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Recent Advances in 3D Time-Resolved Contrast-Enhanced MR Angiography

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Abstract

Contrast-enhanced MR angiography (CE-MRA) was first introduced for clinical studies approximately 20 years ago. Early work provided 3 to 4 mm spatial resolution with acquisition times in the 30 sec range. Since that time there has been continuing effort to provide improved spatial resolution with reduced acquisition time, allowing high resolution three-dimensional (3D) time-resolved studies. The purpose of this work is to describe how this has been accomplished. Specific technical enablers have been: improved gradients allowing reduced repetition times, improved k-space sampling and reconstruction methods, parallel acquisition particularly in two directions, and improved and higher count receiver coil arrays. These have collectively made high resolution time-resolved studies readily available for many anatomic regions. Depending on the application, approximate 1 mm isotropic resolution is now possible with frame times of several seconds. Clinical applications of time-resolved CE-MRA are briefly reviewed.

Keywords

MRA; contrast-enhanced MRA; time-resolved studies; fast imaging; parallel imaging

INTRODUCTION

There are many clinical situations in which time-resolved contrast-enhanced MR angiography (CE-MRA) may be valuable. The multiple images of a time-resolved acquisition may improve diagnostic accuracy by depicting arterial anatomy without venous contamination in cerebral, extremity, and renal and visceral artery imaging. In some cases timing a single-phase CE-MRA scan may be difficult such as in the setting of inflow disease or hyperemia causing asymmetric flow in the lower extremities, or because of the rapid transit time and small contrast bolus volume in children. Time resolved imaging may be useful in visualizing unusual vascular flow patterns, such as retrograde filling of a vessel distal to an occlusion.

One of the initial demonstrations of three-dimensional (3D) CE-MRA was Prince's work in the mid-1990s (1) which showed the ability to image the vasculature in 3D but with coarse spatial resolution. Shortly thereafter there were efforts to simultaneously improve the speed of acquisition and provide improved spatial resolution. In the quest to perform time-resolved imaging one is confronted by a fundamental tradeoff in MRI between acquisition time and spatial resolution. In general, unless some separate physics principle is newly applied or the underlying pulse sequence is altered, reduction of the acquisition time in MRI must be accompanied by a reduction in the spatial resolution.

In the last 20 years the discovery, exploitation, and implementation of various MRI physics principles and the development of improved MRI hardware has radically altered the temporal vs. spatial resolution tradeoff of CE-MRA of the mid-1990s, allowing more than an order of magnitude reduction in the acquisition time. In numerous vascular territories time-resolved CE-MRA is now possible with spatial resolution superior to that which was previously available only in single phase imaging. In this article we provide an overview of this evolution, review methods contributing to this, and describe applications. This work can be considered a companion to recent review articles which have covered parallel imaging (2), MRI temporal acceleration techniques (3), and time-resolved angiography (4). Non-contrast-enhanced (NCE) MRA as reviewed recently (5) constitutes a significant sub-field of its own and is beyond the scope of this article. As the name implies NCE-MRA methods do not require contrast administration. They generally depend on pulsatile or steady arterial blood flow, are subject to signal loss from multiple RF excitations, and for various reasons require acquisition times of minutes. For CE-MRA these first issues are less problematic, and the short T1 relaxation time of enhanced blood allows significantly shorter acquisition times.

THE EVOLUTION OF TIME-RESOLVED MRA

Perhaps the principal take-home point of this review article is that multiple technical advances over the last 20 years have allowed significant, indeed radical, reduction in acquisition time, providing marked improvement in the spatial-temporal resolution of time-resolved 3D CE-MRA. In this section we trace the evolution in performance by selecting a specific anatomic region, the vasculature of the calves, and observing how performance has improved. Table 1 shows this evolution, and is explained as follows.

Assume that bilateral CE-MRA of the calves is to be performed with 3D Fourier Transform (3DFT) acquisition in coronal format: frequency encoding in the superior/inferior (S/I) direction, phase encoding left/right (L/R), and slice encoding anterior/posterior (A/P), with corresponding field-of-view (FOV) of $400 \times 320 \times 132 \text{ mm}^3$. Several possible $Y \times Z$ spatial resolutions within the axial plane are shown heading the columns of Table 1, resolution improving left to right. The absolute $N_Y \times N_Z$ sampling resolution is shown in the third row. For the typical gradient echo pulse sequence used in CE-MRA with one phase encoding measurement or "view" sampled per repetition interval TR, the number of repetitions used for 3DFT acquisition is $N_Y \times N_Z$. However, for all entries in this table it is assumed that the corners of k_Y - k_Z space are not sampled (6), providing more isotropic resolution within the $Y \times Z$ plane, allowing a 25% reduction in this $N_Y \times N_Z$ product. Shown in the fourth row is the

resultant total number of necessary repetitions. The acquisition time is equal to this number multiplied by TR.

Just as moving left to right in Table 1 corresponds to improved spatial resolution, moving downward in the table from the fifth row on corresponds to progressive reduction in acquisition time, with entries for specific repetition times (TR, in msec) shown within groupings for specific acceleration factors $R = 1$ (no acceleration), 2, 4, and 10, where R is the reduction in scan time allowed by parallel acquisition. The evolution of performance corresponds to migration from the top left to the bottom right of the table. Specific entries to be discussed are highlighted in color. Contrast-enhanced MR angiograms which illustrate the various spatial resolution combinations heading the columns are provided in Figure 1.

As a starting point, the parameters described in Prince's early work in abdominal CE-MRA (7), which used a repetition time of 14.1 msec, are applied here to the calves. As highlighted in the table, this corresponds to an acquisition time of 32 sec for the coarsest spatial resolution considered. After this initial demonstration of CE-MRA, investigators quickly recognized the importance of small TR times for enabling high spatial resolution, and the next row in the table shows results for TR 10 msec.

Shortly after the initial demonstration of 3D CE-MRA the view-sharing TRICKS technique (8) was introduced, also highlighted in Table 1. This permitted a reduced frame time, here shown as 8 sec, but due to more frequent sampling of central k-space caused some increase in temporal footprint (32 sec) vs. single phase imaging (23 sec). Note that the times indicated in the table are based on the parameters shown for each column and row and thus might differ from the values reported within the cited references. A subsequent study of TRICKS (9) for imaging the vasculature of the calves used TR times of 7.8 msec and provided improved performance; i.e. moved the operating point in Table 1 to the right and below that for the highlighted entry.

Another distinct step in the development of contemporary CE-MRA was the incorporation of parallel acquisition. Early work (10-12) used one-dimensional (1D) acceleration factor $R = 2$ as highlighted in Table 1 for Ref. (10), with the parameters used therein for renal MRA converted to the peripheral vasculature for this presentation. The acquisition speed of these studies also benefited from TR times as low as 5 msec.

3DFT acquisition incorporates phase encoding along two directions, allowing two-dimensional (2D) (13) vs. 1D parallel acquisition. An early example of 2D acceleration with $R = 4$ for CE-MRA is highlighted (14). In the ensuing years multiple investigators further developed parallel imaging for the periphery (15-17). Highlighted is the work of Bonel (18) which used acceleration $R = 3$ with the GRAPPA technique (19).

Additional development of 2D acceleration, possibly in combination with other undersampling methods such as partial Fourier acquisition and homodyne reconstruction (20), has allowed net acceleration factors of $R = 10\times$ or more (21,22), also highlighted. Effective use of acceleration factors this high requires appropriate receiver coil arrays as will be discussed.

VIEW SHARING

Virtually all time-resolved methods in 3D CE-MRA use view sharing (23) to provide a frame time shorter than the intrinsic acquisition time for a single image. MRI data are measured in the spatial frequency domain of the image, referred to as “k-space,” rather than in the image domain itself. Because individual spatial frequencies are measured serially, images can be generated more frequently than the intrinsic acquisition time for sampling all of k-space.

Although non-Cartesian implementation is possible, this section assumes 3D Cartesian acquisition as in the previous section. In this case analysis of k-space sampling is facilitated by focusing on the k_Y - k_Z plane. All points are located on a rectilinear grid, an individual point (k_Y, k_Z) in this plane is sampled each TR interval, and because sampling along k_X for such a point occurs virtually instantaneously within the several-msec-long echo, effects along k_X can be ignored. Figure 2a schematically shows the possible sampled points of k_Y - k_Z space as well as a central sampling region (orange) and three colored sectors in peripheral k-space (red, green, blue) extending out to some maximum radius k_{MAX} . As assumed in Table 1, the corners of k_Y - k_Z space are not sampled. Time-resolved 3D acquisition is performed by sampling the points within each region according to some prescribed time order and frequency.

The use of sectors in (a) allows clear distinction of individual subsets of peripheral k-space points. With contemporary time-resolved methods alternative decompositions of peripheral k-space are generally used. For example, the “Cartesian Acquisition with Projection Reconstruction-like sampling” (CAPR) technique (24) uses interleaved sets of vanes (b). This tends to better disburse any artifacts due to undersampling vs. the approach in (a). The “Time-resolved angiography With Stochastic Trajectories” (TWIST) method (25) takes this a step further by placing peripheral k-space points into interleaved spiral trajectories based on their distance from the k_Y - k_Z origin. This yields a random appearance as depicted in (c). As a reference the original TRICKS technique (8) used subsets which were progressively located along the $\pm k_Y$ direction (d). Other decompositions are possible, such as “Differential Subsampling with Cartesian Ordering” (DISCO) (26) which has a pattern similar to (c). Once all regions are defined the acquisition proceeds as follows.

Figure 3 shows how view-shared methods are implemented in CE-MRA. As a reference, Fig. 3a shows the typical timing of a CE-MRA exam. Contrast material is injected intravenously and after some delay appears in the targeted arterial vasculature. A short time later the contrast-enhanced blood appears in the companion venous vasculature. The injection-to-arrival time and arterial-venous delay time depend on the arterial vasculature under study and have high patient-to-patient variability. Typical ranges are 5 to 50 sec for the former and 2 to 15 sec for the latter (27,28). The challenge of CE-MRA is to acquire a high spatial resolution image timed to the arterial phase with negligible, potentially confounding enhancement of the companion veins.

A progression of acquisition strategies is shown in the remainder of the figure. In Fig. 3b the “Acq” line shows how the regions of k_Y - k_Z space of any of the methods in Figs. 2a-d might be sampled over time. For purposes of discussion suppose that each colored block is eight

seconds long. The corresponding “Recon” lines show how the acquired data are selected to reconstruct images. Image I1 is reconstructed from the colored data selected within the corresponding bracket, and it is ascribed to time T1, designated with the short vertical arrow. The simplest way to generate a time-resolved image series is to repeat this process multiple times as in (b), yielding Image I2 at time T2, etc. In this case the time between successive images, or the frame time (FT), is equal to the time required to sample all of k_Y - k_Z space, defined as the temporal footprint (TFP) (24). Depending upon when the sequence is initiated, Image I1 or I2 might or might not capture the arterial phase.

With view sharing (c) it is recognized that image reconstruction can be done more frequently than the time required to sample all of k -space. As seen, the data acquisition is identical to (b), and Image I1 of (c) is formed identically to (b). However, Image I2 is reconstructed after the next sub-region of k -space has been sampled (O2). Samplings R1, G1, B1 are “shared” for images I1 and I2. Because the frame time in (c) is four times shorter than in (b), one might think that this method would be more likely to capture the arterial phase. However, the overall appearance of the image is dictated by the sampling of the orange central k -space region. Because the frequency of sampling this region has not changed vs. (b) the performance is not markedly improved. Expressed another way, because images such as I2 and I3 share their central k -space sampling (O2 in this case), they provide redundant information.

View sharing in Cartesian sampling is best exploited if central k -space is sampled more frequently than peripheral k -space (d). In this case the orange central region is sampled prior to each of the three peripheral regions, and an image is reconstructed corresponding to each of these samplings. Compared to (b) the frame time is two times shorter, and because central k -space is sampled more frequently than in (b) the likelihood of capturing the arterial phase is improved. In fact, in (d) the central k -space sampling O_{12} for image I2 is better aligned temporally with the peak arterial signal in (a) than the methods in (b) or (c). However, this is done at the expense of the temporal footprint being longer, six intervals now vs. the four of (b). If one is willing to allow a temporal footprint this long, an alternative acquisition is one which uses this extra time to sample k_Y - k_Z space even more peripherally to obtain a single phase image with even higher spatial resolution. This is shown in the expanded k -space coverage of Figure 2e and the data acquisition and reconstruction in Fig. 3e. In this case one generally uses some kind of timing method (29,30) to initiate this high resolution scan with arterial contrast arrival.

Finally, Figure 3f shows the case when parallel acquisition is applied to the sequence of (d). In this example $R = 3$ is assumed, causing the width of each colored block to be reduced proportionately vs. the previous parts of the figure. Compared to (d), the frame time and the temporal footprint are similarly reduced three-fold. Comparison with the contrast enhancement waveforms in (a) suggests that one or more images of the accelerated sequence in (f) will be temporally aligned with the arterial phase. Also, the reduced temporal footprint reduces the sensitivity of the arterial phase image to venous enhancement.

Modern day time-resolved CE-MRA can be performed with acceleration factors R of 2 to 4 being routine, 4 to 6 being common, and 8 to 16 being possible for certain applications.

Factors $R > 3$ further compress the temporal playout beyond that shown in Fig. 3f. Rather than use such high acceleration factors solely for reduction of frame time and temporal footprint, some of the advantage can be directed toward improved spatial resolution. That is, sampling of more peripheral k-space regions can be incorporated into the sequence while still providing a frame time and temporal footprint smaller than in (d). This usage has allowed migration of the operating point in Table 1 both downward (faster) and rightward (improved spatial resolution).

Prior to the advent of parallel acquisition two other methods used to provide a reduced frame time were keyhole imaging and temporal interpolation. Keyhole imaging (31) repeatedly samples only central k-space during the arterial enhancement period, sampling peripheral k-space only at the end of the scan. This generally results in blurring, akin to that seen in Fast-(FSE) or Turbo-Spin-Echo (TSE) T1-weighted sequences (32). In both cases the peripheral k-space samples have attenuated magnetization levels, in keyhole from waning contrast enhancement and in FSE/TSE from T2 decay. Temporal interpolation was used, for example, in the original TRICKS method (8). Rather than representing the object status at the interpolated time, however, temporal interpolation results in a superposition of the object status at both reference times. The advent of acceleration largely obviates both of these methods, and TRICKS can be used with acceleration (33).

In considering the data sorting for the sequences in (d) and (f) one might question which sampling of central k-space to use in reconstruction. For example, three samplings (O_{12} , O_{13} , O_{21}) are within the temporal footprint for Image I2. Here, the earliest sampling within each footprint was used for illustration. The specific choice is up to the user; considerations are discussed in Ref. (34). Use of an early sampling can cause artifactual, premature enhancement of vessels, called “anticipation artifact.” Use of a late sampling can cause diminished spatial resolution at the leading edge of contrast. Averaging the multiple central samplings blunts the temporal sharpness of only one sampling. Whatever the choice, it is best to maintain it for all images in the series to provide consistent image-to-image portrayal of a changing signal.

Another consideration is the specific time ordering of sampling the phase encoding views within each k-space region. This is arbitrary. Typically, elliptical centric (35), some centric-like sampling such as “Contrast-ENhanced Timing-Robust Angiography” (CENTRA) (36), or a combined centric-in/out sampling of central k_Y - k_Z region is employed (37) to preserve temporal sharpness. Similarly, the number of peripheral k_Y - k_Z regions and the relative size of the central vs. peripheral regions are additional variables. Determination of specific values can depend upon the application, as studied, for example, in renal imaging (37) or fluoroscopic tracking (38).

ACCELERATION IN 3D CE-MRA

Partial Fourier acquisition is performed in MRI for up to two-fold reduction in the number of samples in some direction of k-space (39) and is noted here as a means for reducing scan time in 3D CE-MRA. In one implementation asymmetric sampling is done in k_Y - k_Z space with zero filling (40). In another (41) the sampled and non-sampled regions in peripheral k-space are interleaved and homodyne reconstruction (20) performed. Partial Fourier can also

be done along the readout in both Cartesian (40) and radial acquisition (42) for TR reduction.

Parallel acquisition was first described over a decade ago and has been reviewed various times subsequently, e.g. (2,43,44). This section presents those aspects important for 3D time-resolved CE-MRA. As discussed in these reviews, the measurement of separate coil sensitivity images or autocalibration lines is necessary for SENSE-like (13,45) and GRAPPA-like (19,46) approaches, respectively. This can be done once, either before or after acquisition of the accelerated, time-resolved data sets, or possibly embedded within the accelerated acquisition itself using approaches such as TSENSE (47) or TGRAPPA (48).

3D CE-MRA imaging of the calves is shown schematically in Figure 4a. As assumed for Table 1, frequency encoding (X) is S/I, phase encoding (Y) is L/R, and slice encoding (Z) is A/P. As in Figure 2 the phase encoding (k_Y - k_Z) plane can be isolated for analysis (d). Assume that each open blue circle corresponds to an individual repetition of a reference unaccelerated acquisition. With 3DFT acquisition separate acceleration factors, R_Y and R_Z , can be applied along the respective phase encode directions Y and Z, and the overall acceleration is the product, $R = R_Y \times R_Z$. In k_Y - k_Z space this is implemented by proportionately increasing the sampling distance along the corresponding direction. For example, if the original acquisition in (d) is accelerated by $R_Y = 5.4$ and $R_Z = 2.5$, yielding $R = 13.5$, the resultant sampling pattern corresponds to the orange points shown. This illustrates that: (i) acceleration factors along Y and Z can be different from each other; (ii) acceleration factors need not be integer; and (iii) the actual k-space points sampled can be different in the accelerated vs. the non-accelerated scan. An aliased axial subtraction image from one element of a 16-element array used in acquiring such an accelerated CE-MRA scan of the calves is shown in (e), and the unaliased result in (f). The corresponding MIP image is shown in supplemental Figure S1.

Parallel acquisition requires that the MR signal generated during each repetition of the acquisition be simultaneously detected by each element of a multi-element receiver coil. Thus, the receiver coil array is a critical enabler for effective parallel acquisition.

The receiver coil array must meet several technical specifications. In general the number of coil elements must match or exceed the acceleration factor R. Also, the sizes, positions, and orientations of the coil elements over the FOV must cause the coil sensitivity profiles to be dissimilar from each other. If not, the reconstruction process is prone to severe SNR loss due to mathematically inverting a poorly conditioned matrix. For $R = 2$ SENSE acceleration this is equivalent to solving two equations in two unknowns in which the coefficients of the two equations are nearly identical. The coil sensitivities are analogous to these coefficients. Slight alteration of the two measurements causes major uncertainties in the computed unknowns.

Many coil arrays have been used for parallel acquisition. A progression is illustrated in Figure 4 for the calves. Consider first the four-element case (a) and point P at the surface of the left calf near Coil 1. As one moves from this point through the calf in the posterior direction toward Coil 2, the sensitivity of Coil 1 to the point diminishes owing to the

increased distance, while that of Coil 2 increases because the distance from Coil 2 is decreasing. That is, the sensitivities of Coils 1 and 2 along this Z direction are dissimilar, or more specifically, complementary. Consequently, this coil orientation supports acceleration along Z. Also, Coils 1 and 4 have complementary sensitivities for paths along the L/R direction, supporting acceleration along Y. This configuration has been used for 2D acceleration with $R = 4$ (14). This can be extended to an eight-element array (b) in which the elements are wrapped completely around the calves. Acceleration along the Y (L/R) direction is improved vs. (a) because of the laterally placed elements, and the higher coil count allows higher R values.

The degree to which a coil array provides good quality acceleration due its coil elements having dissimilar responses can be assessed mathematically using the “g-factor” (45). This is a mathematical construct based on the sensitivities of all coil elements and defined to have a numerical value of 1 (unity) or higher. A value of 1 indicates perfect response, i.e. no loss of signal-to-noise ratio (SNR) due to the de-aliasing process. The higher the value, the greater is the SNR loss. For a specific coil array and acceleration factor R, the g-factor can be computed at every point within the FOV. Fig. 4g shows in blue the range of g-factors over the 3D calves volume for various 2D acceleration combinations using the coil shown schematically in (b). Fig. 4h is a CE-MRA image of the calves using this coil and an $R = 7.3$ CE-MRA acquisition. Technical interpretation of these results suggested that the array could be improved if the anterior and posterior elements were adjusted to have a more rapid sensitivity falloff with increased depth in the object and all elements elongated for better S/I coverage. This led to the design of narrower anterior and posterior elements (c), resulting in a different g-factor, shown in the yellow plots in (g), with markedly reduced (improved) values vs. the blue plots. Fig. 4i shows a CE-MRA result with the modified coil of (c) acquired with $R = 8$. In spite of the higher acceleration factor vs. (h), it has higher SNR, consistent with the reduced g-factors.

In the last decade multi-element whole-body coil arrays have been developed as described in e.g. Refs. (49,50), and vendors now offer systems with as many as 100 or more individual coil elements. These are applicable to 2D-accelerated time-resolved CE-MRA. Performance is determined by the number of elements available and deployed for a given FOV, their orientation, and the overall SNR they provide.

3D time-resolved CE-MRA has been one of the most successful applications of parallel acquisition in terms of high ($R = 4$) acceleration factors being common. This is due to several fundamental factors. First, decomposition of the overall acceleration factor R into individual accelerations, $R = R_Y \times R_Z$, as allowed by the 3D acquisition of CE-MRA, is superior in preserving SNR in the final image vs. performing acceleration R solely along one direction. Second, 3D CE-MRA is typically acquired in coronal or sagittal format with the two phase encode directions transverse (A/P and L/R). Because coil elements can be placed facing each other around the body, these directions are more robust for high acceleration factors than the longitudinal (S/I) direction. Third, MRA images in general, and subtraction CE-MRA images in particular (e.g. Fig. 4f), are sparse, in that the percentage of voxels within the 3D FOV with non-zero signal is small. This reduces the overall signal power and objectionable artifact due to any errors in the mathematical correction for the k-space

undersampling. Finally, with its increased speed of covering k-space, accelerated CE-MRA better maps the high, arterial phase, enhanced signal over the entirety of k-space. This provides an overall SNR enhancement effect compared to CE-MRA without acceleration (51) and improved vessel sharpness even with the same sampling resolution as an unaccelerated scan (52). These benefits do not apply when imaging non-time-varying signals. 2D parallel acquisition can likely be further improved by better dispersal of aliasing (53) or parameter optimization on a patient-specific basis (54).

Figure 5 is a plot of temporal footprint vs. frame time for CE-MRA of the calves and shows the dependence of performance on the levels of view sharing and acceleration.

NON-CARTESIAN AND NON-TRADITIONAL METHODS

The discussion thus far has focused on Cartesian k-space sampling methods. In this section alternative acquisition strategies are considered. There can be a number of fundamental reasons for doing this. Non-Cartesian acquisition strategies such as projection reconstruction (PR) disperse artifacts more uniformly across the image, possibly making them less objectionable than the discrete ghosts characteristic of Cartesian methods. Also, dispersed incoherent undersampling artifact is one of the desired conditions for compressed sensing.

Lauterbur's initial description of MRI in 1973 (55) used PR acquisition, and some of the differentiators of PR with Cartesian were noted subsequently (56). An early description of PR application to CE-MRA (57) pointed out the relatively benign nature of the artifact in angularly undersampled MRA. This work combined 2D radial sampling in the k_x - k_y plane with Cartesian sampling in the k_z direction, a so-called "stack of stars" sampling of k-space. Time-resolved CE-MRA results were shown using view sharing. Subsequent work built on this (58-60). Multiple echoes can be sampled per repetition for increased speed in PR (42). Other non-Cartesian k-space sampling strategies can be performed in the k_x - k_y plane such as spiral acquisition (61,62). Rather than the subtle low-signal spoke artifacts of 2D PR, spiral further disperses the artifact in a less coherent fashion due to the further complexity of the spiral trajectory.

PR can be extended to three dimensions (63,64), providing an additional dimension for dispersal of undersampling artifact, potentially allowing greater undersampling than 2D-based PR. One implementation has been dubbed Vastly Undersampled Isotropic Projection Reconstruction or VIPR. 3D PR is particularly compatible with imaging an FOV which is nominally the same size in all three dimensions, such as the brain. With both 2D and 3D PR the low spatial frequency content associated with the overall image contrast is collected at each projection and allows images to be reconstructed on a projection-by-projection basis, providing considerable flexibility in the specific times, temporal footprints, and spatial resolution for which images in a time series are reconstructed. To best exploit this, the ordering of projections is generally done quasi-randomly (65). In contrast to this, the times at which Cartesian-based sampling is reconstructed are linked to the times at which central k-space is sampled, as was noted in Fig. 3. However, the temporally compact sampling of central k-space with centrally-ordered Cartesian methods better freezes transient behavior (34).

Almost a decade ago the method of compressed sensing (CS) was described for reconstruction of a function using reduced sampling of its frequencies (66,67) and was first adapted for MRI by Lustig (68). CS applied to MRI encompasses a broad framework for performing k-space undersampling and image reconstruction, continues to be studied extensively, and a review is beyond the scope of this article. CS techniques can be viewed as another means for acceleration and used synergistically with methods such as view sharing and parallel acquisition. As one example, CS was applied to time-resolved Cartesian acquisitions to restore SNR otherwise lost due to the 20× to 40× acceleration of 2D SENSE and zero filling (69). In another example (70) CS was used with multi-echo 3D radial acquisition and a 12-element receiver coil to provide time-resolved images of the brain with the 1 sec frame time, an approximate 100× acceleration vs. unaccelerated Cartesian sampling.

Another approach used for time-resolved acquisition is highly constrained projection reconstruction (HYPR) (71) with variations (72,73). As applied to CE-MRA this method acquires a high-resolution composite image by sampling the entire or an extended duration of contrast bolus transit. Next, a time series of individually undersampled “raw” frames is formed, typically by sub-sampling the composite image over short time intervals. Each raw frame is then multiplied by the composite to yield the final time series. This step reduces the artifact level, noise, and blurring characteristic of the individual raw frames. Due to the composite being shared in the formation of each final image, the final image series has image-to-image correlation (74) characteristic of view sharing. Although used extensively with PR, HYPR can also be used with Cartesian sampling (75). In one variant the composite image is formed with one type of sequence, such as non-contrast-enhanced phase-contrast (76) or time-of-flight (77), and the HYPR time frames acquired with a coarsely-sampled, contrast-enhanced run.

MR IMAGING OF TIME-VARYING SIGNALS

Time-resolved techniques are generally used in MR angiography because of the desire to portray the arrival or transit of the contrast-enhanced blood. In the ideal, non-attainable limit, all k-space data would be acquired instantaneously at the desired times of imaging. In practice, the mapping of the time-varying signal to k-space as dictated by the speed and k-space trajectory of the pulse sequence profoundly affects the portrayal of the contrast bolus in the image series. Here several aspects of this are described.

Consider Figure 6, taken from Ref. (78), a schematic representation of the arteries in one of the calves. Suppose that contrast-enhanced blood is entering from above the field-of-view shown, is flowing downward, and has an instantaneously abrupt leading edge. At time $T = 0$ this edge is positioned at level (a), and a time-resolved MRA acquisition is initiated in which all of the central portion and some fraction of the periphery of k_Y - k_Z -space are measured using random sampling similar to Fig. 2c. This is represented in the leftmost k-space diagram of the top row. After this data has been accumulated an image is reconstructed. Because not all of peripheral k-space has been sampled with contrast present in the vessel, the signal level in the vessel is not at full value and the vessel is not portrayed with full sharpness.

Next assume the process continues: at time $t = T$ the bolus edge has advanced to level “b,” and the next cycle of the acquisition is repeated, starting with a second sampling of the k-space center followed by a different subset of the periphery. These newly sampled data are used to update the orange central k-space of the first image and provide increased peripheral k-space sampling as depicted in the second from left group in the top row. Upon reconstruction this results in improved signal and sharpness in the image at level “a”. At level “b” however, the vasculature is portrayed with only one subset of peripheral k-space sampled with contrast-enhanced signal, similar to level “a” for time $t = 0$. The process continues for additional cycles until all of k-space is fully sampled, yielding full signal and spatial resolution.

Figure 7 shows these effects experimentally. In both (a) and (b) the spatial resolution is 1 mm isotropic and the frame time is 2.5 sec. However, extensive view sharing was used in (a), with eight frame times required before all of k-space was fully sampled. In the first few images the vessel edges are blurred and artifacted, and it is only until the fourth or fifth image in the series that the quality approaches that required for diagnosis. A more accelerated acquisition was used in (b) in a different volunteer in which only five frame times were required for complete k-space filling. The third or even second image in this series adequately shows the vessels.

These examples show that when the true signal level at a pixel changes over time, it takes several time frames for the signal and spatial resolution in the reconstructed image to “catch up” to the new true level and is a consequence of the sampling of all of k-space not being instantaneous. One might attempt to compensate for this by sampling more of peripheral k-space after central k-space is sampled. However, if an image is ascribed to some time T because that is the time at which central k-space is measured, then the use of any data measured after that specific time in forming that image will cause artifactual non-zero signal in blood vessels in advance of the contrast material actually arriving there. This is called “anticipation artifact” and has been described in Cartesian (34) as well as non-Cartesian (79,80) MRA acquisition. A tell-tale sign is the premature appearance of slight but non-zero signal in venous structures several frame intervals prior to significant enhancement. This can be reduced by reducing the temporal footprint and by sampling central k-space near the end of the footprint for any time frame.

Similar effects were observed a decade ago in studies of injection rate (81). For optimum performance the temporal footprint of the MR acquisition approximately matches the duration of high contrast enhancement.

The above considerations can have consequences in depicting a time-varying contrast bolus in blood vessels. Figure 8a shows a plot of a reference curve (blue) in a presumed S/I-oriented enhancing vessel and approximations of this curve as determined from four assumed acquisition techniques. The vessel diameter is relatively large, 32 mm within an assumed $320 (Y) \times 132 (Z) \text{ mm}^2$ FOV. A view-shared unaccelerated ($R = 1$) scan (green) provides the least accurate estimate of the reference. All three other acquisitions provide high fidelity. In (b) a much smaller vessel diameter is assumed, 6 mm. In this case improved match to the reference is evident in going from unaccelerated $R = 1$ (green) to $R = 4$ (purple)

to $R = 8$ (cyan) cases. For $R = 4$, as view sharing is increased from four peripheral k-space subsets (N4, purple) to eight (N8, orange), although the frame time is reduced, the fidelity is degraded because the temporal footprint increased per the behavior in Fig. 5. As expected, speeding up any sequence, Cartesian or non-Cartesian, provides improved fidelity in portrayal of the contrast bolus due to reduction of the time necessary to sample all of k-space. This is usually traded off with reduced SNR. These and other effects are studied in detail in the references (24,34,79,80,82).

APPLICATIONS

Clinical applications of time-resolved CE-MRA have been widely discussed in the literature. This section provides a sampling of this. The specific application and the choice of time-resolved vs. single phase CE-MRA vs. non-contrast-enhanced MRA vs. use of another modality such as digital subtraction angiography (DSA) or computed tomography angiography (CTA) is highly variable due to both patient-specific factors and radiologist preference.

Head, Neck, and Brain—Imaging of the carotid arteries and the intracranial vasculature is a major application of MRA, and multiple kinds of sequences can be used. Applications of time-resolved CE-MRA include capturing a clear arterial phase of the carotid (25,83-85) and intracranial arteries (24,86-90). Time-resolved acquisitions can simplify existing clinical protocols by avoiding need for a timing bolus (91). Large field of view (FOV) time-resolved acquisitions of the supraaortic vasculature have been developed for covering the great vessel origins and neurovascular anatomy in a single scan (92-94) but sometimes with a need to reduce A/P coverage or resolution to permit adequate temporal or spatial resolution. Single-phase CE-MRA using 2D SENSE with $R = 9$ and 16 has been demonstrated using a 16-element coil (95). A common target for time-resolved CE-MRA is to capture the rapid transit of contrast material in arterial-venous malformations (AVMs) (24,33,76,86,96-101), as shown with the HYPR technique in Figure 9.

Thorax—Imaging the vasculature of the thorax is particularly challenging due to respiratory and cardiac motion. Time-resolved techniques have been used to visualize the transit of contrast material through the pulmonary circulation (102-104). By using a stack of spirals (62) or stack of stars (105) very short, 1.0 sec or less, frame times can be achieved. An example of pulmonary imaging is shown in Figure 10. Time-resolved techniques can also be useful for visualizing the aorta and great vessels, e.g. (15), and improved thoracic coil designs and parallel imaging strategies have allowed portrayal of perfusion of lung parenchyma over the large FOV (106,107). Another application of time-resolved MRA is the evaluation of congenital cardiovascular disease (108,109), and for pediatric cases in particular the exposure-free aspect of MR makes it preferable vs. CT (110). A low-contrast-dose, time-resolved CE-MRA run (40) can be performed in conjunction with a high spatial resolution, single phase study to provide comprehensive interpretation. An example is shown in Figure 11.

Abdomen—The abdomen is also a challenging region to image with MR because of respiratory motion and the large FOV. A common study in abdominal MRA is to rule out

renal artery stenosis. Non-contrast-enhanced methods (5) have been applied but typically require several minutes of acquisition time. Renal CE-MRA has been an area for application of acceleration techniques for both improved single-phase imaging (111-113) and time-resolved techniques (37,114). The former can be challenging due to the need for accurate timing. In Ref (37) the authors adjusted the TWIST sampling to match the anticipated enhancement pattern of the kidney. In Ref. (114) the authors adjusted the CAPR sampling to allow sub-second frame times for timing as well as single phase, high resolution imaging in a split-dose protocol.

Lower Extremity—In imaging the lower extremity one distinction to be made at the outset is whether the study is limited to a single position or “station” of the patient table or if it requires multiple stations, such as for a complete runoff study. The latter imposes additional technical complexity.

Consider single-station imaging first. Compared to thoracic and abdominal imaging, the lower extremity is less prone to motion artifact. However, timing a single phase scan after injection is subject to a greater range of delay times due to the distal nature of the vasculature. Also, the proximity of veins to a companion artery makes proper timing to the arterial phase critical. Yet another factor to be sensitive to is the presence of unusual cross filling or retrograde filling patterns (17,59). Provided the spatial resolution is adequate, time-resolved CE-MRA can well match these factors. As portrayed in Figure 1, the performance has improved due to many techniques previously identified in this work (14,17,21,22,49,54,64,75,115-118). Acquisition parameters for a number of these recent reports are listed in Table 2, and performance noted in Figure 12. A number of these works compare CE-MRA with DSA (116,118) or CTA (119).

The proximal vessels of the feet are often included in multi-station imaging, but in some instances it may be necessary to focus on one or both feet. A recent review of CE-MRA of the hands and feet (120) described how the improved speed of acquisition has facilitated bilateral FOV, isotropic sub-mm resolution, and frame times of several seconds. Such a study allows distinction of arterial vs. venous vasculature in the presence of differential L vs. R contrast arrival. The improvements in spatial resolution allow improved identification of possible sites of anastomosis for surgery (117,121,122).

Next consider multi-station imaging. To keep pace with the advancing contrast bolus from station to station while still dwelling at an individual station long enough to obtain adequate spatial resolution is technically challenging. Since the initial descriptions of multi-station MRA (123,124) considerable effort has been spent to simply get high spatial resolution, single phase images across the entire FOV, e.g. (125,126). Acceleration methods (127,128) and improved coil arrays (49,129) have helped advance this. However, these acquisitions can still be limited with inadequate resolution or venous contamination in the distal-most station. A common way to mitigate this is to use a hybrid approach in which a time-resolved acquisition is done of the calves using a first contrast injection, followed by a dual-station single-phase scan of the abdomen and thighs using a second injection (16,130,131) as illustrated in Figure 13. Recently “fluoroscopic tracking” has been described in which a single injection is used, high (R 8) acceleration is applied at all stations, 3D images are

reconstructed in real time, and table movement is controlled by the operator to match the advancing contrast bolus (38,132).

Upper Extremity—MRA of the hand is a specialized application and also one in which NCE methods have been developed (5). Similar to the feet, the vasculature is relatively distal, making timing uncertain. High spatial resolution is required, and asymmetric L vs. R contrast arrival adds uncertainty. Multi-element receiver coil arrays can be devised for this region, allowing high acceleration (120). Recent literature shows how technical performance has been improving (120,133-137), allowing 1 mm³ acquired resolution and good venous suppression. Applications of time-resolved CE-MRA in the upper extremity include imaging vascular malformations (138) and surveillance of dialysis shunts (139).

DISCUSSION AND SUMMARY

Since its inception almost two decades ago the spatiotemporal resolution of 3D CE-MRA has improved markedly. This advance has been due to increases in the speed of acquisition, as facilitated with improved gradients, reduction of the number of sampled k-space points, as allowed with various acceleration methods, and further permitted with view sharing, which allows a frame time shorter than the intrinsic acquisition time of a single frame. However, when taken to an extreme each has its limits, such as degraded SNR, aliasing artifacts, temporal blur, or premature portrayal of contrast arrival.

Perhaps more than any other application in MRI, time-resolved CE-MRA has benefited from the reduced acquisition times allowed by parallel acquisition.

Due to incomplete filling of k-space, the first image of a time series portraying contrast arrival in an FOV will generally be subject to less than full signal as well as some loss of spatial resolution. If possible, second and subsequent images should also be used in interpretation.

Although outside the scope of this article, acquisition techniques for time-resolved CE-MRA can be adapted to dynamic contrast-enhanced (DCE) perfusion imaging, and this is an area of growing interest (26,140,141).

Time-resolved CE-MRA can be used in many anatomic regions.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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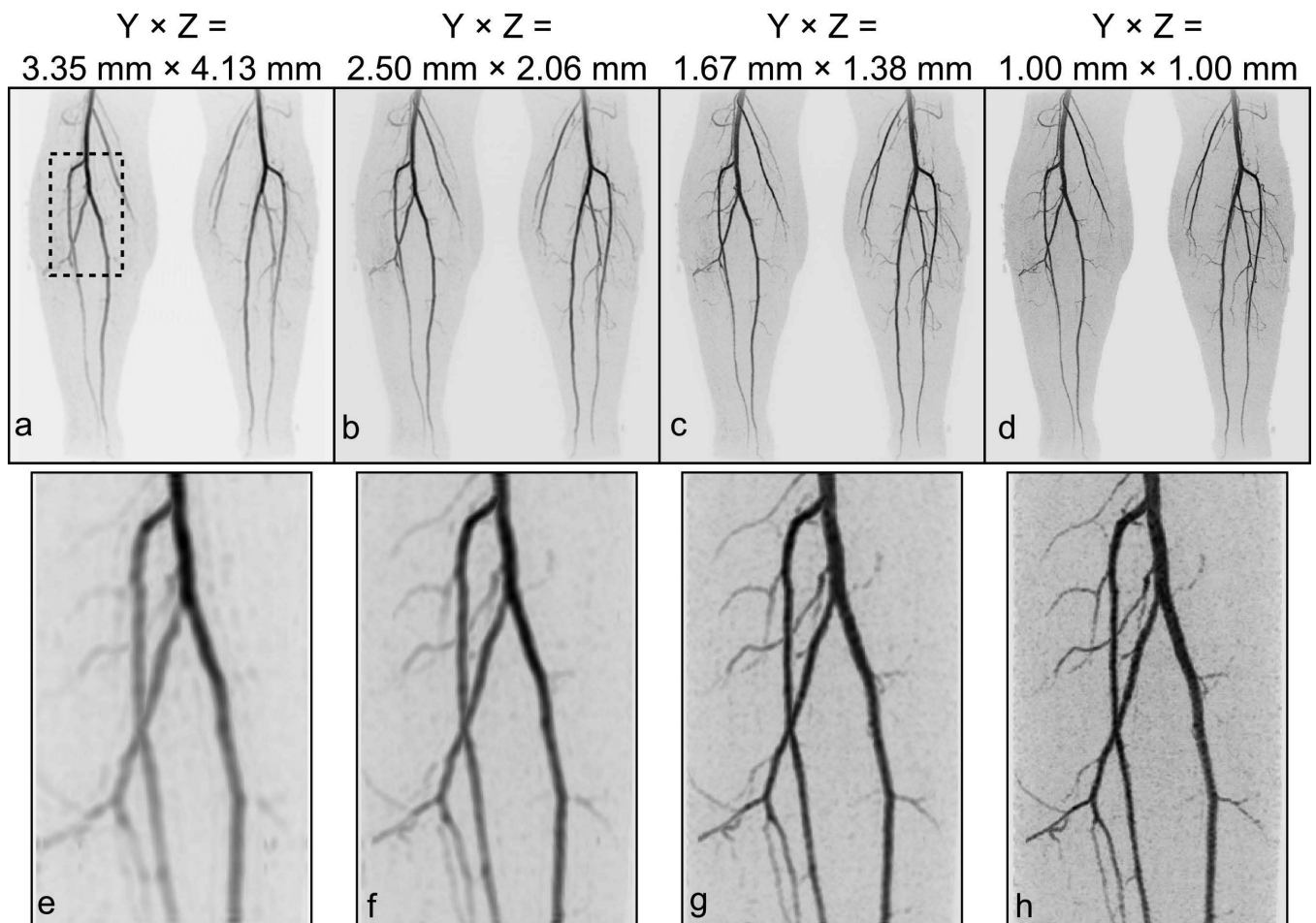


Figure 1. Illustration of the evolution of spatial resolution of contrast-enhanced MRA. Shown for each resolution is a coronal maximum intensity projection (MIP) of the full FOV (a-d) and a targeted MIP (e-h) at a slight angle of a small region of the right calf (identified in the dashed box in (a)). The progressions of images from (a) to (d) and from (e) to (h) correspond to the progression of axial resolution indicated in the column headings of Table 1.

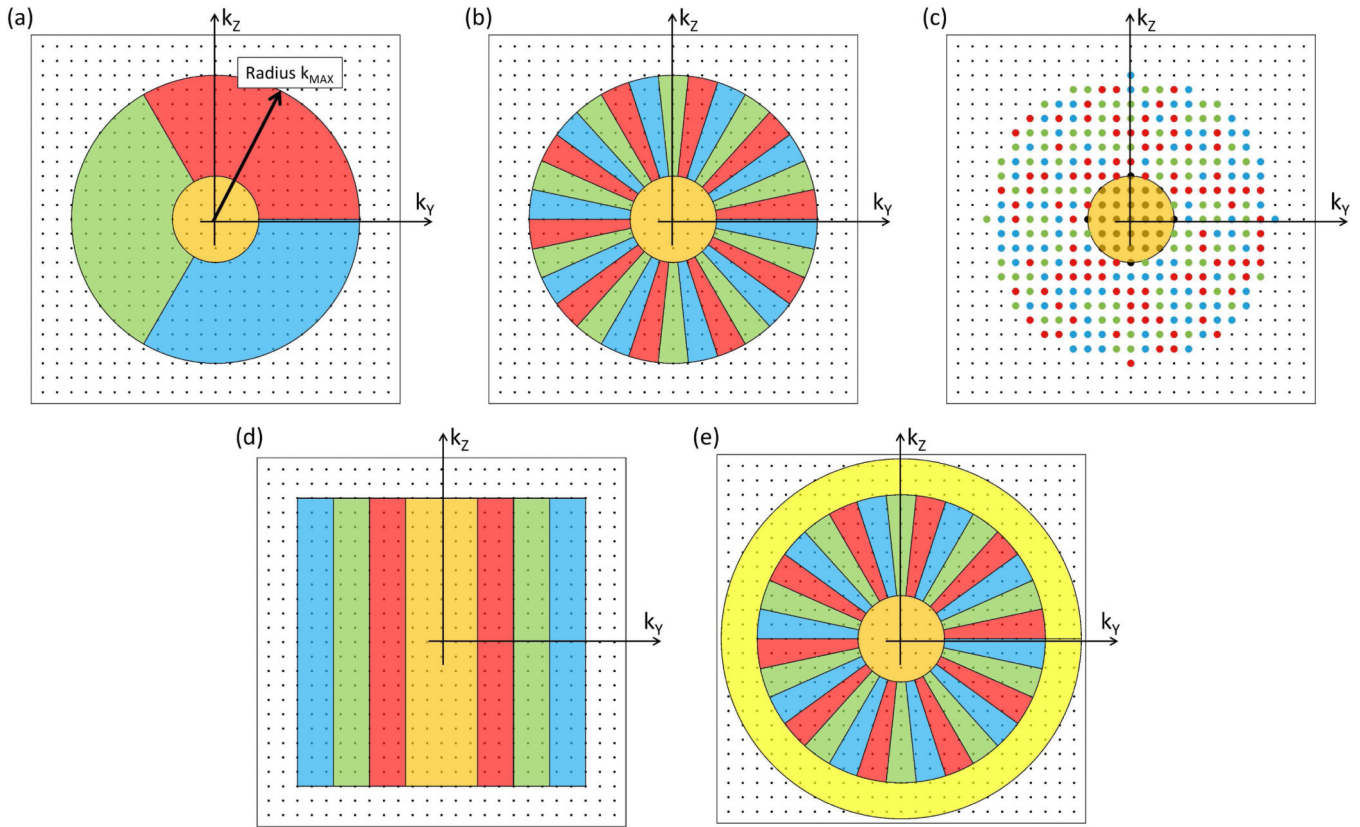


Figure 2. k -space sampling methods for view-shared 3D CE-MRA. Shown for (a) – (e) is a plot of k_y - k_z space. Contemporary methods generally first identify a central region which encompasses the k_y - k_z origin (orange). Next, sampling is defined for peripheral k -space, extending outward to some radius, k_{MAX} , and subdivided into groups. One straightforward way for subdividing is to use sectors (a) with three shown here (red, green, blue). To better disperse artifacts when only central k -space and a limited number of subsets of peripheral k -space are sampled, the points among groups are more interspersed. With the CAPR method (b) this is done using slender vanes. For even better dispersion peripheral k -space can be apportioned into three random-appearing sets (c). For reference the original TRICKS technique used grouping along the k_y direction (d). For improved spatial resolution one can adjust the acquisition to sample out to larger k -radii (e).

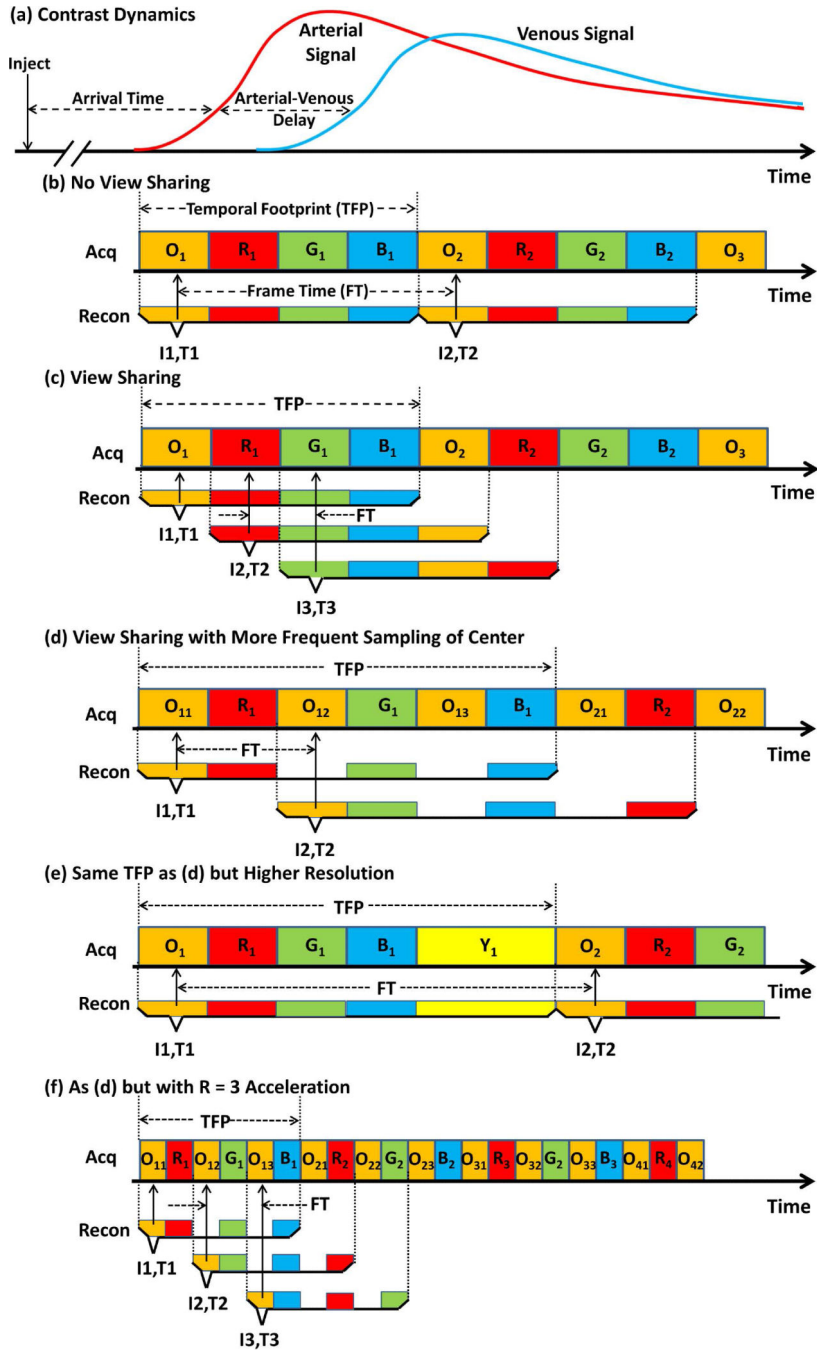


Figure 3. View sharing temporal readouts. (a) Typical timing in CE-MRA. After intravenous contrast injection there is some patient-specific Arrival Time delay after which contrast-enhanced blood arrives at the target artery. After some subsequent Arterial-Venous Delay the companion veins enhance. (b), (c), (d) and (f) show the temporal payout using any of the sampling methods of Fig. 2a-d. (b) Temporal sampling of k_y - k_z space with no view sharing. After all desired samples are first acquired (all four colored groups in the Acq line), those data are used for reconstruction, and the process is repeated. Frame time (FT) matches

temporal footprint (TFP). (c) Simple view sharing. Same acquisition as (b) but more frequent reconstruction with each reconstruction using the designated colored blocks within the corresponding bracket. FT is reduced 4× vs. (b) but TFP remains the same. (d) Acquisition with 3× increased frequency of sampling central (orange) k-space and view sharing. Reconstruction is done in synchrony with sampling central k-space. FT is half that of (b) but TFP is increased by 50%. (e) Alternative acquisition with same TFP as (d) but with improved spatial resolution and increased FT vs. (b) and (d). (f) Temporal sampling of all k-space sectors assuming acceleration factor $R = 3$. Note that image I3 and I4 (not shown) are temporally well aligned with peak arterial signal in (a). Also the temporal footprint has been reduced 3× vs. (d) so that almost all k-space sampling is performed prior to venous enhancement.

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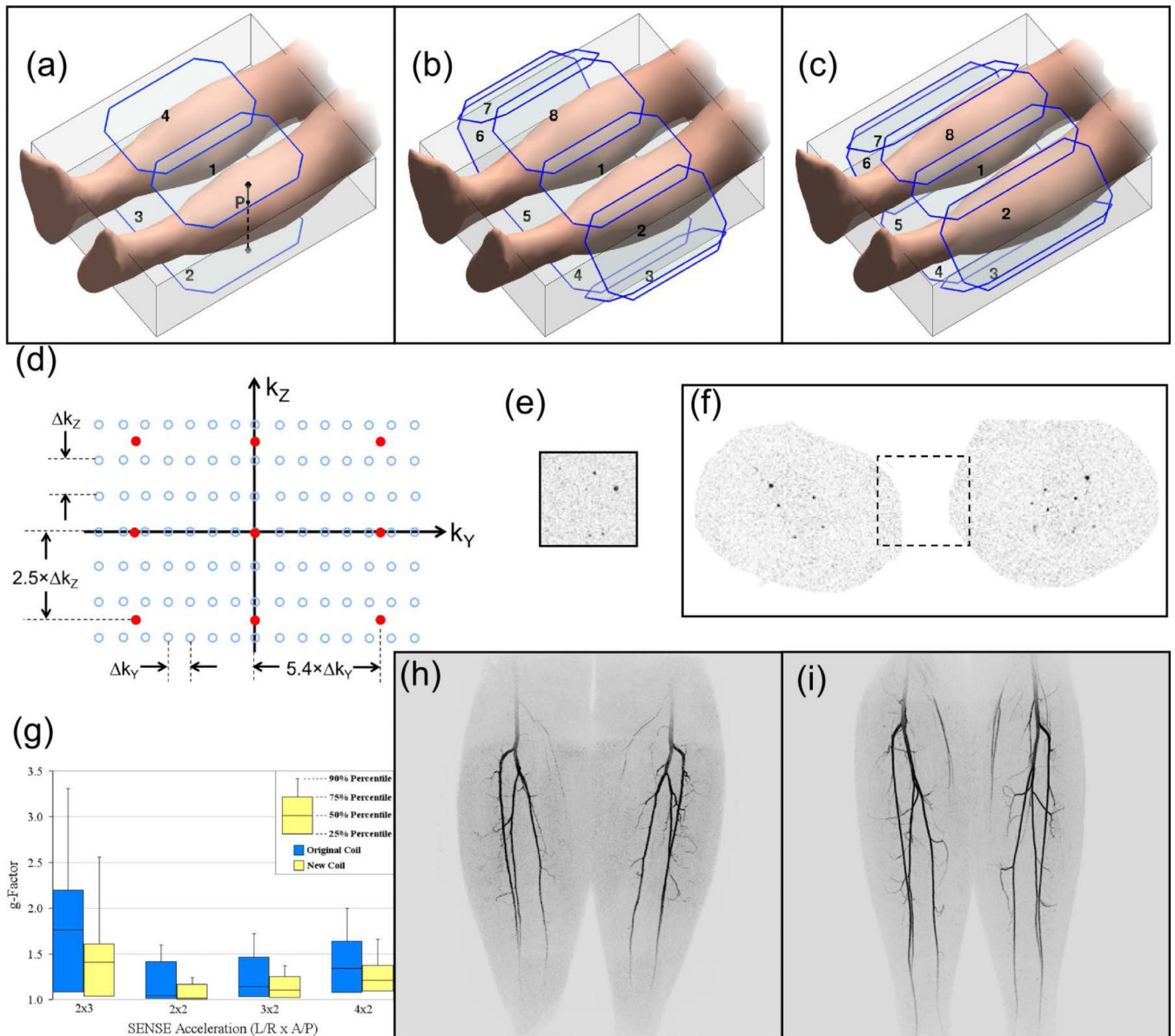


Figure 4.

2D Parallel Acquisition in CE-MRA. (a) Schematic of calves with 3D FOV and simple four-element coil. Point P is described in the text. (b) Similar to (a) but with eight-element coil with all elements of equal area. (c) Similar to (a) and (b) but with further adapted coil with more slender anterior and posterior elements (1,4,5,8) and longer elements overall than in (b). (d) Plot of k_Y - k_Z space with samplings for reference unaccelerated (blue) and accelerated scans (orange). The assumed acceleration is $R_Y \times R_Z = 5.4 \times 2.5 = 13.5$. (e) Axial aliased image from accelerated 3D CE-MRA exam of the calves using the acceleration of (d). (f) Unaliased image from same study as (e). The reduced FOV of the accelerated acquisition is shown in the dashed box. (g) Plot of g-factors of 3D volume of the calves using the receiver coil arrays of (b) (blue) and (c) (yellow) for various acceleration factors. Note the overall reduced g-factors of the latter. (h) MIP made from CE-MRA exam of

calves using coil array of (b) with $R = 7.3$. (i) MIP made from CE-MRA of calves of different volunteer using coil array of (c) with $R = 8$.

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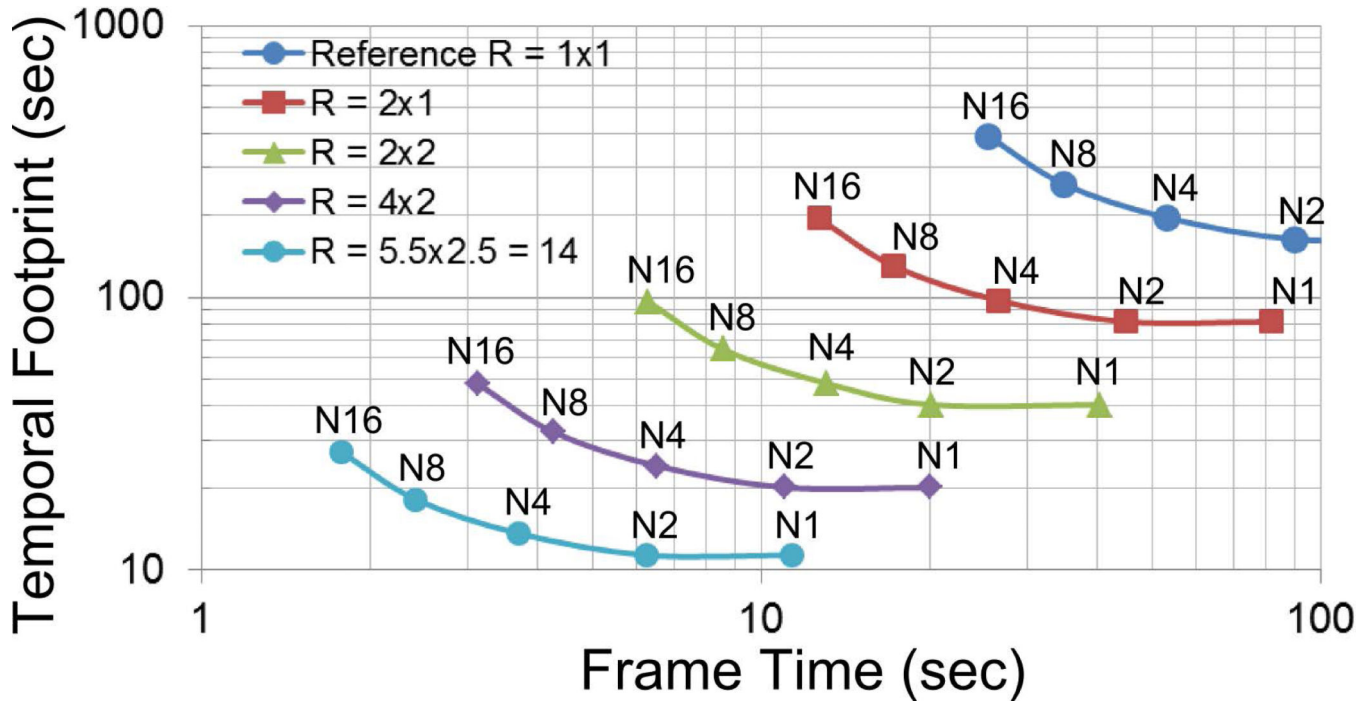


Figure 5. Plot of temporal footprint vs. frame time for different levels of view sharing (N1, N2, etc.) and acceleration ($R = 1 \times 1, 2 \times 1$, etc.) for 3D CE-MRA of the calves with $1 \text{ mm} \times 1 \text{ mm}$ spatial resolution in the $320 \times 132 \text{ mm}^2$ FOV of the axial (L/R \times A/P) plane. “N” is the number of sectors into which peripheral k-space is decomposed as in Figure 2. N1 corresponds to no view sharing in which case the frame time is equal to the temporal footprint. N2 provides reduced frame time but no increase in footprint. Increased levels of view sharing ($N > 2$) provide reduced frame time but larger temporal footprint along a given curve. Acceleration R is expressed as $R_Y \times R_Z$. Increased levels of acceleration (larger R) shift the entire curve to smaller frame times and smaller temporal footprints. For this plot the diameter of central k-space (the orange zone of Fig. 2a) was 30% of the full k-space diameter sampled.

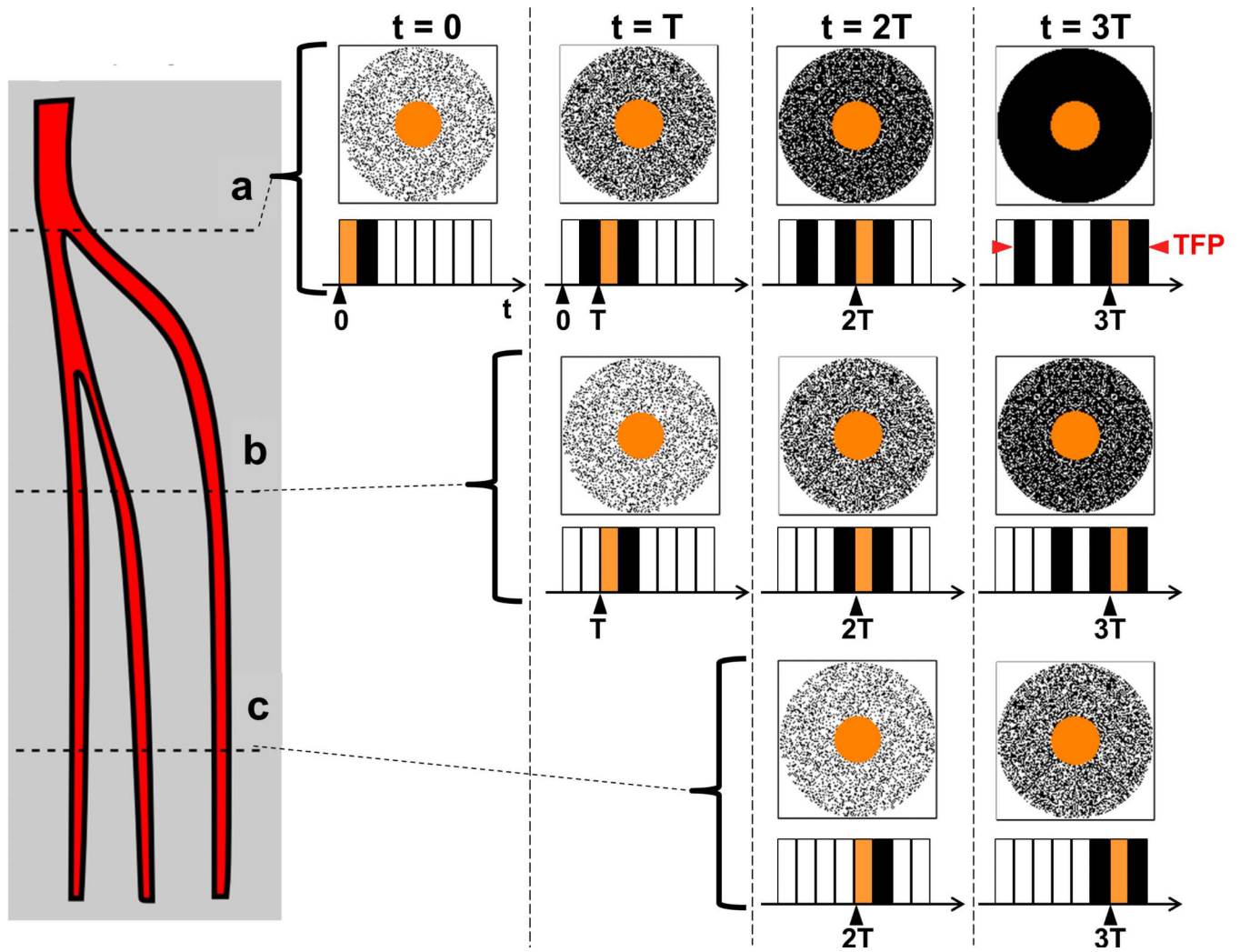


Figure 6. Schematic of the principal arteries of the calf and the degree of k-space filling of k_Y - k_Z space at various times for an assumed view-shared acquisition with sampling pattern similar to that in Fig 2c but with four sectors in peripheral k-space. Contrast bolus is assumed to enter from above and steadily progress to levels (a-c). Images are reconstructed at times $t = 0, T, 2T, 3T$. The temporal footprint (TFP) of the sequence is indicated. For additional details see the text.

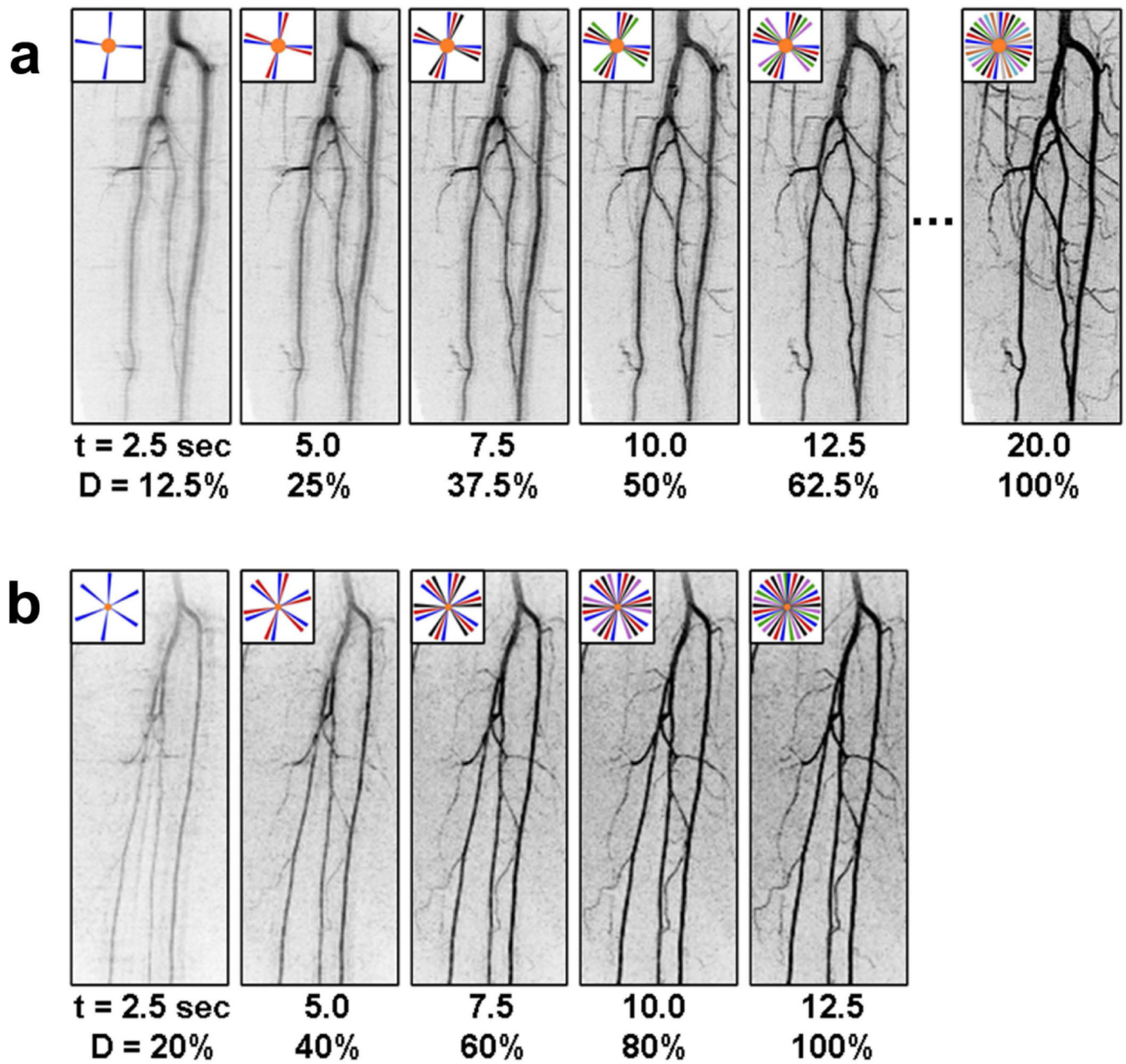


Figure 7. Comparison of the buildup of signal level and spatial resolution in a time-resolved sequence with relatively high level of view sharing (a), (N8, 20 sec temporal footprint) and moderate view sharing (b), (N5, 12.5 sec temporal footprint). In both (a) and (b) the spatial resolution is 1 mm isotropic and the frame time is 2.5 sec. (Taken from Ref. (78)).

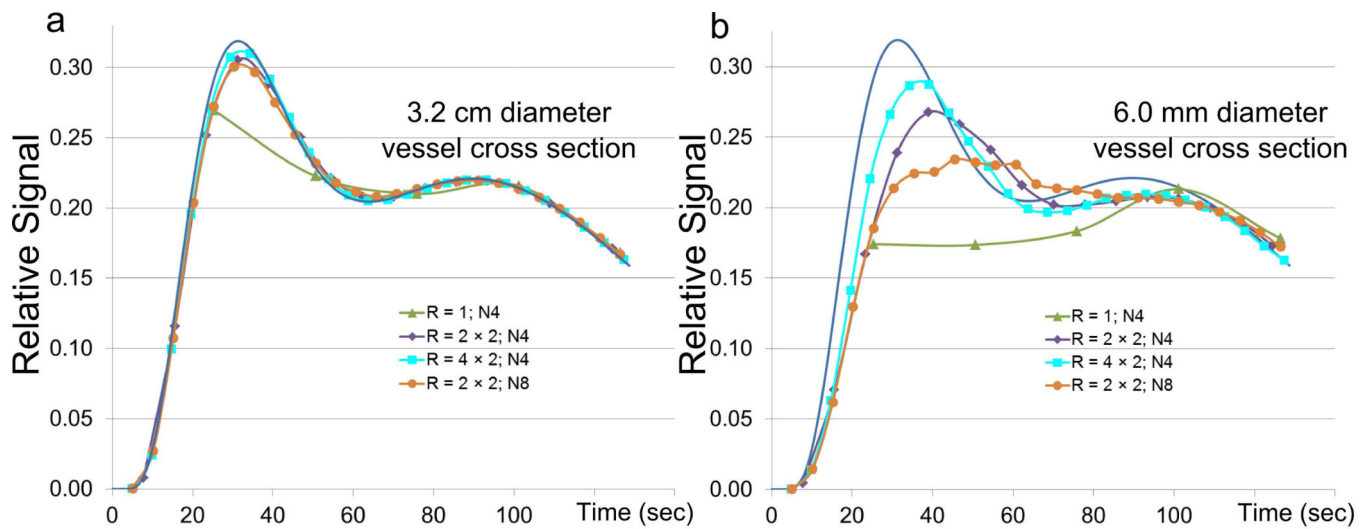


Figure 8.

Illustration of temporal fidelity vs. various acquisition strategies for two different assumed vessel diameters for assumed $Y \times Z$ FOV of $320 \times 132 \text{ mm}^2$. In both (a) and (b) the reference curve, representing actual arterial signal enhancement, is shown in blue. (a) Assumed diameter of vessel in the axial (Y-Z) plane is 32 mm. Other than the unaccelerated acquisition (green), all strategies provide a good match of the reconstructed signal with true assumed reference signal. (b) Assumed vessel diameter of 6 mm. As acceleration increases from $R = 1$ to 4 to 8 the match of the reconstructed signal with the reference improves and the frame time also decreases. For fixed acceleration $R = 4$, as additional view sharing is included, going from N4 (purple) to N8 (orange), although the frame time is reduced, the fidelity decreases owing to the increased temporal footprint

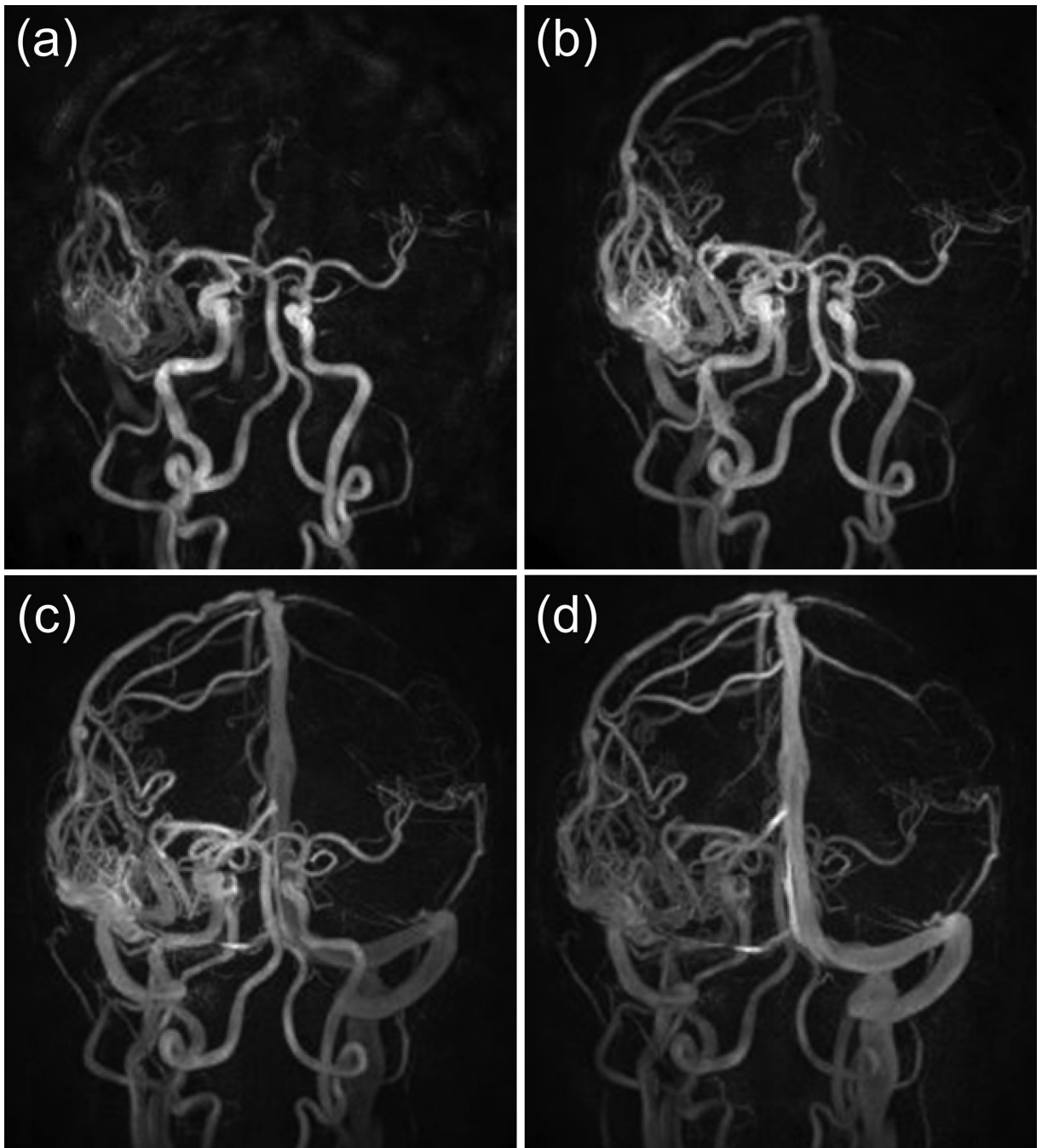


Figure 9.

Image of an arteriovenous malformation using the HYPR-Flow method (76) formed from a time-resolved contrast-enhanced run and a 3D phase-contrast-based composite image.

Spatial resolution 0.68 mm isotropic, 0.75 sec frame time. (Courtesy of Yijing Wu, Ph.D., Kevin Johnson, Ph.D., Patrick Turski, M.D., and Charles Mistretta, Ph.D.)

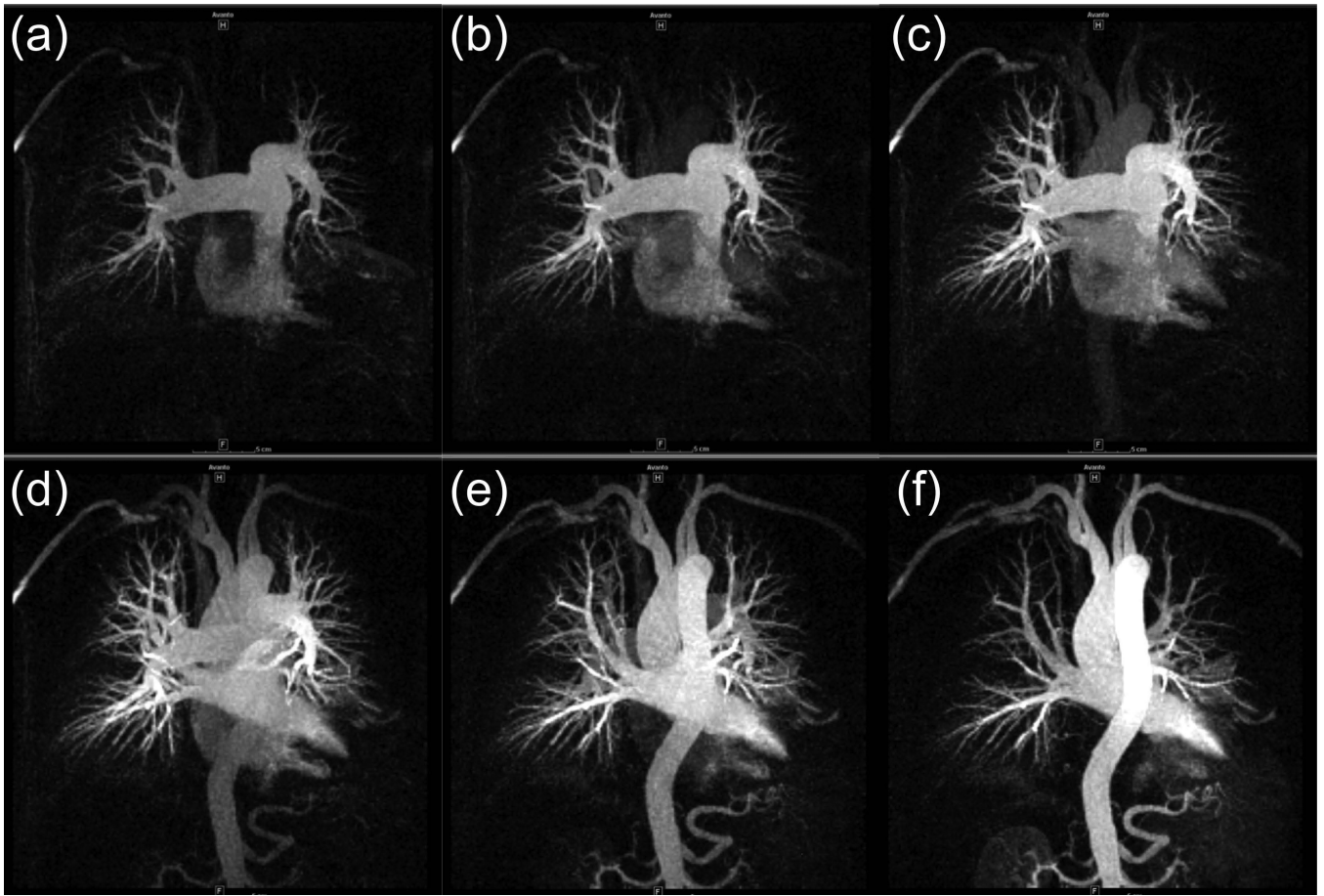


Figure 10. Time-resolved 3D pulmonary CE-MRA acquired using the method reported in Ref. (105). Spatial resolution $1.5 \times 1.5 \times 3.0 \text{ mm}^3$; 0.7 sec frame time. (Courtesy of Timothy J. Carroll, Ph.D.)

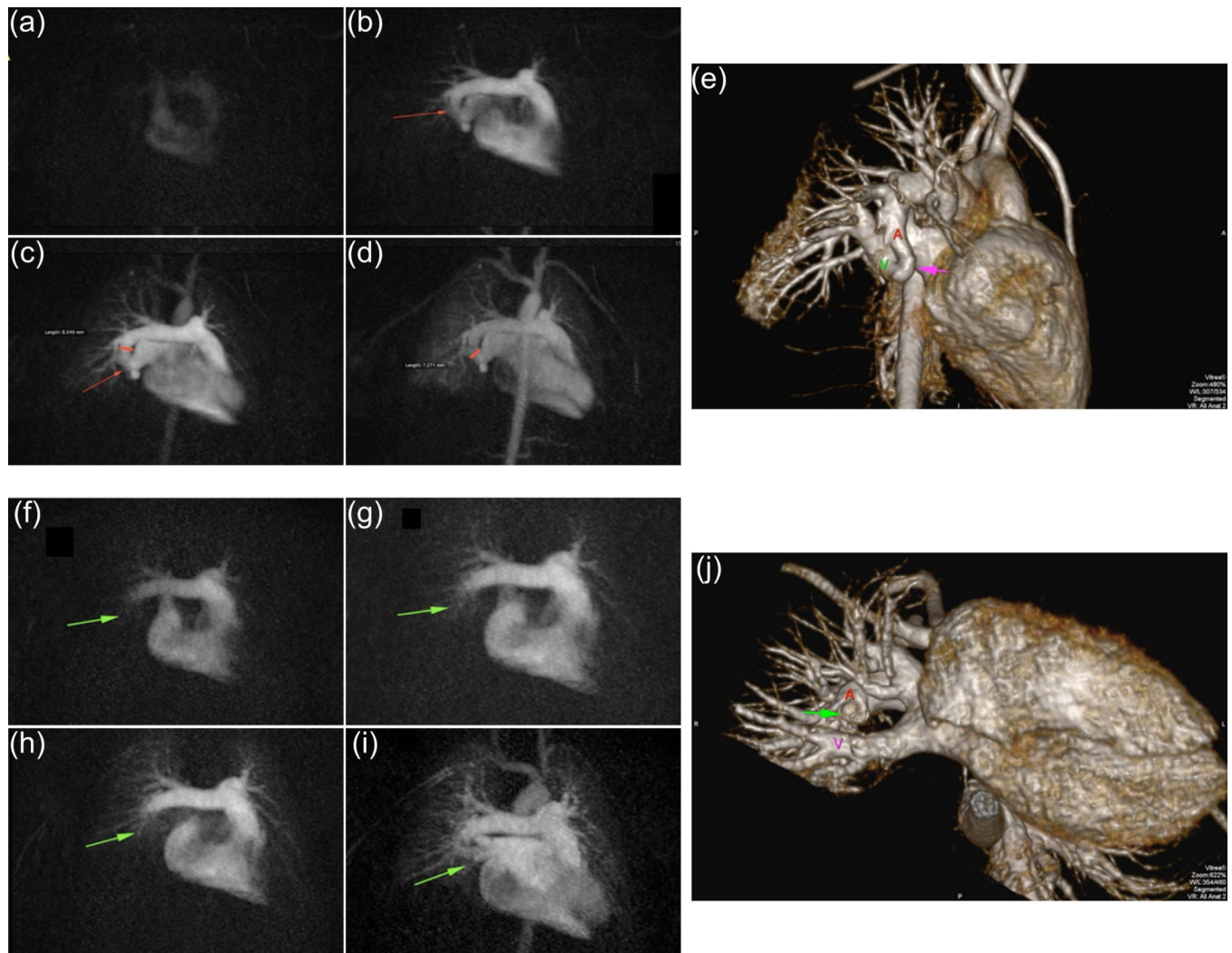


Figure 11.

Example of time-resolved CE-MRA of pulmonary arteriovenous fistula of the right lower lobe in 6 y/o male. (a-d) Sequence of time-resolved 3D CE-MRA images pre-treatment showing the fistula (arrow). Approximate 1 sec frame time; 1 ml contrast material. (e) High resolution single-phase image showing fistula (red arrow) and the abnormal vascular continuity of the pulmonary artery (A) and vein (V). (f-i) Sequence of time-resolved images seven months later, post-treatment, showing normal progression of contrast and absence of fistula. (j) Companion high resolution single phase image confirming occluded fistula (green arrow) and anatomic separation between artery (A) and vein (V). (Courtesy J. Paul Finn, M.D.)

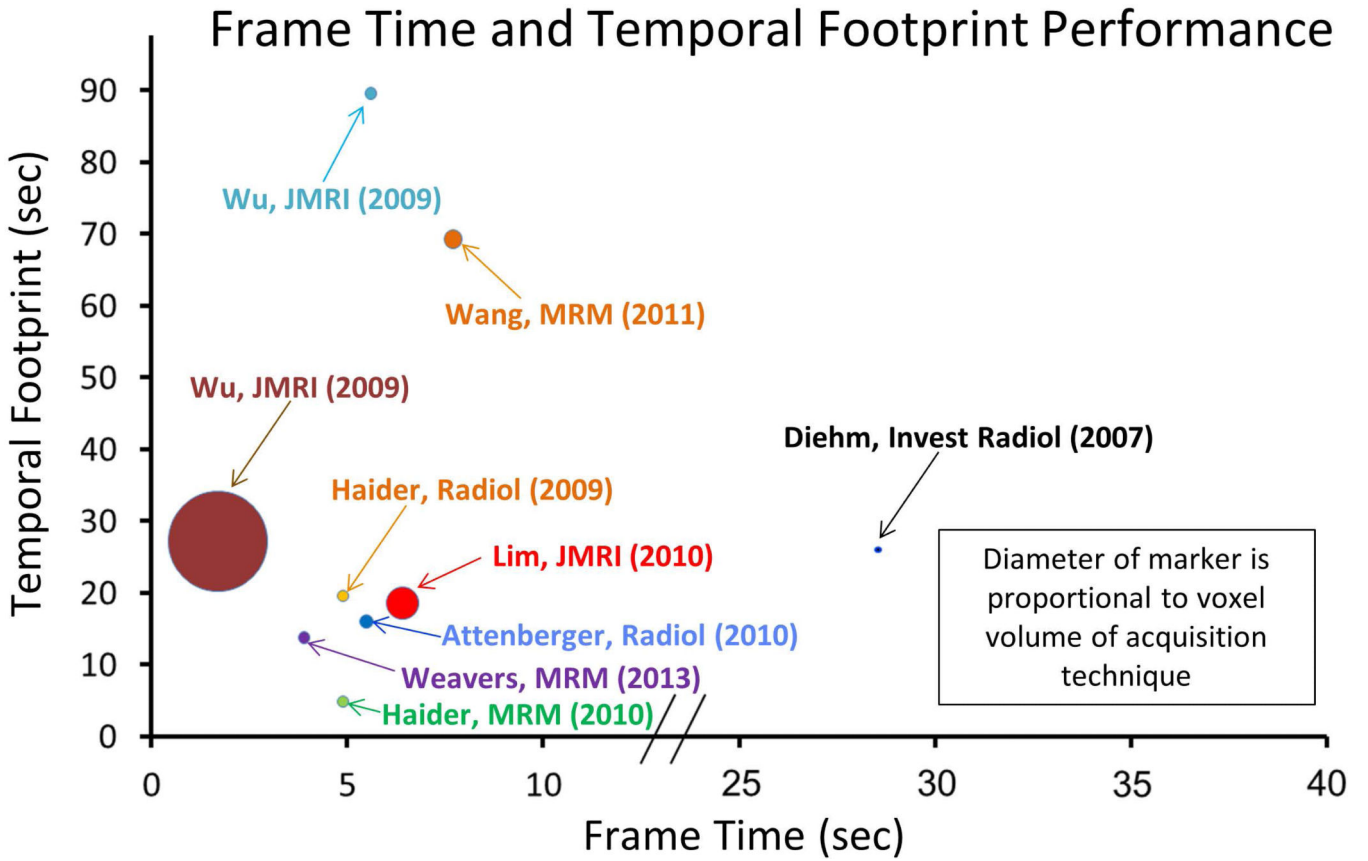


Figure 12. Plot of temporal footprint vs. frame time for time-resolved 3D CE-MRA techniques for imaging the calves listed in Table 2. Diameter of marker for each technique is proportional to the voxel volume of the acquisition. Smaller marker size reflects improved spatial resolution.

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Figure 13. Radiologic study of the calves. (a) Still frame from a time-resolved image series showing a stenosis in the mid-region of the right anterior tibial artery. (b) Companion digital subtraction angiography (DSA) result verifying the finding of (a). (Courtesy of Holger Haubenreisser, M.D. and Stefan Schoenberg, M.D.)

Table 1

The evolution of spatial resolution and acquisition time for contrast-enhanced MRA of the calves.

Spatial Resolution (Y × Z)		3.35 × 4.13 mm ²	2.50 × 2.06 mm ²	1.67 × 1.38 mm ²	1.00 × 1.00 mm ²
Figure		1a, e	1b, f	1c, g	1d, h
Sampling Resolution (Y × Z)		96 × 32	128 × 64	192 × 96	320 × 132
Total No. Views Without k-Space Corners		2304	6144	13824	31680
Times (sec)					
No Acceleration R = 1	TR 14.1	32	87		
	TR 10	23 8.0 / 32	61 21.5 / 86		
	TR 5	11.5	30 10.7 / 43	69 24 / 96	
Acceleration R = 2	TR 10	11.5	30	69	
	TR 5	5.8	15 5.3 / 21	34 12.1 / 48	79 27 / 110
Acceleration R = 4	TR 10	5.7	15	34	79
	TR 5	2.8	7.6 2.7 / 10.7	17 9.6 / 24	39 13.8 / 55
	TR 3	1.7	4.6	10.4 3.6 / 14	23 8.3 / 33
Acceleration R = 10	TR 5	1.1	3.0 1.0 / 4.3	6.9 2.4 / 9.7	15 5.5 / 22
	TR 3	0.7	1.8 0.7 / 2.6	4.1 1.4 / 5.8	9.5 3.3 / 13

References:		
Prince, Ref. (7)	Korosec, Ref. (8)	Weiger, Ref. (10)
Hu, Ref. (14)	Bonel, Ref. (18)	Haider, Ref. (21)

Columns left to right indicate improved axial spatial resolution. Top rows indicate sampling parameters. Subsequent rows from top to bottom indicate reduced times due to TR reduction and increased acceleration (R) from parallel acquisition. Body of table indicates acquisition time (in sec) for single phase imaging and frame time / temporal footprint (assumed to be 4× frame time) for time-resolved acquisition with view sharing. Assumed field of view is 40 (S/I) × 320 (L/R) × 132 (A/P) mm³. Images corresponding to the resolutions indicated are presented in Figure 1. TR times are indicated in msec.

Table 2

Summary of contemporary MRA of the calves. Performance is shown in Figure 12.

First Author	Year	Ref.	X (mm)	Y (mm)	Z (mm)	Volume (mm ³)	Frame Time (sec)	Temporal Footprint (sec)
Diehn	2007	(116)	0.45	0.55	1.00	0.25	28	28
Haider	2009	(21)	1.00	1.00	1.00	1.00	4.9	19.6
Wu	2009	(117)	0.80	0.80	1.60	1.02	5.6	89.6
Wu	2009	(117)	1.70	1.70	3.00	8.67	1.7	27.2
Attenberger	2010	(118)	1.10	1.10	1.10	1.33	5.49	16
Lim	2010	(17)	1.00	1.60	1.70	2.72	6.4	18.6
Haider	2010	(22)	1.00	1.00	1.00	1.00	4.9	4.9
Wang	2011	(75)	0.94	0.94	1.80	1.59	7.7	69.3
Weavers	2013	(54)	1.00	1.00	1.00	1.00	3.9	13.8

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