Chemical shift encoding using asymmetric readout waveforms

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Purpose: To describe a new method for encoding chemical shift using asymmetric readout waveforms that enables more SNR-efficient fat/water imaging.

Methods: Chemical shift was encoded using asymmetric readout waveforms, rather than conventional shifted trapezoid readouts. Two asymmetric waveforms are described: a triangle and a spline. The concept was applied to a fat/water separated RARE sequence to increase sampling efficiency. The benefits were investigated through comparisons to shifted trapezoid readouts. Using asymmetric readout waveforms, the scan time was either shortened or maintained to increase SNR. A matched in-phase waveform is also described that aims to improve the SNR transfer function of the fat and water estimates. The sequence was demonstrated for cervical spine, musculoskeletal (MSK), and optic nerve applications at 3T and compared with conventional shifted readouts.

Results: By removing sequence dead times, scan times were shortened by 30% with maintained SNR. The shorter echo spacing also reduced $T_2$ blurring. Maintaining the scan times and using asymmetric readout waveforms achieved an SNR improvement in agreement with the prolonged sampling duration.

Conclusions: Asymmetric readout waveforms offer an additional degree of freedom in pulse sequence designs where chemical shift encoding is desired. This can be used to significantly shorten scan times or to increase SNR with maintained scan time.

KEYWORDS
chemical shift, Dixon, MRI

1 | INTRODUCTION

Fat tissue has special signal characteristics in MRI. In most sequences, fat is hyperintense due to its relatively short $T_1$, and low diffusivity compared to other soft tissues outside the brain. Due to chemical shift, fat is spatially shifted along the readout dimension. This bright and displaced signal can conceal pathology and decreases the dynamic range of the image. Several methods have been proposed to attenuate the fat signal. Although the physical rationale varies there are significant inefficiencies associated with each method.
The RF fat saturation technique employs a frequency selective RF pulse to excite only fat, followed by gradient spoiling to remove the signal prior to the main pulse sequence playout. The fat saturation module is compatible with all pulse sequences, but has several drawbacks such as increased minimum TR, prolonged scan time, and decreased SNR efficiency. Besides these inefficiencies, the fat saturation technique fails in regions where the local off-resonance from field inhomogeneities exceeds the chemical shift, where it can instead suppress water signal.

The difference in $T_1$ relaxation times can, in combination with an inversion pulse and by synchronizing the excitation with the longitudinal zero-crossing of fat, be used to null the number of signal averages (NSA). Theoretically, it has been reconstructed fat and water images, measured as an effective (CSE) echoes are, however, contributing to the signal in the longs the scan time. These additional chemical shift–encoded signal. Typically, multiple echoes are acquired which pro-

readouts has also been applied to RARE sequences, where three echo trains are played out, each with a unique shift of the readout gradient. In order to fulfill CPMG conditions, the time between successive refocusing pulses must remain constant, and dead time is introduced to allow the desired CSE between every pair of refocusing pulses. For instance, with 320 readout samples acquired with a receiver bandwidth (rBW) of $\pm 25$ kHz/FOV, the readout window duration is 6.4 ms. To achieve a fat/water phase shift of $\pi$, 4.6/2.3 ms of dead time is required at 1.5/3 T, resulting in sampling occupancy of only 58/74%. Besides the inefficient sampling, the echo spacing is prolonged, resulting in stronger $T_2^*$ blurring.

An alternative to shifting echoes is to acquire multiple echoes between every pair of refocusing pulses by applying readouts with alternating polarity, similar to GRASE, but where the additional echoes encode chemical shift rather than phase encodes. Even if this does not introduce any dead time, the sequence is restricted to large echo spacings and an odd number of echoes in order to fulfill CPMG conditions, while only two echoes are required for fat/water separation. More importantly, in terms of sampling efficiency, all echoes acquired after the refocusing pulse have identical CSE to those prior, differing only in the angular direction of dephasing. This redundant sampling pattern negatively impacts the resulting NSA of the separated images.

The inefficiencies of these RARE fat/water imaging methods might seem unrelated, but are a consequence of the readout waveforms being symmetrical. A GRASE readout block is symmetric around the spin echo, while the shifted readout is symmetric around its center of mass. In order to remove the redundant sampling of the GRASE readout block, at least one of the gradient lobes must differ from the rest, thus breaking the block symmetry. The shifted design requires additional echo spacing that cannot be used to acquire data since a prolonged readout is symmetric unless a partial Fourier acquisition is used, that is, asymmetric sampling. The latter method has been described by Ma et al, but it is incompatible with undersampling along the phase encoding directions. The undersampling factor also varies between echoes, and requires special consideration to combine with fat/water separation in k-space. A limiting factor of asymmetric sampling is also the largest possible dephasing time (time between gradient and spin echo), which is limited to half of the readout window duration. The technique is therefore best suited for low rBW acquisitions.

Recently, we described a two-point single-TR RARE sequence where readout asymmetry is achieved by employing two different readout amplitudes (dual bandwidths). While an in-plane resolution of 0.5 mm was achieved at 3T, with a high-performance gradient system, resolutions beyond that are only feasible with a multi-TR acquisition. In this work, we propose using asymmetric readout waveforms to achieve efficient sampling with chemical shift encoding without dead time or redundancy. This technique allows for flexible echo shifts while still fulfilling CPMG conditions. The concept is applied to a RARE sequence to shorten scan time or to increase SNR. As a proof of concept, we also explore a modified symmetric in-phase waveform to improve chemical shift encoding.
2 | THEORY

2.1 | Designing the asymmetric waveform

The simplest asymmetrical waveform is the asymmetric triangle, described by the durations $T_{a,b}$ and slew rates $s_{a,b} = g/T_{a,b}$ of the two segments $a$ and $b$ shown in Figure 1A. The time-to-center, $t_c$, available acquisition time $T_{a,b} = T_{a} + T_{b}$, and area $M$ of the asymmetric triangle are given by the desired shift, echo spacing, and resolution, respectively. With the introduction of the auxiliary variable $T_1 = t_c - t_{as}$, the first segment duration $T_a$ is equal to $2T_1^2/T_{tot}$ in the case of a positive shift. The triangle amplitude is $g = 2M/T_{as}$. For a negative shift, the waveform is mirrored by swapping $a$ and $b$.

The asymmetric triangle achieves a maximum positive shift when $T_b = 0$. Ignoring the infinite slew rate violation, a shift of only $T_{tot} \left( \sqrt{\frac{1}{2} - \frac{1}{2}} \right) = 0.2T_{tot}$ is achieved. This geometrical limitation causes the asymmetric triangle waveform to only be viable in sequence with long echo spacing.

To enable larger shifts, we designed a cubic spline-based waveform, shown in Figure 1B. As defined in the figure, the waveform is split into a trapezoidal base with amplitude $g_0$ and an asymmetric component. Sampling is started at $t_{as}$ which is the instant at which the moment $M_p$ is equal to the ramp area of the corresponding trapezoidal waveform. This is also the starting point of the first spline, $p_1$. The presampling duration $t_{as}$ is known from the phase encoding or crusher gradient duration, whichever is longer. The base amplitude then follows as $g_0 = 2M_p/t_{as}$. With $T_2 = t_{as} - t_c$, the base moments are simply $M_{b1} = g_0T_1$ and $M_{b2} = g_0T_2$.

Three knots are located at $t_{as}$, $t_c$, and $t_{as}$, and the waveform is described by the two interval polynomials:

$$p_1(t) = \sum_{n=1}^{3} a_n t^n$$
$$p_2(t) = \sum_{n=0}^{3} b_n t^n$$

Further, $p_1$ and $p_2$ meet with $C^2$ continuity. In line with natural splines, $p''_2(T_2) = 0$. Given the desired center of mass at $t_c$ the following must hold:

$$\int_0^{T_1} p_1(t) dt = M_1 = \frac{M}{2} - M_{b1} - M_p$$
$$\int_0^{T_2} p_2(t) dt = M_2 = \frac{M}{2} - M_{b2} - M_p$$

Finally, $p_2(T_2) = 0$, forming the exactly determined linear problem which can be written in matrix form as:

$$\begin{bmatrix}
T_1 & T_1^2 & T_1^3 & -1 & 0 & 0 & 0 & a_1 & 0 \\
1 & 2T_1 & 3T_1^2 & 0 & -1 & 0 & 0 & a_2 & 0 \\
0 & 2 & 6T_1 & 0 & 0 & -2 & 0 & a_3 & 0 \\
0 & 0 & 0 & 0 & 2 & 6T_2 & b_0 & b_1 & M_1 \\
T_1^2/2 & T_1^4/3 & T_1^6/4 & 0 & 0 & 0 & 2 & b_2 & b_2 \\
0 & 0 & 0 & T_2 & T_2^2/2 & T_2^3/3 & T_2^4/4 & b_3 & M_2 \\
0 & 0 & 0 & 1 & T_2 & T_2^2 & T_2^3 & b_3 & 0
\end{bmatrix} = 0$$

The coefficients are available in the appendix. A consequence of enforcing the second derivative of $p_2$ to be zero at its endpoint is an undershoot in the waveform for positive shifts. This is avoided by calculating the waveform for a negative shift and then mirroring the result.

2.2 | Matching the in-phase echo

A property of the asymmetric readouts, resulting from the dynamic bandwidth, is the non-uniform k-space sampling density. We propose to counterbalance this with a matched in-phase readout to achieve a more balanced total k-space sampling density in a two-point RARE sequence. Ideally, the combined fat/water estimates will have an improved SNR transfer function with this approach (described below).

Although constrained to be symmetrical about the k-space origin, the sampling density of the matched readout can be varied. Instead of a conventional trapezoidal readout with fixed amplitude $\lambda$, we investigated a symmetric quadratic readout waveform which matches the CSE readout, illustrated
in Figure 2. $Q(t) = q_0 + q_2(t - t_{c,IP})^2$. The constant $q_0$ matches the asymmetric spline such that the combined effective dwell time at the k-space center equals that of two trapezoidal in-phase readouts. That is, the total time spent sampling the k-space center is the same: $\frac{1}{q_0} = \frac{2}{h_0} - \frac{1}{h_0}$. The quadratic coefficient $q_2$ is defined by the gradient moment $M$, the center point $t_{c,IP}$ coinciding with the spin echo, and the constant $q_0$: $q_2 = \frac{3M}{2t_{c,IP}^2} - \frac{2q_0}{t_{c,IP}^2}$.

A comparison between the sampling densities for trapezoidal and quadratic in-phase readouts is shown in Figure 3. It is clear that while the density is non-uniform, the area is consistent between the spline, quadratic, and trapezoidal waveforms because the total readout durations are equal. In contrast, the shifted trapezoidal case shows a uniform distribution with a smaller area. This is a consequence of less time spent sampling the signal. In Figure 4D, it is evident that the combined sampling time at the k-space center is increased and a more uniform density is achieved with the quadratic in-phase waveform. We hypothesize that the asymmetry of the sampling density can be counterbalanced by using real-valued k-space fat/water separation, described below.

3 | METHODS

3.1 | Pulse sequence

A RARE sequence was implemented that alternates the acquisition of in and opposed phase echoes between excitations, as shown in Figure 4. All waveform calculations were calculated on-the-fly, without delay, according to

\[
S_1 = e^{i\phi} \begin{bmatrix} \sqrt{\tau_1} & 0 & 1 & a_1 \\ 0 & e^{i\varphi_1} & \sqrt{\tau_2} & 1 & a_2 & W \end{bmatrix} x
\]

where $\phi$ describes the initial phase after excitation, calculated as described by Berglund et al.\(^{20}\) Equation (4) accounts for multiple fat peak resonances $\alpha$ with relative weights $\alpha^{21}$: $\alpha_n = \sum_{n=1}^{N} \alpha_n e^{i\phi_n}$. Non-chemical shift off-resonances $\varphi$ are determined with a multiscale graph cut technique.\(^{22}\)

Since $S$ is acquired with different gradient strengths, not only between samples but also between echoes, the chemical shift displacement can lead to unfamiliar artifacts. This frequency-dependent blurring can be corrected for by solving for $x$ in k-space.\(^{17}\)

By modeling $x$ as real-valued in image space, fat and water estimation enforces conjugate symmetry in k-space.\(^{20}\) Although Equation (4) is exactly determined (two complex measurements $S_1$, $S_2$, and four real-valued estimates, $\phi$, $\varphi$, $W$, and $F$), estimating and demodulating only low resolution $\phi$- and $\varphi$-maps leaves an over-determined inverse problem. The additional degrees of freedom can be used to take advantage of the known differences in signal uncertainties $A$ for an improved estimation:
Here, $F()$ is the Fourier transform operator, and $S_{\phi\psi} = Se^{-i(\hat{\phi}+\hat{\psi})}$ denotes an acquired echo after image space demodulation with the estimated $\hat{\phi}$ and $\hat{\psi}$. The * superscript indicates the complex conjugate. The encoding matrix $(\Lambda\Lambda)_k$ for sample $k$ is stacked with that of its conjugate sample $c$, that is, the point mirrored around the k-space center. A Tikhonov regularization is applied to the pseudo inverse to reduce colored noise, indicated by $\ddag$. The regularization term is chosen to equalize the noise power spectrum (NPS) of the combined fat and water estimates as previously described. This forms the noise-whitening weighted least squares estimator $E^\ddag$.

The NPS of water and fat can be extracted from the diagonal elements of $E^\ddag$, where $(E^\ddag)^H$ is the conjugate transpose of $E^\ddag$. The modulation transfer function (MTF) describes the frequency loss in the estimates, and can be calculated from the diagonal elements of $E^\ddag$. A theoretical SNR transfer function for $\hat{x}_k$ can be calculated from the ratio between the corresponding MTF and NPS.

3.3 | Data acquisition

Informed consent was obtained from all three volunteers (male, age 29, 33, and 48) in accordance with the institutional review board policy. All images were acquired on a 3T Signa Premier (GE Healthcare). Phase encoding line synthesis was implemented using GRAPPA. Relevant scan parameters are listed in Table 1.

High-resolution PD-weighted images of the foot and knee were acquired with dedicated Rx and Tx/Rx coils, respectively. The in-phase echo was acquired with the matched quadratic waveform.

Sagittal $T_2$-weighted images of the cervical spine were acquired with the conventional shifted approach. The sampling duration was kept constant but the echo spacing was shortened by employing the spline waveform for an opposed phase CSE. The echo train length was increased to achieve the same effective echo time, thereby reducing the scan time. We also acquired images using the matched quadratic in-phase waveform.

The recently described single-TR dBW-RARE technique was compared against the proposed multi-TR RARE sequence. As the multi-TR sequence has twice as long
acquisition time, which also benefits the SNR, the scan
time was matched using in-plane GRAPPA acceleration.
However, as GRAPPA requires reference lines, the scan
time was slightly longer with the proposed technique. The partial
Fourier fraction with dBW-RARE was 0.75.

The use of an asymmetric readout gradient can also be used
to allow more sampling for a fixed echo spacing. We therefore
compared the SNR impact on this increased sample duration
and compared it to the shifted approach. The SNR impact of
a matched quadratic in-phase waveform was also investigated.
$T_2$-weighted images of the optic nerve were also acquired. SNR maps were generated as described previously, with regions of interests placed to avoid bias from flow artifacts. The apparent SNR was measured by acquiring each phase encoding line twice, thus doubling the scan time. Note that these apparent SNR measurements are almost entirely determined by the SNR at the center of k-space, as the majority of the signal originates there and the method for measuring it involves smoothing. A measurement of apparent SNR is therefore highly dependent on the effective dwell time of the k-space center, which varies significantly between the different waveforms. Additionally, with the decision to whiten noise in order to facilitate apparent SNR measurements, the in- and opposed phase images are filtered differently. Compared to a uniform signal resampling, the matched quadratic in-phase signal is in general more low-pass filtered and the opposed phase signal is more high-pass filtered. This does not affect the estimates as this is incorporated in Equation (4). For the full picture, the SNR transfer function describes the expected SNR for all spectral components.

### 3.4 | Supplementary material

An online GitHub repository is provided with interactive plots of the proposed waveforms at henricryden.github.io/
asymreadout, where the user can see set scan parameters and the waveforms and sampling density are updated interactively. A complete interactive plot of the RARE pulse sequence is also available.

4 | RESULTS

Sagittal $T_2$-weighted fat/water images of the cervical spine are shown in Figure 5. In this case, shifted trapezoids prolong the echo spacing by 38%. This is avoided with asymmetric readouts. The proposed matched quadratic in-phase waveform is also shown. Fat and water is successfully separated in both cases, and similar image quality is obtained.

The PD-weighted sagittal knee images in Figure 6 reveal fine anatomical details in both the acquired echoes and the estimated fat and water images. Chemical shift displacement can be seen, which manifests more strongly in the quadratic in-phase echo, due to the lower bandwidth when acquiring the center of k-space. The tibial bone marrow fat is displaced,

**FIGURE 6** Sagittal knee images acquired with quadratic (A) and spline (B) readout waveforms. Signal pile-up from chemical shift displacement is indicated by white arrows, which are more pronounced in (A) compared to (B). This artifact is removed in (C) and (D) by performing the separation in k-space and modeling the k-space trajectory.
superimposing on the cartilage which makes it indiscernible at some regions in the native images. However, the cartilage is clearly seen in the water image, and easily distinguished from the adjacent synovial fluid which appears brighter. The displacement correction is achieved by accounting for the dephasing times of each sample in Equation (5). The opposed phase image acquired with the spline waveform has a sampling density that emphasizes high spatial frequencies. While a higher noise level can be seen in homogeneous regions such as in the muscles, the trabecular structures, and remnant of epiphyseal plates are more strongly depicted.

Enhanced bone structures are also seen in the opposed phase image in Figure 7. Successful suppression of the fat allows a clear depiction of the cartilage across the entire field of view.

The comparison between dBW-RARE and the asymmetric encoding is shown in Figure 8.

Results from the apparent SNR measurements are shown in Figure 9. Although affected by eye motion induced artifacts, fat and water were successfully separated in all acquired slices. Compared to shifted trapezoids, an improved apparent SNR of 20% was obtained with the prolonged trapezoid, and a smaller improvement of 2-3% with the spline. The matched quadratic in-phase waveform has an improved apparent SNR of 34%. The separated fat and water estimates have higher apparent SNR when using the quadratic in-phase waveform. The SNR transfer function derived from the encoding matrices is also shown. At the k-space center (vertical dashed line), the theoretical SNR is higher than with a shifted acquisition, in-line with the apparent SNR measurements. The overall improvement in SNR across k-space is an effect from the increase in sampling duration.

5 | DISCUSSION

Here, we describe a new way of encoding chemical shift with variable density along the readout dimension. More specifically, the encoding is asymmetric to a degree required to achieve the desired CSE. Variable density sampling is commonly used in non-Cartesian sequences, for example, in spiral imaging to reduce aliasing artifacts or for the reduction of artifacts.
Even with a high-performance gradient system, readout resolutions higher than about 1 mm are not achievable with conventional fTED at 3T. By only encoding two echoes with different bandwidths and partial Fourier sampling, we recently reported readout resolutions of 0.5 mm at 3T.

Multi-TR acquisitions, on the other hand, do not have a resolution restriction as only one echo is acquired during $T_{\text{tot}}$, allowing much larger k-space coverage. The downside of multi-TR is the increased scan time, and dead times arising from the shifts, resulting in a more than twofold increase in scan time compared to single-TR acquisitions. The additional excitations contribute to the signal and can be compensated by other acceleration techniques, including reduced phase field of view, decreased NSA, partial Fourier, or parallel imaging. Dead times, however, increase the scan time without SNR benefits. By using asymmetric readout waveforms, these dead times are avoided, thereby reducing scan time and echo spacing. A 38% scan time penalty is reported with the shifted readouts in Figure 5, and would be twice as much at 1.5 T-given the same scan parameters—since the chemical shift dephasing period and hence the dead times for the shifted acquisitions are doubled. It should be noted that non-Cartesian sequences such as time offset spirals do not suffer from these large dead times as the readout is much longer.

The comparison with dBW-RARE in Figure 9 shows a sharper fat image with the proposed technique and less artifacts. These artifacts in dBW-RARE are likely attributed to the partial Fourier sampling. This comparison is, however, somewhat unfair as the dBW-RARE sequence struggles with such short acquisition windows and with the resolution pushed to its limit.

Chemical shift displacement artifacts are seen in all acquired echoes in Figure 9, but less pronounced in the spline images than in the quadratic in-phase images. An apparent SNR improvement is seen with the quadratic waveform. These effects are expected since, by design, the quadratic waveform has a lower bandwidth at the acquisition of k-space center. The reported apparent SNR increase of 14% is in line both with the expected SNR gain of 17% from the prolonged sampling duration and that calculated from the SNR transfer function at the k-space center. This SNR improvement would require 30% longer scan time with shifted readouts. It should be emphasized that the gain in SNR of the proposed technique is sourced from the additional sampling duration, which results in an SNR gain for almost all spectral components as evident from the SNR transfer function. The measured apparent SNR is almost entirely governed by the effective dwell time at the k-space center. A comparison of apparent SNR between acquired echoes is therefore difficult to make due to different resampling filters. The full picture is given by the SNR transfer function, and the apparent SNR measurements should be considered as an in-vivo validation at the center frequency.
FIGURE 9  Apparent SNR comparison of the spline, prolonged trapezoid, and quadratic echoes together with their water estimates. The first column shows images from the three in-phase waveforms. The second column shows opposed phase images. Reconstructed water images are seen in the third column. All images have a colored region showing the measured apparent SNR, with its average value reported in the figure. The bottom rows show the theoretical SNR transfer function of each k-space index, derived from the encoding matrices.
The spline waveform was designed to achieve the desired shift for a given available acquisition time $T_{\text{tot}}$. This typically results in a decreased effective dwell time at the k-space center and more time spent sampling higher spatial frequencies. Even though the apparent SNR does not increase even though the total sample duration does, the high frequency content should be more well resolved. This is supported by the clearer depiction of finer structures in Figures 6 and 7 compared to the in-phase images.

The set of dephasing times for k-space center acquisition is determined by the desired shift, while the first and last sample pair will always have identical CSE. Consequently, $A$ is singular at the edges of k-space, and while Hermitian symmetry will improve conditionality in general, the conjugate operator in Equation (5) results in $E$ being rank deficient. This en passant discovery is intuitive in hindsight, because there is no CSE resolved in the four samples.

Another interesting encoding property arises from the varying sample density (Figure 4), which in combination with varying dephasing times, results in a non-uniform SNR transfer function (Figure 9). The quadratic waveform increases the SNR at the k-space center, which is reflected in the apparent SNR measured in vivo. At the outer samples, the encoding is inferior but this is partly alleviated, for some indices, by the conjugate sample. The real-valued estimates also make the SNR transfer function symmetric. Moreover, water is less affected than the combined estimates in terms of the SNR transfer function and both proposed sequences outperform the shifted alternative in this work.

In Dixon imaging, the fat image can serve as a $T_1$-weighted surrogate that discriminates between tissue types. The improved sharpness in the opposed phase image reported here enhances the trabecular bone structures, and complements the tissue-classifying fat/water images. The opposed phase image might therefore be of value in evaluating fractures and trabecular destruction, similar to previous works. With the correction of chemical shift displacement, we expect simpler evaluation of cartilage thickness.

Chemical shift encoding can be achieved by any asymmetric readout gradient. Other readout waveforms such as a piece-wise linear shape, or an asymmetric triangle stacked on a rectangular base, would be of interest to explore. The waveforms described in this work were designed by only considering the conditioning of the k-space center, and it is expected that other more precisely tailored waveforms can achieve a further improved SNR transfer function. The quadratic waveform shows that the SNR transfer function can be modified by also considering the in-phase waveform, although it must be symmetric. More work can also be done on the regularization to yield optimized fat and water images, as opposed to the current technique where the noise amplification factor considers the sum of $W$ and $F$.

6 | CONCLUSION

Asymmetric readout waveforms can be used as an alternative to shifted trapezoids to encode chemical shift, enabling more efficient Dixon imaging. This allows for significant scan time reduction, increased SNR, or a combination thereof in RARE sequences. By removing sequence dead times, the echo spacing is also shortened, reducing $T_2^*$ blurring. The variable sampling density can be accounted for in the fat/water separation, and further compensated by also modifying the in-phase waveform to achieve a more uniform SNR transfer function.

CONFLICT OF INTEREST

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**APPENDIX A**

**SPLINE COEFFICIENTS**

\[
a_1 = \frac{4 (3M_1T_3^1 - M_2T_3^1 + 6M_1T_2^1T_3^2 - 6M_2T_2^1T_3^2 + 10M_1T_1T_3^3 - 6M_2T_1T_3^3)}{T_1^2T_3^2 (2T_1 + 3T_2) (T_1 + T_2)}
\]  

\[a_2 = \frac{12 (M_2T_1^3 + 4M_2T_1^3T_2^2 - 4M_1T_1T_2^2 - M_1T_2^3)}{2T_1^4T_2^2 + 3T_1^2T_2^3}
\]  

\[a_3 = \frac{4 (M_1T_2^4 - 2M_1T_2^4 + 6M_1T_2^4T_3^2 - 4M_2T_2^4T_3^2 + 5M_1T_1T_2^3 - 8M_2T_1T_2^3)}{T_1^4T_2^2 (T_1 + T_2) (3T_1^2 + 2T_1T_2)}
\]

\[\begin{bmatrix}
    b_0 \\
    b_1 \\
    b_2 \\
    b_3
\end{bmatrix} =
\begin{bmatrix}
    a_1 T_1^2 + a_2 T_2^2 + a_1 T_1 \\
    3a_1 T_1^2 + 2a_1 T_1 + a_1 \\
    3a_1 T_1 + a_2 \\
    - (3a_1 T_1 + a_2) / 3T_2
\end{bmatrix}
\]