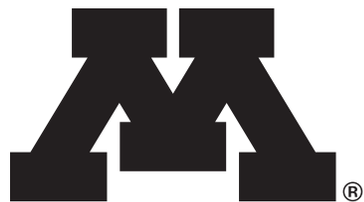


Application Guide & Release Notes

Multi-Band EPI C2P

**Release 017pre13
23 September 2025**



Center for Magnetic
Resonance Research

UNIVERSITY OF MINNESOTA

Driven to DiscoverSM

Conditions of Use

This package is provided to support collaborative research projects between the University of Minnesota and users of Siemens MRI equipment, in accordance with a current C2P agreement.

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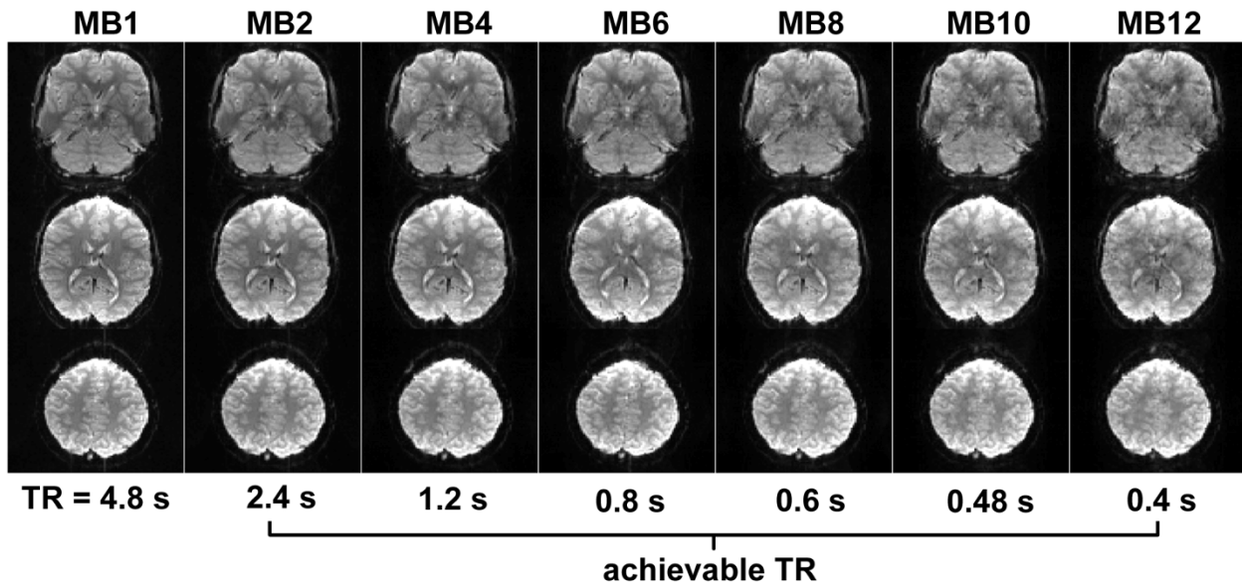
Table of Contents

Overview	3
Sequence Description	4
Hardware and Software Requirements	6
Installation Procedure	8
Usage.....	10
Routine Card	10
Contrast/Common Card.....	11
Resolution/iPAT Card (VB/VD/VE)	12
Resolution/Acceleration Card (Numaris X).....	12
Sequence/Part 1 Card	14
Sequence/Special Card.....	15
Important Issues (not bugs)	21
Known Bugs	22
Version Changes.....	23

Overview

Multi-banded RF pulses can be used to accelerate volume coverage along the slice direction by simultaneously exciting and acquiring multiple slices and subsequently unaliasing them using parallel imaging principles and the spatial information available in multi-channel RF array coils.

This allows for a direct reduction in the volume T_R by the number of simultaneously excited slices (i.e., the multiband (MB) factor or the slice acceleration factor). For example, the figure below shows the minimum achievable T_R for a 64-slice, 2 mm isotropic resolution GRE-EPI acquisition using various MB factors:



Such a significant reduction in minimum T_R is very useful in studies of the brain. In particular, the ability to sample more rapidly while preserving whole-brain coverage is of great benefit to resting-state fMRI. It is also useful to reduce the otherwise long total scan times required for diffusion-weighted imaging techniques such as HARDI and DSI.

Sequence Description

The pulse sequences in this package are based on the product ep2d_bold, ep2d_se, and ep2d_diff sequences from Siemens.

Multi-banded RF pulses are calculated by the sequence using linear combinations of the same RF pulses used in the product sequences. To improve the MB unaliasing performance, partial FoV shifts are applied using gradient blips along the slice direction during the EPI readout. The optimum shift factor is selected automatically by the sequence.

More detailed information about the pulse sequence and image reconstruction implementation can be found in the following references:

Moeller S, Yacoub E, Olman CA, Auerbach EJ, Strupp J, Harel N, Uğurbil K. Multiband multislice GE-EPI at 7 tesla, with 16-fold acceleration using partial parallel imaging with application to high spatial and temporal whole-brain fMRI. *Magn Reson Med*. 2010;63(5):1144–1153. [[PubMed](#)]

Setsompop K, Gagoski BA, Polimeni JR, Witzel T, Wedeen VJ, Wald LL. Blipped-controlled aliasing in parallel imaging for simultaneous multislice echo planar imaging with reduced g-factor penalty. *Magn Reson Med*. 2012;67(5):1210–1224. [[PubMed](#)]

Xu J, Moeller S, Auerbach EJ, Strupp J, Smith SM, Feinberg DA, Yacoub E, Uğurbil K. Evaluation of slice accelerations using multiband echo planar imaging at 3T. *Neuroimage*. 2013 Dec;83:991-1001. [[PubMed](#)]

Auerbach EJ, Xu J, Yacoub E, Moeller S, Uğurbil K. Multiband accelerated spin-echo echo planar imaging with reduced peak RF power using time-shifted RF pulses. *Magn Reson Med*. 2013 May;69(5):1261-7. [[PubMed](#)]

Sotiropoulos SN, Moeller S, Jbabdi S, Xu J, Andersson JL, Auerbach EJ, Yacoub E, Feinberg D, Setsompop K, Wald LL, Behrens TEJ, Uğurbil K, Lenglet C. Effects of image reconstruction on fiber orientation mapping from multichannel diffusion MRI: Reducing the noise floor using SENSE. *Magn Reson Med*. 2013;70(6):1682-1689. [[PubMed](#)]

Cauley SF, Polimeni JR, Bhat H, Wald LL, Setsompop K. Interslice leakage artifact reduction technique for simultaneous multislice acquisitions. *Magn Reson Med*. 2014 Jul;72(1):93-102. [[PubMed](#)]

Setsompop K, Cohen-Adad J, Gagoski BA, Raij T, Yendiki A, Keil B, Wedeen VJ, Wald LL. Improving diffusion MRI using simultaneous multi-slice echo planar imaging. *NeuroImage*. 2012 Oct 15;63(1):569-80. [\[PubMed\]](#)

Polimeni JR, Bhat H, Witzel T, Benner T, Feiweier T, Inati SJ, Renvall V, Heberlein K, Wald LL. Reducing sensitivity losses due to respiration and motion in accelerated echo planar imaging by reordering the autocalibration data acquisition. *Magn Reson Med*. 2016 Feb;75(2):665-79. [\[PubMed\]](#)

Hardware and Software Requirements

Versions of the multi-band accelerated sequences are currently provided for multiple Siemens software versions, supporting various system types. It may additionally be possible to use the sequences with other systems not listed (e.g. 1.5T systems), but compatibility and coil performance has not been tested.

VB17A: MAGNETOM Trio, a Tim System 3T, MAGNETOM Verio 3T, MAGNETOM 7T/9.4T/10.5T/11.7T

VB19A: MAGNETOM Trio, a Tim System 3T

VB19B-SP02: MAGNETOM Trio, a Tim System 3T

VB20P: Biograph mMR

VD11D: MAGNETOM ConnectomA 3T

VD13A-SP4: MAGNETOM Skyra 3T

VD13B: MAGNETOM Skyra Fit 3T

VD13C-SP2: MAGNETOM Skyra 3T with TimTX TrueShape (pTX) option

VD13D: MAGNETOM Prisma/Prisma Fit 3T

VE11A: MAGNETOM Skyra 3T

VE11B: MAGNETOM Prisma 3T

VE11C: MAGNETOM Skyra/Prisma 3T

VE11E: MAGNETOM Skyra/Prisma 3T

VE11P: Biograph mMR

VE11P-SP04: Biograph mMR

VE11K/R: MAGNETOM Terra 7T (1-channel Tx/clinical mode)

VE11U: MAGNETOM Terra 7T (pTX/research mode)

VE12U: MAGNETOM Terra 7T

VE12U-SP01: MAGNETOM Terra 7T

VE12U-AP01: MAGNETOM UHF Plus

VE12U-AP02: MAGNETOM Terra 7T Impulse Edition

VE12U-AP04: MAGNETOM Terra 7T, UHF Plus

NXVA20A: MAGNETOM Vida 3T

NXVA30A: MAGNETOM Skyra/Prisma 3T

NXVA31A: MAGNETOM Vida 3T

NXVA50A: MAGNETOM Vida/Vida Fit/Skyra Fit BioMatrix 3T
NXVA60A: MAGNETOM Skyra/Prisma/Vida 3T, Terra.X 7T
NXVA61A: MAGNETOM Cima.X/Connectom.X/Skyra Fit 3T
NXVA61A-SP01: MAGNETOM Cima.X/Connectom.X/Skyra Fit 3T

For 3T, the **32-channel Head** coil is highly recommended. The **Head/Neck 20** coil can be used, but image quality will be diminished at higher acceleration factors.

The **Head/Neck 64** coil can also be used for 3T, but imaging performance in the brain is typically not improved over the 32-channel coil while reconstruction time can be substantially longer. Enabling “**Matrix Optimization**” on the System/Miscellaneous card can improve reconstruction time when using the Head/Neck 64 coil (note: this option is not available when generating phase images).

For 7T (Terra/Terra.X), the **Nova Medical 32-channel** head coil is recommended, in both the 1T/32R and 8T/32R (pTX) configurations.

Installation Procedure

For Numaris X software versions **NXVA*** please see the included document **Install_Instructions_Numaris_X.pdf**. A multi-step process to install a security certificate must be performed prior to installing the sequence binaries. This only needs to be done once per scanner unless software is upgraded/reloaded.

For versions **VE11C**, **VE11E**, **VE12U**, **VB19A**, **VB19B**, and **VB20P**, the host uses McAfee Embedded Control (Solidifier) security software. It is necessary to switch Embedded Control into UPDATE mode before installing any custom sequences. Please see the included document **Embedded_Control_Mode.pdf** for instructions on how to switch the Embedded Control mode.

The standard install/upgrade procedure is as follows:

1. Numaris X: deactivate Kiosk mode. Other versions: login as “Advanced User”. *If you do not know the password, contact Local Service.*
2. If the system has Embedded Control installed, start the MrEmbeddedControlGui and switch the Embedded Control system into UPDATE mode. *If you do not know the Administrator account name or password, contact Local Service.*
3. Extract the .zip file to a temporary directory.
4. Run the installer program. The installer will automatically restart various processes and reboot the imager if necessary to unlock shared objects.
5. If the installer reports any error, please power cycle the system and try again.
6. If a sample protocol is available for your system, it will appear at the end of the USER tree in the Exam Explorer. If a sample protocol is

not available, create a default protocol in Exam Explorer by selecting Insert Sequence, USER, then cmrr_mbep2d_* for the desired sequence variant.

7. If the system has Embedded Control installed, re-enable host security and switch the Embedded Control system into the ENABLED mode.

Usage

These sequences are based on the Siemens product ep2d_* family of sequences; most of the protocol parameters are the same.

The most important changes appear on the following measurement cards:

Routine Card

The Routine Card interface displays the following parameters:

Slice group	1	FoV read	192	mm
Slices	72	FoV phase	100.0	%
Dist. factor	0	Slice thickness	2.00	mm
Position	Isocenter	TR	818	ms
Orientation	Transversal	TE	31	ms
Phase enc. dir.	A >> P	Multi-band accel. factor	8	
AutoAlign	---	Filter	None	
Phase oversampling	0	Coil elements	HEA;HEP	

At the bottom, a Multi-band accel. factor slider is shown with a scale from 2 to 12, with a green marker at 8. Below the slider is a navigation bar with tabs: Routine, Contrast, Resolution, Geometry, System, Physio, BOLD, and Sequence.

Routine Card (all versions)

Multi-band accel. factor: This parameter is only editable if the number of slices is greater than 1. The number of slices must be an integer multiple of the multi-band acceleration factor. The maximum acceleration factor is 16 (BOLD/SE) or 8 (diffusion) in the current release if iPAT is not used, otherwise the product of the MB accel. factor and the iPAT factor may not exceed 18. Note that image quality is not likely to be acceptable at the highest allowed acceleration factors for most protocols.

Slices: The number of slices must be an integer multiple of the multi-band acceleration factor. It may be necessary to reset the multi-band acceleration factor to 1 in order to select an arbitrary number of slices; a new multi-band acceleration factor can then be re-selected.

Averages: Online averaging has been disabled due to incompatibility with the online reconstruction.

Dist. factor: There must be a minimum gap of 100% of the slice thickness between adjacent bands in the RF pulse. Therefore, in a case where the number of bands is equal to the number of slices, a minimum distance factor of 100% will be required. In the usual case where slices \gg bands, however, a distance factor of 0% can be used.

Contrast/Common Card

The image shows a software interface for the Contrast/Common Card. It has two tabs: 'Common' and 'Dynamic'. The 'Dynamic' tab is active. The interface contains several input fields and controls:

- TR: 3650 ms
- TE: 95 ms
- Fat suppression: Fat sat.
- Grad. rev. fat suppression: Disabled
- MTC:
- Magn. preparation: None
- Flip angle: 78 deg
- A TR slider bar is visible at the bottom, ranging from 3646 to 30000, with a green bar indicating the current value of 3650.

At the bottom of the interface, there are several tabs: Routine, Contrast, Resolution, Geometry, System, Physio, Diff, and Sequence. The 'Contrast' tab is currently selected.

Contrast/Common Card (all versions)

Grad rev. fat suppression (SE/diffusion only): If enabled, the polarity of the slice select gradients will be reversed between excitation and refocusing. This can provide effective fat suppression at no cost in time or SAR. It can

also result in signal losses in areas of B0 inhomogeneity. Typically it is most useful at 3T.

Contrasts (*Numaris X only*): The number of readout trains to acquire can be specified here (“echoes” for multi-echo EPI, *aka* ME-EPI). If two or more echoes are specified, the TE parameter on this card will change to an array displaying the TEs for each echo. Only the first TE can be modified; subsequent TEs will automatically be minimized unless “Inter-TE delay” is specified on the Sequence/Special card. *Note: This parameter is located on the Sequence/Part 1 card in VB/VD/VE versions.*

Resolution/iPAT Card (VB/VD/VE) Resolution/Acceleration Card (Numaris X)

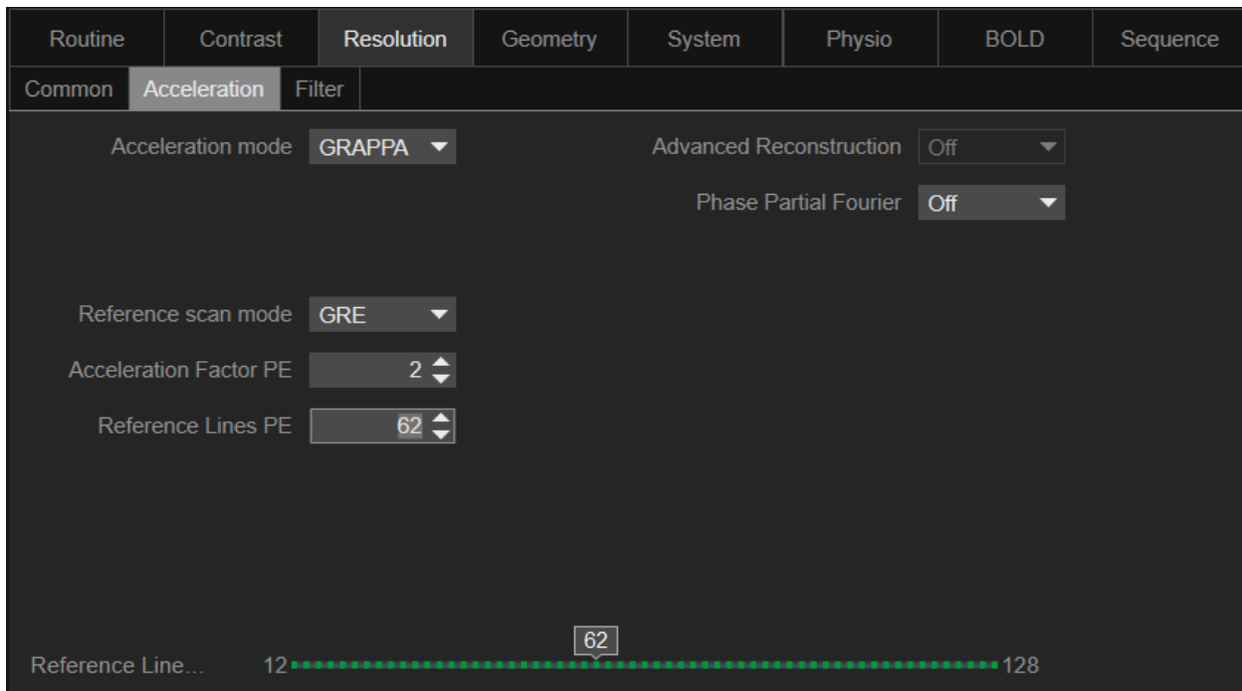
The screenshot displays the Resolution/iPAT Card (VB/VD/VE versions) with the following parameters:

- PAT mode:** GRAPPA
- Accel. factor PE:** 2
- Ref. lines PE:** 24
- Matrix Coil Mode:** Auto (Triple)
- Reference scan mode:** GRE

A slider for Accel. factor PE is shown at the bottom, with a value of 2. The slider has a range from 2 to 4, with a green bar indicating the current value.

The card is part of a sequence of cards: Routine, Contrast, Resolution, Geometry, System, Physio, Diff, and Sequence.

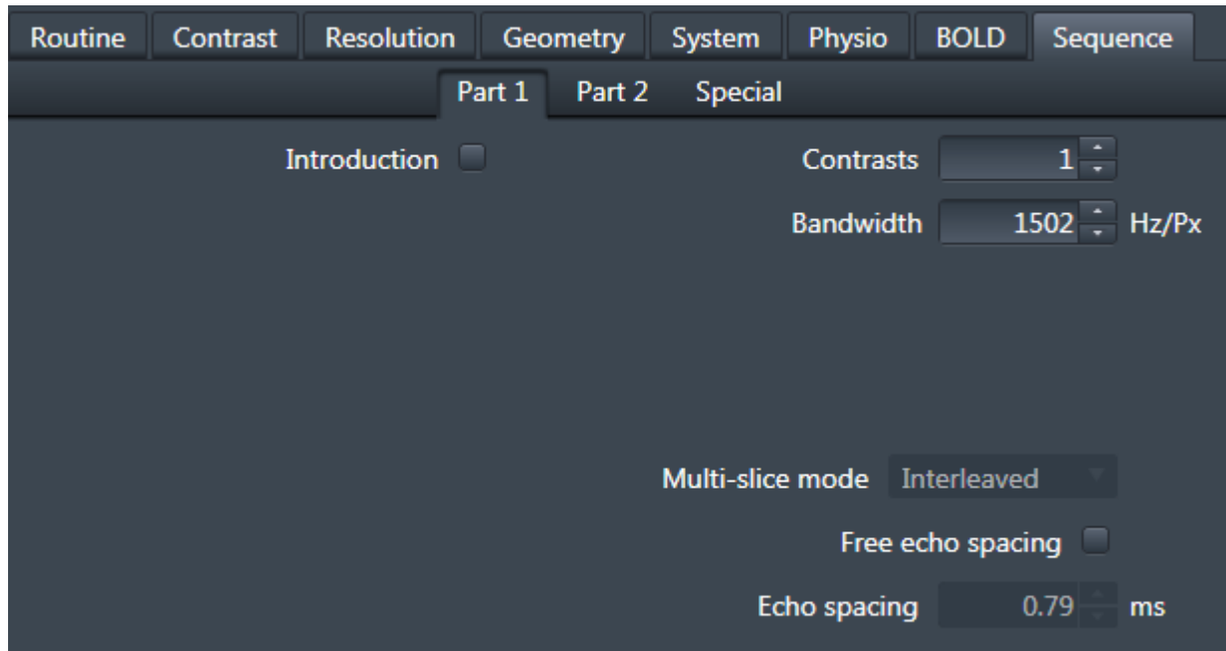
Resolution/iPAT Card (VB/VD/VE versions)



Resolution/Acceleration Card (Numaris X versions)

Reference scan mode: If iPAT is enabled, the reference scan acquisition scheme can be selected here. “Single-shot” EPI (Siemens default for accel. factor 2) is the fastest but limits the number of reference lines. “Segmented” EPI (Siemens default for accel. factor >2) allows for more reference lines, but increases motion sensitivity. “GRE” offers high SNR and is insensitive to motion, but may result in other artifacts (this option is currently experimental). “FLEET” also offers motion insensitivity with the speed of a segmented EPI acquisition, but comparatively low SNR. Note that steady-state preparation scans can be specified on the Sequence/Special card for GRE or FLEET modes.

Sequence/Part 1 Card



Sequence/Part 1 Card (VB/VD/VE versions)

Contrasts (VB/VD/VE only): The number of readout trains to acquire can be specified here (“echoes” for multi-echo EPI, *aka* ME-EPI). If two or more echoes are specified, the TE parameter on the Contrast/Common card will change to an array displaying the TEs for each echo. Only the first TE can be modified; subsequent TEs will automatically be minimized unless “Inter-TE delay” is specified on the Sequence/Special card. *Note: This parameter is located on the Contrast/Common card in Numaris X versions.*

Sequence/Special Card

The screenshot shows the 'Special' tab of the Sequence/Special Card. The interface is divided into several sections:

- Excite pulse duration:** A numeric input field set to 2560, followed by a unit dropdown menu set to 'us'.
- Physio recording:** A dropdown menu set to 'Off'.
- Single-band images:** A checkbox that is currently unchecked, with left and right arrow buttons below it.
- Online multi-band recon.:** A dropdown menu set to 'Online'.
- FFT scale factor:** A numeric input field set to 1.00, with up and down arrow buttons.
- Triggering scheme:** A dropdown menu set to 'Standard'.

At the bottom of the main area, there is a horizontal bar representing the excite pulse duration range. The bar is divided into green and red segments. The green segment starts at 600 and ends at 2560. The red segment starts at 2560 and ends at 16380. The value 2560 is marked above the bar.

At the very bottom, there is a navigation bar with the following tabs: Routine, Contrast, Resolution, Geometry, System, Physio, BOLD, and Sequence. The 'Sequence' tab is currently selected.

Sequence/Special Card (all versions)

Excite pulse duration: The required RF pulse voltage increases linearly with the multi-band acceleration factor. Here the pulse length of the excitation RF pulse can be manipulated manually at the cost of T_E and bandwidth.

Verify on the System/Transmitter/Receiver card that the voltage of all pulses, especially those with names similar to “MBExc” and “MBRef” (SE/diffusion only) are below the system maximum value. Otherwise, the pulse(s) will be clipped and the flip angle(s) will be less than expected. A warning message will pop up on the host if this occurs.

Refocus pulse duration (*SE/diffusion only*): This parameter will appear arrayed beneath the Excite pulse duration parameter in SE or diffusion sequences. It allows for manually controlling the duration of the refocusing pulse(s) only.

Min. prep scans: This parameter can be used to specify a minimum number of dummy scans to acquire (and discard) prior to the main data acquisition. Note: this parameter specifies a floor; more may be acquired automatically by the sequence.

Min. prep scans SB: If multi-band acceleration is enabled, this parameter can be used to specify a minimum number of dummy scans to acquire (and discard) prior to the single-band reference scan acquisition. Note: this parameter specifies a floor; more may be acquired automatically by the sequence.

MI-EPI shift factor (2D BOLD only): Note: This parameter will only appear if “Magn. Preparation: Non-sel. IR” is first selected on the Contrast/Common card, which may itself require first manually setting TR to a large value. This parameter is used to enable MI-EPI acquisition and select the block shift factor. Also known as “IR with rolling slices”; ref. Renvall, et al. *NeuroImage* 134 (2016) 338-354.

HS inv. pulse dur. / HS inv. pulse N / HS inv. pulse R (2D BOLD only): These parameters can be used to modify the nonselective inversion pulse, if “Magn. Preparation: Non-sel. IR” is selected on the Contrast/Common card.

Pause after Meas. (3D BOLD only): For 3D time series, this pause will be inserted between each measurement (i.e., between full 3D volumes). Use to insert a silent gap between measurements e.g., for auditory studies, equivalent to how “Delay in TR” is used for 2D.

3D slab overlap factor (3D only): This factor can be used to modify the thickness of the 3D slab selective RF to intentionally overlap or separate adjacent slabs (to address slab profile issues).

iPAT ref. prep scans: This parameter can be used to specify a number of dummy segments to acquire (and discard) prior to each slice of the GRE or FLEET iPAT reference scan acquisition (if selected), to allow time for the magnetization to approach a steady state.

FLEET ref. min. TR: This parameter can be used to specify a minimum repetition time between segments of the FLEET iPAT reference scan acquisition. If this value is not achievable (or the default of 0 is used), the minimum possible T_R will be used.

Inter-TE delay: If a multi-echo acquisition is specified (“Contrasts” greater than 1), this parameter can be used to add a delay between readout trains, which will increase the effective TE of all echoes after TE1. Normally this parameter is 0, such that TE2 and above are automatically minimized.

Delay before PC scans: Optional delay between the end of the pre-phasing gradients and the start of the first phase correction readout. Can be used in special cases where unusual eddy currents are present. Normally this parameter is 0.

Single-band images: Select this checkbox to send the single-band reference images to the database in a separate series (e.g. for quality control). NOTE: this option is not compatible with FMRI post-processing and motion correction (the single-band images will be quietly discarded in this case).

MB LeakBlock kernel: Select this checkbox to replace the default slice-GRAPPA kernel optimization with LeakBlock kernel optimization (ref. *MRM* 2014;72:93-102). This approach is designed to minimize inter-slice correlations, but may come at the cost of somewhat higher overall noise and within-slice artifacts. NOTE: this option may cause the reconstruction of the first image to be delayed due to increased initial computational requirements.

MB dual kernel: Select this checkbox to calculate separate kernels for odd and even lines (ref. *NeuroImage* 2012;63:569-580). This approach is designed to reduce ghosting, but it has not been carefully vetted. It is provided for experimental use.

MB RF phase scramble: Select this checkbox to optimize inter-band phase offsets in the RF pulse shape to reduce the required peak RF power (ref. *Proc. ISMRM* 2012, #2209). Overall SAR is unchanged. This option is currently only defined for MB acceleration factors 3-8.

Time-shifted MB RF (SE/diffusion only): Select this checkbox to temporally offset the bands in the RF pulse shape to reduce the required peak RF power (ref. *MRM* 2013;69:1261-7). Overall SAR is unchanged. For SE and monopolar diffusion, the “aligned- T_E ” scheme described in the reference is used. Refer to the tooltip for specific shift offset information.

SENSE1 coil combine: Select this checkbox to use a SENSE R=1 algorithm for SNR-optimized coil combination (ref. *MRM* 2013;70(6):1682-9). Coil combine mode “sum of squares” must also be selected on the System/Miscellaneous card.

Invert RO/PE polarity: Select this checkbox to automatically apply a 180-degree in-plane rotation to the acquisition parameters (e.g. to use for eddy current correction in post-processing).

Sparse TR acq.: Select this checkbox to disable readout and phase encoding gradients for selected measurements, e.g. to allow for “silent” periods during auditory scans (while maintaining RF steady state). Note: the on/off periods must be defined using the **Triggering scheme** option; *this checkbox will only appear if a triggering paradigm is active.*

PF omits higher k-space (BOLD only): When phase partial Fourier is active, select this checkbox to fully acquire the early k-space lines (before the k-space center) and omit the latest lines. This option can be used to reduce minimum T_R when long T_E is used. This is the opposite of the product (default) approach, which always omits the early k-space in order to reduce minimum T_E .

Disable freq. update: This option will disable dynamic transmitter frequency updating, which is intended to correct for frequency drift over the course of long scans (caused by e.g. gradient heating). This update adds a 20 ms dead period at the end of every T_R . Image drift in the phase encoding direction will still be corrected in post-processing if this option is selected.

Suppress 16-bit DICOM: This option will revert the image output to the traditional standard 12-bit DICOM format. This may be needed for

compatibility with some postprocessing software. Note that dynamic range is limited with this option, and care will need to be taken to adjust the **FFT scale factor** to ensure clipping does not occur.

Force equal slice timing: This option will force the inter-slice T_R fill delay to be exactly equal throughout the acquisition. Any remaining delay that cannot be evenly distributed between the slices (remainder of the division of the T_R by the number of slices) will simply be discarded, which means the actual T_R of the acquisition may be shorter than the protocol T_R . The tooltip will report the actual T_R . The **Disable freq. update** option must also be selected (if present).

Online multi-band recon.: *Online* will reconstruct the multi-band accelerated images in ICE and send them to the database. *Aliased* will send the aliased images directly to the database without reconstruction (for debugging). *Quick* will reconstruct using the fastest reconstruction options without regard to image quality (for rapid validation). *None* is equivalent to EmptyIce (e.g. for later reconstruction offline).

If the optional remote reconstruction service is configured, the option to enable it will be found here.

FFT scale factor: This parameter may be used to adjust the image brightness, i.e. to optimize dynamic range. Note that on VD11D and newer platforms 16-bit DICOM format is used.

Inversion pulse FA: This parameter can be used to correct the nominal flip angle of the nonselective inversion pulse (default = 180 deg), only if “Magn. Preparation: Non-sel. IR” is selected on the Contrast/Common card.

Fat saturation FA: This parameter specifies the flip angle to use for fat saturation RF pulses (default = 110 deg).

Fat sat. offset: This parameter specifies a frequency offset to apply to fat saturation RF pulses (0 = use the system calculated fat frequency).

GRE iPAT ref. FA: This parameter specifies the flip angle to use for the GRE iPAT reference acquisition.

FLEET iPAT ref. FA: This parameter specifies the flip angle to use for the FLEET iPAT reference acquisition.

Sinc exc. pulse BWTP (*BOLD only*): This parameter specifies the bandwidth*time product (*aka* R factor) of the sinc RF excitation pulse. Lowering this parameter from the default will reduce SAR, but at the cost of a less sharp slice profile.

Physio. recording: Control recording of physiological signals (cardiac, respiration, ECG, external) to text or encoded DICOM files. Text files will be stored in C:\MedCom\Log\Physio\ on the host computer (MRC). *Legacy* uses the classic CPmuSequence/IdeaCmdTool logging facility (this is the only option available on V_{Axx}/V_{Bxx} systems and VD11x; it is known to be unreliable on VD11x/VD13x/VE11x). *DICOM* and *File* use the new online/ICE logging available since VD13A, which is recommended for use when available. *File* writes the log data to individual text files for each signal (this option was named *Online* in previous versions). *DICOM* embeds these logs in a special DICOM “image” stored in the database (sample Matlab code for reading the special DICOM files is provided in the [GitHub repository](#)). *Multiple* enables both *Legacy* and *File* options (not recommended for routine use).

Triggering scheme (*BOLD/SE only*): This parameter, and the arrayed parameters below it, can be used to control the sending of the external trigger signal (EXT_TRIG) per slice, per volume, and/or controlled by a custom paradigm. Refer to the tooltips for details. The paradigm definition is similar to that of the BOLD card. If scheme “standard” is selected, the external trigger will be sent once at the start of each repetition (same behavior as the product sequences).

Important Issues (not bugs)

This release will expire on 10 February 2026!

Please refer to <https://github.com/CMRR-C2P/MB> for the latest information.

- Only diffusion modes “MDDW” and “Free” are compatible with multi-band. The other modes have been disabled. The MDDW license is therefore required to insert the MB diffusion sequence.
- It is not possible to use fMRI post-processing and/or motion correction and also save the single-band reference images. If both are selected, the ICE program will silently discard the single-band reference images. It is generally not recommended to use these post-processing steps since they slow down the online reconstruction and can very easily be performed offline (often with better results).
- Demanding diffusion acquisitions with high b values may fail with gradient duty cycle errors if the highest gradient amplitudes are requested on the same axis over several consecutive T_{RS} . This can be avoided by using e.g. a randomized vector table (in “Free” mode) to distribute the gradient demand more evenly over time. (This also applies to the product diffusion sequence.)
- Monopolar+ and bipolar+ diffusion schemes are not available. Unfortunately, Siemens does not make the code to enable these modes available for customer sequences.
- Reconstruction is sometimes slow, especially with iPAT enabled. This is unavoidable due to the complexity of the unaliasing calculations, but it is something that we continually aim to improve. It is recommended to have the fastest available MRIR/MaRS from Siemens. Using the [remote reconstruction service](#) can offer dramatic performance improvements with older MRIR/MaRS hardware.

Known Bugs

Please refer to <https://github.com/CMRR-C2P/MB> to read and submit bug reports.

Version Changes

Release 017pre13, 23 September 2025

- Fixed issue where an incorrect refocusing RF pulse was being used for single-band reference acquisitions in the diffusion sequence, resulting in increased artifacts (likely introduced ca. R017pre10).
- VE12U/Numaris X: The pop up messages indicating clipped RF pulses or an expired license now use the native syngo message service and should be impossible to be hidden behind the syngo window.
- The maximum allowed b-value for diffusion has been increased from 16000 to 50000 s/mm² for systems with high performance gradient sets.
- XA60A: Improved stimulation prediction for diffusion sequences on Terra.X systems.
- VE12U: Increased maximum Mosaic size from 4K×4K to 8K×8K. This may cause problems with DICOM viewers for very large mosaics.
- Added “Limit grad. capability” and “Disable new stim. pred.” checkbox options to closely match Prisma sequence parameters on Cima.X systems and Terra parameters on Terra.X W60. Please consult the UI tooltips for more information about the usage of these parameters (intended for testing/troubleshooting only!).
- Added ini file option to enable manual control of diffusion δ/Δ (for advanced users only; please consult the Wiki for usage).
- iPAT² protocols should no longer stop when magn/phas reconstruction is selected.

Release 017pre12, 23 April 2025

- Improved image database storage performance for XA60A-SP02.
- Added “NOISE” label to DICOM image type field for EPI noise scans.
- Added UI option to optionally split echoes into separate series (may improve image database storage performance on XA versions).
- Fixed issue where GRE iPAT reference scan type was not selectable when editing stored protocols on XA30A and newer versions.
- The “Fat saturation FA” parameter now has a minimum value of 30°. If a lower value is found, it will be reset to the default (110°). This

addresses an issue where protocols converted from VE to XA sometimes would have this parameter inadvertently set to 0°.

Release 017pre11, 10 February 2025

- Fixed logging of physiology data for XA61A and newer versions.
- UHF systems: Fixed UI issues restricting settings of RF pulse lengths in diffusion and SE sequences.
- Fixed timing calculation error when using Bipolar diffusion mode.
- Additional fixes for calculation of change proposals when gradient stimulation limits are exceeded.
- Corrected the placement of optional SB prep scans (all are now acquired before any iPAT reference scans).
- The in-plane iPAT factor is no longer restricted if MB acceleration is not enabled. If MB acceleration is enabled, the in-plane iPAT factor is limited to 4, and the total iPAT*MB acceleration is limited to 18 (for sanity).
- Slight reduction of maximum diffusion gradient for UHF Plus systems with SC72 gradient to avoid possibility of overflow errors.

Release 017pre10, 8 January 2025

- Fixed some cases where GRE or FLEET iPAT reference scans could not be selected.
- XA50A/XA60A: Fixed crash when attempting to log physiology data.
- XA61A: Added full support for high performance Cima.X and Connectom.X gradient systems in the diffusion sequence.
- ICE GPU code is now split into a separate binary and support scripts have been removed to simplify sequence installation and avoid missing driver issues on legacy systems unequipped with a GPU in the MaRS.
- The DisableGPABalance ini file option now works as intended (see Wiki for details).
- Additional improvements to SENSE1 phase reconstruction: Siemens coil compression and noise decorrelation are now disabled by default to avoid “black hole” artifacts at the site of phase discontinuities.

Release 017pre9, 15 May 2024

- The “MB RF phase scrambling” option is now enabled by default for new protocols. It is always recommended to use this option when it is available.
- Disabled the B_1 control loop in the BOLD sequence for all versions prior to XA61A due to stability/oscillation issue, pending Siemens fix.
- VE12U Terra systems: Added optional Skope synchronization and triggering.
- Improved the consistency and quality of phase correction for SENSE1 when output of phase images is selected (namely, interslice phase discontinuities should be eliminated). The use of this improved algorithm is indicated by “SENSE1++” in the image comment.
- Maximum diffusion gradient amplitudes are now reported in the diffusion scheme tooltip and saved in the DICOM header (see Wiki for details).
- “Sparse T_R ” mode should be more silent.

Release 017pre8, 23 February 2024

- Added support for XA60A, XA61A.
- Fixed the case where SENSE1 could be selected when using a single-channel receive coil, which would result in an image reconstruction error.
- Reduced the minimum b-value step size to 5 (to restore previous functionality in R016 and earlier versions).

Release 017pre7, 3 October 2023

- XA30A/XA31A: Updated the warning in the installation guide about certificate installation to clarify that security patch SD05 must be installed to prevent the system becoming unbootable.
- XA31A/XA50A: Fixes for logging of physiology data.
- XA versions: Performance improvements for logging of physiology data in DICOM format.
- XA versions: Restored the option for "legacy" physiology data logging.
- Fixed issue where the stimulation checker would always fail when attempting to calculate a change proposal in the BOLD sequence.

Release 017pre6, 27 March 2023

- Added support for VE11E, VE12U-AP02, XA20A, XA50A.
- XA versions: Removed restrictions on the use of 16-bit DICOM format and DICOM format physiology following the issuance of the XA30A-SP02 patch which increases system stability.
- VE11x/VE12x only: Fixed issue preventing use of the “Free” diffusion mode (for versions with build date of 25 April 2023).
- XA versions: Added warning to installation instructions about certificate installation causing systems to become unbootable.
- SBRef phase images are now stored in a separate series; phase series are labeled with the suffix “_Pha”.
- For protocols with multiple echoes and/or phase images, only the magnitude images from the first contrast will now appear in the inline display to reduce flickering.

Release 017pre5, 27 June 2022

- Added support for VE11C, VE12U-SP01, VE12U-AP01, XA31A.
- XA only: Temporarily disabled option to output physiology in DICOM format due to crashes on XA systems.
- Fixed issue with improper normalization of user-defined diffusion vectors in “Free” mode.

Release 017pre4, 3 June 2022 (first public release)

- Initial public release for XA30A
- XA only: Temporarily disabled the output of 16-bit DICOM format by default due to crashes on XA systems.

Release 017pre3, 11 April 2022 (limited release)

- Added option to force 12-bit DICOM format.
- Improved FLEET iPAT reference scan performance.

Release 017pre2, 11 January 2022 (limited release)

- Enabled UI for “diffusion averaging”, allowing for acquisition of multiple “averages” per diffusion b-value.
- Added UI options to manually set a minimum number of preparing scans for single-band reference scan and/or main acquisition.

Release 017pre, 7 December 2021 (internal release)

- Added support for Numaris X software versions.

- Added 3D EPI acquisition option for all sequence variants.
 - Note: sequences will continue to be named cmrr_mbep2d* for historical and sentimental reasons.
 - Options: single slab, multiple slabs, multiple slabs with multi-band acceleration; slice partial-Fourier; iPAT2 acceleration; manually adjustable slab excitation thickness/overlap factor.
- Added multi-TI T₁ mapping option (aka MIR, MI-EPI, multiple TI IR with rolling slices) with user configurable hyperbolic secant inversion pulse parameters (BOLD sequence only).
- Increased maximum allowed repetitions from 4096 to 32768 (BOLD & SE sequences).
- Added UI options to manually configure fat saturation frequency offset and nominal flip angle.
- Added UI option to configure preparing scans before GRE iPAT reference scan.
- Increased range of flip angles selectable in UI.
- Reduced minimum slice gap to -50%, minimum bandwidth to 30 (matches gre sequence).
- Reduced minimum allowed slice thickness.
- Added “Quick” MB image reconstruction option (can be used for testing/validation; automatically selects fastest reconstruction options, which may degrade image quality).
- Improved reliability of UI for selecting number of iPAT reference lines when using GRE or FLEET reference scans.
- Added UI option to configure sinc excitation RF pulse bandwidth (BOLD sequence only).
- Updated calculation of spoiler b-value displayed in monopolar diffusion UI to account for all axes.
- Removed nonfunctional phase scrambling option for MB=2, corrected description in UI.
- Improved consistency of SAR look-ahead calculation for MB protocols and/or when using alternative reference scan types (applies to VE11A and newer versions).
- Fixed bookkeeping issue with physiology log data recorded from the diffusion sequence (should now report correct total number of diffusion volumes).
- Fixed crash when attempting to use FLEET iPAT reference scan in a single-band only protocol (applies to VD13A and newer versions).

- Fixed RF spoiling for FLEET and GRE iPAT reference scan types when (legacy) slice multiplier option is used.

Release 016b, 8 January 2020

- VE11E/VE12U only: correct an issue where GPU acceleration could have been disabled for image reconstruction even when an appropriate GPU was present in the MaRS.
- VD11D only: adds support for GPU accelerated reconstruction on systems which have been upgraded with the "128-channel (VD13)" MaRS option.

Release 016a, 19 December 2017

- Added .ini-file override parameter to restore the option of using base resolutions < 64 (the limit was raised in R016 because it previously invoked a serious Siemens UI bug which could change the FOV phase % factor in existing protocols unknown to the user). Use this override at your own risk; see the [GitHub wiki](#) for details.
- Fixed issue on VE11A/B/C pTX systems where the "Excitation mode" was not being set to a valid default when upgrading protocols from previous C2P versions; this could prevent protocols from being printed.
- Fixed crash when selecting "Invert RO/PE" option in combination with FLEET iPAT reference scan.
- Fixed various minor issues where the reference scan UI could reset the reference scan type unexpectedly in response to enabling/disabling iPAT and/or changing the iPAT factor.
- Minor update to installation script to hopefully (but unlikely) address issues where for unknown reasons ICE updates on some systems occasionally are not recognized properly without a full power cycle.

Release 016, 20 November 2017

- The installation launcher has been updated for VE11C and newer versions with improved error handling.
- The C2P license has been moved into a separate file so that in the future it will be possible to update the license without needing to reinstall all sequence and ICE binaries.

- FLEET iPAT reference scan option added. *Note: the configurable parameters for FLEET (dummy scans, min. T_R , flip angle) have not been optimized.*
- All advanced shimming modes are now selectable for VB19A and VB20A.
- Various default gradient parameters have been updated to match the product defaults for Terra systems (VE11K/R/U).
- The MB RF phase scrambling offsets and various diffusion gradient parameters (Δ , δ , spoiler b-value) are now stored in the DICOM images (Phoenix protocol) for posterity. Information about how to retrieve these parameters will be available on the [CMRR MB-EPI wiki](#).
- “High dynamic range” (16-bit DICOM) mode is now enabled for Magn./Phas. reconstruction mode on VD13x/VE11x platforms.
- The phase only reconstruction mode option has been removed. For consistency, only two reconstruction modes are supported. Magnitude reconstruction is the default, and fully supports the standard Siemens EPI reconstruction pipeline. For Magn./Phas. reconstruction, “Maxwell” correction is automatically disabled. The “Maxwell” correction is an image-space dewarping which cannot be applied to phase images at this time. It is disabled to allow for coregistration of magnitude and phase images.
- The SENSE1 coil combination option now properly supports the calculation of phase images. This is the only recommended scheme for acquiring phase images with multi-channel receive coils.
- Fixed a UI issue where the Phase FOV sometimes could not be set to 100%.
- Patient-specific and volume-selective B_1 shimming modes are now selectable for all pTX-capable systems on VD13C/D and VE11x platforms.
- “Force equal slice timing” option added.
- Fixed scan time calculation and display when GRE/FLEET reference scans are enabled.

Release 015a, 11 April 2017

- Stability improvements for GPU reconstructions.
- Support multi-echo acquisitions when using “online” physiology logging.

- Triggering fix for “Sparse T_R ” mode.
- Added version check and file integrity verification to install script.

Release 015, 3 March 2017

- The SMS license is now required to enable slice acceleration on versions VE11C and newer.
- Added new option to embed “Online” physiology log data in a special DICOM file (VD13x/VE11x and newer versions).
- The scale of respiratory traces should no longer change during an acquisition when using “Online” physiology logging.
- ECG traces should now be stored at the highest temporal resolution (400 Hz) when using “Online” physiology logging.
- “Learning phase” physiology samples will no longer be stored when using “Online” physiology logging.
- Inactive/unused signal sources will not produce log files when using “Online” physiology logging.
- If a K10 GPU is detected in the MaRS, both logical devices will be used properly for improved reconstruction performance.
- The excitation pulse voltage is now displayed correctly on VD13x/VE11x.
- Various “common” ICE errors will now cause friendly/informative popup messages to appear on the host.
- The installation script will report an error if an encrypted file is encountered (this can occur inadvertently when extracting .zip archives to NTFS volumes on macOS systems).
- Additional triggering paradigm options added.

Release 014, 20 July 2016

- VE11C version added.
- Multi-echo support fully enabled.
- GPU support for accelerated image reconstruction enabled for VE11x versions.
- Maximum allowed MB factor limit raised from 4 to 8 for diffusion variant; maximum combined MB*x*iPAT factor limit raised from 12 to 18 for all variants.
- Partial GPU support added for systems with K10 GPUs.
- “Sparse T_R ” acquisition option added to allow for the prescription of “quiet” periods during a scan, e.g. for auditory studies.

- Protocol option added to disable frequency feedback, e.g. for timing-dependent studies.
- Dual kernel reconstruction option added for evaluation.
- Fixed UI and protocol conversion issues related to hidden “SB fake slice bands” option.
- Image reconstruction stability improved for very small matrix sizes.

Release 013a, 7 January 2016

- Fixed bug in VB17A/VB19A/VB20P diffusion variant which prevented existing protocols from being loaded without being “fixed” and reset to default parameters.

Release 013, 14 December 2015

- VE11A/VE11B versions added.
 - Note: GPU support is currently not available for VE11A/VE11B versions.
- VD13x/VE11x: New ‘Online’ physiology logging option. This should be much more reliable than the scheme used in previous versions (still available as ‘Legacy’ mode). Example MATLAB code for reading the new file format is available in the [GitHub repository](#).
- New protocols created with this version will now default to using the LeakBlock kernel.

Release 012d, 14 December 2015

- VD13C/D: The pop up message indicating clipped RF pulses was not being displayed; this has been fixed.
- VD13A/B/C/D: Eliminated some harmless but severe-looking error messages that could appear in the logs at the start of sequence execution (particularly with physiology logging enabled).

Release 012c, 13 October 2015

- A pop up message will now be displayed on the host at runtime to clearly indicate when the sequence has expired; protocols will still be visible and editable after expiration.
- VD13x: Corrected 10 μ s per T_R timing error when real-time frequency feedback is enabled.
- VD13x/VD11D: If online motion correction is enabled, 16-bit DICOM encoding (“high dynamic range”) will be disabled due to

incompatibility with the Siemens online motion correction routines. Using the online motion correction is not recommended.

- If magnitude+phase reconstruction is selected, a new option to force enabling of the Maxwell correction is available. Otherwise, by default, Maxwell correction is disabled for phase image reconstructions.
- Minimum T_E has been slightly reduced for diffusion acquisitions (allows for replicating the HCP/Lifespan 3T protocol exactly on Prisma systems).
- The maximum number of repetitions allowed is no longer affected by the number of receiver coils selected (e.g. the Head/Neck 64).
- Fixed preparation failure when selecting a GRE iPAT reference scan in combination with time-shifted RF pulses.
- The default RF time shift has been adjusted to match the published 7T HCP protocol (for 5.12 ms excitation pulse, 1.92 ms shift).
- On Tx-Array systems, the clipped RF pulse check has been disabled since it can not be made compatible with amplitude correction factors (CP_mode.ini) $\neq 1$ on arbitrary transmit channels.
- Due to poor performance, GPU acceleration will be automatically disabled if a Tesla K8, K10, or similar device is detected.
- If the remote reconstruction service is configured, an option is now available to save images directly to disk on the server for real-time processing with minimal latency.
- Fixed reconstruction failure for multi-echo, single-band acquisitions.
- Various UI fixes for multi-echo protocols (generally improved handling of arrayed T_E values).
- Fixed reconstruction error for multi-echo + multi-band acquisitions when using GPU acceleration on VD13x.
- Reduced ICE log chatter...

Release 012b, 18 May 2015

- VD13D: On systems with dual GPUs, reconstruction should no longer occasionally fail with kernel errors.

Release 012, 19 November 2014

- An option has been added to use LeakBlock (aka Split slice-GRAPPA, ref. Cauley *MRM* 2014;72:93-102) kernel optimization in place of the default slice-GRAPPA kernel. This option may offer reduced leakage artifacts (inter-slice crosstalk) at the cost of overall

reduced SNR, increased within-slice aliasing artifacts, and increased reconstruction time.

- The slice acquisition timing reported in the DICOM files (e.g. MosaicRefAcqTimes in the shadow header) should now be physically correct for all repetitions as well as for the single-band reference. *This time it should really be true...*
- The default CAIPIRINHA PE shift has been changed from FoV/3 to FoV/4 for iPAT R=2 protocols to improve unaliasing performance.
- When time-shifted RF pulses are being used in combination with iPAT using the GRE reference scan option, the reference scan acquisition should now be shorter and offer improved performance.
- The single-band reference images are now always acquired with linear slice order to reduce the possibility of banding artifacts that may appear during interleaved acquisitions (the MB acquisition is unchanged).
- Minimum allowed T_R may be shorter for certain protocols, especially on systems with >40 mT/m body gradient sets (Prisma, Connectom).
- An internal hard limit on maximum allowed RF pulse bandwidth has been removed. This will allow for more coverage in the slice direction, but B_1 transmit homogeneity over large bandwidths has not been carefully evaluated on all systems (warnings will be written to the trace log when the expected limit is exceeded).
- The popup message that appears when the requested RF flip angle cannot be achieved due to insufficient RF peak power should no longer appear repeatedly if a SAR or stimulation change proposal is being calculated by the system.
- VD13: Image reconstruction should no longer fail at the end of a scan (with no data loss) if SENSE1 coil combination is used with DTI postprocessing enabled.
- Image reconstruction should no longer fail at the start of a scan if an Image Numbering scheme (defined on System/Miscellaneous card) other than the default is used.
- VD13: When the “log physiology to file” option is selected, the EXT_2 signal will now also be logged.
- An option to use RF spoiling is now available (possibly useful for extremely short T_R).
- Optional flow compensation in the slice direction is now available in the BOLD variant.

- The “free echo spacing” option is no longer disabled.
- Other bug fixes and UI performance improvements.

Release 011a, 9 June 2014

(eja: vb17a/d7f82ea, vb19a/da328c9, vd11d/1f6d304, vd13a/de62ca8, vd13c/af3cafa, vd13d/5cd7089)

- Fixed image reconstruction preparation error when selecting GRE-based iPAT reference scan option.
- GRE-based iPAT reference scan can now be used in combination with time-shifted MB RF pulses (SE, diffusion variants).
- Fixed UI issue where diffusion mode “Free” could not be selected in new protocols under VB17A/VB19A/VD11D.
- VD13x: Fixed failure to prepare image reconstruction on systems without a CUDA GPU (48-channel MARS).
- VD13C: Corrected CUDA version; GPU support should now work properly when a GPU is present.

Release 011, 23 May 2014

(eja: vb17a/d9ea75d, vb19a/504763c, vd11d/483fe46, vd13a/5a1c0a3, vd13c/1b34f94, vd13d/c791fc7)

- GRE-based iPAT reference scan option added (note: segmented reference scan was used for MB+iPAT acquisitions in all previous versions).
- MB acquisitions with descending slice order should work properly.
- The slice timing info in the image shadow header under MosaicRefAcqTimes should now be accurate for all MB repetitions (previously the info for the first repetition was incorrect).
- VD13: GPU acceleration has been enabled for VD13A/C/D systems with a 64- or 128-channel MARS2 installed (only newer revisions with Tesla C2070 or better). Most scans using 32 or fewer channels should reconstruct in real time. This unfortunately does not seem to be true for >32 channels...
- VD13: All available advanced B₀ shim modes are now selectable.
- VD13A: “Tx Feedback” is now enabled by default on VD13A Skyra systems for improved stability. If you have a Skyra with an older Tx box and MB scans stop with TxK2304 errors, contact Siemens to have it upgraded, or in the meantime contact us or see the support web page for instructions on how to disable it again.

Release 010c, 28 March 2014

(eja: vd13a/59bad81, vd13c/1b2c8d6, vd13d/4b0f402)

- VD13C: Fixed issue preventing SE sequence from being inserted.
- VD13A: Fixed issue where diffusion mode Free could not be selected in new protocols, related diffusion UI slowness.
- VD13D: New protocols now default to gradient mode Performance if it is available (e.g. for Prisma).
- Removed non-functional diffusion modes (only MDDW and Free are supported).

Release 010b, 13 December 2013

(eja: seqs only all versions; vb17a/41a9b7e, vd11d/R010/dae2076, vd13a/sp2/3451f7c, vd13a/R010/fd129fc, vd13d/R010/7d45ede; added 29 Jan 2014: vb19a/6a5af2e, vd13c/2bfb96d)

- VB17A: Fixed scanner crash and reboot if a non-MB sequence was used following a MB sequence in the same session.
- VD13D: Fixed incorrect orientation labeling of slices with the “invert RO/PE” option selected.
- VD11D/VD13x: Fixed inconsistent synchronization of frequency feedback acquisition. This could result in timing errors of ~20ms per T_R and degraded image quality for certain protocols.
- If the “log physiology to file” checkbox is selected, the directory C:\MedCom\log\Physio will now be created if it does not already exist.
- iPAT reference scan for R=2 is now segmented as it was in previous versions (protocols with larger numbers of reference lines should no longer break on import).
- Descending slice order should now work properly (really).

Release 010a, 22 November 2013

(eja: vb17a/857e1d9 all seqs, vd11d/R010/2638d72 diff seq only)

- VB17A: Fixed display of current measurement in the status bar.
- VB17A/VD11D: Fixed inability to select diffusion mode Free.
- VB17A: RF pulse durations can now be set freely for single-band SE/diffusion protocols on UHF systems.

Release 010, 20 November 2013

(eja: vb17a/r206, vd11d/R010/6fea7e1, vd13a/sp2/49ca23b, vd13a/R010/5b5bc96, vd13d/R010/eacfbce)

- The number of preparation scans acquired at the start of the measurement has been reduced, notably for diffusion protocols and protocols using iPAT.
- VD13A: Transmitter feedback is now specifically disabled for Skyra systems by default. This new feature is currently causing crashes with multiband pulses; Siemens may fix it in a future hardware update.
- Gradient reversal fat suppression is now an independently selectable option. In previous versions it was automatically enabled if fat suppression modes “none” or “strong” were selected (following Siemens default behavior since VD11).
- New options have been added for sending optical trigger signals per slice or repetition.
- New options have been added to use phase scrambling and/or time-shifting approaches to reduce peak RF power.
- Clipped RF pulse warning window should no longer pop under the exam window.
- Descending slice order should now work properly.
- VB17: iPAT>2 should work now on pTX systems.
- VD13: Images should no longer jump in the phase direction.
- Copying sequence parameters from other protocols should be more reliable.
- UI responsiveness and stability should be improved.

Release 009a, 21 May 2013

(eja: seq/ICE VB17A r158/r157, VD11D r236/r234, VD13A-SP2 r22/r20)

- VB17: Fixed inability to load existing diffusion protocols.
- Minimum T_E/T_R should now match previous versions.

Release 009, 15 May 2013

(eja: VB17A r157, VD11D r234, VD13A-SP2 r20)

- First “development” release.
- Added VD13A-SP2 version.
- Dramatic image quality improvements. All images should appear significantly sharper, and MB+iPAT performance is especially improved.

- The maximum allowed MB factor has been increased from 8 to 16.
- Allowed MB*iPAT factors have been increased to 12.
- Many user interface improvements. The user interface should be more responsive and more stable in general.
- All diffusion modes aside from MDDW and Free have been disabled. Previously the other modes were selectable but would crash image reconstruction.
- Saving single-band images in combination with FMRI postprocessing and/or motion correction will no longer crash image reconstruction (the single-band images will be discarded).
- Minimum T_E/T_R /thickness increments have been reduced.

Release 008, 13 February 2013

(eja: seq/ICE VB17A r121/r122; VD11D r206/r205)

- First “stable” release.
- Added options for reconstructing phase images
- MB+iPAT reconstruction improvements

Release 007, 21 November 2012

(eja: VB17A r113; VD11D r195)

- Reconstruction speed improvements
- Fixed occasional occurrence of intensity variations between bands in some (mostly diffusion) protocols
- VD11: Siemens product frequency feedback/update enabled
- VD11: Checkbox added to invert readout and phase encoding gradient polarity (useful for post-processing eddy current corrections)
- SENSE1: image comment now indicates whether SENSE1 has been applied
- SENSE1: susceptibility-induced signal loss reduced

Release 006b, 21 June 2012

(eja: VB17A r92)

- VB17: fixed cases in monopolar diffusion where spoiler gradients failed to be played out for single-band $b=0$ reference scan

Release 006a, 23 May 2012

(eja: VB17A r91; VD11D r154)

- Fixed memory leak which caused recon to eventually crash for long series

- VB17: fixed FFT scaling factor for SENSE1
- Minor improvements to exported physiological log data

Release 006, 18 May 2012

(eja: VB17A r88; VD11D r151)

- Expiration date extended to 1 December 2012
- Added spin-echo EPI sequence variant
- Fixed clipped RF pulse dialog false alarms (for real this time...)
- VD11: fixed crash when physiological logging is enabled
- VB17: improved EPI ghost correction for BOLD variant
- VB17: fixed DTI recon preparation failure for certain syngo patch levels
- Limited max. MB factor to 3 for diffusion
- Removed non-working “free echo spacing” checkbox
- Misc. UI improvements

Release 005, 6 April 2012

(eja: VB17A r75; VD11D r125)

- Image quality improvements for MB7 and MB8
- Added SENSE1 coil combination option
- Restored option to save single-band reference images to a separate series
- Fixed clipped RF pulse dialog for VB17 diffusion refocusing pulses
- Further reconstruction speed improvements

Release 004a, 20 March 2012

(eja: VB17A r72; VD11D r111)

- Fixed incompatibility with protocols created with earlier revisions
- Reconstruction speed improvements

Release 004, 20 March 2012

(eja: VB17A r71; VD11D r110)

- Fixed DTI recon crash when postprocessing was explicitly not selected
- Added warning dialog on host if (MB) RF pulses will be clipped
- Fixed crash when creating new DTI protocols
- Fixed gradient errors on systems with lower slew rates (e.g., Verio)
- UI responsiveness, stability improvements

- Removed “single-band images” option due to unresolved ICE crashes; will restore in a later release

Release 003, 27 January 2012

(eja: VB17A r64; VD11D r78)

- Reconstruction speed improvements
- MB+iPAT image quality improvements
- Added option to save single-band reference images to a separate series
- Reduced number of prep scans at start of BOLD measurement
- Adaptive coil combine now compatible with MB (VD11D only)

Release 002a, 4 January 2012

(eja: VB17A r60; VD11D r61)

- VB17A ICE program expiration date fixed

Release 002, 18 November 2011

(eja: VB17A r58; VD11D r61)

- Diffusion variant (ep2d_diff) released
- iPAT (GRAPPA only) enabled
- Option to log physiological data to file added
- Additional FFT scale factor UI parameter added (VB17 only)
- Bug fixes

Release 001, 26 September 2011

(eja: VB17A r50; VD11D r36)

- Initial public release