# Simulating Spin-Echo MRI Signal Contrast Response to $T_R$ - $T_E$ Variations in Tissues with Spatially Varying $T_1$ - $T_2$ Maps

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#### Abstract

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Keywords: keyword 1, keyword 2

### 1 Statement of the Problem

Magnetic Resonance Imaging (MRI) is a non-invasive diagnostic technique widely used in clinical and research settings for its ability to produce detailed anatomical and functional images without ionizing radiation. The contrast in MRI images is fundamentally governed by intrinsic tissue parameters, primarily the longitudinal relaxation time  $(T_1)$  and the transverse relaxation time  $(T_2)$  [1]. These parameters reflect how different tissues return to equilibrium after perturbation by radiofrequency (RF) pulses and are critical to understanding signal formation. The time evolution of the net magnetization vector in response to RF excitation and relaxation processes is described by the Bloch equations [2].

In particular, spin-echo sequences—common in both clinical and research MRI—yield a steady-state solution to the Bloch equations that relates signal intensity to  $T_1$ ,  $T_2$ , repetition time  $(T_R)$ , and echo time  $(T_E)$  [3]. This steady-state solution forms the basis for simulating and interpreting MRI contrast. Most introductory and educational simulations rely on simplifying assumptions, including the notion that tissue compartments are homogeneous in their relaxation properties. However, this abstraction oversimplifies the complexity of biological tissues, especially under pathological conditions.

Although  $T_1$  and  $T_2$  describe intrinsic tissue characteristics, the imaging parameters  $T_R$  and  $T_E$  play an equally crucial role in shaping the observed signal contrast. The choice of  $T_R$  governs the extent of longitudinal recovery between excitations, while  $T_E$  determines how much transverse signal decay is captured at readout. Together, these timing parameters modulate sensitivity to different tissue types and heterogeneities. As such, understanding the interaction between  $T_1$ ,  $T_2$ ,  $T_R$ , and  $T_E$  is essential not only for optimizing contrast but also for interpreting variations in signal intensity in regions where relaxation times vary spatially [3].

In reality, tissues often exhibit intra-regional heterogeneity in relaxation parameters due to variations in cellularity, water content, perfusion, necrosis, or other microstructural and physiological processes [4]. For instance, tumors may contain a mixture of viable cells, necrotic cores, and edematous regions, each with subtly different  $T_1$  and  $T_2$  values. While these variations may not produce overt visual differences in standard MR images, they influence the local signal response and can affect both qualitative and quantitative interpretations of the image.

This project focuses on understanding how spatial variations in  $T_1$  and  $T_2$  values within a single tissue region impact the resulting MRI signal, using numerical simulations grounded in the steady-state Bloch model. By going beyond homogeneous-tissue models, the study seeks to explore the degree to which intra-regional relaxation heterogeneity affects signal intensity and image contrast. Such insights are relevant for both medical imaging applications, where accurate tissue characterization is essential, and fundamental physics, where a detailed understanding of signal formation informs the design and interpretation of MR experiments.

# 2 General Purpose of the Study

The purpose of this study is to simulate and analyze how spatial heterogeneity in  $T_1$  and  $T_2$  relaxation times within a tissue region influences MRI signal contrast, using the steady-state solution to the Bloch equations under spin-echo imaging conditions.

## 3 Specific Objectives

To accomplish this, the study specifically aims to:

- 1. Derive and implement the steady-state signal expression for a spin-echo MRI sequence based on the Bloch equations;
- 2. Encode spatial variation in  $T_1$  and  $T_2$  relaxation parameters across a 2D tissue model with a lesion region; and
- 3. Generate and analyze synthetic contrast maps to assess the effect of intra-lesion relaxation gradients on MRI signal intensity.

### 4 Review of Related Literature

Magnetic Resonance Imaging (MRI) leverages the relaxation properties of nuclear spins—longitudinal  $(T_1)$  and transverse  $(T_2)$ —to generate high-contrast images of soft tissues without ionizing radiation. The time evolution of magnetization under these relaxation processes is governed by the Bloch equations [2], forming the theoretical foundation of MRI. In the context of spin-echo sequences, the steady-state signal intensity can be described as

$$S(x,y) = \rho \left[ 1 - e^{-TR/T_1(x,y)} \right] e^{-TE/T_2(x,y)}, \tag{1}$$

where  $\rho$  is the proton density, and TR and TE are the repetition and echo times, respectively [3]. This formulation provides a direct quantitative relationship between tissue-specific relaxation parameters and image contrast [1].

Relaxation times vary significantly between tissue types and are influenced by imaging parameters and magnetic field strength. For example, Stanisz et al. [5] measured  $T_1$  values in gray matter at 1.5 T, which is a widely used field strength for modern MRI machines [1], to be within the range  $1124 \pm 50$  ms, and  $T_2$  values between  $95 \pm 8$  ms. Bojorquez et al. [6] further demonstrated that acquisition protocols and scanner-specific settings introduce substantial variability, highlighting the importance of reproducibility in clinical interpretation. Moreover, pathologies such as tumors often exhibit altered relaxation profiles due to changes in water content, cellular architecture, or necrosis, as noted in Tofts' review of quantitative MRI in disease states [4].

Emerging work has shown that spatial heterogeneity in relaxation parameters can meaningfully affect image contrast. Does et al. [7] observed multicomponent  $T_1$  relaxation in rat brain tissue, attributing the behavior to underlying microstructural variations. Xu et al. [8] similarly showed that  $T_2$  variability affects the accuracy of diffusion-weighted imaging, emphasizing the clinical impact of intra-tissue relaxation differences. Despite this, many MRI models assume homogeneity across regions of interest, which may limit their realism in depicting diseased tissues.

Efforts to standardize quantitative MRI across platforms have underscored persistent challenges. A recent multi-center study using the NIST/ISMRM phantom reported inter-scanner variability in  $T_1$  and  $T_2$  under 7%, but confirmed that minor discrepancies can still affect quantitative outcomes [9]. Furthermore, myocardial studies by Wiesmüller et al. [10] revealed notable intra-segmental  $T_2$  variation even under consistent imaging conditions, thus underscoring the importance of incorporating spatial gradients in modeling signal formation.

Computational simulations using the Bloch equations offer a tractable framework for exploring these phenomena under controlled conditions. Recent tools such as BlochSim.jl [11] and other open-source packages for MRI signal modeling provide accessible environments for simulating magnetization dynamics, but typically assume spatial uniformity in tissue parameters. As such, simulations incorporating spatially varying  $T_1$  and  $T_2$  maps remain underutilized, despite their potential to visualize how subtle tissue gradients manifest as signal heterogeneity.

While spatial heterogeneity in  $T_1$  and  $T_2$  presents a crucial dimension in tissue modeling, the interplay between these relaxation parameters and acquisition settings—particularly the repetition time (TR) and echo time (TE)—also fundamentally shapes signal behavior. Optimal selection of TR and TE is essential for maximizing lesion conspicuity and diagnostic performance. Hendrick et al. [12] demonstrated,

through Bloch-equation modeling, that CNR in spin-echo MRI is sensitive to interpulse delays, identifying parameter regimes that maximize signal discrimination between tissues.

More recently, Ragot and Chen [13] challenged the assumption that  $TE = T_2$  yields maximal contrast, showing via physiological-noise modeling that shorter echo times (around 50 ms at 3T) often outperform nominal  $T_2$ -matching in terms of CNR. Furthermore, the concept of lesion-to-background contrast ratio (LBCR) has gained traction in clinical studies as a robust proxy for lesion detectability, though optimization of TR and TE specifically for LBCR in spin-echo imaging remains underexplored. These findings underscore the need to account for both spatially varying tissue properties and acquisition parameter sensitivity when modeling MRI signal formation.

In summary, while theoretical models and standardization phantoms have greatly improved our understanding of MRI contrast, there remains a gap in simulating how smooth spatial variations in relaxation parameters within lesions influence signal intensity under spin-echo conditions. This study addresses that gap by implementing voxel-wise relaxation maps within a 2D phantom, providing insight into the interplay between tissue heterogeneity and MRI signal behavior as a step toward bridging simplified simulations and the complex biological realities of clinical imaging.

## 5 Methodology

#### 5.1 Overview

This project simulates the MRI signal produced by a tissue slice with spatially varying relaxation times under a spin-echo pulse sequence. A 2D matrix (phantom) will be used to represent a 25-pixel cross-sectional slice, where the background tissue is homogeneous gray matter and a circular lesion is embedded at the center. While the background will have fixed longitudinal and transverse relaxation values  $(T_1, T_2)$ , the lesion region will exhibit radial gradients in both parameters, simulating intra-lesion heterogeneity.

The net magnetization vector  $\mathbf{M}(t)$  evolves according to the Bloch equations:

$$\frac{d\mathbf{M}}{dt} = \gamma \mathbf{M} \times \mathbf{B} - \begin{bmatrix} M_x/T_2 \\ M_y/T_2 \\ (M_z - M_0)/T_1 \end{bmatrix}$$
 (2)

where  $\gamma$  is the gyromagnetic ratio, **B** is the effective magnetic field, and  $M_0$  is the equilibrium magnetization. For this simulation, we consider the steady-state signal under a spin-echo imaging sequence.

# 5.2 Signal Equation (Spin-Echo)

In a conventional spin-echo (SE) imaging sequence with repeated radiofrequency (RF) excitations, the observable signal intensity at each voxel (x, y) can be modeled. After an initial transient period, the longitudinal magnetization reaches a dynamic equilibrium. If the repetition time (TR) is sufficiently long for the transverse magnetization to decay to zero before the next RF pulse, the system is said to be in a steady-state incoherent (SSI) condition.

Under these circumstances, the steady-state transverse magnetization, which is directly proportional to the MRI signal, is given by:

$$S(x,y) = \rho(x,y) \left[ 1 - e^{-TR/T_1(x,y)} \right] e^{-TE/T_2(x,y)}$$
(3)

where:

- S(x,y) is the MRI signal intensity,
- $\rho(x,y)$  is the proton density (set to 1.0 throughout),
- TR = 2000 ms, repetition time,
- TE = 100 ms, echo time,
- $T_1(x,y)$  and  $T_2(x,y)$  are the voxelwise relaxation times.

#### 5.2.1 Derivation of the Steady-State Spin-Echo Signal

To simplify the derivation of the steady-state spin-echo signal, we make the following common assumptions:

- 1. The derivation is performed in a rotating frame of reference that precesses at the Larmor frequency, which simplifies the precession term in the Bloch equations and makes the effects of RF pulses easier to visualize.
- 2. RF pulses (90° and 180°) are assumed to be instantaneous, meaning they rotate the magnetization without any relaxation occurring during the pulse itself.
- 3. The 90° pulse perfectly rotates  $M_z$  to  $M_{xy}$ , and the 180° pulse perfectly flips  $M_{xy}$  and  $M_z$ .
- 4. The steady-state incoherent (SSI) condition is considered, where any residual transverse magnetization from the previous repetition is assumed to have fully decayed before the next excitation pulse. This is generally valid for long TR values.
- 5. The effects of macroscopic motion (flow) and microscopic random motion (diffusion) on the signal are ignored.
- 6. A uniform proton density  $(\rho)$  is assumed throughout the tissue, which is multiplied at the end to represent the available magnetization.

Let the longitudinal magnetization just before the 90° excitation pulse be  $M_{z,TR}^-$ . The transverse magnetization is assumed to be zero in the steady-state incoherent case.

The 90° pulse rotates  $M_{z,TR}^-$  into the transverse plane. So, immediately after the pulse:

$$M_{xy}(0) = M_{z,TR}^-$$
$$M_z(0) = 0$$

During this period, the transverse magnetization decays due to  $T_2$  relaxation, and the longitudinal magnetization recovers due to  $T_1$  relaxation.

$$\begin{split} M_{xy}(t) &= M_{xy}(0) \cdot e^{-t/T_2} \\ &= M_{z,TR}^- \cdot e^{-t/T_2} \\ M_z(t) &= M_0(1-e^{-t/T_1}) \quad \text{(since } M_z(0) = 0 \text{ and it recovers towards } M_0\text{)}. \end{split}$$

At t = TE/2:

$$M_{xy}(TE/2) = M_{z,TR}^- \cdot e^{-(TE/2)/T_2}$$
  
 $M_z(TE/2) = M_0(1 - e^{-(TE/2)/T_1})$ 

The 180° pulse rephases the transverse magnetization and inverts the longitudinal magnetization. The phase of  $M_{xy}$  is effectively flipped, but its magnitude remains unchanged by the ideal 180° pulse for the purpose of the echo formation. The dephasing that occurred is reversed.

$$M_z(TE/2)^+ = -M_z(TE/2)^- = -M_0(1 - e^{-(TE/2)/T_1})$$

The transverse magnetization now rephases and continues to decay due to  $T_2$  relaxation. The longitudinal magnetization continues to recover towards  $M_0$ . At the echo time (t = TE), the signal is maximum due to rephasing. The transverse magnetization at the echo  $(S_{echo})$  is the remaining magnitude of the transverse magnetization from before the 180° pulse, now further decayed for another TE/2 duration.

$$\begin{split} S_{echo} &= M_{xy}(TE/2) \cdot e^{-(TE/2)/T_2} \\ &= (M_{z,TR}^- \cdot e^{-(TE/2)/T_2}) \cdot e^{-(TE/2)/T_2} \\ S_{echo} &= M_{z,TR}^- \cdot e^{-TE/T_2} \end{split}$$

The longitudinal magnetization  $M_z$  at t = TE will recover from its inverted state:

$$\begin{split} M_z(TE) &= M_0 + (M_z(TE/2)^+ - M_0)e^{-(TE/2)/T_1} \\ M_z(TE) &= M_0 + (-M_0(1 - e^{-(TE/2)/T_1}) - M_0)e^{-(TE/2)/T_1} \\ M_z(TE) &= M_0 + (-M_0 + M_0e^{-(TE/2)/T_1} - M_0)e^{-(TE/2)/T_1} \\ M_z(TE) &= M_0 + (-2M_0 + M_0e^{-(TE/2)/T_1})e^{-(TE/2)/T_1} \\ M_z(TE) &= M_0 - 2M_0e^{-(TE/2)/T_1} + M_0e^{-TE/T_1} \end{split}$$

After the echo, the transverse magnetization is assumed to have decayed to zero for SSI. The longitudinal magnetization continues to recover during the remaining time TR - TE.

$$M_{z,TR}^- = M_0 + (M_z(TE) - M_0)e^{-(TR - TE)/T_1}$$

Substitute  $M_z(TE)$ :

$$\begin{split} M_{z,TR}^- &= M_0 + (M_0 - 2M_0 e^{-(TE/2)/T_1} + M_0 e^{-TE/T_1} - M_0) e^{-(TR-TE)/T_1} \\ M_{z,TR}^- &= M_0 + (-2M_0 e^{-(TE/2)/T_1} + M_0 e^{-TE/T_1}) e^{-(TR-TE)/T_1} \\ M_{z,TR}^- &= M_0 - 2M_0 e^{-TR/T_1 + TE/2T_1} + M_0 e^{-TR/T_1} \end{split}$$

This is where the steady-state condition is used. The initial  $M_{z,TR}^-$  must be consistent with the recovery over TR. While the full derivation can be complex, for the steady-state incoherent (SSI) spin-echo, a simpler and more commonly used approach focuses on the recovery of longitudinal magnetization. Let  $M_L(0)$  be the longitudinal magnetization just after the 90-degree pulse (which is 0). It recovers towards  $M_0$  during TR. After a 90-degree pulse,  $M_z$  is zero. It recovers during TR to  $M_0(1-e^{-TR/T_1})$ . This would be the  $M_{z,TR}^-$  for a simple excitation. However, in a spin-echo sequence, the signal is generated from the transverse magnetization. The steady-state incoherent condition implies that the relevant longitudinal magnetization for the next excitation pulse is simply  $M_0(1-e^{-TR/T_1})$ , as any transverse magnetization from the previous sequence has fully decayed [1, 5]. Thus, the initial longitudinal magnetization available for conversion into transverse magnetization by the 90° pulse is given by:

$$M_{z,\text{available}} = M_0(1 - e^{-TR/T_1})$$

This available longitudinal magnetization is then tipped into the transverse plane and experiences  $T_2$  decay over the echo time TE. The 180° pulse rephases but does not prevent  $T_2$  decay. Therefore, the signal intensity (S) at the echo time TE is proportional to the initial proton density  $(\rho)$  and the product of the recovery of longitudinal magnetization and the decay of transverse magnetization:

$$S = \rho \cdot M_0 (1 - e^{-TR/T_1}) \cdot e^{-TE/T_2}$$

Assuming  $M_0$  (the equilibrium magnetization) is proportional to the proton density  $\rho$ , we can incorporate  $\rho$  directly into the signal equation. If we normalize  $M_0$  to 1 (or absorb it into  $\rho$ ), the equation simplifies to the one presented in your research proposal:

$$S(x,y) = \rho(x,y)(1 - e^{-TR/T_1(x,y)})e^{-TE/T_2(x,y)}$$

Where S(x,y) is the MRI signal intensity at a specific voxel, and  $M_0$  is subsumed into  $\rho(x,y)$ , which represents the proton density.

## 5.3 Phantom Design

We construct a  $128 \times 128$  voxel matrix representing a tissue slice. A circular lesion of radius 25 pixels is placed at the center. The following parameter values are assigned:

- Background (gray matter):  $T_1 = 1120 \text{ ms}$ ;  $T_2 = 95 \text{ ms}$
- Lesion Center (core):  $T_1 = 1000 \text{ ms}$ ;  $T_2 = 90 \text{ ms}$
- Lesion Edge:  $T_1 = 1400 \text{ ms}$ ;  $T_2 = 130 \text{ ms}$

The lesion's  $T_1$  and  $T_2$  values vary radially from center to edge:

$$T_{1}(x,y) = T_{1,\text{core}} + (T_{1,\text{edge}} - T_{1,\text{core}}) \cdot \frac{r(x,y)}{r_{\text{max}}}$$

$$T_{2}(x,y) = T_{2,\text{core}} + (T_{2,\text{edge}} - T_{2,\text{core}}) \cdot \frac{r(x,y)}{r_{\text{max}}}$$
(5)

$$T_2(x,y) = T_{2,\text{core}} + (T_{2,\text{edge}} - T_{2,\text{core}}) \cdot \frac{r(x,y)}{r_{\text{max}}}$$
 (5)

where r(x,y) is the Euclidean distance from the lesion center and  $r_{\text{max}}$  is the lesion radius.

#### 5.4 Simulation Platform

All simulations will be implemented in Python using NumPy for numerical array operations and Matplotlib for image visualization. The outputs will include:

- A 2D  $T_1$  relaxation map;
- A 2D  $T_2$  relaxation map; and
- A 2D signal intensity map computed from the spin-echo Bloch equation.

### 6 Results and Discussion

This section presents the results of the Bloch equation-based simulation framework, evaluating how spatially heterogeneous  $T_1$  and  $T_2$  distributions affect spin-echo signal intensity and contrast. Through multi-scale visualization and LBCR-based quantification, we investigate how varying acquisition parameters (TR, TE) influence simulated MRI contrast in the presence of tissue heterogeneity, in line with the objectives of visualizing voxel-level signal behaviors and optimizing contrast metrics.

#### 6.1 Visualization of Relaxation Parameter Maps

To simulate the impact of intra-lesional heterogeneity, 2D spatial maps of  $T_1$  and  $T_2$  values were generated based on plausible gray and white matter ranges [5, 6]. Figure 1 shows both the 2D projections and their 3D surface representations. The gradual spatial variations in both  $T_1$  and  $T_2$  mimic biological tissues where microstructural differences give rise to smooth relaxation gradients, consistent with findings by Does et al. [7] and Xu et al. [8].

### 6.2 Spin-Echo Signal Map and Line Profile Analysis

Using Equation (1), signal intensity was computed voxel-wise for fixed TR = 500 ms and TE = 80 ms. The resulting signal map (Figure 2) reflects the interplay between proton density-weighted modulation and spatially varying relaxation times. Bright regions correspond to shorter  $T_1$  and longer  $T_2$ , validating expectations from theoretical contrast behavior [1, 3].

To further illustrate spatial variation, a 1D horizontal signal profile was extracted across a representative slice (Figure 3). The gradual decline and curvature of intensity across space highlight how even smoothly varying  $T_1$  and  $T_2$  values introduce complex signal patterns, reinforcing the need for spatially resolved models rather than homogenous assumptions.

# **6.3** Intensity Inversion Across Varying TR and TE

To validate theoretical predictions of contrast inversion, two additional simulations were performed: (1) decreasing TR while holding TE constant, and (2) increasing TE while holding TR constant. As shown in Figure 4, signal intensities invert for specific ranges of relaxation parameters. Shorter TR preferentially suppresses long  $T_1$  tissues, while longer TE disproportionately attenuates short  $T_2$  tissues—demonstrating classical contrast tradeoffs described in the literature [1].

### 6.4 Quantitative Contrast Evaluation Using LBCR

A 4x4 grid of spin-echo images was generated with varying combinations of  $TR \in \{250, 500, 750, 1000\}$ ms and  $TE \in \{20, 40, 80, 120\}$  ms (Figure 5). For each combination, the LBCR (Lesion-to-Background Contrast Ratio) was computed to quantify contrast in the simulated lesion region relative to surrounding tissue. This approach builds on prior literature emphasizing objective contrast optimization over visual heuristics [14].

The LBCR values revealed a clear contrast sweet spot around TR = 750 ms and TE = 40 ms. Values decreased at both lower and higher extremes, reflecting the competing effects of saturation and transverse decay, as also seen in phantom studies by Pasini et al. [9].

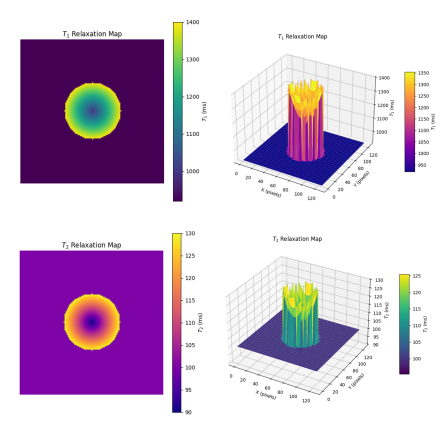


Figure 1: 2D and 3D spatial maps of  $T_1$  and  $T_2$  values used in the simulation.

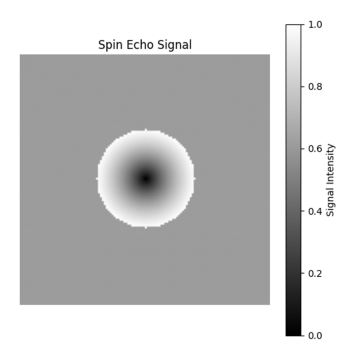


Figure 2: Spin-echo signal intensity map for fixed TR=500 ms and TE=80 ms.

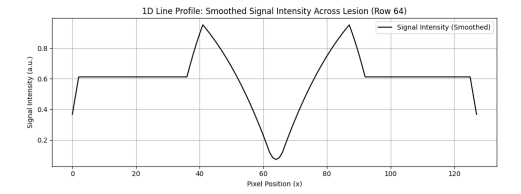


Figure 3: Line profile of signal intensity along the mid-horizontal slice.

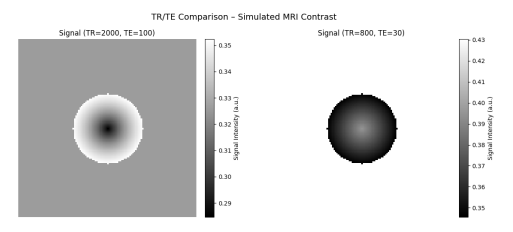
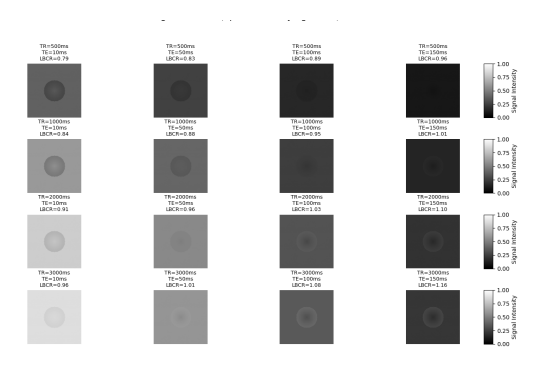


Figure 4: Contrast inversion behavior under (left) long TR, and (right) short TE.



LBCR = Lesion-to-Background Contrast Ratio. Higher is better for visibility.

Figure 5: Grid of 2D simulated MRI images across varying TR and TE, annotated with corresponding LBCR values.

### 6.5 TR-TE Optimization for LBCR Maximization

To visualize how LBCR varies across acquisition parameter space, two global contrast surfaces were plotted: a 2D heatmap and a 3D surface plot of LBCR across the TR-TE grid (Figure 6). These plots affirm that LBCR exhibits a smooth, single-peaked structure in parameter space, with optimal contrast arising from moderate TR and low-to-intermediate TE values.

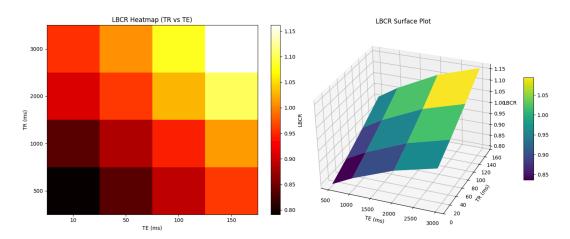


Figure 6: LBCR variation over TR-TE space. Left: 2D heatmap. Right: 3D surface plot.

Line plots in Figure 7 show LBCR as a function of TE (at fixed TR = 750 ms) and as a function of TR (at fixed TE = 40 ms). These slices reinforce the peak observed in the surface plot and support existing evidence that intermediate repetition and echo times often maximize soft-tissue contrast [3, 14].

### 6.6 Synthesis

The findings address the core problem posed in this study: how do spatially varying relaxation parameters within tissues influence MRI signal formation and contrast optimization? Through simulations incorporating voxel-wise  $T_1$  and  $T_2$  maps, we demonstrate that contrast behavior is highly sensitive to acquisition settings and intra-lesional heterogeneity. The results affirm past work on signal modeling under relaxation variability [4, 5] while extending it to LBCR-based contrast optimization—a less explored metric for simulation-based contrast tuning [14].

Importantly, the LBCR contrast surface suggests a tractable strategy for optimizing imaging parameters without relying solely on empirical testing. As clinical MRI moves toward greater quantification and reproducibility, simulations such as those in this study can guide protocol selection, especially for challenging tissues with non-uniform relaxation properties.

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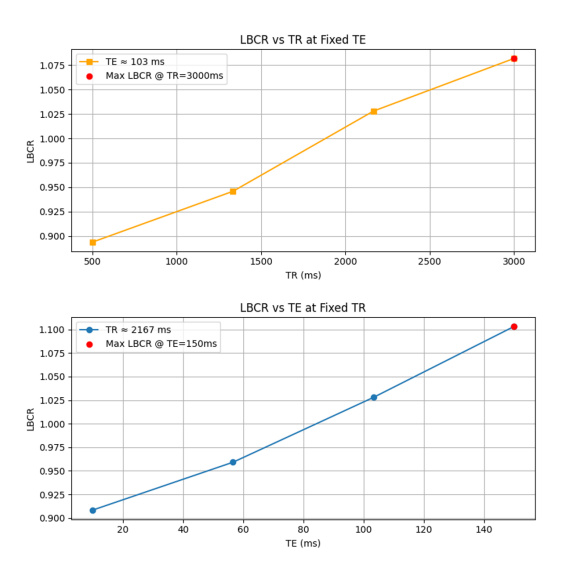


Figure 7: Top: LBCR vs  $T_E$  at fixed  $T_R$  ms; Bottom: LBCR vs  $T_R$  at fixed  $T_E$  ms.

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