DPPP (the Default Preprocessing Pipeline, previously NDPPP for New Preprocessing Pipeline) is the LOFAR data pipelined processing program. It can be used to do all kind of operations on the data in a pipelined way, so the data are read and written only once.

DPPP started as a new and faster version of IDPPP. The original differences can be seen here.

DPPP preprocesses the data of a LOFAR observation by executing steps like flagging or averaging. Such steps can be used for the raw data as well as the calibrated data by defining the data column to use. One or more of the following steps can be defined as a pipeline. DPPP has an implicit input and output step. It is also possible to have intermediate output steps.

DPPP comes with quite some predefined steps, but it is possible to plugin arbitrary steps, either implemented in C++ or Python.

The following steps are possible:

- Flagging and Filtering
 - **AOFlagger** for automatic flagging in time/freq windows using Andre Offringa's advanced aoflagger.
 - **Preflagger** to flag given baselines, time slots, etc.
 - **UVWFlagger** to flag based on UVW coordinates, possibly in the direction of another source.
 - **MADFlagger** for automatic flagging in time/freq windows based on median filtering.
 - **Filter** to filter on baseline and/or channel (only the given baselines/channels are kept). The reader step has an implicit filter.
- Averaging
 - Averager to average data in time and/or freq.
- Phase Shifting
 - **PhaseShift** to shift data to another phase center.
- **Demixing** to remove strong sources (A-team) from the data.
 - **Demixer** to demix in the old way.
 - **SmartDemixer** to demix in a new, smarter way.
- Station summation
 - StationAdder to add stations (usually the superterp stations) forming new station(s) and baselines.
- Counter to count the number of flags per baseline, frequency, and correlation. A flagging step also counts how many visibilities it flagged. Counts can be saved to a table to be plotted later using function plotflags in python module lofar.dppp.
- Data calibration and Data scaling
 - ApplyCal to apply an existing calibration to a MeasurementSet.
 - GainCal to calibrate gains using StefCal.
 - **DDECal** to calibrate direction dependent gains.
 - **Predict** to predict the visibilities of a given sky model.
 - **H5ParmPredict** to predict visibilities corrupted by an instrument model (in H5Parm)
 - **ApplyBeam** to apply the LOFAR beam model, or the inverse of it.
 - ScaleData to scale the data with a polynomial in frequency (based on SEFD of LOFAR stations).
 - **Upsample** to upsample visibilities in time
 - **Out** to add intermediate output steps

- Interpolate for improving the accuracy of data averaging.
- User defined steps provide a plugin mechanism for arbitrary steps implemented in C++.
- **Python defined** steps provide a plugin mechanism for arbitrary steps implemented in Python.

The input is one or more (regularly shaped) MeasurementSets (MSs). The data in the given column are piped through the steps defined in the parset file and finally written (if needed). It makes it possible to, say, flag at the full resolution, average, flag on a lower resolution, average further, and finally write the data.

Regularly shaped means that all time slots in the MS must contain the same baselines and channels. DPPP can handle only one spectral window. If the MS has multiple spectral windows, one has to be selected.

If multiple MSs are given as input, their data are combined in frequency. It means that the time, phase direction, etc. of the different MSs have to be the same. Note that other steps (like averaging) can still be used.

When combining MSs (thus combining subbands), it is possible that one or more of them do not exist. Flagged data will be inserted for them. The missing frequency info is deduced from the other subbands. Note that in order to insert missing subbands in the data, the names of the missing MSs have to be given at the right place in the list of MS names. Otherwise DPPP does not know that subbands are missing.

The output can be a new MeasurementSet, but it is also possible to update the flags if the input is a single MS. If averaging or phase-shifting to another phase center is done, the only option is to create a new MeasurementSet.

At the end the run time is shown. Note that on a multi-core machine the user time can exceed the elapsed time (user time is counted per core). By default the percentage of time each step took is also shown.

The AOFlagger, MADFlagger, and Demixer, by far the most expensive parts of DPPP, can run multithreaded if DPPP is built with OpenMP. It is possible to define the number of threads to use by the global key numthreads. Is that is not set, it uses the environment variable OMP_NUM_THREADS. If also that variable is undefined, an DPPP run uses as many threads as there are CPU cores. Thus if multiple DPPP runs are started on a machine, the default total number of threads will exceed the number of CPU cores.

MeasurementSet Access

- The 'msin' step defines which MS and which DATA column to use. It is possible to specify multiple MSs using a glob-pattern or a vector of MS names.
- If multiple MSs are given, they will be concatenated in frequency. It means that all MSs must have the same times, baselines, etc. Flagged data can be inserted for MSs that are specified, but do not exist.
- It is possible to select baselines and/or a band (spectral window) and/or skip leading or trailing channels. This is the same for each input MS.
- Optionally proper weights can be calculated using the auto-correlation data.
- It sets flags for invalid data (NaN or infinite).
- Dummy, fully flagged data with correct UVW coordinates will be inserted for missing time slots in the MS. This can only be done if a single input MS is used.

- Missing time slots at the beginning or end of the MS can be detected by giving the correct start and end time. This is particularly useful for the imaging pipeline where BBS requires that the MSs of all subbands of an observation have the same time slots. When updating an MS, those inserted slots are temporary and not put back into the MS.
- The 'msout' step step defines the output. If a band is selected, the output MS (including its SPECTRAL_WINDOW subtable) contains that band only (its id is 0). The input MS is updated if no output name is given or if the output name is equal to the input name or equal to a dot.

The calculation of the weights is done as follows.

```
Weight[ANT1_POL1, ANT2_POL2] = N / (autocorr[ANT1_POL1] *
autocorr[ANT2_POL2])
N = EXPOSURE * CHAN_WIDTH * WGHT
```

where WGHT is the weight put in by RTCP (number of samples used / total number of samples). This note discusses weighting in some more detail.

Flagging

It is important to realize that a MeasurementSet contains columns FLAG and FLAG_ROW to indicate if data are flagged. If FLAG_ROW is set, all data in that row are flagged. DPPP will set FLAG_ROW if all FLAG are set (and vice-versa).

When clearing the flags manually, it is important to realize that both columns have to be cleared. For example:

taql 'update my.ms set FLAG=F, FLAG_ROW=F'

DPPP flagging behaviour is as follows.

- If one correlation is flagged, all correlations will be flagged (e.g. XX,YX,YY are flagged if XY is flagged).
- The msin step flags data containing NaNs or infinite numbers or if FLAG_ROW is set.
- An AOFlagger step can be used to flag using Andre Offringa's rficonsole code. Because DPPP always reads entire time slots, the flagging can be done on limited time windows only (depending on the available memory). An overlap can be defined to reduce boundary effects. By default QUALITY subtables will be created containing statistical flagging quality information. They can be inspected using tools like aoqplot.

The default strategy works well for HBA data, but not for LBA data. The strategy LBAdefault should be used for it.

• A Preflagger step can be used to flag (or unflag) on time, baseline, elevation, azimuth, simple uv-distance, channel, frequency, amplitude, phase, real, and imaginary. Multiple values (or ranges) can be given for one or more of those keywords. A keyword matches if the data matches one of the values. The results of all given keywords are AND-ed. For example, only data matching given channels and baselines are flagged.

Keywords can be grouped in a set making it a single (super) keyword. Such sets can be OR-ed or AND-ed. It makes it possible to flag, for example, channel 1-4 for baseline A and channel 34-36 for baseline B. Here it is explained in a bit more detail.

• A UVWFlagger step can be used to flag on UVW coordinates in meters and/or wavelengths. It is possible to base the UVW coordinates on a given phase center. If no phase center is given, the

UVW coordinates in the input MS are used.

• A MADFlagger step can be used to flag on the amplitudes of the data. It flags based on the median of the absolute difference of the amplitudes and the median of the amplitudes. It uses a running median with a box of the given size (number of channels and time slots). It is a rather expensive flagging method with usually good results.

The flagging parameters can be given as an expression to make them dependent on baseline length.

It is possible to specify which correlations to use in the MADFlagger. Flagging on XX only, can save a factor 4 in performance.

Furthermore it is possible to only flag the auto-correlations and apply the results to the crosscorrelations with a baseline length within optionally given limits.

Averaging

- Unflagged visibility data are averaged in frequency and/or time taking the weights into account. New weights are calculated as the sum of the old weights. Some older LOFAR MSs have weight 0 for unflagged data points. These weights are set to 1.
- The UVW coordinates are also averaged (not recalculated).
- It fills the new column LOFAR_FULL_RES_FLAG with the flags at the original resolution for the channels selected from the input MS. It can be used by BBS to deal with bandwidth and time smearing.
- Averaging in frequency requires that the average factor fits integrally. E.g. one cannot average every 5 channels when having 256 channels.
- When averaging in time, dummy time slots will be inserted for the ones missing at the end. In that way the output MeasurementSet is still regular in time.
- An averaged point can be flagged if too few unflagged input points were available

Demixing

Demixing (or Smart Demixing explained below) is a faster and more flexible way of the old demixing python script to demix and subtract strong sources (A-team). Jones matrices can be estimated for the direction of the subtract-sources, model-sources, and the optional target-source.

- It is possible to have different averaging for the demix and subtract step.
- Selected (e.g. shorter) baselines can be demixed (others will be averaged only). By default only the cross-correlations are used.
- Four different direction types can be given:
 - The subtract-sources are subtracted from the data. They must have a source model.
 - The model-sources can be given to take the contribution of other strong sources into account when solving for the gains. They must have a source model as well. The target source should NOT be part of this list.
 - $\circ\,$ The other-sources directions are taken into account when demixing. They are projected away when solving for the gains.
 - If the target source is given, it must have a source model and no other-sources can be given. If no target source is given, the target direction can be projected away like the extra-sources. Weak target sources should not be projected away.
- A source model mentioned above is the patch name in the SourceDB (e.g. CasA). At the moment only point and Gaussian sources are supported. The direction used for demixing is the

- It is important to note that the target source model must NOT be given using the subtractsources or model-sources. If it has to be used, give it using the targetsource parameter.
- The Jones matrices will be estimated jointly for all directions, so better results are expected if the sources are close to the target. However, joint estimation of the Jones matrices for all directions is slower than estimating the Jones matrices for each direction separately. In the near future an option will be added to estimate the Jones matrices for each direction separately like the old demixing script is doing.

Smart Demixing

Smart Demixing does demixing as above, but in a smarter way using a scheme developed by Reinout van Weeren. For each time chunk (say 2 minutes) it is decided how to demix.

It needs three source models, which are made from a text file using makesourcedb. Note that for performance it is best to run makesourcedb with parameter outtype=blob.

- A detailed model of the A-team sources used in the solve and subtract steps.
- A coarse model of the A-team sources used in the estimate step. If not given, the detailed model will be used.
- A model of the target field. Usually the user can create it from the GSM using gsm.py.

Smart demixing works as follows:

- If an A-team source is at about the same position as a source in the target model, the source is removed from the A-team list and its detailed model replaces the source in the target model used in the solve step (not for the estimate step).
- Using the coarse A-team model, the visibilities are estimated per baseline for each A-team source. By default the beam model is applied to get the apparent visibilities. The sources and baselines are selected for which the maximum amplitude exceeds a given threshold. A source/station will be solved for if the station appears in at least N of the selected baselines for that source. A detailed source model is used in that step to get as accurate gains as possible.
- The visibilities of the target are estimated in a similar way using the target model. The target is included in the solve if its maximum amplitude exceeds a threshold or if the amplitude ratio Target/Ateam exceeds a threshold. The target is also included if it is close to an A-team source and the ratio exceeds another (smaller) threshold. Otherwise, the target is ignored (if close) or deprojected.

A detailed decision tree that the smart demixing algorithm follows is available here.

When solving for the complex gains of the selected A-team sources, the detailed A-team model is used to get the correct gains. Note that by default the sources/stations not solved for are still used in the solve step. There Jones matrices will have a small gain value on the diagonal and zeroes for the off-diagonal values.

At the end a log is produced showing how the demixing behaved. It shows:

- percentage of converged solves and the average number of iterations used for them.
- percentage of times the target was included, deprojected, and ignored.
- percentage of times a source/station was solved for (thus matched the threshold/ratio criteria).

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• average and standard deviation of percentage amplitude subtracted per source per baseline

Phase shifting

- Data can be shifted to another phase center.
- A shift step can shift back to the original phase center (by giving an empty center). If that is done by the last shift step, no new MS needs to be created.

Upsample

Upsampling data can be useful for at least one use case. Consider data that has been integrated for two seconds, by a correlator (the AARTFAAC correlator) that sometimes misses one second of data. The times of the visibilities will then look like [0, 2, 4, 7, 9, 12], each having integration time 2 seconds. DPPP will automatically fill missing time slots, which will lead to times [0, 2, 4, 6, 7, 9, 11, 12]. This is still a nonuniform time coverage, which is not desirable. Calling the upsample step with timestep=2 on this data will create times [0, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13] (it will remove the inserted dummy time slots that overlap, i.e. at 7 and 12). This data is then useful for further processing, e.g. averaging to 10 seconds.

Station summation

- One or more new stations can be defined from a list of existing stations. An existing station can occur in only one new station.
- The data of baselines containing only one of the stations are added to form a new baseline.
- Optionally the auto-correlations can be added to form a new auto-correlation 'baseline'.
- The data can be added with or without weight.
- Optionally averaging instead of summing can be done.

Data scaling

- The data can be scaled with a polynomial in frequency to correct for the SEFD of the LOFAR stations.
- The default coefficients have been determined empirically. It is possible to specify them per station.
- It can take the number of used dipoles/tiles into account when scaling (e.g. for remote/international or for failing ones).

Filtering

- Similar to the msin step a filter makes it possible to keep only the given channels and/or baselines.
- By default, a station is always kept in the ANTENNA table, even if all its baselines are removed. This can be changed with the key remove.

Flag statistics and plotting

Several steps show statistics about flagged data points.

- A MADFlagger and AOFlagger step show the percentage of visibilities flagged by that flagging step. It shows:
 - $\circ\,$ The percentages per baseline and per station.
 - The percentages per channel.
 - The number of flagged points per correlation, i.e. which correlation triggered the flagging. This may help in determining which correlations to use in the MADFlagger.
- A UVWFlagger and PreFlagger step show the percentage of visibilities flagged by that flagging step. It shows percentages per baseline and per channel.
- The msin step shows the number of visibilities flagged because they contain a NaN or infinite value. It is shown which correlation triggered the flagging, so usually only the first correlation is really counted.
- A Counter step can be used to count and show the number of flagged visibilities. Such a step can be inserted at any point to show the cumulative number of flagged visibilities. For example, it can be defined as the first and last step to know how many visibilities have been flagged in total by the various steps.
- Each step giving flagging percentages can save the percentages per frequency and per station to a table. The extension .flagfreq is used for the table containing the flags per frequency; the extension .flagstat for the flags per station. The full basename of the table is the main part of the MS followed by _<stepname> followed by the extension. The path for these tables can be specified in the parset file.
- The plotflags function in the Python module lofar.dppp can be used to plot those tables. It can plot multiple subbands by giving it a list of table names. The flags per station will be averaged for those subbands.

Intermediate output step

The step out can write data to disk at an intermediate stage. It takes the same arguments as the 'msout' step. As an example, the following reduction will flag, save flagged data at high resolution, then average and save the result in another measurement set. On the averaged data, it will also apply a calibration table and save that in the CORRECTED_DATA column.

msin=L123.MS
steps=[aoflag,out1,average,out2,applycal]
Write out flagged data at full resolution
out1.type=out
out1.name=L123-flagged.MS
average.timestep=4
Write out averaged data
out2.type=out
out2.name=L123-averaged.MS
out2.datacolumn=DATA

applycal.parmdb=instrument.parmdb

Write the corrected data to CORRECTED_DATA
msout=L123-averaged.MS
msout.datacolumn=CORRECTED_DATA

User defined step

Besides the predefined DPPP steps like AOFlagger, etc., it is possible to use any user-defined DPPP step implemented in C++ or Python.

If implemented in C++ such a step has to reside in a shared library, that will dynamically be loaded by DPPP. The name of such a shared library has to be the step type name. DPPP will try to load the library libdppp_xxx.so (or .dylib on OS-X) for a step type xxx.

To make this a bit more flexible it is possible to define multiple steps in a single shared library. In such a case the step type name has to consist of 2 parts separated by a dot. The first part is the library name, the second part the step type in that library.

For example:

```
steps=[averager, mystep1, mystep2]
mystep1.type = mystep.stepa
mystep2.type = mystep.stepb
```

defines two user steps. Both step implementations reside in library libmystep.so. A description and example of a dynamically loaded step can be found in the LOFAR source code repository in LOFAR/CEP/DPPP/TestDyDPPP.

Python defined step

The mechanism described above is used to make it possible to implement a user step in Python. The step type has to be pythoDPPP and the name of the Python module and class containing the code have to be given. DPPP will load the library libdppp_pythonDPPP.so, which will start an embedded Python shell, load the module, and instantiate an object of the class. A detailed description is available.

ParSet File

Similar to most LOFAR programs, the parameters for the DPPP program are given in a so-called parset file. Note that it is possible to add parameters or overwrite parameters, defined in the parset file, using command line arguments. For example:

DPPP DPPP.pset parm1=value1 parm2=value2 ...

The steps to perform have to be defined in the parset file. They are executed in the given order, where the data are piped from one step to the other until all data are processed. Each step has a name to be used thereafter as a prefix in the keyword names specifying the type and parameters of the step.

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The most basic parset is as follows. It copies the DATA column of the MS and flags NaN and infinite data.

msin = ~/SB0.MS
msout = SB0_DPPP.MS
steps=[]

The following example is more elaborate. It flags (using a median flagger), averages all channels, flags the result of the average, and finally averages in time.

Note that 'msin' and 'msout' can be seen as an implicit first and last step.

```
msin = \sim /SB0.MS
msin.startchan = 8
msin.nchan = 240
msin.datacolumn = DATA
                           # is the default
msout = "SB0 DPPP.MS"
                           # if empty, the input MS is updated and
                           # no averaging steps can be done
msout.datacolumn = DATA
                           # is the default
steps = [flag1,count,avg1,flag2,avg2,count]
flag1.type=madflagger
flag1.threshold=1
flag1.freqwindow=31
flag1.timewindow=5
flag1.correlations=[0,3]
                           # only flag on XX and YY
flag1.count.save = true
                           # save flag percentages
flag1.count.path = $HOME
                           # to a table in $HOME
avgl.type = average
avg1.freqstep = 240
                           # is the default
avgl.timestep = 1
flag2.type=madflagger
flag2.threshold=2
flag2.timewindow=51
avg2.type = average
avg2.timestep = 5
```

Plotting the flag percentages, saved by the first MADFlagger step, could be done in python like:

```
import lofar.dppp as ld
ld.plotflags ('$HOME/SB0_flag1.flagfreq') # step name was flag1
```

Description of all parameters

The parameters in the parset are divided into several groups like input (msin), output (msout), madflagger, average, preflagger, and uvwflagger. Because multiple flagging and averaging steps can be specified, their parameters have to be prefixed with the step name as shown in the example above.

Parameter	type	default	description
General		•	
steps	string vector		Names of the steps to perform. Each step has to be defined using the step name as a prefix. The step type parameter defines the type of step (averager, madflagger, preflagger, uvwflagger, counter). The step type defaults to the name of the step, which is especially handy for count steps. msin and msout are implicit steps which should not be given here. An empty vector [] means that the input MS is copied to the output MS while flagging NaN and infinite numbers. Note that a step name can be used more than once meaning that the same step will be executed multiple times (e.g., multiple times count).
numthreads	int	\${OMP_NUM_THREADS}	Maximum number of threads to use.
showprogress	bool	true	Show a progress bar?
showcounts	bool	true	Show flagging statistics?
showtimings	bool	true	At the end the percentage of elapsed time each step took can be shown; the overall time is always shown.
checkparset	integer	0	 What to do if parameters in the ParSet file are not used. -1 means ignore. 0 means give a warning showing those parameters. In this way misspelled parameters can be detected. 1 means give an error and stop. For backward compatibility False (0) and True (1) can also be given.
uselogger	bool	false	If false, all DPPP messages are written on stdout. If true, the logging framework is used.

Counter

<step>.type</step>	string		Case-insensitive step type; must be 'counter' (or 'count'). Note that the type defaults to the step name, so if step name count is used, nothing more needs to be specified.
<step>.showfullyflagged</step>	bool	false	If true, all fully flagged baselines are shown in the baseline selection format using their antenna indices (not names). For example: 0&1; 3&7

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<step>.save</step>	bool		If true, the flag percentages per frequency are saved to a table with extension .flagfreq and percentages per station to a table with extension .flagstat. The basename of the table is the MS name (without extension) followed by the stepname and extension.
<step>.path</step>	string	<i>u n</i>	The directory where to create the flag percentages table. If empty, the path of the input MS is used.
<step>.warnperc</step>	double	0	If > 0, print an extra message for each baseline or channel with a percentage flagged higher than this value. Such a message line can be easily grep-ed.
<step>.flagdata</step>	bool	false	If COUNT is the only step in an DPPP run, the data won't be read, so unflagged invalid data (NaN. infinite) won't be noticed and counted as flagged. Setting this flag forces DPPP to read and check the data.

Input

msin msin.name	string		Name of the input MeasurementSets. If a single name is given, it can be a glob-pattern (like L23456_SAP000_SB*) meaning that all MSs matching the pattern will be used. A glob-pattern can contain *, ?, [], and {} pattern characters (as used in bash). If multiple MSs are to be used, their data are concatenated in frequency, thus multiple subbands are combined to a single band. In principle all MSs should exist, but if 'missingdata=true' and 'orderms=false' flagged zero data will be inserted for missing MS(s) and their frequency info will be deduced from the other MSs.
msin.sort	bool	false	Does the MS need to be sorted in TIME order?
msin.orderms	bool	true	Do the MSs need to be ordered on frequency? If true, all MSs must exist, otherwise they cannot be ordered. If false, the MSs must be given in order of frequency.
msin.missingdata	bool	false	true = it is allowed that a data column in an MS does not exist. In that case its data will be 0 and flagged. It can be useful if the CORRECTED_DATA of subbands are combined, but a BBS run for one of them failed. If 'orderms=false', it also makes it possible that a MS is specified but does not exist. In such a case flagged data will be used instead. The missing frequency info will be deduced from the other MSs where all MSs have to have the same number of channels and must be defined in order of frequency.
msin.baseline	string		Baselines to be selected (default is all baselines). See Description of baseline selection parameters. Only the CASA baseline selection syntax as described in this note can be used.
msin.band	integer	-1	Band (spectral window) to select (<0 is no selection). This is mainly useful for WSRT data.

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msin.startchan	integer	0	First channel to use from the input MS (channel numbers start counting at 0). Note that skipped channels will not be written into the output MS. It can be an expression with `nchan` (nr of input channels) as parameter. E.g. nchan/32 will be fine for LOFAR observations with 64 and 256 channels.
msin.nchan	integer	0	Number of channels to use from the input MS (0 means till the end). It can be an expression with `nchan` (nr of input channels) as parameter. E.g. 15*nchan/16
msin.starttime	string	first time in MS	Center of first time slot to use; if < first time in MS, dummy time slots are inserted. A date/time must be specified in the casacore MVTime format, e.g. 19Feb2010/14:01:23.817
msin.starttimeslots	int	0	Starting time slot. This can be negative to insert flagged time slots before the beginning of the MS.
msin.endtime	string	last time in MS	Center of last time slot to use; if > last time in MS, dummy time slots are inserted.
msin.ntimes	integer	0	Number of time slots to use (0 means till the end).
msin.useflag	bool	true	Use the current flags in the MS? If false, all flags in the MS are ignore and the data (except NaN and infinite values) are assumed to be good and will be used in later steps.
msin.datacolumn	string	DATA	Data column to use, i.e. the name of the column in which the visibilities are written.
msin.weightcolumn	string	WEIGHT_SPECTRUM or WEIGHT	Weight column to use. Defaults to WEIGHT_SPECTRUM if this exists, otherwise the WEIGHT column is used.
msin.modelcolumn	string	MODEL_DATA	Model data column. Currently only used in gaincal
msin.autoweight	bool	false	Calculate weights using the auto-correlation data? It is meant for setting the proper weights for a raw LOFAR MeasurementSet.
msin.forceautoweight	bool	false	In principle the calculation of the weights should only be done for the raw LOFAR data. It appeared that sometimes the autoweight switch was accidently set in a DPPP run on already dppp-ed data. To make it harder to make such mistakes, the forceautoweight flag has to be set as well for MSs containing dppp-ed data.

Output

DPPP

msout msout.name	string		Name of new output MeasurementSet; if empty, the input MS is updated. The other msout parameters are not applicable (apart from countflag). Normally an update is only done if a step is given that can change the data (e.g. PreFlagger). However, a name '.' or a name equal to the name of the input MS means that the input MS will always be updated, even if no step is given. This is useful if only flagging of NaN-s in the MS needs to be done. Note that when doing averaging, the input MS cannot be updated.
msout.overwrite	bool	false	When creating a new MS, overwrite if already existing?
msout.datacolumn	string	DATA	The column in which to write the data. When creating a new MeasurementSet, only column DATA can be used. When updating the input MeasurementSet, any column can be used. If not existing, it will be created first.
msout.weightcolumn	string	WEIGHT_SPECTRUM	The column in which to write the weights. When creating a new MeasurementSet, only WEIGHT_SPECTRUM can be used. When updating the input Measurementset, any column can be used. If not existing, it will be created first.
msout.writefullresflag	bool	true	Write the full resolution flags?
msout.tilesize	integer	1024	For expert user: tile size (in Kbytes) for the data columns in the output MS.
msout.tilenchan	integer	8	For expert user: maximum number of channels per tile in output MS.
msout.clusterdesc	string	un	If not empty, create the VDS file using this ClusterDesc file.
msout.vdsdir	string	<i>un</i>	Directory where to put the VDS file; if empty, the MS directory is used.

msout.storagemanager msout.storagemanager.name	string		What storage manager to use. When empty (default), the data will be stored uncompressed. When set to "dysco", the data will be compressed. Settings below will set the compression settings; see the Dysco wiki and the paper for more info. The default settings are reasonably conservative and safe.
msout.storagemanager.databitrate	integer	10	Number of bits per float used for columns containing visibilities. Can be set to zero to compress weights only.
msout.storagemanager.weightbitrate	integer	12	Number of bits per float used for WEIGHT_SPECTRUM column. Can be set to zero to compress data only. Note that compressing weights will set all polarizations to the same weight (determined by the minimum weight over the polarizations).
msout.storagemanager.distribution	string	"TruncatedGaussian"	Assumed distribution for compression; "Uniform", "TruncatedGaussian", "Gaussian" or "StudentsT".
msout.storagemanager.disttruncation	double	2.5	Truncation level for compression with the Truncated Gaussian distribution.
msout.storagemanager.normalization	string	"AF"	Compression normalization method: AF, RF or Row.

Filter

<step>.type</step>	string		Case-insensitive step type; must be 'filter'
<step>.startchan</step>	integer	0	First channel to use from the input MS (channel numbers start counting at 0). Note that skipped channels will not be written into the output MS. It can be an expression with `nchan` (nr of input channels) as parameter. E.g. nchan/32 will be fine for LOFAR observations with 64 and 256 channels.
<step>.nchan</step>	integer	0	Number of channels to use from the input MS (0 means till the end). It can be an expression with `nchan` (nr of input channels) as parameter. E.g. 15*nchan/16
<step>.baseline</step>	string	""	Baselines to keep. See Description of baseline selection parameters.
<step>.blrange</step>	double vector	<i>u </i>	Baselines to keep. See Description of baseline selection parameters.
<step>.corrtype</step>	string	""	Correlation type to match? Must be auto, cross, or an empty string.

Upsample

<step>.type</step>	string	Case-insensitive step type; must be 'upsample'
<step>.timestep</step>	integer	Number of times into which each timestep will be expanded

AOFlagger

<step>.type</step>	string		Case-insensitive step type; must be 'aoflagger' (or 'aoflag').
<step>.count.save</step>	bool	false	If true, the flag percentages per frequency are saved to a table with extension .flagfreq and percentages per station to a table with extension .flagstat. The basename of the table is the MS name (without extension) followed by the stepname and extension.
<step>.count.path</step>	string	""	The directory where to create the flag percentages table. If empty, the path of the input MS is used.
<step>.strategy</step>	string		The name of the strategy file to use. If no name is given, the default strategy is used which is fine for HBA. For LBA data the strategy LBAdefault should be used. A strategy file is looked up as given. If not found, it is looked up in \$LOFARROOT/share/rfistrategies that contains the standard strategies.
<step>.memoryperc</step>	integer	0	If >0, percentage of the machine's memory to use. If memorymax nor memoryperc is given, all memory will be used (minus 2 GB (at most 50%) for other purposes). Accepts only integer values (LOFAR v2.16). Limiting the available memory too much affects flagging accuracy; in general try to use at least 10 GB of memory.
<step>.memorymax</step>	double	0	Maximum amount of memory (in GB) to use. $\leftarrow 0$ means no maximum. As stated above, this affects flagging accuracy.
<step>.timewindow</step>	integer	0	Number of time slots to be flagged jointly. The larger the time window, the better the flagging performs. 0 means that it will be deduced from the memory to use. Note that the time window can be extended with an overlap on the left and right side to minimize possible boundary effects.
<step>.overlapperc</step>	double	0 or 1	If >0, percentage of time window to be added to the left and right side for overlap purposes (to minimize boundary effects). If overlapmax is not given, it defaults to 1%.
<step>.overlapmax</step>	integer	0	Maximum overlap value (0 is no maximum).
<step>.autocorr</step>	bool	true	Flag autocorrelations?
<step>.pulsar</step>	bool	false	Use flagging strategy optimized for pulsar observations?
<step>.pedantic</step>	bool	false	Be more pedantic when flagging?
<step>.keepstatistics</step>	bool	true	Write the quality statistics?

MADFlagger

<step>.type</step>	string		Case-insensitive step type; must be 'madflagger' (or 'madflag').
<step>.count.save</step>	bool	false	If true, the flag percentages per frequency are saved to a table with extension .flagfreq and percentages per station to a table with extension .flagstat. The basename of the table is the MS name (without extension) followed by the stepname and extension.
<step>.count.path</step>	string	<i>u </i>	The directory where to create the flag percentages table. If empty, the path of the input MS is used.
<step>.threshold</step>	float	1	The flagging threshold that can be baseline dependent. It can be any (TaQL-like) expression that evaluates to a float. In the expression the variable 'bl' can be used which is the baseline length (in meters). In this way the value can be made baseline dependent. For example: iif(bl<100, 0.5, iif(bl<500, 0.75, iif(bl<1000, 0.9, 1))) defines the threshold between the baseline lengths 100, 500, and 1000 meter.
<step>.timewindow</step>	integer	1	Number of times in the median box. If not odd, 1 is subtracted. It is silently reduced if exceeding the actual number of time slots. In a way similar to 'threshold' it can be made baseline length dependent.
<step>.freqwindow</step>	integer	1	Number of channels in the median box. If not odd, 1 is subtracted. It is silently reduced if exceeding the actual number of channels. In a way similar to 'threshold' it can be made baseline length dependent.
<step>.correlations</step>	integer vector	[]	The correlations to use in the flagger; an empty vector means all. They are handled in the order given; if the flagging criterium holds for one correlation, the other correlations are not tested anymore. So if one knows that most RFI is found in YY, then in XX and finally some in XY and YX, the vector should be [3,0,1,2] because it makes the program run faster. Note that the statistics printed at the end show how many flagged data points have been found per correlation.
<step>.applyautocorr</step>	bool	False	True means that the MADFlagger is used on the auto- correlations only. The resulting flags are applied to the cross- correlations, thus data are flagged where the corresponding auto-correlations are flagged. An error is given if set to True, while the MS does not contain auto-correlations.
<step>.blmin</step>	integer	-1	Minimum baseline length (in meters). Only baselines with a length >= this minimum are flagged. If applyautocorr=true, the autocorrelations are applied to the matching baselines only.
<step>.blmax</step>	integer	1e30	Maximum baseline length (in meters). It is similar to minimum.

PhaseShift

<step>.type</step>	string	Case-insensitive step type; must be 'phaseshifter' (or 'phaseshift').
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<step>.phasecenter string vector</step>	The RA and DEC (in J2000) of the new phase center. If an empty vector (i.e. []) is given, the original phase center is used. The RA and DEC can be given in sexagesimal format or as a value followed by a unit (default rad). For example, [12h31m34.5, 52d14m07.34] or [187.5deg, 52.45deg]
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Demixer

<step>.type</step>	string		Case-insensitive step type; must be 'demixer' (or 'demix').
<step>.baseline</step>	string	un	Baselines to demix. See Description of baseline selection parameters.
<step>.blrange</step>	double vector	[]	Baselines to demix. See Description of baseline selection parameters.
<step>.corrtype</step>	string	cross	Baselines to demix. Correlation type to match? Must be auto, cross, or an empty string.
<step>.timestep</step>	integer	1	Number of time slots to average when subtracting. It is truncated if exceeding the actual number of times. Note that the data itself will also be averaged by this amount.
<step>.freqstep</step>	integer	1	Number of channels to average when subtracting. It is truncated if exceeding the actual number of channels. Note that the data itself will also be averaged by this amount.
<step>.demixtimestep</step>	integer	timestep	Number of time slots to average when demixing. It is truncated if exceeding the actual number of times. It defaults to the averaging used for the subtract.
<step>.demixfreqstep</step>	integer	freqstep	Number of channels to average when demixing. It is truncated if exceeding the actual number of channels. It defaults to the averaging used for the subtract.
<step>.ntimechunk</step>	integer	#cores	Number of demix time slots (after averaging) that are processed jointly in as much a parallel way as possible. If subtract uses different time averaging, it has to fit integrally.
<step>.skymodel</step>	string	sky	The name of the SourceDB to use (i.e., the output of makesourcedb).
<step>.instrumentmodel</step>	string	instrument	The name of the ParmDB to use. The ParmDB does not need to exist. If it does not exist it will be created.
<step>.subtractsources</step>	string vector		Names of the sources to subtract. If none are given, demixing comes down to averaging. The sources must exist as patches in the SourceDB.

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<step>.modelsources</step>	string vector	[]	Names of sources with models to take into account when solving. the sources must exist as patches in the SourceDB. Note that the target should NOT be part of this parameter. If a model of the target has to be used, it has to be given in parameter targetsource.
<step>.targetsource</step>	string	un	It can be used to specify the name of the source model of the target. If given, the target source model (its patch in the SourceDB) is taken into account when solving; in this case parameter othersources cannot be given. It cannot be given if ignoretarget=true. If not given, the target is projected away or ignored (depending on parameter ignoretarget).
<step>.ignoretarget</step>	bool	false	false = project the target source away; true = ignore the target
<step>.othersources</step>	string vector	[]	Names of sources of which the direction is taken into account when demixing by projecting the directions away. The direction needs to be specified if the source is unknown (which is usually the case). It can be done using parameters <step>.<sourcename>.phasecenter.</sourcename></step>
<step>.<sourcename>.phasecenter</sourcename></step>	string vector	Taken from SourceDB	The J2000 direction [ra,dec] of a source given above.
<step>.propagatesolutions</step>	bool	true	If set to true, solutions of a time slot are used as initial values for the next time slot. If set to false, the diagonal elements of the Jones matrix are initialized to one and the off-diagonal elements to zero.
<step>.defaultgain</step>	double	1.0	The default and initial gain for the directional gains that are computed internally.
<step>.maxiter</step>	int	50	Maximum number of iterations used in the LM solve

SmartDemixer

<step>.type</step>	string		Case-insensitive step type; must be 'smartdemixer' (or 'smartdemix').
<step>.baseline</step>	string	<i>u n</i>	Baselines to demix. See Description of baseline selection parameters.
<step>.blrange</step>	double vector	<i>u n</i>	Baselines to demix. See Description of baseline selection parameters.
<step>.corrtype</step>	string	cross	Baselines to demix. Correlation type to match? Must be auto, cross, or an empty string.

<step>.target.baseline</step>	string	"CS*&"	Baselines to use in prediction of median target amplitude. See Description of baseline selection parameters.
<step>.target.blrange</step>	double vector	un	Baselines to use in prediction of median target amplitude. See Description of baseline selection parameters.
<step>.target.corrtype</step>	string	cross	Baselines to use in prediction of median target amplitude. Correlation type to match? Must be auto, cross, or an empty string.
<step>.timestep</step>	integer	1	Number of time slots to average when subtracting. It is truncated if exceeding the actual number of times. Note that the data itself will also be averaged by this amount.
<step>.freqstep</step>	integer	1	Number of channels to average when subtracting. It is truncated if exceeding the actual number of channels. Note that the data itself will also be averaged by this amount.
<step>.demixtimestep</step>	integer	timestep	Number of time slots to average when demixing. It is truncated if exceeding the actual number of times. It defaults to the averaging used for the subtract.
<step>.demixfreqstep</step>	integer	freqstep	Number of channels to average when demixing. It is truncated if exceeding the actual number of channels. It defaults to the averaging used for the subtract.
<step>.chunksize</step>	integer	demixtimestep	Number of time slots in a chunk for which it is decided how to demix (which sources/stations to use and how to deal with the target). It has to be a multiple of parameter 'demixtimestep'.
<step>.ntimechunk</step>	integer	#cores	Number of time chunks that are processed jointly in as much a parallel way as possible. Preferably it is a multiple of the number of cores. Note that for a typical LOFAR observation the data of a single time slot is about 4 MB. A typical chunk size can be 2 minutes, thus 120 time slots per core. For 24 cores this amounts to about 11 GB!!
<step>.ateam.skymodel</step>	string		The detailed sky model of the A-team sources used to solve for the complex gains. It is the name of the SourceDB to use (i.e., the output of makesourcedb).
<step>.estimate.skymodel</step>	string	<i>u</i> "	The coarse sky model of the A-team sources used to estimate the visibilities when deciding how to demix a chunk. It is the name of the SourceDB to use (i.e., the output of makesourcedb outtype=blob). If no name is given, the detailed A-team model will be used. The SourceDB must contain the same sources as the detailed model at about the same position. The order can be different though.
<step>.target.skymodel</step>	string		The sky model of the target. It is the name of the SourceDB to use (i.e., the output of makesourcedb).

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<step>.target.delta</step>	double	60	Angular distance uncertainty (in arcsec) to determine if an A-team source is at the same position as a target source.
<step>.instrumentmodel</step>	string	instrument	The name of the ParmDB to use. The ParmDB does not need to exist. If it does not exist it will be created. Note that the ParmDB is created after the output MS is created, so it can be a subdirectory of the output MS.
<step>.sources</step>	string vector	<i>u n</i>	Names of the A-team sources to use. If none are given, all sources in the A-team sky model will be used.
<step>.ateam.threshold</step>	double	50 for LBA 5 for HBA	Take a source/baseline into account if its maximum estimated amplitude > threshold.
<step>.minnbaseline</step>	integer	6	Solve a source/station if the station occurs in at least 'minnbaseline' baselines with amplitude > ateam.threshold.
<step>.minnstation</step>	integer	5	Solve a source if at least 'minnstation' stations are solvable for the source.
<step>.target.threshold</step>	double	200 for LBA 100 for HBA	Include the target in the solve if its maximum estimated amplitude > threshold.
<step>.ratio1</step>	double	5	Include the target in the solve if the estimated amplitude ratio Target/max(Ateam) > ratio1.
<step>.distance.threshold</step>	double	60	Distance threshold (in degrees). The target is close to the A-team if the angular distance (scaled with freq) < threshold for any A-team source (thus angdist*obsfreq/reffreq < threshold).
<step>.distance.reffreq</step>	double	60e6	The 'reffreq' frequency used above.
<step>.ratio2</step>	double	0.25	Include the target in the solve if the target is close to the A-team and the estimated amplitude ratio Target/min(Ateam) > ratio2.
<step>.maxiter</step>	integer	50	Maximum number of iterations to use in the solve.
<step>.propagatesolutions</step>	bool	true	If set to true, solutions of a time slot are used as initial values for the next time slot. If set to false, the diagonal elements of the Jones matrix are initialized to one and the off-diagonal elements to zero. However, solutions will not be transferred between chunks processed in parallel.
<step>.defaultgain</step>	double	1e-3	The default gain to use for the real part of the diagonal Jones elements for the unsolvable sources/stations. Take into account that the scale of the raw visibilities changed when COBALT was adopted. In the case of data correlated with BG/P, this parameter should be tuned down (1e-8).

<step>.verbose</step>	int	0	0 = only show basic demix statistics 1 = show for each time chunk how target is handled, which sources are solvable, and how many stations. >10 = various levels of debugging output.
<step>.solveboth</step>	bool	false	Mainly for test purposes. True means that in the solve only the baselines are used for which both stations are solvable. Usually this gives worse results.
<step>.targethandling</step>	integer	0	Mainly for test purposes. It enforces the target handling. 1=include, 2=deproject, 3=ignore, else=use smart way.
<step>.applybeam</step>	bool	true	Mainly for test purposes. Apply the station beam in the estimate, solve, and subtract steps?
<step>.subtract</step>	bool	true	Mainly for test purposes. False means that the subtract step is not done, thus only a solve of the gains is done.

Averager

<step>.type</step>	string		Case-insensitive step type; must be 'averager' (or equivalent 'average' or 'squash').	
<step>.timestep</step>	integer	T	Number of time slots to average. It is truncated if exceeding the actual number of times.	
	integer		Number of channels to average. It is truncated if exceeding the actual number of channels.	
<step>.minpoints</step>	integer	0	If number of averaged unflagged input points < minpoints, the averaged point is flagged.	
<step>.minperc</step>	float	0	Like minpoints, but expressed as a percentage of timestep*freqstep.	
<step>.timeresolution</step>	float	0 Target time resolution, in seconds. If this is given, and bigger than zero, it overrides <step>.timestep</step>		
<step>.freqresolution</step>	float		Target frequency resolution, in Hz (or append "MHz" or "kHz" to specify it in those units). If this is given, and bigger than zero, it overrides <step>.freqstep</step>	

StationAdder

<step>.type</step>	string	Case-insensitive step type; must be 'stationadder' (or equivalent 'stationadd').
<step>.stations</step>	record	<pre>One or more names of new stations each followed by the list of stations it consists of. A station name in the list can be a glob-like pattern. Optionally such a pattern can be negated by a ! or ^ meaning that names matching that pattern are excluded from the selection so far. For example: stations={ST6:'CS00[2-7]*'} can be used to form the superstation from all superterp stations. {ST6:['CS00[2-7]*','!CS005*']} is similar, but excludes CS005. {ST001:[CS001,CS002,CS003], ST002:[CS004,CS005,CS006]} defines 2 new stations ST001 and ST002 consisting of the stations in the lists following their names.</pre>

<step>.minpoints</step>	int		Flag a new data point if number of unflagged data points added is less than minpoints.
<step>.useweights</step>	bool		Use the input data weights? False means all input visibilities have weight 1.
<step>.average</step>	bool	true	Is a visibility of a new station the weighted average of its input visibilities and its UVW the weighted average of the input UVWs?
<step>.autocorr</step>	bool	false	Form new auto-correlations?
<step>.sumauto</step>	bool	true	Sum auto- or cross-correlations to form new auto-correlations?

ScaleData

<step>.type</step>	string		Case-insensitive step type; must be 'scaledata'.
<step>.stations</step>	string vector	[]	Zero or more glob-like patterns defining the stations for which the corresponding coefficient vector has to be used. The coefficients of the first matching pattern are used. Default coefficients (determined by Adam Deller for LBA and HBA) are used for stations not given. For example: stations=[CS*, RS*, *]
<step>.coeffs</step>	double vector	[]	Zero or more vectors of coefficients defining a polynomial in frequency (MHz). For example: coeffs=[[1.5, 0.7, 0.04], [1.7, 0.65], [1.2, 0.8]] The first vector results in a scale factor of 1.5 + 0.7*f + 0.04*f*f where f is the channel frequency in MHz. Note that an extra scaling can be applied taking into account the number of used dipoles/tiles of a station (see next parameter).
<step>.scalesize</step>	bool		This parameter determines if an extra scaling has to be applied to correct for the number of tiles/dipoles actually used in a station. By default this will be done for the stations using the default coefficients, because those coefficients have been determined for an LBA station with 48 dipoles and HBA station with 24 tiles. By default it will not be done for explicitly given coefficients, because it is supposed they are determined specifically for that station. Note that giving stations=* coeffs=1 scalesize=true will correct for station size only.

PreFlagger

<step>.type</step>	string		Case-insensitive step type; must be 'preflagger' (or 'preflag').
<step>.count.save</step>	bool	false	If true, the flag percentages per frequency are saved to a table with extension .flagfreq and percentages per station to a table with extension .flagstat. The basename of the table is the MS name (without extension) followed by the stepname and extension.
<step>.count.path</step>	string	""	The directory where to create the flag percentages table. If empty, the path of the input MS is used.

<step>.mode</step>	string	set	Case-insensitive string telling what to do with the flags of the data matching (or not matching) the selection criteria given in the other parameters. 'set' means set the flags for the matching data. This is the default mode. 'clear' means clear the flags for the matching data. However, flags of invalid data (NaN or zero) are always set. 'setcomplement' or 'setother' means set flags for NON- matching data. 'clearcomplement' or 'clearother' means clear flags for NON- matching (valid) data.	
.expr	string	[]	Expression of preflagger keyword sets (see above). Operators AND, OR, and NOT are possible (or their equivalents &&,&, , , and !). Parentheses can be used to change precedence order. For example: c1 and (c2 or c3) Take care that the name of the set is used as an extra prefix in the PreFlagger parameter names.	
.timeofday	time vector	[]	Ranges of UTC time-of-day given as stend or val+-delta. Each value must be given as 12:34:56.789, 12h34m56.789, or as a value followed by a unit like h, min, or s.	
.abstime	date/time vector	[]	Ranges of absolute UTC date/time given as stend or val+- delta. Each value (except delta) must be given as a date/tir in casacore MVTime format, for instance 12- Mar-2010/11:31:00.000. A delta value must be given as a t (for instance 1:30:0 or 20s).	
.reltime	time vector	[]	Ranges of times (using or $+$ -) since the start of the observation. A time can be given like 1:30:0 or 20s.	
.timeslot	integer vector	[]	Time slot sequence numbers. First time slot is 0. stend means end inclusive.	
.lst	time vector	[]	Ranges of Local Apparent Sidereal Times like 1:30:0 +- 20min. The LST of a time slot is calculated for the array position, thus not per antenna.	
.azimuth	direction vector	[]	Ranges of azimuth angles given as stend or val+-delta. Each value has to be given as a casacore direction like 12:34:56.789 or 12h34m56.789, 12.34.56.789 or 12d34m56.789, or a value followed by a unit like rad or deg.	
.elevation	direction vector	[]	Ranges of elevation angles (similar to azimuth). For example: 0deg10deg	
.baseline	baseline vector	un	See Description of baseline selection parameters.	
.corrtype	string	<i>u </i>	Correlation type to match? Must be auto, cross, or an empty string.	
.blmin	double	-1	If blmin > 0, baselines with length < blmin meter will match.	
.blmax	double	-1	If $blmax > 0$, baselines with length > $blmax$ meter will match.	
.uvmmin	double	-1	If uvmmin > 0, baselines with UV-distance < uvmmin meter will match. Note that the UV-distance is the projected basel length.	
.uvmmax	double	-1	If uvmmax > 0, baselines with UV-distance > uvmmax meter will match.	

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.freqrange	string vector	[]	Channels in the given frequency ranges will match. Each value in the vector is a range which can be given as startend or start+-delta. A value can be followed by a unit like KHz. If only one value in a range has a unit, the unit is also applied to the other value. If a range has no unit, it defaults to MHz. For example: freqrange=[1.2 1.4 MHz, 1.8MHz+-50KHz] flags channels between 1.2MHz and 1.4MHz and between 1.75MHz and 1.85MHz. The example shows that blanks can be used at will.	
.chan	string vector	[]	The given channels will match (start counting at 0). Channels exceeding the number of channels are ignored. Similar to msin, it is possible to specify the channels as an expression nchan. Furthermore, can be used to specify ranges. For example: chan=[0nchan/32-1, 31*nchan/32nchan-1] to flag the first and last 2 or 8 channels (depending on 64 or 256 channels in the observation).	
.amplmin	float vector	-1e30	Correlation data with amplitude < amplmin will match. It can be given per correlation. For example, amplmin=[100,,,100] matches data points with XX or YY amplitude < 100. The non-specified amplitudes get the default value. It is also possible to give a single value (without brackets) meaning that it is used as the minimum for all correlations.	
.amplmax	float vector	1e30	Correlation data with amplitude > amplmax will match.	
.phasemin	float vector	-1e30	Correlation data with phase < phasemin (in radians) will match.	
.phasemax	float vector	1e30	Correlation data with phase > phasemax (in radians) will match.	
.realmin	float vector	-1e30	30 Correlation data with real complex part < realmin will matc	
.realmax	float vector	1e30	Correlation data with real complex part > realmax will match.	
.imagmin	float vector	-1e30	Correlation data with imaginary complex part < imagmin will match.	
.imagmax	float vector	1e30	Correlation data with imaginary complex part > imagmax will match.	

ApplyCal

<step>.type</step>	string	Case-insensitive step type; must be 'applycal' (or 'correct').
<step>.parmdb</step>	string	Path of parmdb in which the parameters are stored. This can also be an H5Parm file, in that case the filename has to end in '.h5'
<step>.solset</step>	string	In case of applying an H5Parm file: the name of the solset to be used. If empty, defaults to the name of one solset present in the H5Parm (if more solsets are present in an H5Parm and solset is left empty, an error will be thrown))

DPPP

<step>.correction</step>	string	gain	Type of correction to perform, can be one of 'gain', 'tec', 'clock', '(common)rotationangle' / 'rotation', '(common)scalarphase', '(common)scalaramplitude' or 'rotationmeasure' (create multiple ApplyCal steps for multiple corrections). When using H5Parm, this is for now the name of the soltab; the type will be deduced from the metadata in that soltab, except for full Jones, in which case correction should be 'fulljones'.
<step>.soltab</step>	string vector		The name or names of the H5 soltab. Currently only used when correction=fulljones, in which case soltab should list two names (amplitude and phase soltab).
<step>.direction</step>	string	<i>u n</i>	If using H5Parm, the direction of the solution to use
<step>.updateweights</step>	bool	false	Update the weights column, in a way consistent with the weights being inverse proportional to the autocorrelations (e.g. if 'autoweights' was used before).
<step>.interpolation</step>	string	nearest	If using H5Parm, the type of interpolation (in time and frequency) to use, can be one of 'nearest' or 'linear'.
<step>.invert</step>	bool	true	Invert the corrections, to correct the data. Default is true. If you want to corrupt the data, set it to 'false'
<step>.timeslotsperparmupdate</step>	int	100	Number of time slots to handle after one read of the parameter file. Optimization to prevent spurious reading from the parmdb.
<step>.steps</step>	list	[]	<pre>(new in version 3.1) ApplyCal substeps, e.g. [myApplyCal1, myApplyCal2]. Their parameters can be specified through e.g. <step>.myApplyCal1.correction=tec. If a parameter is not given for the substep, it takes the value from <step></step></step></pre>

GainCal

<step>.type</step>	string	Case-insensitive step type; must be 'gaincal' or 'calibrate'.
<step>.caltype</step>	string	The type of calibration that needs to be performed, can be one of 'fulljones', 'diagonal', 'phaseonly', 'scalarphase'. Experimental values are 'amplitude' or 'scalaramplitude', 'tec', 'tecandphase'
<step>.parmdb</step>	string	Path of parmdb in which the computed parameters are to be stored. If the parmdb already exists, it will be overwritten. Note : You cannot use this parmdb in an applycal step in the same run of DPPP. To apply the solutions of the gaincal directly, use 'gaincal.applysolution' (see below). New in LOFAR 3.1: if the parmdb name ends in .h5, an H5Parm will be written.

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<step>.blrange</step>	vector		Vector of baseline lengths to use for calibration. See Description of baseline selection parameters. New in version 2.20
<step>.uvlambdamin</step>	double	0	Ignore baselines / channels with UV < uvlambdamin wavelengths. Note : also all other variants of uv flagging described in UVWFlagger (uvmmin, uvmrange, uvlambdarange, etc) are supported (New in 3.1)
<step>.baseline</step>	string		Baseline selection filter for calibration. See Description of baseline selection parameters. New in version 2.20
<step>.applysolution</step>	bool	false	Apply the calibration solution to the visibilities. Note that you should always also inspect the parmdb afterwards to check that the solutions look reasonable.
<step>.solint</step>	int	1	Number of time slots on which a solution is assumed to be constant (same as CellSize.Time in BBS). 0 means all time slots. Note that for larger settings of solint, and specially for solint = 0, the memory usage of gaincal will be large (all visibilities for a solint should fit in memory).
<step>.nchan</step>	int	0	Number of channels on which a solution is assumed to be constant (same as CellSize.Freq in BBS). 0 means all channels. When caltype = 'tec' or 'tecandphase', the default is 1, meaning that a TEC will be fitted through a phase for each channel.
<step>.usemodelcolumn</step>	bool	false	Use model column. The model column name can be specified with msin.modelcolumn (default MODEL_DATA)
<step>.applybeamtomodelcolumn</step>	bool	false	Apply the beam model (at the phase center) to the visibilities in the model column. If this option is true, all options from applybeam are valid as well (except .invert, since the model data will always be corrupted for the beam)
<step>.propagatesolutions</step>	bool	true	Use solutions of one time interval as a starting value for the next time interval
<step>.maxiter</step>	int	50	Maximum number of iterations of stefcal
<step>.detectstalling</step>	bool	true	Detect if the iteration does not converge anymore and then stop iterating even if maxiter is not reached
<step>.tolerance</step>	float	1.e-5	Tolerance to which the model should match the data
<step>.minblperant</step>	int	4	If an antenna has less than minblperant unflagged data points for a given solution slot, it is not used for calibration
<step>.timeslotsperparmupdate</step>	int	500	Number of solution intervals after which the parmdb is updated
<step>.debuglevel</step>	int	0	Debugging. If debuglevel==1, then a file debug.h5 is created containing all iterands. This file will be very large; you can use it to check the convergence speed etc.
<step>.sourcedb</step>			Same as in Predict step

<step>.sources</step>	Same as in Predict step
<step>.usebeammodel</step>	Same as in Predict step
<step>.operation</step>	Same as in Predict step
<step>.applycal.*</step>	ApplyCal sub-step, same as in Predict step
<step>.onebeamperpatch</step>	Same as in ApplyBeam step
<step>.usechannelfreq</step>	Same as in ApplyBeam step
<step>.beammode</step>	Same as in ApplyBeam step

DDECal

<step>.type</step>	string		Case-insensitive step type; must be 'ddecal'.
<step>.sourcedb</step>	string		Sourcedb (created with `makesourcedb`) with the sky model to calibrate on.
<step>.directions</step>	list	[]	List of directions to calibrate on. Every element of this list should b a list of facets. Default: every facet is a direction.
<step>.usemodelcolumn</step>	bool	false	Use model data from the measurement set. This implies solving for one direction, namely the pointing of the measurement set. If you specify usemodelcolumn to be true, directions and sourcedb are not required
<step>.maxiter</step>	int	50	Maximum number of iterations.
<step>.detectstalling</step>	bool	true	Stop iterating when no improvement is measured anymore (after a minimum of 30 iterations).
<step>.stepsize</step>	double	0.2	stepsize between iterations.
<step>.h5parm</step>	string		Filename of output H5Parm (to be read by e.g. losoto). If empty, defaults to instrument.h5 within the measurement set.
<step>.solint</step>	int	1	Solution interval in timesteps.
<step>.usebeammodel</step>	bool	false	use the beam model. All beam-related options of the Predict step are also valid.
<step>.mode</step>	string	diagonal	Type of constraint to apply. Options are scalarcomplexgain, scalarphase, scalaramplitude, tec, tecandphase. Modes in development are fulljones, diagonal, phaseonly, amplitudeonly, rotation, rotation+diagonal.
<step>.tolerance</step>	double	1e-5	Controls the accuracy to be reached: when the normalized solutions move less than this value, the solutions are considered to be converged and the algorithm finishes. Lower values will cause more iterations to be performed.
<step>.minvisratio</step>	double	0	Minimum number of visibilities within a solution interval, e.g. 0.6 for at least 60% unflagged vis. Intervals with fewer vis will be flagged.
<step>.propagatesolutions</step>	bool	false	Initialize solver with the solutions of the previous time slot.
<step>.propagateconvergedonly</step>	bool	false	Propagate solutions of the previous time slot only if the solve converged. Only effective when propagatesolutions=true.
<step>.flagunconverged</step>	bool	false	Flag unconverged solutions (i.e., those from solves that did not converge within maxiter iterations).

<step>.flagdivergedonly</step>	bool	false	Flag only the unconverged solutions for which divergence was detected. At the moment, this option is effective only for rotation+diagonal solves, where divergence is detected when the amplitudes of any station are found to be more than a factor of 5 from the mean amplitude over all stations. If divergence for any one station is detected, all stations are flagged for that solution interval. Only effective when flagunconverged=true and mode=rotation+diagonal.
<step>.approximatetec</step>	bool	false	Uses an approximation stage in which the phases are constrained with the piece-wise fitter, to solve local minima problems. Only effective when mode=tec or mode=tecandphase.
<step>.maxapproxiter</step>	int	maxiter/2	Maximum number of iterations during approximating stage.
<step>.approxchunksize</step>	int	0	Size of fitted chunksize during approximation stage in nr of channels. With approxchunksize=1 the constraint is disabled during the approx stage (so channels are solved for independently). Once converged, the solutions are constrained and more iterations are performed until that has converged too. The default is approxchunksize=0, which calculates the chunksize from the bandwidth (resulting in 10 chunks per octave of bandwidth).
<step>.approxtolerance</step>	double	tolerance*10	Tolerance at which the approximating first stage is considered to be converged and the second full- constraining stage is started. The second stage convergences when the tolerance set by the 'tolerance' keyword is reached. Setting approxtolerance to lower values will cause more approximating iterations. Since tolerance is by default 1e-5, approxtolerance is by default 1e-4.
<step>.nchan</step>	int	1	Number of channels in each channel block, for which the solution is assumed to be constant. The default is 1, meaning one solution per channel (or in the case of constraints, fitting the constraint over all channels individually). O means one solution for the whole channel range. If the total number of channels is not divisable by nchan, some channelblocks will become slightly larger.
<step>.coreconstraint</step>	double	0	Distance in meters. When unequal to 0, all stations within the given distance from the reference station (0) will be constraint to have the same solution.
<step>.antennaconstraint</step>	list	[]	A list of lists specifying groups of antennas that are to be constrained to have the same solution. Example: "[[CS002HBA0,CS002HBA1],[CS003HBA0,CS003HBA1]]" will keep the solutions of CS002HBA0 and 1 the same, and the same for CS003.
<step>.smoothnessconstraint</step>	double	0	Kernel size in Hz. When unequal to 0, will constrain the solutions to be smooth over frequency by convolving the solutions with a kernel of the given size (bandwidth). The default kernel is a Gaussian kernel, and the kernel size parameter is the 3 sigma point where the kernel is cut off.

string	un	File to write the step-sizes to. Form of the file is: " <iterationnr> <normalized-stepsize> <unnormalized-stepsize>", and all solution intervals are concatenated. File is not written when this parameter is empty.</unnormalized-stepsize></normalized-stepsize></iterationnr>
double	0	Ignore baselines / channels with UV < uvlambdamin wavelengths. Note : also all other variants of uv flagging described in UVWFlagger (uvmmin, uvmrange, uvlambdarange, etc) are supported (New in 3.1).
bool	false	Subtracts the corrected model from the data. NOTE This may not work when you apply a uv-cut.
bool	false	Do image-based prediction using IDG.
string	un	Name of fits file for IDG prediction. Should have the phase direction of the measurement set.
string	un	DS9 regions file describing the facets for IDG prediction.
bool	false	Write out each facet as a fits file (named facet <n>.fits). Only useful when useidg=true.</n>
bool	false	Instead of solving, output the predicted visibilities instead. This is useful for testing, although when doing faceted prediction with IDG, it might be fast for certain cases.
		ApplyCal sub-step, same as in Predict step. One can pass an h5parm with as many directions as set in "directions" and each direction model is corrupted accordingly.
	double bool bool string string bool	Stringdouble0boolfalseboolfalsestring""stringboolboolfalse

Predict

<step>.type</step>	string		Case-insensitive step type; must be 'predict'
<step>.sourcedb</step>	string		Path of sourcedb in which a sky model is stored (the output of makesourcedb)
<step>.sources</step>	string vector	[]	Patches to use in the predict step of the calibration
<step>.usebeammodel</step>	bool	false	Use the LOFAR beam in the predict part of the calibration
<step>.operation</step>	string	replace	Should the predicted visibilities replace those being processed (replace, default), should they be subtracted from those being processed (subtract) or added to them (add)
<step>.applycal.*</step>			Set of options for applycal to apply to this predict. For this applycal-substep, .invert is off by default, so the predicted visibilities will be corrupted with the parmdb
<step>.onebeamperpatch</step>			Same as in ApplyBeam step
<step>.usechannelfreq</step>			Same as in ApplyBeam step
<step>.beammode</step>			Same as in ApplyBeam step

H5ParmPredict

<step>.type</step>	string	Case-insensitive step type; must be 'h5parmpredict'
<step>.sourcedb</step>	string	Path of sourcedb in which a sky model is stored (the output of makesourcedb)
<step>.applycal.parmdb</step>	string	Path of the h5parm in which the corruptions are stored

<step>.applycal.correction</step>	string		SolTab which contains the directions to be predicted, or "fulljones". The names of the directions need to look like [dir1,dir2], where dir1 and dir2 are patches in the sourcedb.
<step>.directions</step>	string vector	[]	List of directions to include. Each of those directions needs to be in the h5parm soltab. If empty, all directions in the soltab are predicted.
<step>.usebeammodel</step>	bool	false	Use the LOFAR beam in the predict part of the calibration
<step>.operation</step>	string	replace	Should the predicted visibilities replace those being processed (replace, default), should they be subtracted from those being processed (subtract) or added to them (add)
<step>.applycal.*</step>			Set of options for applycal to apply to this predict. For this applycal-substep, .invert is off by default, so the predicted visibilities will be corrupted with the parmdb
<step>.onebeamperpatch</step>			Same as in ApplyBeam step
<step>.usechannelfreq</step>			Same as in ApplyBeam step
<step>.beammode</step>			Same as in ApplyBeam step

ApplyBeam

<step>.type</step>	string		Case-insensitive step type; must be 'applybeam'
<step>.direction</step>	string vector	[]	A RA/Dec value specifying in what direction to correct the beam. See phaseshift.phasecenter for syntax. If empty, the beam is corrected in the direction of the current phase center.
<step>.onebeamperpatch</step>	bool	false	Compute the beam only for the center of each patch (saves computation time, but you should set this to false for large patches). In the ApplyBeam step, this setting does not make sense (but it does if the applybeam is part of predict, ddecal, gaincal, h5parmpredict, etc.). Generally, FALSE is the right setting for this option. The default has changed to false in a recent (Nov 2018) version.
<step>.usechannelfreq</step>	bool	true	Compute the beam for each channel of the measurement set separately. This is useful for merged / concatenated measurement sets. For raw LOFAR data you should set it to false, so that the beam will be formed as in the station hardware. Also, setting it to false is faster.
<step>.updateweights</step>	bool	false	Update the weights column, in a way consistent with the weights being inverse proportional to the autocorrelations (e.g. if 'autoweights' was used before).
<step>.invert</step>	bool	true	Invert the beam. When applying the beam to transfer calibration solutions, this should be true. In other words: invert=true means correcting for the beam, invert=false means corrupting with the beam. When using the beam in a predict (or gaincal) step, this option defaults to false (so it will corrupt for the beam).

<step>.beammode</step>	string	Beam mode to apply, can be "array_factor", "element" "default" or "default". Default is to apply both the element beam
		and the array factor.

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UVWFlagger

<step>.type</step>	string		Case-insensitive step type; must be 'uvwflagger' or 'uvwflag'.
<step>.count.save</step>	bool	false	If true, the flag percentages per frequency are saved to a table with extension .flagfreq and percentages per station to a table with extension .flagstat. The basename of the table is the MS name (without extension) followed by the stepname and extension.
<step>.count.path</step>	string		The directory where to create the flag percentages table. If empty, the path of the input MS is used.
<step>.uvmrange</step>	string vector	[]	Flag baselines with UV within one the given ranges (in meters). Delimiters and +- can be used to specify a range. E.g., uvmrange = [2030, 40+-5] flags baselines with UV in range 20-30 meter and 35-45 meter.
<step>.uvmmin</step>	double	0	Flag baselines with UV < uvmmin meter.
<step>.uvmmax</step>	double	1e15	Flag baselines with UV > uvmmax meter.
<step>.umrange</step>	string vector	[]	Flag baselines with U within one of the given ranges (in meters).
<step>.ummin</step>	double	0	Flag baselines with U < ummin meter.
<step>.ummax</step>	double	1e15	Flag baselines with $U > ummax$ meter.
<step>.vmrange</step>	string vector	[]	Flag baselines with V within one of the given ranges (in meters).
<step>.vmmin</step>	double	0	Flag baselines with V < vmmin meter.
<step>.vmmax</step>	double	1e15	Flag baselines with $V > vmmax$ meter.
<step>.wmrange</step>	string vector	[]	Flag baselines with W within one of the given ranges (in meters).
<step>.wmmin</step>	double	0	Flag baselines with W < wmmin meter.
<step>.wmmax</step>	double	1e15	Flag baselines with $W > wmmax$ meter.
<step>.uvlambdarange</step>	string vector	[]	Flag baselines/channels with UV within one the given ranges (in wavelengths). Delimiters and +- can be used to specify a range. E.g., $uvlambdarange = [2030, 40+-5]$ flags baselines/channels with UV in range 20-30 wavelengths and 35-45 wavelengths.
<step>.uvlambdamin</step>	double	0	Flag baselines/channels with UV < uvlambdamin wavelengths
<step>.uvlambdamax</step>	double	1e15	Flag baselines/channels with UV > uvlambdamax wavelengths
<step>.ulambdarange</step>	string vector	[]	Flag baselines/channels with U within one the given ranges (in wavelengths).
<step>.ulambdamin</step>	double	0	Flag baselines/channels with U < ulambdamin wavelengths
<step>.ulambdamax</step>	double	1e15	Flag baselines/channels with $U > u$ lambdamax wavelengths
<step>.vlambdarange</step>	string vector	[]	Flag baselines/channels with V within one the given ranges (in wavelengths).
<step>.vlambdamin</step>	double	0	Flag baselines/channels with V < vlambdamin wavelengths
<step>.vlambdamax</step>	double	1e15	Flag baselines/channels with V > vlambdamax wavelengths

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<step>.wlambdarange</step>	string vector	[]	Flag baselines/channels with W within one the given ranges (in wavelengths).
<step>.wlambdamin</step>	double	0	Flag baselines/channels with W < wlambdamin wavelengths
<step>.wlambdamax</step>	double	1e15	Flag baselines/channels with W > wlambdamax wavelengths
<step>.phasecenter</step>	string vector	[]	If given, use this phase center to calculate the UVW coordinates to flag on. The vector can consist of 1, 2 or, 3 values. If one value is given, it must be the name of a moving source (e.g. SUN or JUPITER). Otherwise the first two values must contain a source position that can be given in sexagesimal format or as a value followed by a unit. The third value can contain the direction type; it defaults to J2000. Possible types are GALACTIC, ECLIPTIC, SUPERGAL, J2000, B1950 (as defined in the casacore Measures system).

Split

<step>.type</step>	string		Case-insensitive step type; must be 'split' or 'explode'
<step>.steps</step>	string vector	[]	List of next steps; each step will run after this step. E.g. [average, msout]
<step>.replaceparms</step>	string vector	[]	The substep keys that should be different for each of the next steps. Instead of their default type, they should now be a list of those things. E.g. [average.timestep, msout.name]

Interpolate

The interpolate step replaces flagged values by interpolating them using "neighbouring" samples (samples close in time and frequency). It calculates the Gaussian weighted sum over non-flagged samples, with a sigma parameter of one timestep/one channel. The flags are removed after interpolation. This is in particular useful in combination with averaging; by replacing flagged values before averaging, the output visibilities will more accurately represent the true sky. This step was aimed to solve frequency structure from flagging/averaging for the EoR experiment, but might be useful in other cases as a more accurate averaging step. Details are published in Offringa, Mertens and Koopmans (2018).

<step>.type</step>	string		Case-insensitive step type; must be 'interpolate'.
<step>.windowsize</step>	int	15	Size of the window over which a value is interpolated. Should be odd.

Description of baseline selection parameters

Parameters to select on baseline can be used in the steps preflagger and filter. The step msin only supports .baseline. The parameters are described in the table below.

Parameter	type	default	description
.corrtype	string		Correlation type to match? Must be auto, cross, or an empty string (= all).

DPPP

Parameter type d	efault description
.blrange double []	Zero or more ranges of physical baseline lengths (in m). A baseline matches if its length is within one of the ranges. E.g., blrange=[0,10000, 100000, 1e30]
.baseline baseline "	 Names of baselines to be matched. It can be given as either a vector of vectors or as a casacore MSSelection string. These two methods are mutually exclusive. When in doubt, use the second syntax. I. If given as a vector, a vector element can be a vector of two names giving the stations forming a baseline. For example: baselines [[CS001,RS003], [CS002,RS005]] selects baselines CS001-RS003 and CS002-RS005. Each name can be a shell-type pattern (with wildcards * ? [] or {}). Thus baseline=[[CS*,RS*]] selects all baselines between core and remote stations. Note that the wildcard characters {} mean OR. They can be used to pair groups of stations (uotes are needed). For example: baseline=[[(CS001,CS002)", "(RS003,RS005,CS002-RS003, and CS002-RS005. Besides giving a baseline, it is also possible to give a single station name (possibly wildcarded) meaning that all baselines containing that station will be selected. For example: baselines. RCS*] selects all baselines containing remote or cor estations. Please note that an extra bracket pair is needed to specify baselines between RS and CS inke in baseline=[[RS*, CS*]] It is a bit hard to select international stations using this syntax. The casacore note 263. The advantage of this syntax is that it is more concise and that besides a station name pattern, it is possible to give a station number. The examples above can be expressed as: baseline=CS001, CS002RS003, RS005 for CS001-RS003, and CS002-RS005 baseline=RS*&&CS* baseline=ES12 baseline between station number 8 and 12. Note that & means cross-correlations, & means cross and auto, &&& means auto only. International stations. baseline=fCR]S*&& to select all baselines containing an international station. use baseline=fCR]S*&& to select all baselines containing an international station. use baseline=cCR]S*&& to select all baselines containing an international station. use baseline=cCR]S*&&

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