the data.

Consequently, these programs must not be used for clinical applications of MEG data. Datasets created by writeCTFds and files created by writeCTFMRI and writeCTFhdm must not be used later in clinical applications, even with software that has received clearance for clinical applications.

New in version 1.1:

- (1) getCTFdata, writeCTFds handle data arrays exceeding 2 GB.
- (2) addCTFds allows addition of trials to an existing dataset.
- (3) readCTFhom and writeCTFhom handle v5 and v6 head-model files.

New in version 1.2:

- (1) readCTFds and writeCTFds have been changed to accommodate fMEG .hc files.
- (2) A minor change is made to handle text strings when writing CPerist objects.
- (3) **readCPersist** has been changed so it will not give an error when a CPersist file has extra bytes after the finale '**EndofParameters**' field.

Installation



2. Installation and Help

2.1 Installation

The codes are provided as a .zip file, on a floppy disk, or on a CD (PN530-0036 MATLAB Import/Export Routines: Program Files). To make them available for calls from MATLAB, follow the standard procedure:

- (1) Create a folder on the computer that is running MATLAB. For example, in Windows the folder could be C:\CTF-MATLAB or C:\MATLAB\toolbox\CTFDataSet. On a Linux system, you might create /usr/local/matlab/toolbox/CTFDataSet.
- (2) Copy the MATLAB programs to this folder.
- (3) Start MATLAB and use the Set Path command available in the Files Menu to add the folder to the path.

2.2 Help

Typing the name of any of the programs documented here gives a brief summary of the program and a list of input and output arguments. For example typing

```
getCTFdata;
```

produces the response

Reading Data



3. Reading CTF data into MATLAB

3.1 readCTFds - reading the dataset description

The user issues the MATLAB command

ds=readCTFds (datasetname) ;

where datasetname is the name of the dataset (i.e. the directory containing the dataset). readCTFds creates structure ds which holds the information that describes the dataset. Table 1 (page 5) lists the fields of ds and the corresponding dataset files. For analysis in MATLAB, usually only the .res4, and possibly the .hc, fields are used, but the additional fields are present to supply information for dataset files when subsequently creating a dataset using writeCTFds.

ds.res4 holds the parameters that describe the data including sample rate, trial size, channel names, and sensor parameters (sensor positions, gains and gradiometer balancing). Appendix A describes the fields of structure ds in detail and Appendix B describes ds.res4 in detail.

For a 275-channel CTF system, the size of structure **ds** is about 4 MB, of which 3.6 MB is the .res4 field.

3.1.1 No clinical use of readCTFds

On the first call to **readCTFds** in a MATLAB session (or on the first call since issuing the commands **clear all** or **clear readCTFds**), the following message is displayed on the monitor:

readCTFds: You are reading CTF data for use with a software-application tool that is not manufactured by VSM MedTech Ltd. and has not received marketing clearance for clinical applications. If CTF MEG data are processed by this tool, they should not be later employed for clinical and/or diagnostic purposes.

Table 1. List of fields of structure ds produced by readCTFds

readCTFds always produces fields .baseName, .path, .res4 and .meg4. The other fields are created only if the files listed are actually present in the dataset. getCTFdata uses fields baseName, .path and .res4, and file baseName .meg4. The Table shows which fields are required by writeCTFds. Appendix A describes the fields.

ds field	Dataset file	Required by writeCTFds	Information/Description	
.baseName		x	Dataset is directory	
.path		X	[path,baseName,'.ds']	
.res4	baseName.res4	x	Description of the data including sensor gains, balancing data, and trial structure. Central to any analysis of the dataset.	
.meg4	baseName.meg4	X	Data size (bytes) and 8-character header	
.infods	baseName.infods	X	Dataset information. If .infods is not present, writeCTFds will create one.	
.newds	baseName.newds		Obsolete dataset information file. Included for compatibility with older CTF software.	
.acq	baseName.acq		Acquisition parameters.	
.hist	baseName.hist		Text file describing acquisition and processing history.	
.hc	baseName.hc		Head coil positions	
.eeg	baseName.eeg		EEG electrode channels, names and positions.	
.mrk	MarkerFile.mrk		Event markers. Generated by Acq or by DataEditor.	
.TrialClass	ClassFile.cls		Trial classes. Generated by Acq or by DataEditor.	
.badSegments	bad.segments		Marks bad segments of data. Generated by DataEditor.	
.Virtual	VirtualChannels		Virtual channels specified by DataEditor.	
.BadChannels	BadChannels		List of channels known to be bad.	
proceesing	processing of		DataEditor processing parameters (filtering	

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3.2 getCTFdata - reading the data

Data are read from the dataset's .meg4 file and converted to user-specified units by getCTFdata. The command is

data=getCTFdata(ds,data list,chan_list,unit,prec);

where the input arguments are:

ds: The structure produced by readCTFds. (Only the first three fields in Table 1 are used by getCTFdata.)

data list: Tells getCTFdata which data to read.

- (1) data_list<=0 or [] or not defined. Read the complete dataset.
- (2) size (data list) = [1 N]. Read the N trials listed in row vector data list.
- (3) size(data_list)=[3 1]. Read points data_list(1):data_list(2) from trial data list(3).

chan_list: Optional list of channels to read. If not specified, or if chan_list=[], then all
 channels are read. chan_list may be specified as

- (1) A list of channel numbers (i.e. a vector of doubles).
- (2) As a list of character channel names (i.e. a character array).

 chan_list(n,:) is the name of the nth channel to be returned in array data.

 getCTFdata compares the characters preceding the first blank or '-' character to the rows of ds.res4.chanNames.

unit: Controls the conversion from data in file baseName.meg4 to user-specified units.

- 'fT', [] or not defined: SQUIDs in fT, EEGs in μV, ADC's and DACS in V, HLC channels in m, clock channels in s.
- 'T': SQUIDs in T, EEGs in V, ADC's and DACS in V, HLC channels in m.
- phi0': SQUIDs in ϕ_0 , EEGs in μV , ADC's and DACS in V, HLC channels in m.
- int': No conversion. All channels read as signed 32-bit integers.

prec: Allows the user to specify 'single' in order to reduce memory requirements. If not specified or if set to anything other than 'double' or 'single', it returns an error message and sets data=[].

All except the first input argument are optional.

getCTFdata reads the .meg4 file of a dataset, and closes it. The data are produced as a 3-index array: size(data) = [no_samples no_channels no_trials].

getCTFdata does not alter the balance of the signals recorded in the dataset. Field
grad_order_no of structure array ds.res4.senres gives the balance of the recorded data. Function
setCTFDataBalance described below is used to change data balance.

3.2.1 Examples of the application of getCTFdata:

(1) Read the entire meg4 file into a double array, converting the SQUID channels to fT, use the command

```
data=qetCTFdata(ds);
```

(2) Read points 1:10000 of trial 1 of the MEG channels in ϕ_0 units. Usually, MEG channels are distinguished by starting with 'M'.

```
MEGlist=strmatch('M',ds.res4.chanNames);
MEGdata=getCTFdata(ds,[1 10000 1]',MEGlist,'phi0');
```

3.3 setCTFDataBalance - balancing the data

External interference in the MEG gradiometers is reduced by applying balancing to the raw MEG signals. Before applying balancing to a dataset, check the balance of the recorded data by examining ds.res4.senres(k).grad_order_no where k is a MEG channel. If grad_order_no=3, then the MEG signals are already balanced.

The complete set of balance coefficients for a MEG sensor is contained in structure array ds.res4.scrr which is produced by readCTFds. setCTFDataBalance takes an array of data, reads the balance table which is stored as structure array ds.res4, and changes the data balance.

The possible balance states of the MEG sensors are

```
'none' : Unbalanced data (grad_order_no=0)
'G1BR' : 1st order balancing (grad_order_no=1)
'G2BR' : 2nd order balancing (grad_order_no=2)
'G3BR' : 3rd order balancing (grad_order_no=3)
```

The possible balance states of the reference gradiometers are

```
'none': Unbalanced data,
'G1BR': 1st order balancing
```

There are no situations where balanced reference gradiometers are needed in MEG analysis and CTF software assumes unbalanced references. Therefore, to prevent passing erroneous reference gradiometer information to other datasets, writeCTFds will not create datasets with balanced reference gradiometers.

The MATLAB command to change data balance is

[data2,ds2]=setCTFDataBalance(data1,ds1,bal2,unit,chan list,messages);

Inputs: data1: The current data (e.g. the array produced by getCTFdata).

- **ds1: ds1** is a structure produced by **readCTFds** that provides the balance coefficients for rebalancing the data and also what is the balance state of **data1**.
- bal2: Character array specifying the new MEG and reference gradiometer balance. Example: If the user wants order-3 MEG balance and unbalanced reference gradiometers, then bal2=strvcat('G3BR', 'none').
 - If the reference-gradiometer balance is omitted from **bal2** (e.g. **bal2='G2BR'**), then **setCTFDataBalance** assumes reference-gradiometer balance='none'.
 - If bal2=[], then it is set to strvcat('none', 'none') (i.e. data2 will be unbalanced).
- unit: Optional input. Options: 'fT', 'T' 'phi0', 'int'. If unit is omitted from the
 input arguments, or unit=[], it will be set to the default: unit='fT'.
- chan_list: Optional input. List of dataset channels included in array data1 (numbering
 referred to the channel numbering in structure ds). If chan_list=[] or <0 or
 chan_list is omitted from the list of inputs, it is assumed that array data1 has dataset
 channels 1:size(data,2).(i.e. channel numbered consecutively from 1). If
 chan_list is included, length(chan_list)=Nchan where size(data1)=[Npt Nchan
 Ntrial]. chan list must include the reference sensors required for balancing.</pre>

messages: Optional input. If omitted or messages=[], set messages=1.

messages=0: Do not print tracing message.

=1: Print a message when the requested balance changes.

=2: Always print a message.

Outputs: data2: Rebalanced data.

ds structure for the rebalanced data. It is the same as ds1, but grad_order_no in ds.res4.senres is set to the new balance order.

setCTFDataBalance assumes that all of the MEG sensors also have the same balance (usually grad_order_no=0 or 3). However, it is possible for the coefficients for some of the MEG channels to be missing from the balance table. When this occurs, it is not possible to balance the missing channel. If setCTFDataBalance is requested to balance a channel missing from the table, it sets data for the channel to 0, and prints a message when messages=1 or 2.

If some of the MEG channels or reference gradiometers have a different <code>grad_order_no</code> than the majority of the class of channels, these channels are considered bad, and <code>setCTFDataBalance</code> sets the corresponding part of <code>data2</code> to zero, and adds the channels to <code>ds.BadChannels</code>.

3.3.1 Examples of the application of setCTFDataBalance:

(1) Array data3 is in fT units and has 3rd-order balanced MEGs, and unbalanced reference gradiometers. To convert to unbalanced MEG data, while keeping a copy of the balanced data,

```
[data0,ds0]=setCTFDataBalance(data3,ds3,'none');
```

(2) Array data is in ϕ_0 units and has unbalanced SQUID data. To convert to 3rd-order balanced MEGs and unbalanced reference gradiometers,

```
[data,ds]=setCTFDataBalance(data,ds,'G3BR','phi0');
```

This replaces data with balanced MEGs, and updates structure ds so it shows the new balance (G3BR).

(3) A more complicated example is calculating the balanced response of MEG sensors to modeled sources. To model the SQUID response, start with the sensor descriptions in structure array ds.res4.senres. MEGlist and Reflist are lists of the MEG and reference sensors referred to the dataset channel numbering. Field grad_order_no of ds.res4.senres must be 0 for all MEG sensors and reference gradiometers.

Let LO(k,n) be the calculated unbalanced response of sensor MEGlist(k) to source n (k=1,..,length(MEGlist), size(LO)=[nMEG nSource]). R(p,n) is the response of sensor Reflist(p) to source n. Then the balanced MEG response L3 would be calculated with the following MATLAB instructions:

```
chan_list=[Reflist MEGlist];
[L3,ds3]=setCTFDataBalance([R L0]',ds,'G3BR',[],chan_list,0);
L3=L3(:,length(Reflist)+[1:length(MEGlist)],:)';
```

It is necessary to use the chan_list option in this case because the forward solution [R;L0] does not include the non-SQUID channels of the dataset (e.g. clock, trigger, EEG, and ADC channels). Note the matrix transpositions that are used to make the channel become the second index in the argument of setCTFDataBalance.

3.4 getCTFBalanceCoefs - Balancing data using the table of balance coefficients

Example 3 of the application of **setCTFDataBalance** in Section 3.3 showed how to calculate the response of balanced MEG sensors to model sources. An alternative approach is to read the balance table from structure **ds.res4.scrr** using function **getCTFBalanceCoefs** and then balance the model solution directly.

To read the balance coefficients for the MEG channels:

```
[alphaMEG,MEGlist,Refindex]=getCTFBalanceCoefs(ds,balance type,unit);
```

To read the balance coefficients for the MEG channels and the reference gradiometers.

[alphaMEG,MEGlist,Refindex,alphaGref,Greflist,Gbalanceindex] = getCTFBalanceCoefs(ds,balance type,unit);

There are no situations where balanced reference gradiometers are needed in MEG analysis and CTF software assumes unbalanced references. To prevent passing erroneous reference gradiometer information to other datasets, writeCTFds will not create datasets with balanced reference gradiometers.

Inputs: - ds is the structure that describes the dataset (output of readCTFds).

- balance_type is a character array that specifies the balance type. The balance options are 'none','G1BR','G2BR','G3BR'.
 - size (balance_type) = [1 4]. If the reference-gradiometer balance table is requested by giving 6 output arguments, getCTFBalanceCoefs extends balance_type by adding balance_type(2,:)='none'.
 - balance_type(1,:): MEG-gradiometer balance table. The options are 'none','G1BR','G2BR','G3BR'.
 - balance_type(2,:): Reference-gradiometer balance table. The options are 'none','G1BR'.
- unit: Options: 'fT', 'T' 'phi0', 'int'. If unit is omitted from the input arguments, or unit=[], it will be set to the default unit='fT'.

Outputs: The MATLAB command must specify either 3 or 6 output arguments.

- alphameg, meglist, Refindex. The MEG gradiometer balance table. Meglist is the list of the MEG sensor channels that appear in the balancing table (N_{MEG} channels) referred to the dataset channel numbering (i.e. the channel numbering in structure ds). Refindex is the list of N_{BAL} reference SQUID channels that are used for balancing MEG channels. Refindex is a subset of the list of all reference channels ($N_{BAL} \le N_{REF}$) and is referred to the dataset channel numbering. alphameg is an $N_{BAL} \times N_{MEG}$ array of balance coefficients. alphameg(k,m) is the coefficient of reference sensor Refindex (k) for balancing MEG gradiometer Meglist (m).
- alphaGref, Greflist, Gbalanceindex. The reference gradiometer balance table. Greflist is a list of all reference gradiometers (N_{GREF} channels). Gbalanceindex is the list of N_{GBAL} reference channels that are used for balancing reference gradiometers. Gbalanceindex is a subset of the list of all reference channels. alphaGref is the N_{GBAL} × N_{GREF} array of balance coefficients for the reference gradiometers. alphaGref(k,m) is the coefficient of reference sensor Gbalanceindex(k) for balancing reference gradiometer Greflist(m). Channel numbers are referred to the channel numbering in structure ds.
- When the input balance_type(k,:)='none' (k=1 or 2) is selected, getCTFBalanceData returns lists of sensors, but the balance matrix and balance-reference list are empty:

3.4.1 Examples of application of getCTFBalanceCoefs

(1) To perform the forward-solution balancing described in Example 3 in Section 3.3 above where R is the reference-sensor response, LO is the unbalanced MEG response, Reflist is a list of reference sensors referred to dataset channel numbering, and the sensor is described by structure ds, use the following commands:

```
[alphaMEG,MEGlist,Refindex]=getCTFBalanceCoefs(ds,'G3BR');
[Cx,Refx]=intersect(Reflist,Refindex);
L3=L0-alphaMEG'*R(Refx,:);
```

Note that the transpose of alphaMEG is used since size(alphaMEG)=[nBal nMEG] while size(LO)=[nMEG nSource].

(2) Suppose array data is a single trial of unbalanced data read by getCTFdata with option unit='int' and chan_list=[] (i.e. all of the dataset channels are present in array data). Then the MEG channels could be converted to order-3 balance by the following commands:

```
[alphaMEG,MEGlist,Refindex]=getCTFBalanceCoefs(ds,'G3BR','int');
data(:,MEGlist)=data(:,MEGlist)-data(:,Refindex)*alphaMEG;
```

Writing Data



4. Writing data from MATLAB to CTF format

Datasets created by writeCTFds must not be used for clinical applications.

4.1 No clinical use

The MATLAB programs supplied by VSM have been verified for correct operation, but it is not possible to know how data have been manipulated in MATLAB before being written as a CTF dataset by writeCTFds adds "NOT FOR CLINICAL USE" messages to text fields in the .res4, .infods, .hist, and if present .newds files of the new dataset.

In addition writeCTFds prints the following message on the monitor the first time it is called in a session, or after a clear all or clear writeCTFds command:

writeCTFds: The data you are writing have been processed by software not manufactured by VSM MedTech Ltd. and that has not received marketing clearance for clinical applications. These data should not be later employed for clinical and/or diagnostic purposes.

4.2 Data preparation for writeCTFds

It is expected that the user will start by reading a dataset using readCTFds and getCTFdata (Sections 3.1 and 3.2). This will create the ds structure for the dataset including the res4, and meg4 and infods fields, plus a field for each of the other files found in the dataset as listed in Table 1.

The user will manipulate the data in MATLAB. This may include balancing the data (Section 3.3), adding test signals to existing data, adding or removing channels or trials from the dataset, and possibly filtering and subsampling the data. The output data may be either single or double precision.

Before writing the dataset, the ds.res4 information must be updated as listed below:

(a) Filtering. Increase ds.res4.num_filters to the correct number of stages. As an example, if a single filter was applied to the data, the res4 structure would be changed as follows:

The only allowed value of fClass is 1. Set fType=1 for lowpass, 2 for highpass and 3 for notch filtering. This assumes that the filters defined in the .res4 file have no extra parameters, and that only Butterworth filters are applied to the data.

CTF's DataEditor does not make use of this information, but it is a simple way to record the data in the .res4 file of the dataset, and it is consistent with the way that the data-acquisition software records the filtering information. writeCTFds uses the information in ds.res4.filters to update the bandwidth parameters in the .newds and .infods files of the output dataset, and also makes a notation in the .hist file.

If ds.res4.filters is not defined, then the .infods file of the dataset will have bandwidth set to 0 to (sample rate)/4.

- (b) Subsampling. Adjust parameter ds.res4.sample_rate to match the sample rate in the data array.
- (c) Truncating data. Simply remove trials or samples as required. writeCTFds will set

```
[no_samples no_channels no_trials]=size(data)
```

and will adjust the relevant fields of the .res4 structure.

(d) Removing channels. If a channel is removed from the dataset, then remove the corresponding channel from ds.res4.chanNames and ds.res4.senres. For example if channels 100:200 have been removed from the dataset, make the following changes:

```
keeplist=setdiff(1:ds.res4.no_channels,[100:200]);
ds.res4.chanNames=ds.res4.chanNames(keeplist,:);
ds.res4.senres=ds.res4.senres(keeplist);
ds.res4.no_channels=size(ds.res4.chanNames,1);
```

writeCTFds will detect the absence of these channels, and they will be removed from the balance coefficient arrays in the output dataset. If writeCTFds detects a ds.res4.chanNames or ds.res4.senres table that is longer than the number of channels in array data, it issues an error message since writeCTFds does not know which channels have been removed. It is not really necessary to adjust ds.res4.no_channels since writeCTFds reassigns it on the basis of the size of the data array.

(e) Adding channels. Suppose one new channel is created and that it is added at the end of the array data. Before adding channel,

```
size(data)=[Ndat Nchan Ntrial]
size(newdata)=[Ndat 1 Ntrial]
```

To add data,

```
data(:,Nchan+1,:)=newdata;
ds.res4.chanNames=[ds.res4.chanNames; (new 32-character name)];
ds.res4.senres(Nchan+1)=...
```

The new channel name must include a null byte char (0) to terminate the string.

If a user does not provide names for new channels or entries in the **senres** table, **writeCTFds** will notice a difference between the lengths of array **ds.res4.chanNames** and the data array and will add names **MXT001**, **MXT002**,... at the end of **ds.res4.chanNames** (i.e. it will assume that the new channels are at the end of array data). **writeCTFds** also extends the **ds.res4.senres** table for the new channels by creating channels with **sensorTypeIndex=4**, a nominal gain of 3×10^9 ϕ_0/T to the channel, and a position 21 cm above the centre of the MEG helmet (i.e. typical MEG gradiometer gains, but the system treats it as a magnetometer, and it's position clearly marks it as different).

writeCTFds does not add balancing coefficients for the new channels, so DataEditor will not adjust the new channels when different gradient-balancing orders are requested.

Generally, it will be better if users do not depend on this default approach, but rather supply a description for added channels by making entries in ds.res4.chanNames and ds.res4.senres.

(f) Saving balanced data. The user tells CTF software (including DataEditor) the balance of channel k (a type=5 MEG sensor) by setting

```
ds.res4.senres(k).grad order no=p;
```

where p=0,1,2,3 is the gradiometer balancing order. All MEG channels (i.e. gradiometers with sensorTypeIndex=5) must have the same balance order, and a MEG channel with a different balance order is considered a bad channel and is added to the list of bad channels in ds.BadChannels. All of the reference gradiometers must have grad order no=0.

If balancing is done using **setCTFDataBalance** (Section 3.3) with the **ds** structure as an output argument, then **grad order no** will be set to the correct value by **setCTFDataBalance**.

(g) Changing text information. The following text strings appear in the .res4 file and structure ds.res4. If they are not changed, the strings read from the original dataset will be written into the output .res4 file. writeCTFds checks the lengths of the strings before writing them in order to maintain the correct res4 file format. If a string is too long the trailing characters will be removed except for fields marked with * where leading characters are removed in order to ensure that the message "NOT FOR CLINICAL USE" is not removed from the field. Four of the .res4 fields are also transferred to the infods file.

res4 field	max. chars	infods tag
* ds.res4.appName	256	
* ds.res4.dataOrigin	256	
* ds.res4.dataDescription	256	
ds.res4.data_time	255	
ds.res4.data_date	255	
ds.res4.nf_run_name	32	
* ds.res4.nf_run_title	256	_DATASET_PROCSTEPTITLE
ds.res4.nf_instruments	32	
ds.res4.nf_collect_descrip	otor 32	_DATASET_PROCSTEPDESCRIPTION
* ds.res4.nf_subject_id	32	_DATASET_PATIENTID
ds.res4.nf_operator	32	_DATASET_OPERATORNAME
ds.res4.nf_sensorFileName	56	
ds.res4.run_description	no limit	

In all cases, res4 strings are terminated by a null byte (char (0)).

In addition, text changes in the .newds, .hist and .infods fields of ds (including the message "NOT FOR CLINICAL USE") will appear in the output dataset.

4.3 Creating datasets with writeCTFds

The MATLAB command is

newds=writeCTFds (datasetname, ds, data, unit);

Inputs:

datasetname: Name of dataset (i.e. directory) to be created including the complete path. The

extension .ds is optional: if it is missing, writeCTFds will add it.

ds : The structure that describes the dataset and is created by **readCTFds**. It must

include the fields shown in Appendix A. Section 4.2 describes how to change

the ds.res4 structure for a modified dataset.

data : The array of data to be written to the new .meg4 file. It may be a single or

double precision array.

size(data)=[Ndat Nchan Ntrial]

unit : Described in the documentation for getCTFdata. This controls the conversion

of array data into 32-bit integers which are written to file datasetname.meg4. If not included in the call, or if unit=[], writeCTFds assumes unit='fT'. The

options are

(i) 'ft' or 'fT'

(ii) 't' or 'T'

(iii) 'int'

(iv) 'phi0'

The first three inputs arguments (datasetname, ds, data) must be included. Argument unit may be omitted, if the SQUID data are in fT units.

Outputs:

- **newds**: The **ds** structure for the new dataset.
- CTF dataset datasetname.

The new CTF-format dataset contains one file for each field of structure ds as listed in Table 1. If ds does not include a .hist field, it is added and indicates that the dataset was created by writeCTFds rather than CTF MEGTM software. The bandwidth information in files *.infods and *.newds is adjusted to reflect new filters and subsampling specified by the user in ds.res4. The new dataset will not include hz.ds or hz2.ds subdirectories which contain the signals recorded during pre- and post-run head localization. writeCTFds also adds messages indicating that the dataset was produced by writeCTFds and not by CTF MEGTM software and that the data must not be used for clinical purposes.

Version 1.2 of writeCTFds will create multiple meg4 files if the total data size exceeds 2 GB.

4.4 Adding trials to existing datasets with addCTFtrial

It is not uncommon for the size of a dataset to exceed the memory available for MATLAB in a computer running Windows. As an example, with a 2 GB machine memory. MATLAB will encounter memory problem if the dataset exceeds ~100 million points. In this case, if a user wants to analyze a dataset and the create a new output dataset of the same size, it will be necessary to process the data in segments, and then to add the results of later segments to the dataset created from the earlier ones. addCTFtrial adds one or more trials to a dataset previously created by writeCTFds. addCTFtrial will not add trials to datasets that were not created by writeCTFds.

The MATLAB command is

cntrl=addCTFtrial(datasetname,data,unit,mrk,cls,badSegments);

Inputs:

datasetname: Name of dataset (i.e. directory) to be created including the complete path. The

extension .ds is optional.

data : The array of data to be added to the existing .meg4 file. It may be a single or

double precision array.

size(data)=[Ndat Nchan Ntrial]

where Ntrial is the number of trials being added.

unit : Described in the documentation for getCTFdata and writeCTFds.

mrk : Markers for the trials being added. Trial=1 is the first trial being added.

cls : Classes for to the trials being added. Trial=1 is the first trial being added.

badSegments: Bad segments markings for the trials being added. Trial=1 is the first trial being

added. See description of ds.BadSegements in Appendix A.15.

Output: cntrl: 0 = failure.

1 = success.

MRI and Head Model Files



5. Reading and creating MRI and head-model files

Analysis of MEG data often requires analysis of the MRI of a subject to create a head model as part of the calculation of the magnetic field of point dipole in the brain (the "forward solution"), or to establish the anatomical location of a source determined by MEG analysis. Programs have been prepared to allow users to read MRIs and head models into MATLAB and to create files that are compatible with CTF MEGTM software (e.g. MRIViewer, DipoleFit, and the SAM beamformer programs). These files may be especially useful for users who need to convert MRIs in other formats to CTF MRI format, or who wish to do multiple-local-spheres forward solutions in MATLAB and need access to the model parameters calculated by Linux program localSpheres.

The formats of MRI and head-model files are described in document "CTF MEG™ File Formats", PN900-0088.

5.1 Reading CTF MRIs with readCTFMRI

A CTF MRI can be read into MATLAB by the following command:

[MRItags, MRIdata] = readCTFMRI (MRIfile);

where

MRIfile is the name of a v4 CTF-format MRI including the full path and extension '.mri'. CTF MRI files (version 4) are in the CPersist format (like .infods and .acq)

MRItags is a structure array listing the names, types and contents of the tags found in the file.

The tags are listed in the reference.

MRIdata is a 256x256x256 int16 array containing the MRI. MRIdata(j,k,n) refers to coronal slice j, axial slice k, sagittal slice n. j=1: anterior, j=256: posterior, k=1: superior, k=256: inferior, n=1: left, n=256: right.

Tag 'CTFMRI_TRANSFORMMATRIX' is a text string giving the 4×4 matrix that connects positions in CTF Head Coordinates and MRI pixel coordinates. The matrix elements are separated by a '\' character:

string='M11\M12\M13\...\M43\M44'.

Let d = pixel size. Then the relation between physical positions (x,y,z) in CTF Head Coordinates and MRI pixel coordinates (j,k,n) is

$$\begin{bmatrix} M_{11} & M_{12} & M_{12} & M_{14} \\ M_{21} & M_{22} & M_{23} & M_{24} \\ M_{31} & M_{32} & M_{33} & M_{34} \\ 0 & 0 & 0 & 1 \end{bmatrix} \times \begin{bmatrix} x/d \\ y/d \\ z/d \\ 1 \end{bmatrix} = \begin{bmatrix} j \\ k \\ n \\ 1 \end{bmatrix}$$

(In this matrix $M_{41} = M_{42} = M_{43} = 0$, $M_{44} = 1$ and $[M_{14} \ M_{24} \ M_{34}] =$ pixel coordinates of the coordinate-system origin). CTF Head Coordinates are defined in Appendix A of the document "CTF MEGTM File Formats", PN900-0088.

5.1.1 No clinical use of readCTFMRI

readCTFMRI prints the following message on the monitor the first time it is called in a session, or after a clear all or clear readCTFMRI command:

readCTFMRI: You are reading a CTF-format MRI for use with a software-application tool that is not manufactured by VSM MedTech Ltd. and has not received marketing clearance for clinical applications. If the MRI is processed by this tool, it should not be later employed for clinical and/or diagnostic purposes.

5.2 Creating CTF MRIs with writeCTFMRI

writeCTFMRI reverses the operation of readCTFMRI and produces an MRI file compatible with MRIViewer. The call is

writeCTFMRI(MRIfile,MRItag,MRIdata);

where

MRIfile is the name of the file to be created including the full path and extension '.mri'.

MRItags is a structure array in the format produced by readCTFMRI listing the names, types and contents of the tags found in the file. The tags are listed in the MRI format section of document "CTF MEGTM File Formats", PN900-0088.

MRIdata is a 256x256x256 array containing the MRI. MRIdata(j,k,n) refers to coronal slice j, axial slice k, sagittal slice n. j=1: anterior, j=256: posterior, k=1: superior, k=256: inferior,. n=1: left, n=256: right. MRIdata is converted to int16 when it is written to file MRIfile.

This program will be useful for users who need to modify the CTF MRI (for example, by expanding or shrinking it, or by processing it to remove artifacts in the MRI), or who need the MRI to prepare plots in MATLAB. Alternatively, writeCTFMRI provides a way to convert MRIs in other formats (e.g. AFNI) to CTF MRI format if the other formats can be read into MATLAB.

5.2.1 No clinical use of writeCTFMRI

The message "NOT FOR CLINICAL USE" is added to the patient identification and comment tags in the output MRI file. In addition writeCTFMRI prints the following message on the monitor the first time it is called in a session, or after a clear all or clear writeCTFMRI command:

writeCTFMRI: The MRI you are creating has been processed by software not manufactured by VSM MedTech Ltd. and that has not received marketing clearance for clinical applications. This MRI should not be later employed for clinical and/or diagnostic purposes.

5.3 Reading CTF head-model files with readCTFhdm

CTF MEGTM program **DipoleFit** and the SAM beamformer software use single or multiple local sphere models of the subject's head. These models are based on the positions of head coils and the shape of the head determined from the MRI. Command-line program **localSpheres** produces a text file with extension .hdm (usually the filename is default.hdm) containing the centre and radius of the local sphere appropriate for each MEG sensor.

readCTFhdm reads the head-model file into a structure **hdm**. The head-model file is a text file in the CTF "Config Reader" format (see "CTF MEGTM File Formats", PN900-0088). The command is

hdm=readCTFhdm(hdmFile)

where

hamfile is the name of the head-model file including the path and .ham extension

hdm is a structure containing the information read from hdmFile. It has a field for each of the Config Reader classes identified in file hdmFile, and fields named note contain comment lines read from the file.

The most important part of structure hdm is hdm.MultiSpheres_Data which contains the list of SQUID sensor names and the local sphere for each. Version 5 head-model files have the following sub-fields:

```
hdm.MultiSphere Data =
```

note: [2x34 char]

SEARCH RADIUS: 7

HEADSHAPE FILE: 'MRI/s2 20051209 matched fids head.shape'

SQUIDname: [301x5 char] sphereOrigin: [3x301 double] radius: [1x301 double]

Release 5.4 of the CTF software assumes version 6 head-model files in which two sub fields are added to **MultiSphere Data**:

```
hdm.MultiSphere_Data.SURFACE_TYPE: 'SCALP' OF 'BRAIN"
```

.HEADPOS.HEADPOS: List of head coil positions in the dewar reference

frame (same information as ds.hc.dewar)

.HEADPOS.NOMINAL: 'TRUE' Or 'FALSE'

The SEARCH_RADIUS parameter is the radius (cm) of the head patch that is used to estimate the centre and radius of the local sphere. Character array SQUIDname is a list of all the SQUID sensors in the dataset that was used to create the head-model file. sphereOrigin and radius are the sphere parameters in cm for each of the SQUID sensors. The coordinate system is CTF Head Coordinates. The sensor positions in this coordinate frame are given in the pos field of the ds.res4.senres structure array (see Appendix B.2).

Command-line program **localSpheres** is described in more detail in documents "Command Line Programs Guide", PN900-0016, and SAMsuite Guide, PN900-0037.

5.3.1 No clinical use of readCTFhdm

readCTFhdm prints the following message on the monitor the first time it is called in a session, or after a clear all or clear readCTFhdm command:

readCTFhdm: You are reading a CTF head model for use with a software-application tool that is not manufactured by VSM MedTech Ltd. and has not received marketing clearance for clinical applications. If CTF MEG data are processed by this tool, they should not be later employed for clinical and/or diagnostic purposes.

5.4 Creating CTF head-model files with writeCTFhdm

writeCTFhdm allows creation of a single- or multiple-local-spheres head model that can be used by DipoleFit and the SAM programs to generate forward solutions for MEG analysis. The command is

```
writeCTFhdm(hdmFile,hdm);
```

where the arguments have the same meaning as for the complementary program readCTFhdm:

hamfile is the name of the head-model file including the path and .ham extension

hdm is a structure in the form produced by **readCTFhdm** containing the information required to create a head-model file. It has a field for each of the Config Reader classes that will be created in **hdmFile**.

To create a multiple-local-sphere model, structure hom must have a field MultiSphere_Data in the format discussed in Section 5.3 (readCTFhom).

A single-sphere MEG model can be defined by field **MEG Sphere**. An example is:

```
hdm.MEG_Sphere =
note: ''
ORIGIN_X: -2.0030
ORIGIN_Y: -0.1110
ORIGIN_Z: 4.6580
RADIUS: 9.0850
```

This defines the origin in cm in CTF Head Coordinates.

writeCTFhom produces both version 5 and version 6 hdm files depending on the presence of the new subfields in hom.MultiSphere Data.

writeCTFhdm requires fields Model and one or both of MEG_Sphere and MultiSphere_Data (see "CTF MEGTM File Formats", PN900-0088 for a description of the classes in .hdm files). Without these fields it is not a head-model file.

5.4.1 Converting head-model files from v5 to v6

v6 head-model files have two tags added to the MultiSphere_Data class and the version number changed in the File_Info class. This extra information is required by v5.4 Linux command <code>DipoleFit</code>, and if it is not present an error message will be printed. An example of a v5 text file (Config Reader format) is

```
File_Info
{
    VERSION:    CTF_HEAD_MODEL_FILE_VERSION_5.0
    DATE:    18-Oct-2006 16:05
    PATIENT:    none
    STATUS:
}
...
MultiSphere_Data
{
    SEARCH_RADIUS: 7.000
    HEADSHAPE_FILE:    MRI/head.shape

    // Multiple Sphere locations in cm
    //    X    Y    Z    Radius
    BG1:    -2.950 -0.038 5.544 8.365
    BG2:    -2.950 -0.038 5.544 8.365
```

The v6 file has tags for head-coil positions (in dewar coordinates) and a keyword to indicate whether this is a scalp or brain surface:

The simplest way to change from 5 to v6 is to regenerate the head-model file using the v5.4 **localSpheres** command. A second way is to edit the v5 head model file with a Linux text editor to add the extra information as shown above. Note that Config Reader format separates fields by tabs. Finally, the following MATLAB script could be used where the head-model file name is **default.hdm**:

5.4.2 No clinical use of writeCTFhdm

The message "NOT FOR CLINICAL USE" is added to the comment section at the start of the file. In addition writeCTFhcm prints the following message on the monitor the first time it is called in a session, or after a clear all or clear writeCTFhcm command:

writeCTFhdm: The head-model data you are writing have been processed by software not manufactured by VSM MedTech Ltd. and that has not received marketing clearance for clinical applications. These data should not be later employed for clinical and/or diagnostic purposes.

Appendix A: Data Structure



Appendix A: The data structure produced by readCTFds

ds is the structure produced by readCTFds:

ds=readCTFds (dsName)

The fields of ds are listed in Table 1 (page 5). This Appendix describes the structure and information in each field of ds while Appendix B describes ds.res4 in detail. The format and information in the dataset files are described in the document "CTF MEGTM File Formats", PN900-0088. The function of many of the fields is simply to store information that can later be used by writeCTFds to create a CTF-format dataset. For operations within MATLAB, only the baseName, path and res4 fields are required.

A.1 ds.baseName

Character string containing the name of the dataset, without path information and without the .ds extension. E.g.: ds.baseName='AbsCal Noise 20050301 04'.

A.2 ds.path

Character string containing the complete path for the dataset, including the final file-separator character. E.g.:

Windows:ds.path= S:\CHL\NIMH_20060606\

Linux: ds.path=/home/hwilson/data/CHL/NIMH 20060606/

A.3 ds.res4

The contents of the file [baseName,'.res4']. This field contains the description of the system sensors in substructure ds.res4.senres, and the balance tables in ds.res4.serr. ds.res4 is a complicated field, and Appendix B discusses ds.res4 in detail. It is the most important part of ds. For a 275-channel CTF MEG system, structure ds has a size of about 4 MB, of which 3.6 MB is the .res4 field.

A.4 ds.meg4

This gives the 8-bytes MEG4 file header and the size of the MEG4 file in bytes

```
E.g.: ds.meg4=

fileSize: 961920008

header: 'MEG41CP'
```

A.5 ds.infods

This is a structure array containing all of the information listed in the file baseName.infods. The file structure (CPersist format) is listed in Appendix B of "CTF MEGTM File Formats", PN900-0088. Each element of ds.infods contains the name, type and data of one of the fields in the infods file:

An example of one element of the structure array is

The CPersist reading function also creates two data types in addition to the types listed in the reference:

type=0: The string 'ws1' that marks the start of a CPersist object.

For data processing, the important .infods fields are the bandwidth fields

```
DATASET LOWERBANDWIDTH and DATASET UPPERBANDWIDTH
```

and the fields that describe head motion when Continuous Head Localization is applied:

```
_DATASET_HZ_MODE,
_DATASET_MOTIONTOLERANCE,
_DATASET_MAXHEADMOTION,
_DATASET_MAXHEADMOTIONTRIAL,
DATASET_MAXHEADMOTIONCOIL.
```

A.6 ds.newds

ds.newds is a simple character image of the file baseName.newds. writeCTFds updates the bandwidth information in ds.newds, indicates that the dataset was created by writeCTFds, and adds the no-clinical-use message "NOT FOR CLINICAL USE" in the appName field.

ds.newds will have no application within MATLAB. If newds is not a field of ds, writeCTFds will not indicate that ds.newds is missing. The newds file is obsolete, and is now replaced by the infods file.

A.7 ds.acq

This is a structure array containing all of the information listed in the file baseName.acq which describes the electronics and acquisition setup. The file structure (CPersist) format is listed in Appendix B of "CTF MEGTM File Formats", PN900-0088. Each element of ds.acq contains the name, type and data of one of the fields in the baseName.acq.

```
ds.acq=
1x1319 struct array with fields:
name
type
data
```

The acq file is read simply to enable writeCTFds to create an .acq file in the output dataset. It is unlikely that MATLAB users will need to access ds.acq.

A.8 ds.hist

ds.hist is a simple character image of file baseName.hist. writeCTFds adds a record of filter information from ds.res4, indicates that the dataset was created by writeCTFds, and adds the message "NOT FOR CLINICAL USE".

ds.hist will have no application within MATLAB. If hist is not a field of ds, writeCTFds will create it.

A.9 ds.hc

ds.hc contains the information from the head-coil position file baseName.hc. It records the head coil positions in three frames: "standard", "measured" and "head". "measured" gives the head coil positions in cm in the dewar coordinates (origin at the centre of the reference-sensor array, 21 cm above the centre of the MEG helmet. "head" gives head coil positions in CTF head coordinates. If there are 3 head coils in the .hc file then

```
ds.hc =
    names: [3x6 char]
    standard: [3x3 double]
    dewar: [3x3 double]
    head: [3x3 double]
```

ds.hc.dewar(:,k) is the position of coil k (name=ds.hc.names(k,:)), in the dewar reference frame.

readCTFds and writeCTFds will also handle .hc files with 4 head coils.

$A.10\,ds.eeg$

If file baseName.eeg exists, ds.res4.eeg is a list of EEG channels (channel number in the dataset), channel names and electrode positions in cm in CTF Head Coordinates. If 32 EEG sensors were defined, then

A.11ds.mrk

ds.mrk is a structure array with one element for each marker set found in MarkerFile.mrk. An example of an element of ds.mrk is

```
ds.mrk(2) =
   ClassGroupId: 0
        Name: 'Tr18'
   Comment: 'Trigger Marker corresponding to bit 18'
        Color: 'Red'
   Editable: 'No'
   ClassId: 2
   BitNumber: 18
   Polarity: '-'
        Source: 'Source is not defined'
   Threshold: 0
        trial: [1x83 double]
        time: [1x83 double]
```

These field names are taken directly from the keywords in MarkerFile.mrk (see document "CTF MEGTM File Formats", PN900-0088, for the file format), but note the addition of BitNumber, Polarity, Source and Threshold for ClassGroupID=0 markers. These four fields are empty for ClassGroupID=3 markers.

Inside MarkerFile.mrk, trial numbering starts at 0, but in ds.mrk it starts at 1. This means it will be easy to match ds.mrk(k).trial to data(:,:,k) when the whole dataset is read, but if only a few

trials are read by getCTFdata (e.g. data_list=2), then it will be necessary to edit the trial and time arrays within ds.mrk.

A.12ds.TrialClass

readCTFds forms ds.TrialClass from ClassFile.cls, if it exists. ds.TrialClass is a structure array with one element for each class set found in ClassFile.cls. ds.TrialClass has 7 fields:

In ClassFile.cls, trial numbering starts at 0, but in ds.TrialClass it starts at 1. This means it will be easy to match ds.TrialClass (k).trial to data(:,:,k) when the whole dataset is read, but if only a few trials are read by getCTFdata (e.g. data_list=2), then it will be necessary to edit the trial arrays within ds.TrialClass.

A.13 ds.badSegments

If file bad.segments exists, readCTFds stores information about segments of data marked BAD in ds.badSegments. bad.segments is usually generated by DataEditor. An example is

Trial numbering starts at 1. If writeCTFds finds this field in structure ds, it will create file bad.segments. This field can be used to mark artefacts or bad segments found during MATLAB analysis of the data, and then to pass the information to programs such as DataEditor or the SAM beamformer programs.

A.14ds.Virtual

DataEditor allows the creation of virtual channels as weighted sums of real channels, and stores the definitions in file **VirtualChannels**. **readCTFds** creates **ds.Virtual** as a way of passing the definitions to **writeCTFds** for creation of an output dataset. It is a structure array with one element for each virtual channel. An example is

It is not likely that ds. Virtual will be used within MATLAB.

A.15 ds. BadChannels

The BadChannels file is a text file produced by the data-acquisition program or by DataEditor that lists SQUID channels that should not be used in data analysis. readCTFds simply copies this file. ds.BadChannels is a character array with one line for each bad channel. An example is

```
ds.BadChannels = size(ds.BadChannels)=[2 5]
Q11
MRO21
```

This provides a simple way to pass back to DataEditor a list of bad channels. An alternative would be to remove the bad channels from data, ds.res4.chanNames, and ds.res4.senres.

A.16 ds.processing

ds.processing is a simple character image of the file processing.cfg which records filter and balancing commands that have been executed by DataEditor. writeCTFds copies ds.processing to file processing.cfg in the output dataset.

ds.processing will have no application within MATLAB. If processing is not a field of ds, writeCTFds will not indicate that ds.processing is missing.

CTF | MEG[™]

Appendix B: Structure ds.res4



Appendix B: Structure ds.res4

Reference: "CTF MEG™ File Formats", PN900-0088.

B.1 Fields of structure ds.res4.

Here is an example of a MATLAB listing of ds.res4 for a 275-channel CTF MEG system. The names of most fields indicate their meaning.

```
ds.res4 =
                   header: 'MEG42RS '
                  appName: ''
               dataOrigin: ''
          dataDescription: ''
           no trials avgd: 0
                data time: '16:37'
                data date: '06-Jun-2006'
               no_samples: 720000
              no channels: 334
              sample rate: 2400
               epoch_time: 300
                no trials: 1
               preTrigPts: 0
           no trials done: 0
        no trials display: 0
              save trials: 0
           primaryTrigger: -65536
      triggerPolarityMask: 0
             trigger_mode: 0
       accept reject Flag: 0
         run time display: 0
           zero Head Flag: 0
            artifact mode: 0
              nf_run_name: '
             nf run title: 'CHL test'
           nf instruments: '
    nf collect descriptor: 'SEF no motion
            nf subject id: 'Anonymous
              nf operator: 'Staff
        nf_sensorFileName: ''
                    rdlen: 14
          run description: 'SEF no motion '
              num filters: 0
                  filters: []
                chanNames: [334x32 char]
                   senres: [1x334 struct]
```

numcoef: 1393

scrr: [1x1393 struct]

B.2 ds.res4.senres

ds.res4.senres is a structure array containing the channel descriptions and sensor calibration data (it contains the same information as the .sens file). For a typical SQUID channel (a 3rd-order balanced MEG gradiometer in this case) the structure is

sensorTypeIndex: 5 originalRunNum: 0 coilShape: 0 properGain: -2.8982e+009 qGain: 1048576 ioGain: 1 ioOffset: 0 numCoils: 2 grad order no: 3 gain: -0.3291 pos0: [3x2 double] ori0: [3x2 double] area: [2.5472 2.5472] numturns: [2 2] pos: [3x2 double] ori: [3x2 double]

All SQUID channels look like this except that **sensorTypeIndex** is 0 for reference magnetometers and 1 for reference gradiometers, and **numCoils** is 1 for magnetometers. Field **area** is the loop area in cm². The possible values of **sensorTypeIndex** are listed in the RES4 File Format Section of the document "CTF MEGTM File Formats", PN900-0088.

B2.1 Gain parameters in ds.res4.senres

To convert SQUID data from the 32-bit integer format of the dataset to ϕ_0 units, divide the raw data by qGain (= 2^{20} for the current generation of SQUID electronics). The physical unit of qGain is ϕ_0^{-1} .

To convert data from the 32-bit integer format of the dataset to physical MKS units (T), divide by ioGain, properGain and qGain:

data in physical units = (raw integer data)/(ioGain*properGain*qGain)

The physical unit of **properGain** is ϕ_0/T .

To convert to fT, multiply by an extra factor 10^{15} . ds.res4.senres field gain is given by

```
gain = 10<sup>15</sup>/(ioGain*properGain*qGain) (SQUIDs)
gain = 1/(ioGain*properGain*qGain) (non-SQUID channels)
```

For SQUID channels multiplying by gain converts 32-bit integer data to fT and the physical unit of gain is fT. Field gain is calculated by readCTFds and is not part of the res4 file.

If getCTFdata encounters a channel with ioGain*properGain*qGain=0, it sets the channel to 0. This allows properGain=0 to be used to mark bad channels.

B.2.2 Sensor position and orientation information

Arrays **pos0** and **ori0** (size=[3 numCoils]) are the Cartesian vectors for the pickup-loop positions (cm) and orientations (unit vector) in "dewar" coordinates where the reference sensors are located at height z=0 and the coordinate frame is rotated 45° so \hat{x} is 45° to the subject's right. These are the values that would be read from the .sens file. Arrays **pos** and **ori** are the positions and orientations after conversion to the CTF Head Coordinates (or to room coordinates in the case of fMEG sensors). If a Head Localization is done as part of a dataset, then **pos** and **ori** are referred to coordinates established by the head coils: the origin ([0 0 0]) is the midpoint of the left and right coils, and the \hat{x} direction is the line from the origin to the nasion coil. If no Head Localization is done, then **pos** and **ori** refer to an origin 21 cm below the reference sensors, where \hat{z} is the dewar axis, and \hat{x} points forward (toward the subject's nose).

Fields pos0, ori0, area, numturns, pos and ori appear in the structure array ds.res4.senres.

	pos0, ori0	pos, ori (no head zeroing)	pos, ori (with head zeroing)
\hat{x} (coord 1)	45° right of nose	nose	Line from [0 0 0] through the nasion coil
\hat{y} (coord 2)	45° left of nose	left	$\hat{z} \times \hat{x}$
\hat{z} (coord 3)	up (dewar axis)	up (dewar axis)	⊥ to the head-coil plane
Origin	Centre of the reference sensors	21 cm below references	Mid point between left and right head coils

B.2.3 grad order no (field of ds.res4.senres)

Field grad_order_no indicates what is the gradient balancing of data read from the dataset. Function setCTFDataBalance updates grad_order_no when an output data structure is specified in the output-argument list.

It is important that all of the MEG channels have the same <code>grad_order_no</code> because channels with different values of <code>grad_order_no</code> may indicate bad or disabled channels. <code>setCTFDataBalance</code> sets these channels to zero and adds them to the list of bad channels in <code>ds.BadChannels</code>.

B.3 Structure ds.res4.scrr

This structure array contains all of the balancing data for the sensor. The number of records in this structure is much greater that the number of SQUID channels because ds.res4.scrr contains 5 sets of MEG balancing coefficients (G1BR, G2BR, G3BR, G1OI, G2OI; the last two types are not true balance tables), plus G1BR coefficients for balancing the reference gradiometers. A typical element of the structure array is

```
ds.res4.scrr(101) =
```

sensorName: [1x32 uint8] 32-character sensor name
coefType: [1x4 uint8] 4-character balance type
numcoefs: 17 Number of reference sensors

sensor: [31x50 uint8] List of 31-character names of reference sensors

coefs: [50x1 double] The balance coefficients

There is room for coefficients for 50-term balancing in ds.res4.scrr, but only numcoefs terms are used (the remaining elements are zeros). numcoefs is 8 for order-1 and order-2 balancing, and 15-18 for order-3.

The documentation for **getCTFBalanceCoefs** (Section 3.4) shows how to use the balance coefficients.





