

# Reduced-order modelling of Cascadia's slow slip cycles

Yohai Magen<sup>1</sup>, Dave A. May<sup>1</sup>, Alice-Agnes Gabriel<sup>1,2</sup>

<sup>1</sup>Institute of Geophysics and Planetary Physics, Scripps Institution of Oceanography University of California, San Diego, CA, USA

<sup>2</sup>Department of Earth and Environmental Sciences, Ludwig-Maximilians-Universität München, Munich, Germany

## Key Points:

- Our reduced-order models of slow slip cycle simulations are  $\sim 360,000$  times faster than volumetric rate-and-state friction simulations
- ROM-based inversion finds near-lithostatic pore pressure (99.6%) and a  $45 \pm 16$  km wide low-stress zone controlling Cascadia slow-slip events
- Bayesian inversion using reduced-order models constrains both the magnitude and spatial extent of deep slow slip normal stress in Cascadia

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Corresponding author: Yohai Magen, [ymagen@ucsd.edu](mailto:ymagen@ucsd.edu)

## Abstract

Slow-slip events (SSEs) modulate the earthquake cycle in subduction zones, yet understanding their physics remains challenging due to sparse observations and high computational cost of physics-based simulations. We present a scientific machine-learning approach using a data-driven reduced-order modeling (ROM) framework to efficiently simulate the SSE cycle governed by rate-and-state friction in a Cascadia-like 2D subduction setting. Our approach projects fault slip, slip-rate, and state variable trajectories onto a spline-based latent space, which is subsequently emulated using proper-orthogonal decomposition and radial-basis-function interpolation. Achieving a speedup of  $\sim 360,000\times$  compared to volumetric simulations, the ROMs enable comprehensive parameter exploration and Bayesian Markov chain Monte Carlo (MCMC) inversion. Our analysis reveals complex, non-linear dependencies of SSE characteristics on the width and magnitude of the deep, low-effective-normal-stress region. Our MCMC inversion constrained by Northern Cascadia SSEs observations indicates near-lithostatic pore fluid pressure ( $99.6\pm 0.17\%$  lithostatic) and positions the upper frictional transition zone at  $30.4 \pm 2.8$  km depth, consistent with geophysical observations. The inversion resolves the deep SSE-portion of the slab spanning  $45\pm 16$  km with low effective normal stress of  $3.8\pm 1.4$  MPa. This framework provides a new tool for advancing the physics-based understanding of SSEs and subduction zone faulting mechanics. By systematically linking megathrust properties such as fluid pressure and fault strength to rate-and-state friction governed slow slip cycle characteristics, such as recurrence interval, our approach helps to constrain the first- and second-order physics-based controls and the uncertainties of how plate boundaries slip.

## Plain Language Summary

Slow earthquakes, also known as slow slip events (SSEs), are subtle, slow-moving movements that occur deep underground in subduction zones, regions where one tectonic plate slides beneath another. Unlike sudden, destructive earthquakes, slow earthquakes unfold over days to weeks. Although they do not cause shaking, they may influence how and when larger earthquakes happen. Because they occur far below the surface, slow earthquakes are difficult to observe directly. Simulating them with computer models is also challenging and time-consuming, limiting what scientists can explore. In this study, we develop a fast and efficient method to simulate slow earthquakes using scientific machine learning. Our reduced-order model reproduces slow earthquake behavior with over 300,000 times less computational effort than traditional methods. We apply this approach to a model of the Northern Cascadia subduction zone, in the Pacific North-West. Our results show that slow earthquakes are highly sensitive to a deep part of the fault that has unusually low resistance to slip, likely caused by high fluid pressure. This new modeling tool allows us to test a wide range of physical conditions much more efficiently. It provides clearer insight into how slow earthquakes work and how they may influence the broader earthquake cycle.

## 1 Introduction

The conventional view of the earthquake cycle as a simple stick-slip phenomenon, where accumulated tectonic strain is released only through seismogenic earthquakes (Ried, 1911), has evolved dramatically over the past two decades. Advanced seismological and geodetic observation networks have revealed a rich spectrum of fault slip behavior (Gomberg et al., 2016). Among these, slow slip events (SSEs) have emerged as an important component of the seismic cycle, accommodating a large portion of tectonic plate motion and thus contributing to the moment budget at large continental faults (Linde et al., 1996; Rousset et al., 2019) and in many subduction zones worldwide (Behr & Bürgmann, 2021; Schwartz & Rokosky, 2007). For example, SSEs have been documented at the Casca-

64 dia Subduction Zone (CSZ) (Schmidt & Gao, 2010), offshore from Boso Peninsula (Ozawa  
 65 et al., 2007), in the Nankai Trough, Japan (Obara et al., 2004; Araki et al., 2017), and  
 66 along the Hikurangi margin in New Zealand (Wallace et al., 2012). Slow slip events typ-  
 67 ically manifest as transient reversals in the direction of surface deformation, detected by  
 68 high-precision Global Navigation Satellite System (GNSS) networks. SSE displacement  
 69 amplitudes reach several millimeters to centimeters which is 10–100 times faster than  
 70 the tectonic loading and durations range from days to years (Dragert et al., 2001; Peng  
 71 & Gombert, 2010; Bartlow et al., 2011). The CSZ has been particularly instrumental  
 72 for studying SSEs (Schmidt & Gao, 2010; Bartlow et al., 2011). There, transients recur  
 73 quasi-periodically every  $\approx 14$  months and have been monitored for the past  $\sim 25$  years  
 74 (Gombert et al., 2016; Schmidt & Gao, 2010).

75 The physical mechanisms controlling SSEs remain enigmatic, as their potential role  
 76 as precursors of imminent megathrust earthquakes (Obara & Kato, 2016; Ruiz et al., 2014;  
 77 Li & Gabriel, 2024), despite extensive research (Bürgmann, 2018). Their occurrence at  
 78 depths ranging from  $\sim 25$ –40 km introduces uncertainties about the structure, material  
 79 properties, and pressure and temperature conditions at these transitional depths (Behr  
 80 & Bürgmann, 2021). This, combined with the small surface deformation signals produced  
 81 by SSEs and the rapid loss of resolution with depth, limits the constraints that obser-  
 82 vational data alone can place on their governing physics (Liu & Rice, 2007). Consequently,  
 83 numerical simulations incorporating laboratory-derived friction laws have become essen-  
 84 tial for investigating the mechanics of these phenomena (Liu & Rice, 2007; Segall et al.,  
 85 2010). Rate-and-state friction has emerged as the predominant framework for simulat-  
 86 ing sequences of earthquakes and aseismic slip (e.g., Rice & Tse, 1986; Kato, 2002; Bar-  
 87 bot et al., 2012; Jiang et al., 2022; Erickson et al., 2023). These constitutive laws describe  
 88 fault strength as dependent on both slip-rate and a state variable, representing contact  
 89 properties at the fault interface (Dieterich, 1979; Ruina, 1983).

90 Traditionally, two classes of rate-and-state friction models have been used to re-  
 91 produce the key characteristics of SSEs. In both frameworks, SSEs arise spontaneously  
 92 from specific combinations of frictional stability regimes along the fault, particularly, in-  
 93 cluding a conditionally stable region near the transition between fully locked and con-  
 94 tinuously creeping sections. In the first class of models, SSEs emerge from a narrow velocity-  
 95 weakening (VW) region embedded within a velocity-strengthening (VS) zone, where the  
 96 ratio between VW patch width and nucleation size governs SSEs recurrence and prop-  
 97 agation (Rubin, 2008). The second class, introduced by Liu and Rice (2007), focuses on  
 98 transitional frictional stability near the down-dip limit of the seismogenic zone. In this  
 99 approach SSEs nucleate within conditional stable fault portions characterized by VW  
 100 behavior that is stabilized by low effective normal stress and proximity to the VW-VS  
 101 transition. To model spontaneous SSEs, these models incorporate a fault portion with  
 102 reduced effective normal stress (e.g. Liu & Rice, 2005, 2009; Rubin, 2008; Matsuzawa  
 103 et al., 2010; Li & Liu, 2016a; Luo & Ampuero, 2018). In subduction zones, this may re-  
 104 sult from elevated pore fluid pressure at the megathrust interface (Audet et al., 2009;  
 105 Suppe, 2014). The Liu and Rice (2007) framework demonstrates that the ratio between  
 106 the width of this low effective stress zone and the nucleation length scale ( $h^*$ ) critically  
 107 controls both the occurrence and characteristics of SSEs, with higher ratios producing  
 108 more frequent events of smaller magnitude (Liu & Rice, 2009; Cattania, 2019).

109 More recent efforts have focused on incorporating rate-and-state friction fault, along  
 110 with fluid flow, permeability evolution, and changes in pore fluid pressure to produce SSEs  
 111 (Ozawa et al., 2024; Perez-Silva et al., 2023). This aims to provide a more realistic phys-  
 112 ical explanation that aligns with observations. Despite their success in reproducing ob-  
 113 served characteristics of SSEs, physics-based forward models are often simplified, e.g. by  
 114 reducing dimensionality or idealizing fault geometry, and typically focus on limited as-  
 115 pects of the parameter space due to computational challenges. Simulating SSEs requires  
 116 capturing processes across vastly different timescales, from the evolution of individual

slow slip events to the periods where fault slip-rate and slip-rate gradients are low. This multi-scale character necessitates repeated fine temporal resolution to capture SSE nucleation dynamics, while simultaneously requiring long simulation times to capture multiple SSE cycles. The computational costs become problematic when attempting to conduct sensitivity analyses or quantify uncertainties in model predictions, motivating the need for more efficient simulation approaches.

Scientific machine learning (ML) methods have recently advanced the analysis of SSE observations. Deep learning models, in particular, have proven effective in detecting low signal-to-noise ratio signatures of SSEs from geodetic data and their seismic counterparts, low-frequency earthquakes (LFEs) and non-volcanic tremor (Hulbert et al., 2022). For instance, Lin et al. (2024) applied Convolutional Neural Networks (CNNs) to continuous seismic records to build comprehensive LFE catalogs, revealing thousands of previously uncataloged events, suggesting hidden SSEs which were unraveled previously. Münchmeyer et al. (2024) demonstrated that such models can generalize across different subduction zones, suggesting that LFEs may share universal waveforms characteristics that are learnable by deep networks. From a geodetic perspective, Costantino et al. (2023) developed a multi-station deep learning detector that operates on raw GNSS time series to identify subtle deformation transients associated with SSEs. To overcome the scarcity of labeled training data, they used a synthetic training simulation data set. Early efforts to apply ML directly to the governing physics of fault slip include Physics-Informed Neural Networks (PINNs), embedding the governing equations into the loss function of neural networks to estimate fault properties (e.g. Fukushima et al., 2023; Okazaki et al., 2022; Fukushima et al., 2025; Rucker & Erickson, 2024). Recently, Fourier Neural Operators (FNOs) have been used to emulate the nonlinear equations governing dynamic rupture propagation (Tainpakdipat et al., 2025).

Reduced-order models (ROMs) have proven to be an effective method to accelerate geoscience simulations, including modeling slow slip events (Kaveh et al., 2024), seismic shake maps (Rekoske et al., 2023) and wave fields (Rekoske et al., 2025), thermal structures in subduction zones (Hobson & May, 2025a), geothermal geodynamic processes (Degen et al., 2023) and magnetotellurics (Quiaro et al., 2025). By projecting high-dimensional systems onto lower-dimensional subspaces that retain the primary characteristics, ROMs make it possible to capture the overall evolution of complex physical systems at a fraction of the cost of full-order models (FOMs). Reported speedups range from several hundred to over 100,000 times faster, depending on the application. Such reductions in computational cost enable global exploration of the parameter space and facilitate uncertainty quantification that are impractical with physics-based models.

In this study, we introduce and evaluate a new reduced-order modeling framework designed to efficiently simulate SSE cycles governed by rate-and-state friction. We utilize the volumetric, scalable discontinuous Galerkin library (Uphoff et al., 2022) on unstructured meshes to simulate sequences of aseismic slip in a Cascadia-like model setup following Liu and Rice (2009), and expand their exploration of the model parameter space. Our ROM methodology combines an efficient spline-based latent representation of rate-and-state friction SSE cycle models outputs, leveraging their phase-space characteristics, with Proper Orthogonal Decomposition (POD, Bui-Thanh et al. (2003)) and coefficient-interpolation using Radial Basis Functions (RBFs, Audouze et al. (2009)). This approach reduces the computational cost by  $\sim 3.6 \times 10^5$  times compared to performing a physics-based tandem simulation, enabling comprehensive parameter space exploration and uncertainty quantification of SSE characteristics. We perform a full Bayesian Markov chain Monte Carlo (MCMC) inversion constrained by Northern Cascadia SSEs characteristics, constraining the width ( $44.7 \pm 16.2$  km) and magnitude ( $3.8 \pm 1.4$  MPa) of a deep low effective normal stress region. We demonstrate the accuracy and efficiency of our ROM framework and highlight its potential for advancing the physics-based understanding and simulation of complex fault slip behavior.

**Table 1.** Summary of tandem SSE cycle forward simulation parameters.

Symbol	Definition	Value	Reference
$W_l$	Up-dip extent of low effective normal stress zone	varied [152, 189.5] km	
$W_r$	Down-dip extent of low effective normal stress zone	252 km along dip	
$\sigma_0$	Fault normal stress outside of $[W_l, W_r]$	50 MPa	Liu and Rice (2009)
$\sigma_W$	Fault normal stress within $[W_l, W_r]$	varied [1, 6] MPa	See Eq. (3)
$L_0$	Characteristic slip distance outside of $[W_l, W_r]$	13 mm	Liu and Rice (2009)
$L_W$	Characteristic slip distance within $[W_l, W_r]$	0.16 mm	Liu and Rice (2009)
$a$	Direct effect parameter	varies along-fault, see Figure 1	He et al. (2006)
$b$	Evolution effect parameter	0.0045	He et al. (2006)
$\mu$	Shear modulus	33.91 GPa	
$\nu$	Poisson’s ratio	0.25	
$f_0$	Reference coefficient of friction	0.6	
$\dot{s}_0$	Reference slip-rate	$1 \times 10^{-6}$ m/s	

## 2 Full-order models of slow slip cycles in Cascadia

To investigate the time-dependence of repeating SSEs in a Cascadia-like subduction zone, we first describe a full-order model (FOM) simulation framework. The fault stress evolution in this model follows the regularized form of the rate-and-state friction law (Dieterich, 1979; Ruina, 1983; Lapusta et al., 2000; Rice & Ben-Zion, 1996), given by

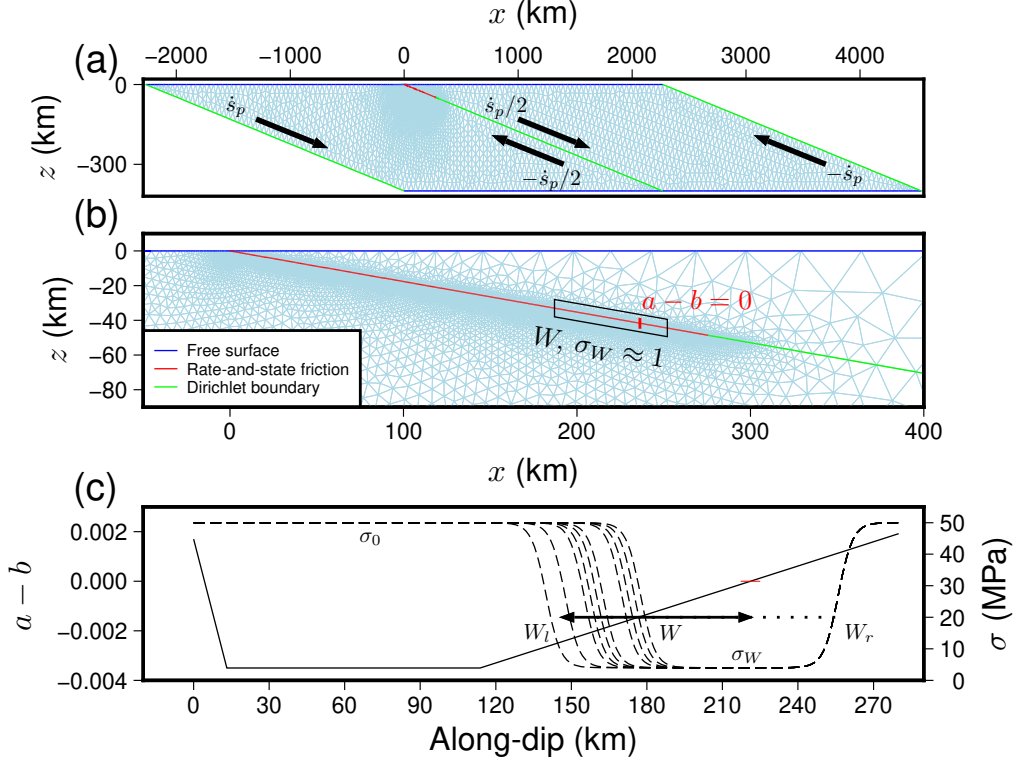
$$\tau(\dot{s}, \psi) = a \operatorname{arcsinh} \left( \frac{\dot{s}}{2\dot{s}_0} \exp \left( \frac{\psi}{a} \right) \right) \quad (1)$$

where the fault shear stress  $\tau$  is a function of the slip-rate  $\dot{s}$  (with  $\dot{s} = \|\dot{\mathbf{s}}\|$ ) and a state variable  $\psi$ , with  $a$  representing an empirical friction parameter to describe the “direct effect” and  $\dot{s}_0$  a reference slip-rate. The state variable evolves temporally according to the Dietrich-Ruina aging law

$$\frac{d\psi}{dt}(\dot{s}, \psi) = \frac{b\dot{s}_0}{L} \left( \exp \left( \frac{f_0 - \psi}{b} \right) - \frac{\dot{s}}{\dot{s}_0} \right) \quad (2)$$

where  $b$  is an empirical frictional parameter describing the time-dependent “evolution” effect,  $L$  denotes the characteristic slip distance, and  $f_0$  is a reference friction coefficient.

The evolution of and interaction between slip-rate and state variable can be analyzed in a two-dimensional phase-space defined by these variables, which offers a simplified framework for examining the system’s inherent instability (Ranjith & Rice, 1999). For VW faults, trajectories in this phase-space may form either closed or open orbits (i.e.,



**Figure 1.** Model setup and unstructured triangular mesh for the full-order SSE cycle models using tandem. (a) View of the computational domain, boundary conditions and applied loading, where  $\dot{s}_P$  is the applied loading rate. Not to scale, note the  $15\times$  vertical exaggeration of the  $y$ -axis. (b) Zoomed view of the rate-and-state friction fault, showing the adaptively refined mesh, with  $W$  and  $\sigma_W$  being the width and magnitude of a low effective normal stress zone respectively and  $a, b$  are rate-and-state friction parameters. (c) Along-fault profiles of  $a - b$  derived from gabbro friction experiments (He et al., 2006) and the initial effective normal stress distribution along the fault where  $W_l$  and  $W_r$  are the up-dip and down-dip extent of the low effective normal stress zone,  $\sigma_0$  is the effective normal stress outside of this zone and red horizontal bar mark the VW-VS transition point. Profiles of normal stress are shown for  $\sigma_W = 4$  MPa.

limit cycles, e.g., (Pranger et al., 2022)), reflecting stable oscillatory behavior or runaway slip, respectively. The geometry of phase-space trajectories provides a clearer, lower-dimensional view of the complex faulting behavior than the time-domain (Viesca, 2016a; Ciardo & Viesca, 2024). We will later utilize this phase-space representation of the rate-and-state friction SSE cycle simulations as a more suitable representation for reduced-order modeling.

All models in this study consist of two-dimensional parallelogram-shaped domains measuring 4500 km in length and 400 km in width, containing a single planar fault that bisects the domain into two geometrically identical regions (Figure 1(a)). This setup follows the configuration of the BP3 SCEC community benchmark (Erickson et al., 2023). To approximate the CSZ, both the fault and lateral domain boundaries are inclined at a shallow dip angle of  $10^\circ$  relative to the upper and lower domain boundaries. The up-dip edge of the fault intersects the upper free surface, while the down-dip edge terminates at the bottom boundary, which is also treated as a free surface.

Loading is prescribed as steady tectonic convergence, (Figure 1(a), Biemiller et al., 2024), imposed via Dirichlet boundary conditions, with velocities of  $\dot{\mathbf{s}}_p = (11.5, 2.0)$  cm/year in the  $x$  and  $z$  components, respectively, to the fault at depths exceeding 280 km along-dip. In addition  $\pm \dot{\mathbf{s}}_p/2$  is applied to the lateral domain boundaries. Fault slip behavior governed by rate-and-state friction can be classified into two regimes based on the difference of the frictional parameters  $a - b$ . When  $a - b < 0$ , the fault exhibits VW behavior, which promotes the release of accumulated stress through rapid seismic slip. In contrast, when  $a - b > 0$ , the fault is VS, favoring stable creep (Scholz, 1998).

SSEs are emerging in our model framework by applying a reduced effective normal stress ( $\sigma_W$ ) to a localized subregion of the rate-and-state friction fault, which encompasses the stability transition zone ( $a - b = 0$ ) between VW (up-dip) and VS (down-dip) frictional behavior (1), following the approach established in previous studies (e.g., Liu & Rice, 2005, 2007, 2009; Rubin, 2008). The width of this region will be denoted as  $W$ .

The initial normal stress along the fault (Figure 1(c)) is given by

$$\sigma(d) = \left( \frac{1}{1 + \exp(-\gamma(d - W_l))} - \frac{1}{1 + \exp(-\gamma(d - W_r))} \right) (\sigma_W - \sigma_0) + \sigma_0, \quad (3)$$

where  $d$  is the distance along the fault in the down-dip direction with  $d = 0$  at the top free surface,  $\sigma_0$  is the normal stress outside the low effective normal stress zone which is taken to be 50 MPa. The parameter  $\gamma$  is a smoothing coefficient controlling the smoothness of the transition between the two normal stress regions and taken to be 0.35, while  $W_l$  and  $W_r$  represent the up-dip and down-dip boundaries of the region of low effective normal stress.

We follow Liu and Rice (2009) and define the width of the low-effective normal stress zone ( $W$ ) as the distance from the stability transition point ( $a - b = 0$ ) to its up-dip termination ( $W_l$ ). In the following, we build reduced order models to represent variations in both the width (by varying  $W_l$ ) and normal stress reduction (varying  $\sigma_W$ ) of this region (Figure 1). Other model parameters remain fixed.

We use frictional parameters resembling experimentally derived values for gabbro (He et al., 2006). The parameter  $b$  is held constant throughout the domain, while  $a$  is varied to achieve the target  $a - b$  values (Figure 1) consistent with the experimental data. As characteristic slip distance, we assume  $L_W = 0.16$  mm within the low effective normal stress zone and  $L_0 = 13$  mm outside of this zone. All model parameters are listed in Table 1.

We employ the open-source SEAS simulation software tandem (Uphoff et al., 2022) to conduct quasi-dynamic simulations of SSEs in this two-dimensional Cascadia-like setting (Liu & Rice, 2009). Each simulation constitutes a full-order model (FOM), where simulations are run to contain at least 20 SSE cycles, after a spin up phase. tandem utilizes a discontinuous Galerkin (DG) finite element method and supports curvilinear elements within an unstructured mesh composed of triangular elements. Preconditioned Krylov methods from the PETSc-TAO library (Balay et al., 2025b, 2025a, 1997) are used to solve the elasticity problem arising from the DG spatial discretization. Time integration of the rate-and-state friction ODEs within tandem uses an embedded sixth-order Runge-Kutta scheme (Abhyankar et al., 2018). We implement time step adaptivity by using an error estimator obtained from the difference between the 5th order and 6th order solution following (Dormand & Prince, 1980).

For all simulations, the on-fault resolution is chosen to resolve the smaller of the characteristic nucleation size,  $h^*$ , and the process zone size,  $\Lambda$ , by a factor of  $20/d_{\text{poly}}$ , where  $d_{\text{poly}} = 6$  is the polynomial degree of DG basis functions used, thereby ensuring adequate spatial resolution (Uphoff et al., 2022). The simulations are initiated by

setting the state variable  $\psi$  to a value that satisfies the governing rate-and-state friction law for the prescribed initial stress conditions and slip-rate, respectively.

In a general setting, the FOM described above is a set of coupled parametric ODEs coupled to a PDE. In any given study designed to explore model parameters, one is required to hold some parameters constant and vary others. To generalize the presentation of the reduced-order model in Section 3 we denote the parameters varied via

$$\boldsymbol{\xi} = \{\xi_1, \dots, \xi_{\dim(\mathcal{P})}\}. \quad (4)$$

In this work we consider a two-dimensional parameter space ( $\dim(\mathcal{P}) = 2$ ) with  $\xi_1 = W \in [30.5, 68]$  km and  $\xi_2 = \sigma_W \in [1, 6]$  MPa. All other parameters such as  $a$ ,  $b$ ,  $L$  (Table 1 for a complete list) are not varied. We will denote our 2D parameter space by  $\mathcal{P}$ .

### 3 SEAS model order reduction

Our reduced-order model (ROM) methodology is designed provide an approximate value of slip, slip-rate and state variable at locations along the fault as a function of time  $t$  and the parameters  $\boldsymbol{\xi} = (W, \sigma_W)$ . An overview of our ROM methodology is shown in Figure 2.

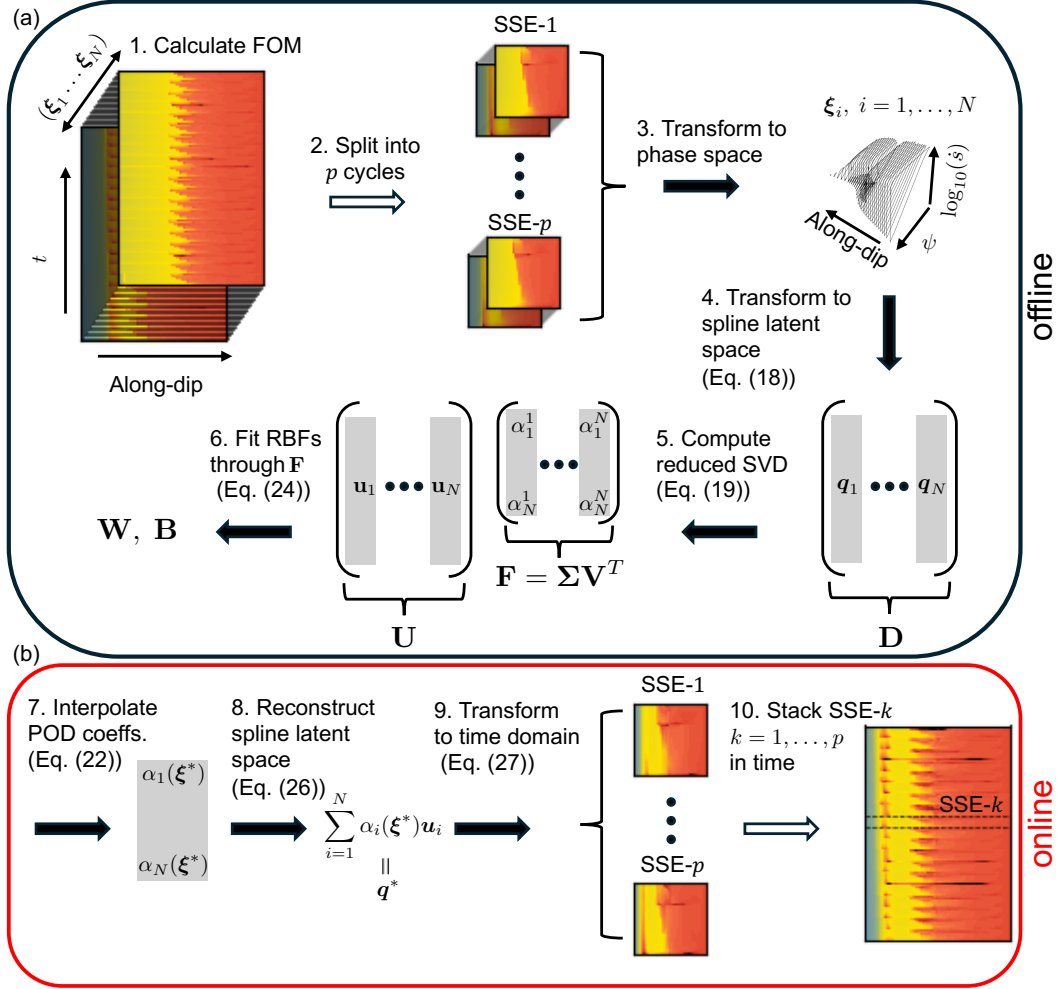
The construction of our ROM starts by uniformly sampling the parameter space within prescribed minimum and maximum values for each  $\xi_i$ . We then evaluate the FOM at different parameter realization  $\boldsymbol{\xi}_i$ ,  $i = 1, \dots, N$  to obtain  $N$  simulation outputs, each being denoted by  $\mathbf{Q}(\boldsymbol{\xi}_i)$  (step 1 in Figure 2). The  $N$  outputs consist of time discrete values of slip, slip-rate and state variable at different locations along the fault, that is  $\mathbf{Q}(\boldsymbol{\xi}_i) = (\mathbf{t}, \dot{\mathbf{S}}, \boldsymbol{\Psi}, \mathbf{S})$ , where  $\mathbf{t}$  is a vector of simulation time stamps of length  $n$ , and  $\dot{\mathbf{S}}, \boldsymbol{\Psi}, \mathbf{S} \in \mathbb{R}^{n \times m}$  are matrices representing slip-rate, state variable, and cumulative slip, respectively, across  $m$  fault observation points. We will refer to the time series of length  $n$  for quantities at observation points  $j = 1, \dots, m$  via the sub-script  $j$ . That is,  $\mathbf{S}_j$  denotes the time series for the slip at the  $j^{\text{th}}$  observation point and thus  $\mathbf{S} = [\mathbf{S}_1, \dots, \mathbf{S}_m]$  and  $\mathbf{Q}_j = (\mathbf{t}, \dot{\mathbf{S}}_j, \boldsymbol{\Psi}_j, \mathbf{S}_j)$ .

A key aspect of our ROM strategy is that, instead of reducing the dimensionality of simulations defined over fixed time intervals, we reduce the dimensionality over each SSE cycle independently. Each trajectory  $\mathbf{Q}(\boldsymbol{\xi}_i)$  is segmented into  $p$  individual cycles  $\mathbf{Q}^k(\boldsymbol{\xi}_i)$ ,  $k = 1, \dots, p$  by applying a slip-rate threshold criterion at a pre-selected location on the fault (step 2 in Figure 2). This location is set to 195 km along-dip, a position chosen to be within the VW region of the low effective normal stress zone ( $W$ ) for all FOM simulations. Whenever the slip-rate at this location exceeds a threshold of  $1 \times 10^{-4}$  m/s, a new cycle boundary is defined, thereby triggering a split in the data sequence. To prevent the clustering of multiple triggers within a single event, we impose an additional temporal constraint which ensures that no new cycle boundary can be defined within a three-month period following the previous one. This period is well below the expected recurrence interval of the SSE and thus ensures that each distinct SSE is identified as a single cycle.

The construction of the ROM corresponding to each individual  $k$  SSE cycle is detailed in Sections 3.1.1 and 3.1.2. Once constructed, each cycle-specific ROM defines a mapping

$$\text{ROM}^k(\boldsymbol{\xi}^*) = \mathbf{Q}^k(\boldsymbol{\xi}^*), \quad \text{for } k = 1, \dots, p, \quad (5)$$

where  $\text{ROM}^k(\cdot)$  represents the reduced-order model for the  $k^{\text{th}}$  cycle,  $\boldsymbol{\xi}^* \in \mathcal{P}$  denotes an arbitrary parameter vector in which  $\boldsymbol{\xi}^* \neq \boldsymbol{\xi}_i$ ,  $i = 1, \dots, N$  and  $\mathbf{Q}^k(\boldsymbol{\xi}^*)$  is the resulting ROM approximation for the complete time series of the slip-rate, state and slip at all fault observation points  $m$ . The superscript  $*$  is used to distinguish ROM inputs from the full-order model inputs  $\boldsymbol{\xi}_i$ ,  $i = 1, \dots, N$  used to construct the ROM. To reconstruct a full multi-cycle simulation sequence for a given parameter  $\boldsymbol{\xi}^*$ , we evaluate



**Figure 2.** Overview of the SEAS ROM methodology. (a) The offline component of the ROM framework, which is performed once during initialization. This computationally intensive setup stage primarily involves data collection by performing  $N$  FOM simulations with parameters vectors  $\xi_i$  (step 1). This is followed by latent space encoding ( $q_i$ ) via spline interpolation (step 4). Subsequently, dimensionality reduction using POD of  $\mathbf{D}$ , the stacked latent vector matrix, via SVD, where  $\mathbf{D} = \mathbf{U} \mathbf{\Sigma} \mathbf{V}^T$  (step 5). The columns of  $\mathbf{U}$  ( $u_i$ ) form the POD basis. The ROM is constructed by interpolating the POD coefficients ( $\alpha_r^i$ ) along the rows of the POD coefficient matrix  $\mathbf{F}$  using RBFs with polynomial terms, defined by RBF weights  $\mathbf{W}$  and polynomial coefficient  $\mathbf{B}$  matrices. (b) The online component of the ROM framework, which enables fast inference of ROM-based simulations for any new parameter vector  $\xi^*$ , where the  $*$  superscript distinguishes ROM inputs and outputs from their FOM counterparts, within the bounds of the training set. Steps 3 through 9 are performed for each  $k = 1, \dots, p$  cycles and marked with solid arrows.

each of the  $p$  individual cycle ROMs to obtain  $\mathbf{Q}^1(\boldsymbol{\xi}^*), \dots, \mathbf{Q}^p(\boldsymbol{\xi}^*)$  and concatenate them to form a continuous approximation of the complete time series for slip, slip-rate and state variable (step 10 in Figure 2).

While the uniform sampling of our parameter space ensures broad coverage, it does not account for the possibility that the underlying function approximated by the ROM may locally exhibit high gradients, or non-smooth behavior. In such regions of the parameter space, the ROM may yield large approximation errors. To address this, we adopt an empirical refinement strategy wherein the uniformly sampled parameter space is iteratively refined in regions exhibiting high ROM approximation errors. Errors are estimated using leave-one-out cross-validation (LOOCV). Regions which are identified as having a high error are resampled, e.g., refined in parameter space, by performing additional FOM simulations, and rebuilding the ROM (Section 3.2). The iterative refinement continues until the ROM leave-one-out errors fall within the range of SSEs recurrence interval and potency ( $P_0$ ) standard deviations of the corresponding FOM simulations. The potency is defined as

$$P_0 = \int_A \|\mathbf{s}^+ - \mathbf{s}^-\|_2 dA, \quad (6)$$

where  $A$  is the fault surface area and  $\mathbf{s}^-$  and  $\mathbf{s}^+$  are the cumulative slip before and after each SSE event respectively (Ben-Zion & Zhu, 2002).

### 3.1 Reduced-order models of slow slip cycles in Cascadia

#### 3.1.1 Simulation latent space representation

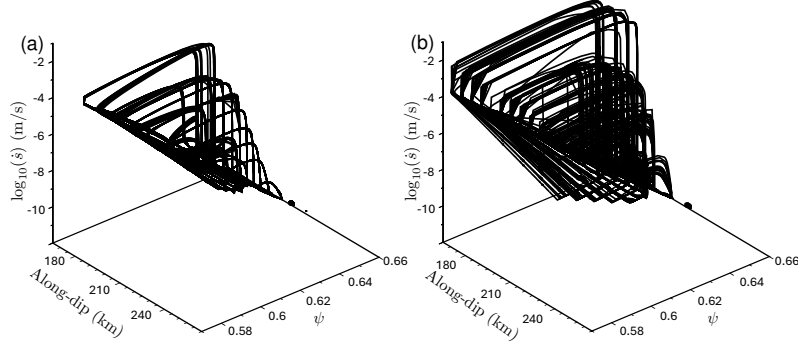
Directly reducing the order of the FOM is challenging for two reasons. First, the simulations are complex, spanning tens of orders of magnitude in slip-rate and involving adaptive time steps ranging more than 6 order of magnitude. Second, each  $N$  simulation output ( $\mathbf{Q}(\boldsymbol{\xi}_i)$ ) amounts to approximately 2 GB of floating-point data, making it impractical to apply a ROM scheme directly to the raw simulation output. Instead, we propose to use a B-spline interpolant (spline latent space hereafter) to represent the simulation output, which in general constitutes a lossy compression framework.

Casting spline interpolation as a compression framework aligns with a broader challenge in scientific computing, where high-performance computing (HPC) applications generate increasingly large datasets, making storage and analysis difficult. Data compression provides a viable solution by reducing the data size. While lossless compression preserves all original information, its compression is usually not efficient for scientific simulation data, which lack the necessary redundant patterns (Lindstrom, 2014). In contrast, lossy compression, such as the spline latent space approach, achieves higher compression factors by selectively discarding less critical information under user-defined error constraints (Tao et al., 2017). This enables efficient representation while preserving essential features for further analysis.

To formalize the lossy compression framework, we define a transformation of simulation data  $\mathbf{Q}_j$  (the parameter vector is omitted for notational clarity) into a latent space representation  $\mathbf{q}_j \in \mathbb{R}^l$  at a single observation point ( $j$ ) along the fault via

$$G(\mathbf{Q}_j) = \mathbf{q}_j, \quad G^{-1}(\mathbf{t}, \mathbf{q}_j) \approx \mathbf{Q}_j \quad \text{such that} \quad \mathcal{F}(\mathbf{Q}_j, G^{-1}(\mathbf{t}, \mathbf{q}_j)) > 1 - \epsilon, \quad (7)$$

where  $\mathcal{F}(\cdot, \cdot)$  is the fidelity criterion with tolerance  $\epsilon \ll 1$ ,  $G(\cdot)$  and  $G^{-1}(\cdot, \cdot)$  are the transformations defined in this subsection (Eqs. (12), (13), (14)) and  $G^{-1}$  is evaluated over  $\mathbf{t} \in \mathbf{Q}_j$ . For this study, this criterion is met if the Pearson Correlation Coefficient (PCC) between its original slip-rate time series  $\dot{\mathbf{S}} \in \mathbf{Q}_j$ , and its corresponding reconstructed slip-rate time series  $\tilde{\dot{\mathbf{S}}}_j \in G^{-1}(\mathbf{t}, \mathbf{q}_j)$  is greater than  $1 - \epsilon$ , where we have used  $\epsilon = 0.001$ .



**Figure 3.** Phase-space  $H$  (Eq. (9)) as a function of the along-dip distance of FOM simulation results. (a) FOM results using  $\xi_1 = (W = 37.375 \text{ km}, \sigma_W = 2 \text{ MPa})$ . (b) FOM results using  $\xi_2 = (W = 48 \text{ km}, \sigma_W = 4 \text{ MPa})$ . Both panels show trajectories plotted for observation points sampled every 6 km. For each observation point, the trajectory forms a nearly closed loop. The time-domain behavior of these results is depicted in Figure 4.

Seismic cycle simulations often span a wide range of time steps and slip velocities, covering several orders of magnitude. This variability makes direct spline placement and coefficient estimation over  $t \rightarrow \|\dot{\mathbf{s}}(t)\|$  particularly challenging. However, the simulation data in phase-space, along the parametric curve

$$H(t) = (\log_{10}(\dot{\mathbf{s}}(t)), \psi(t)) \quad (8)$$

exhibit a well-behaved, loop-like structure (Figure 3). Consequently, SEAS simulation data can be compressed more efficiently via splines when the output is represented in phase-space rather than time. We achieve this by first parametrization time  $t$  with a variable  $\phi$  which is defined by:

$$\phi(t) \in [0, 1] \quad \text{s.t.} \quad \phi(t_1) = 0 \text{ and } \phi(t_n) = 1.$$

such that  $\phi$  represents the time progression along the trajectory in the phase-space, and then

$$\begin{aligned} H(\phi) &= H(\phi(t)) \\ &= \left( \log_{10}(\dot{\mathbf{s}}(\phi(t))), \psi(\phi(t)) \right). \end{aligned} \quad (9)$$

Using spline interpolation, we define an interpolant for  $\phi$  in terms of  $t$ . Similarly, we also can define a spline interpolate for slip-rate, state and slip in terms of  $\phi$ . This two-step procedure results in the construction of four B-splines, constructed via least-squares fitting. These splines will be used to define the forward transformation  $G$ . The inverse transformation  $G^{-1}$  is then defined by first evaluating spline interpolant to approximate  $\phi(t)$ , and this result is used to evaluate the splines approximating the slip-rate, state and slip (as functions of  $\phi$ ).

#### Definition of $G$ and $G^{-1}$

Given a set of data points whose elements consist of paired values of independent ( $x$ ) and dependent variables ( $f(x)$ ) denoted by  $\{\hat{x}_i, \hat{f}_i\}_{i=1}^n = (\hat{\mathbf{x}}, \hat{\mathbf{f}})$  where  $\hat{f}_i = f(\hat{x}_i)$ , then the construction of the B-Spline is defined by

$$\mathcal{B} : \hat{\mathbf{x}}, \hat{\mathbf{f}} \rightarrow \mathbf{k}_x, \mathbf{c}_f, \quad (10)$$

where  $\mathbf{k}_x, \mathbf{c}_f$  are the spline knots and spline coefficients respectively. The B-spline approximation of the function  $f(\cdot)$  is given by

$$f(x) \approx \tilde{f}(x) = \mathcal{B}^{-1}(\mathbf{k}_x, \mathbf{c}_f, x), \quad x_1 \leq x \leq x_n. \quad (11)$$

We note that subscript on the knot vector  $\mathbf{k}$  indicates the independent variable, and the subscript on the coefficient vector  $\mathbf{c}$  indicates the dependent variable.

In this work we use B-spline interpolants to define a latent space transformation for the SSE simulation output. Given  $\mathbf{Q}_j$ , the transformations  $G : \mathbf{Q}_j \rightarrow \mathbf{q}_j$  is defined as:

$$\begin{aligned} G(\mathbf{Q}_j) &= (\mathcal{B}(\mathbf{t}, \Phi_j), \mathcal{B}(\Phi_j, \dot{\mathbf{S}}_j), \mathcal{B}(\Phi_j, \Psi_j), \mathcal{B}(\Phi_j, \mathbf{S}_j)) \\ &= (\mathbf{k}_t, \mathbf{c}_\phi, \mathbf{k}_\phi, \mathbf{c}_{\dot{s}}, \mathbf{c}_\psi, \mathbf{c}_s) \\ &= \mathbf{q}_j, \end{aligned} \quad (12)$$

where  $\Phi_j = (\phi(t_1) \dots \phi(t_n))$  for  $\mathbf{t} = (t_1, \dots, t_n) \in \mathbf{Q}_j$ ,  $\mathbf{k}_t, \mathbf{c}_\phi \in \mathbb{R}^{K_0}$  are the spline knot and coefficient vector for  $\phi(t)$  and  $\mathbf{k}_\phi, \mathbf{c}_{\dot{s}}, \mathbf{c}_\psi, \mathbf{c}_s \in \mathbb{R}^{K_1}$  are the spline knot and coefficient vector for  $\dot{s}(\phi), \psi(\phi), s(\phi)$  respectively. As the same independent variable  $\phi$  is used for the slip-rate, state and slip, the knot vector  $\mathbf{k}_\phi$  is only stored once in Eq. (12). In Section 4.2 we discuss how  $K_0, K_1$  were chosen. Furthermore, in Appendix B a description of how the knot placement is defined within  $\mathbf{k}_t, \mathbf{k}_\phi$  (given  $K_0, K_1$ ) is provided.

For a specific time  $t$ , the spline latent vector  $\mathbf{q}_j$  can be used to reconstruct the slip-rate, state and slip. Defining this as  $g^{-1} : t, \mathbf{q}_j \rightarrow \tilde{s}(t), \tilde{\psi}(t), \tilde{s}(t)$  we have:

$$\begin{aligned} g^{-1}(t, \mathbf{q}_j) &= (\mathcal{B}^{-1}(\mathbf{k}_\phi, \mathbf{c}_{\dot{s}}, \tilde{\phi}(t)), \mathcal{B}^{-1}(\mathbf{k}_\phi, \mathbf{c}_\psi, \tilde{\phi}(t)), \mathcal{B}^{-1}(\mathbf{k}_\phi, \mathbf{c}_s, \tilde{\phi}(t))) \\ &= (\tilde{s}(t), \tilde{\psi}(t), \tilde{s}(t)), \end{aligned} \quad (13)$$

where  $\tilde{\phi}(t) = \mathcal{B}^{-1}(\mathbf{k}_t, \mathbf{c}_\phi, t)$  is the B-spline approximation for  $\phi$ .

Lastly we have the inverse transformation given by

$$G^{-1} : \tilde{\mathbf{t}}, \mathbf{q}_j \rightarrow \tilde{\mathbf{Q}}_j = (\tilde{\mathbf{t}}, \tilde{\dot{\mathbf{S}}}_j, \tilde{\Psi}_j, \tilde{\mathbf{S}}_j), \quad (14)$$

where  $\tilde{\mathbf{t}} \in \mathbb{R}^{n^*}$  is an arbitrary time vector which in general is different to  $\mathbf{t}$ , and  $\tilde{\dot{\mathbf{S}}}_j, \tilde{\Psi}_j, \tilde{\mathbf{S}}_j$  are the length  $n^*$  vectors containing the spline approximations for slip-rate, state and slip at observation point  $j$  for times  $t = \tilde{t}_1, \dots, \tilde{t}_{n^*}$ . The entries of  $\tilde{\dot{\mathbf{S}}}_j, \tilde{\Psi}_j, \tilde{\mathbf{S}}_j$  are computed using  $\tilde{s}(t), \tilde{\psi}(t), \tilde{s}(t)$  obtained from applying  $g^{-1}$  pointwise.

The transformations from simulation output into a spline latent space at a specific observation point could easily be expanded to all simulation fault observation points by applying  $G$  on each specific observation point and letting the latent space vector  $\mathbf{q}$  store the spline knots and coefficients of all observation points. For notation simplicity we will use from here on  $\mathbf{q} \in \mathbb{R}^l$ ,  $G$ ,  $g^{-1}$  and  $G^{-1}$  to represent the transformation of the entire simulation to latent space with

$$G(\mathbf{Q}) = (\mathbf{k}_t, \mathbf{c}_{\phi 1}, \mathbf{k}_{\phi 1}, \mathbf{c}_{\dot{s} 1}, \mathbf{c}_{\psi 1}, \mathbf{c}_{s 1}, \dots, \mathbf{c}_{\phi m}, \mathbf{k}_{\phi m}, \mathbf{c}_{\dot{s} m}, \mathbf{c}_{\psi m}, \mathbf{c}_{s m}) = \mathbf{q} \quad (15)$$

where  $\mathbf{Q} = (\mathbf{t}, \dot{\mathbf{S}}, \Psi, \mathbf{S})$  with  $\dot{\mathbf{S}}, \Psi, \mathbf{S} \in \mathbb{R}^{m \times n}$  as defined in Section 2. We note that all fields (slip-rate, state and slip) and at all observation points use the same vector of time values  $\mathbf{t}$ , hence for efficiency the knot vector  $\mathbf{k}_t$  can be reused in Eq (15) and thus only appears once. Therefore, the length  $l$  of the spline latent space representation  $\mathbf{q}$  is given by:

$$l = (1 + m)K_0 + 4mK_1, \quad (16)$$

where  $K_0$  is the dimension of the knot vector  $\mathbf{k}_t$  and each of the  $m$  coefficient vectors  $\mathbf{c}_{\phi j}$ . Similarly,  $K_1$  is the dimension of each of the  $m$  knot vectors  $\mathbf{k}_{\phi j}$  and their corresponding coefficient vectors  $(\mathbf{c}_{sj}, \mathbf{c}_{\psi j}, \mathbf{c}_{sj})$ .

In the same fashion we will expend  $g^{-1}$  to be

$$g^{-1}(t, \mathbf{q}) = \left( \begin{pmatrix} \tilde{s}_1(t) \\ \vdots \\ \tilde{s}_m(t) \end{pmatrix}, \begin{pmatrix} \tilde{\psi}_1(t) \\ \vdots \\ \tilde{\psi}_m(t) \end{pmatrix}, \begin{pmatrix} \tilde{s}_1(t) \\ \vdots \\ \tilde{s}_m(t) \end{pmatrix} \right) \in \mathbb{R}^{m \times 3} \quad (17)$$

resulting in three column vectors of B-splines evaluations at time  $t$  at all  $m$  observation points.

This spline-based latent space provides a compact representation of complex seismic cycle simulations. In the following, we demonstrate how this latent encoding can be leveraged to construct a ROM that accurately emulates the full-order simulations with orders of magnitude lower computational cost.

### 3.1.2 One SSE cycle reduced-order model

To construct a ROM for an individual SSE cycle, we adopt a strategy similar to the approach of Rekoske et al. (2025), originally developed for seismic wave propagation. Here, we extend this methodology to one cycle of SSE simulation.

Let us consider a collection of  $N$  full-order model simulations, each corresponding to a distinct parameter realization  $\boldsymbol{\xi}_1, \dots, \boldsymbol{\xi}_N$ . For a given SSE cycle  $k$ , these simulations yield full-order outputs  $\mathbf{Q}^k(\boldsymbol{\xi}_1), \dots, \mathbf{Q}^k(\boldsymbol{\xi}_N)$ . Using Eq. (15) we map each simulation output into its spline latent representation to form a matrix  $\mathbf{D} \in \mathbb{R}^{l \times N}$  (step 4 in Figure 2) given by

$$\mathbf{D} = \left( G(\mathbf{Q}^k(\boldsymbol{\xi}_1)) \dots G(\mathbf{Q}^k(\boldsymbol{\xi}_N)) \right) = \left( \mathbf{q}^k(\boldsymbol{\xi}_1) \dots \mathbf{q}^k(\boldsymbol{\xi}_N) \right), \quad (18)$$

where  $\mathbf{q}^k(\boldsymbol{\xi}_i)$  is the spline latent column vector representation of the  $i^{\text{th}}$  simulation in the  $k^{\text{th}}$  cycle. We then compute the Singular Value Decomposition (SVD) of  $\mathbf{D}$ :

$$\mathbf{D} = \mathbf{U} \boldsymbol{\Sigma} \mathbf{V}^T, \quad (19)$$

where  $\mathbf{U}$  contains the left singular vectors (basis functions),  $\boldsymbol{\Sigma}$  is a diagonal matrix of singular values  $\lambda_i$ , and  $\mathbf{V}$  contains the right singular vectors. The SVD provides an optimal low-rank approximation in the least-squares sense and allows each latent vector  $\mathbf{q}^k(\boldsymbol{\xi}_i)$  to be expressed as a linear combination of the orthonormal basis vectors in  $\mathbf{U}$ :

$$\mathbf{q}^k(\boldsymbol{\xi}_i) = \sum_{r=1}^N \lambda_r v_{ir} \mathbf{u}_r = \sum_{r=1}^N \alpha_r^i \mathbf{u}_r, \quad (20)$$

where  $\alpha_r^i = \lambda_r v_{ir}$  are the Proper Orthogonal Decomposition (POD) coefficients. This decomposition is consistent with viewing the SVD as a POD of the matrix  $\mathbf{D}$ , capturing the most energetic modes of variability across the simulations dense representation (step 5 in Figure 2, Berkooz et al., 1993; Bui-Thanh et al., 2003; Druault et al., 2005).

We adopt a radial basis functions (RBFs) interpolation strategy to perform a mapping from an arbitrary  $\boldsymbol{\xi}^* \in \mathcal{P}$  to approximate POD coefficients (e.g., Lazzaro & Montefusco, 2002; Audouze et al., 2009; Xiao et al., 2015; Rekoske et al., 2025). To avoid bias due to differing parameter scales, we first normalize the parameter space:

$$\begin{pmatrix} \bar{\xi}_1 \\ \vdots \\ \bar{\xi}_N \end{pmatrix} = \left[ \begin{pmatrix} \xi_1^1 & \dots & \xi_1^b \\ \vdots & \ddots & \vdots \\ \xi_N^1 & \dots & \xi_N^b \end{pmatrix} - (\bar{\xi}^1 \dots \bar{\xi}^b) \right] \oslash (\text{std}(\xi^1) \dots \text{std}(\xi^b)), \quad (21)$$

where  $\bar{\xi}^r$  and  $\text{std}(\xi^r)$  denote the mean and standard deviation of the  $r^{\text{th}}$  parameter across the  $N$  samples, and  $\oslash$  denotes element-wise division.

Given a new normalized parameter vector  $\bar{\xi}^*$ , the interpolated POD coefficient vector  $\alpha(\bar{\xi}^*)$  (step 7 in Figure 2) is expressed as:

$$\alpha(\bar{\xi}^*) = \sum_{i=1}^N w_i \varphi(\|\bar{\xi}^* - \bar{\xi}_i\|_2) + \sum_{r=1}^{|V|} b_r p_r(\bar{\xi}^*), \quad (22)$$

where  $w_i$  are the RBFs weights and  $\varphi(\cdot)$  is the RBF kernel defined in terms of the Euclidean distance ( $R$ ) between points. In this study, we consider the linear kernel  $\varphi(R) = R$ , which provides a simple yet effective means of interpolating in the reduced space (see Text S?? and Figure ??). In Eq. (22)  $b_r$  are polynomial weights, and  $p_r(\cdot)$  are the monomials of the input parameters up to degree  $d$ :

$$p_r(\bar{\xi}^*) = \xi_1^{a_1} \cdots \xi_{\dim(\mathcal{P})}^{a_{\dim(\mathcal{P})}}, \quad \text{for } a_1, \dots, a_{\dim(\mathcal{P})} \geq 0 \text{ and } \sum_{l=1}^{\dim(\mathcal{P})} a_l \leq d. \quad (23)$$

The interpolation weights are determined by solving the following linear system (Step 6 in Figure 2):

$$\begin{pmatrix} \Phi & \Lambda \\ \Lambda^T & \mathbf{0} \end{pmatrix} \begin{pmatrix} \mathbf{W} \\ \mathbf{B} \end{pmatrix} = \begin{pmatrix} \mathbf{F} \\ \mathbf{0} \end{pmatrix} \quad (24)$$

where

$$\Phi_{ij} = \varphi(\|\bar{\xi}_i - \bar{\xi}_j\|_2), \quad \Phi \in \mathbb{R}^{N \times N} \quad (25a)$$

$$F_{ij} = \alpha_i^j = \lambda_i v_{ji}, \quad \mathbf{F} \in \mathbb{R}^{N \times N} \quad (25b)$$

and  $\Lambda \in \mathbb{R}^{N \times |V|}$  is the matrix of monomial evaluations at each  $\bar{\xi}_i$ .

Following the calculation of the POD coefficients and the RBF interpolant, the coefficient corresponding to a new parameter vector  $\bar{\xi}^*$  can be retrieved by interpolating the POD coefficients corresponding to the training parameter realizations  $\bar{\xi}_1, \dots, \bar{\xi}_N$  used for the FOM simulations. To estimate a spline latent vector (step 8 in Figure 2) using the ROM scheme for a new normalized parameter vector  $\bar{\xi}^*$ , we perform

$$\mathbf{q}^k(\bar{\xi}^*) = \sum_{i=1}^N \alpha_i(\bar{\xi}^*) \mathbf{u}_i, \quad (26)$$

where  $\alpha_i(\bar{\xi}^*)$  is the  $i^{\text{th}}$  interpolated POD coefficient computed using Eq. (22).

The ROM-computed spline latent vector  $\mathbf{q}^k(\bar{\xi}^*)$  can then be mapped back to the full-order output space (step 9 in Figure 2) by applying the transformation  $G^{-1}$  given by Eq. (12) to yield:

$$\begin{aligned} \mathbf{Q}^k(\bar{\xi}^*) &= G^{-1}(\mathbf{t}^*, \mathbf{q}^k(\bar{\xi}^*)) \\ &= \left( \mathbf{t}^*, g^{-1}(t_1^*, \mathbf{q}^k(\bar{\xi}^*)), \dots, g^{-1}(t_k^*, \mathbf{q}^k(\bar{\xi}^*)) \right) \\ &= \left( \mathbf{t}^*, \dot{\mathbf{S}}^{k*}, \Psi^{k*}, \mathbf{S}^{k*} \right), \end{aligned} \quad (27)$$

where  $\mathbf{t}^* \in \mathbb{R}^{n^*}$  yielding  $m \times n^*$  matrices approximating the simulation output corresponding to the new parameter realization  $\bar{\xi}^*$  over the  $n^*$  length time vector  $\mathbf{t}^*$ .

### 3.2 Training and validation: Leave-one-out cross validation

A leave-one-out cross-validation (LOOCV) approach is employed to assess the accuracy of our ROM predictions (Rippa, 1999; Kohavi, 1995). Under this framework, each

parameter realization  $\xi_i$  is systematically excluded from the training set, the model is constructed using the remaining  $N - 1$  simulations, and prediction accuracy is evaluated against the withheld simulation. This process iterates across all  $N$  parameter realizations to obtain an unbiased estimate of generalization error.

For each  $\xi_i$  and each SSE cycle  $k$ , we consider the POD coefficient vector  $\alpha_r$  the  $r^{\text{th}}$  row of the  $\mathbf{F}$  matrix as our validation set and perform the procedure described in Eq. (22) to Eq. (27) with  $\bar{\xi}^* = \bar{\xi}_i$ . This yields the predicted coefficients  $\alpha_r(\bar{\xi}_i^*)$  and the corresponding ROM output  $\mathbf{Q}^k(\bar{\xi}_i^*)$  for the  $k^{\text{th}}$  cycle.

To assess the accuracy of the ROM predictions, we:

1. Compare the recurrence intervals of the full-order and ROM simulations using the average cycle durations:

$$T_c = \frac{1}{p} \sum_{k=1}^p (\max(t_k) - \min(t_k)), \quad t_k \in \mathbf{Q}^k(\xi_i), \quad (28)$$

and similarly for the predicted outputs:

$$T_c^* = \frac{1}{p} \sum_{k=1}^p (\max(t_k^*) - \min(t_k^*)), \quad t_k^* \in \mathbf{Q}^k(\bar{\xi}_i^*). \quad (29)$$

2. Compare the SSEs mean potency  $P_0$  (Eq. (6)). In a 3D context, the potency can be related to the seismic moment by  $M_0 = \mu P_0$ , where  $\mu$  is the shear modulus. In our 2D model the potency can be estimated without further assumptions where  $\int_A$  is a line integral, we can approximate the seismic moment by assuming some fault width of  $W_s$  km along strike, yielding  $M_0 = \mu W_s P_0$ .

## 4 Results

### 4.1 FOM results

We performed a total of 76 FOM evaluations using the computational resources of the Nautilus Kubernetes cluster of the National Research Platform. Our simulation suite was initiated with a relatively coarse sampling grid spanning  $W \in [30.5, 68]$  km and  $\sigma_W \in [1, 6]$  MPa. Subsequently, the sampling density was increased within the range  $W \in [30.5, 53]$  km, and further refined within  $W \in [35.5, 38]$  km. This adaptive refinement strategy aimed to improve the accuracy and robustness of the ROMs, particularly letting the leave-one-out cross-validation error estimation described in Section 3.2 stay within simulation  $T_c$  and  $P_0$  standard deviation.

The computational expense per simulation exhibited considerable variation, depending on the specific values chosen for the  $W$  and the  $\sigma_W$  within the localized subregion around the stability transition zone  $a-b=0$ . As is standard for seismic cycle simulations, an initial spin-up phase was removed to minimize the influence of initial conditions on the subsequent system time-dependence behavior (Rubin & Ampuero, 2005). The computation time required for this spin-up phase proved to be highly sensitive to the model parameters, particularly the normalized fault width

$$\frac{W}{h^*} = \frac{W\pi(1-\nu)(b-a)\sigma_W}{2\mu L}, \quad (30)$$

where  $h^*$  is the characteristic nucleation size,  $\nu$  is the Poisson ratio and  $\mu$  is the shear modulus (Table 1). This is consistent with the results of Liu and Rice (2009), who showed that lower  $W/h^*$  induces more frequent SSE oscillations, necessitating longer spin-up times to allow the system to reach a quasi-steady state before the analysis period.

We establish a uniform simulation duration cutoff of  $t = 75$  years for all model runs. The majority of the computational effort during each simulation was concentrated on resolving the fine-scale time-dependence associated with the SSEs themselves. Individual simulation run times ranged from 19.2 to 302.4 hours on 30 MPI ranks, corresponding to computational costs between 628 and 9125 CPU hours per simulation. The cumulative computational cost for executing the entire suite of 76 FOM simulations amounted to approximately 233,565 CPU hours.

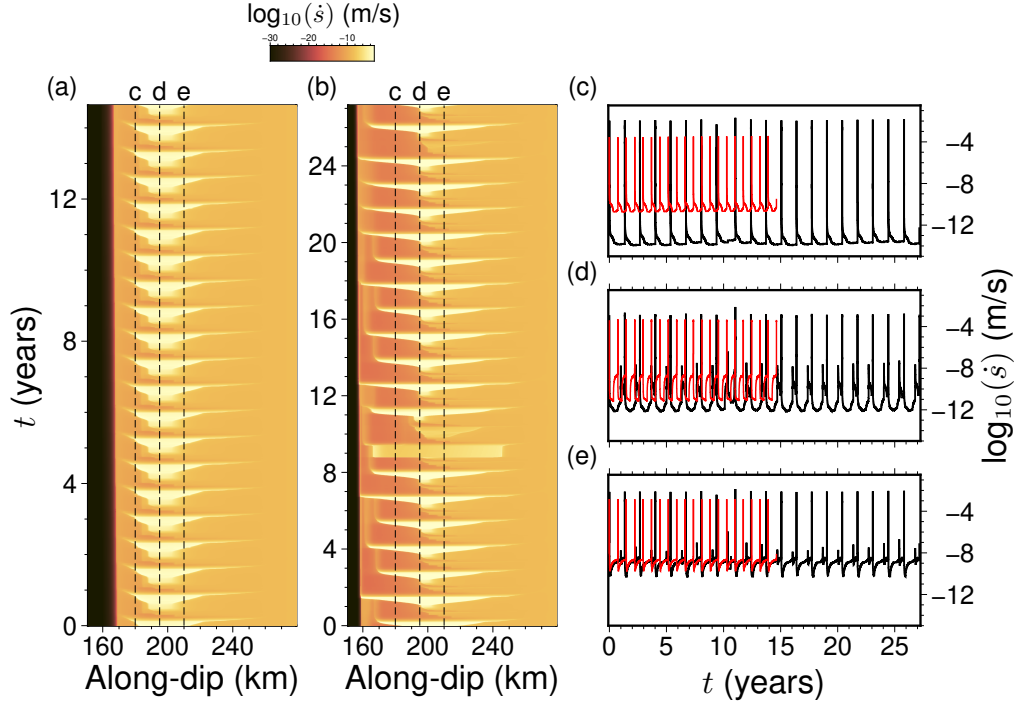
To illustrate the typical time-dependence observed in our FOM simulations, we first examine two representative cases: simulation  $\xi_1$  with parameters ( $W = 37.375$  km,  $\sigma_W = 2$  MPa) and simulation  $\xi_2$  with parameters ( $W = 48$  km,  $\sigma_W = 4$  MPa), yielding  $W/h^*$  values of 5.44 and 16.73, respectively (Figure 4). Both simulations exhibit quasi-periodic SSEs:  $\xi_1$  recurs approximately every  $T_c = 0.73 \pm 0.004$  years with peak slip velocities  $V_{max} \approx 10^{-2.7}$  m/s and accumulating a mean total slip of  $\delta = 0.98 \pm 3.4 \times 10^{-5}$  cm per event, while  $\xi_2$  shows  $T_c = 1.36 \pm 0.07$  years,  $V_{max} \approx 10^{-1.6}$  m/s, and  $\delta = 2.7 \pm 8.8 \times 10^{-3}$  cm.

Analyzing these simulations in their phase-space (Figure 3) shows a simplified view of the underlying rate-and-state friction characteristics. For both cases, as an SSE nucleates, the representative phase-space trajectory rapidly transitions towards higher slip velocities. It then evolves through a characteristic loop, involving stages of rapid state evolution and slip-rate changes that reflect the weakening and subsequent healing phases dictated by Eq. (2). The cycle is completed by the slow evolution during the long inter-seismic period, which, although dominant in the time-domain, represents a relatively small portion of the path length traced in the phase-space during one full cycle. This confirms the utility of the phase-space perspective for capturing SEAS cycles and highlighting the periods of rapid dynamic change during the events themselves, across different parameter regimes.

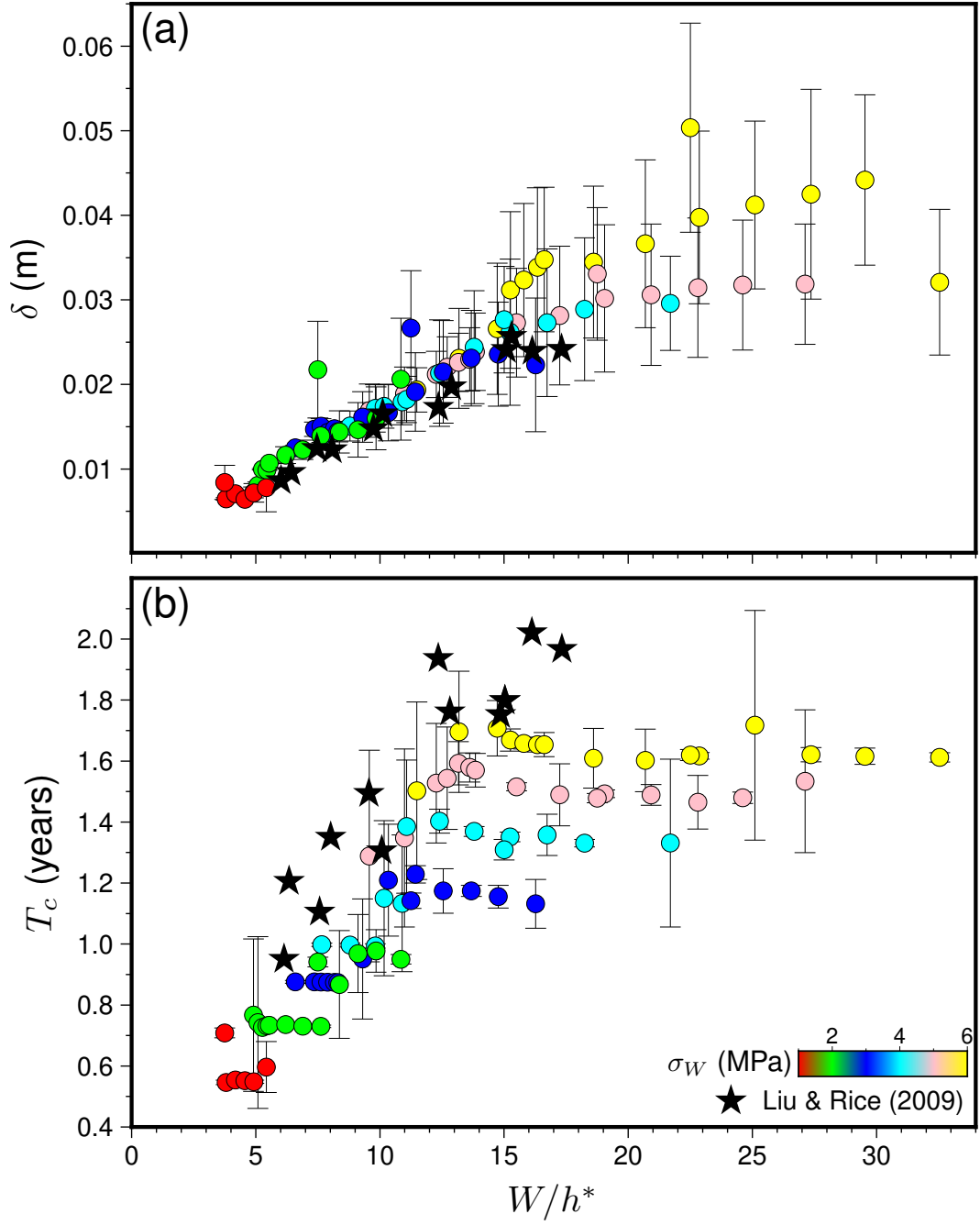
Figure 5 presents the resulting recurrence time ( $T_c$ ) and average SSE slip ( $\delta$ ) as a function of the normalized fault width ( $W/h^*$ ) for all 76 FOM simulations, juxtaposed with the results reported by Liu and Rice (2009) for gabbro rheology. Notably, our results, obtained using the volumetric discontinuous Galerkin finite element code tandem (Uphoff et al., 2022), show excellent agreement with those of Liu and Rice (2009), which were generated using a boundary element method.

The simulations reveal that spontaneous, quasi-periodic SSEs are generated over a wide range of the explored parameter space, corresponding to  $W/h^*$  values ranging from 3.75 to 32.54. For  $W/h^*$  values smaller than 3.75, these simulation settings do not produce distinct SSEs; the system either evolves towards steady creep or exhibits very fast recurrence intervals ( $T_c \ll 1$  year) with the slip-rate oscillating around the tectonic loading rate. Within the approximate range  $3.75 < W/h^* < 18$ , our findings are consistent with Liu and Rice (2009), we observe clear trends where both the recurrence time ( $T_c$ ) and the mean slip ( $\delta$ ) appear to increase approximately linearly with  $W/h^*$ . However, for  $W/h^* \gtrsim 18$ , we observe a deviation from this linear trend, particularly for  $T_c$ , which tends to slightly decrease and then plateau. The specific  $W/h^*$  value where this transition begins appears influenced by  $\sigma_W$ , for instance, the deviation becomes noticeable around  $W/h^* \approx 11$  for  $\sigma_W = 3$  MPa, but closer to  $W/h^* \approx 15$  for  $\sigma_W = 6$  MPa. In contrast, the mean accumulated slip in those events ( $\delta$ ) seems to maintain an approximately linear dependence on  $W/h^*$  across the entire range where SSEs are observed.

These results indicate a complex dependency between the SSE time-dependence behavior and the model parameters  $W, \sigma_W$ . Although the normalized fault width ( $W/h^*$ ) controls the time-dependence behavior at first order, our results suggest second-order independent dependencies on both  $W$  and  $\sigma_W$ , which we will explore further using our ROM approach.



**Figure 4.** FOM results for two parameter vectors:  $\xi_1 = (W = 37.375 \text{ km}, \sigma_W = 2 \text{ MPa})$  and  $\xi_2 = (W = 48 \text{ km}, \sigma_W = 4 \text{ MPa})$ . Each simulation presents 20 SSE cycles, with  $\xi_1$  spanning a shorter time period due to its reduced recurrence interval. (a) Slip-rate as a function of time and along-dip distance for  $\xi_1$ . The locked portion of the subduction interface is not shown. (b) Equivalent slip-rate representation for  $\xi_2$ . (c)-(e) Temporal profiles of slip-rate at specific observation points along the fault (indicated by dashed lines in panels (a) and (b)), with  $\xi_1$  shown by the red curve and  $\xi_2$  by the black curve.



**Figure 5.** FOM results for key SSE characteristics as a function of normalized fault width ( $W/h^*$ ). Panel (a) shows the mean SSE slip ( $\delta$ ) occurring within the low effective normal stress zone versus  $W/h^*$ . Panel (b) depicts the SSE recurrence time ( $T_c$ ) versus  $W/h^*$ . Our FOM results using tandem (Uphoff et al., 2022) (coloured circles) are compared with the BEM results reported in Liu and Rice (2009) (black stars). The volumetric based FOM results obtained from tandem exhibit good agreement with those of Liu and Rice (2009) in their common range of  $W/h^*$ .

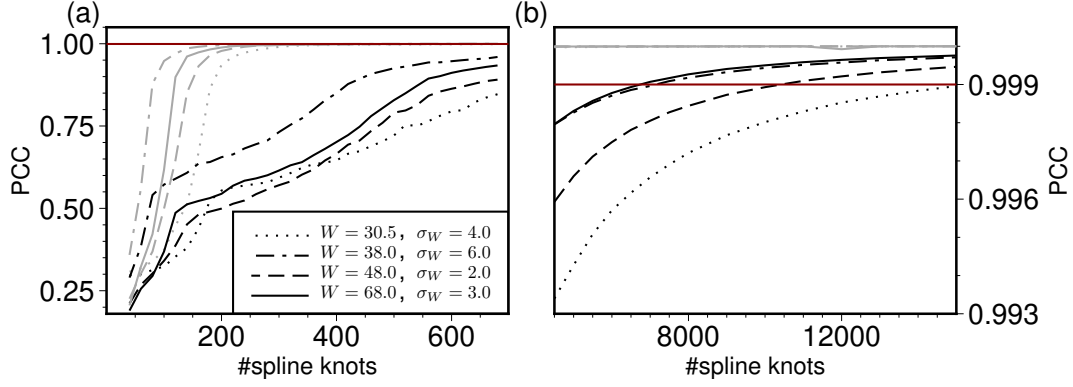
## 4.2 Latent space representation accuracy analysis

The primary goal of employing the spline latent space representation, as detailed in Section 3.1.1, is to achieve a substantial compression of the complex FOM simulation data while preserving the essential characteristics, a necessary requirement to construct an accurate reduced-order model. As our chosen representation employs lossy compression, it is important to quantify the quality of this representation by evaluating how accurately the original simulation variables can be reconstructed from the latent vector  $\mathbf{q}$ . Following Eq. (7), we assess this accuracy primarily using the Pearson Correlation Coefficient (PCC) between the original FOM slip-rate time series  $\hat{\mathbf{S}}$  within the low effective normal stress zone  $W$  and the corresponding reconstruction  $\hat{\mathbf{S}}^*$  obtained via the inverse transformation  $G^{-1}(\mathbf{t}, \mathbf{q})$ . We aim to ensure the PCC remains above a threshold,  $1 - \epsilon$  where  $\epsilon = 0.001$ . The trade-off lies in minimizing the dimensionality  $l$  of the latent vector  $\mathbf{q}$  (i.e., minimizing the number of spline knots and coefficients) while maintaining this high level of reconstruction accuracy.

Figure 6 provides a quantitative comparison validating the efficiency of representing the slip-rate ( $\dot{s}$ ) using our proposed phase domain spline approach versus a conventional time-domain spline, illustrated using four different FOM parameter pairs. The plot shows the number of spline knots required for the B-spline representation to achieve a PCC exceeding 0.999 when reconstructing the slip-rate time series. This comparison considers the full simulation duration after the initial spin-up period, without constraining the number of SSE cycles included. As clearly demonstrated, splines parameterized by the phase progression variable  $\phi$  consistently require considerably fewer spline knots to reach this high accuracy threshold compared to splines based directly on the time-domain. The four example FOM simulations contain approximately  $2 \times 10^5$  to  $4 \times 10^5$  time steps, requiring only 400 - 900 spline knots for satisfactory representation using the phase trajectory approach, compared to 7000 - 15000 spline knots needed for the time-domain approach. This empirically confirms that the simulation characteristics can be represented much more efficiently in the phase-space.

Building on the now established efficiency of phase-space parameterization, Figure 7 further assesses the performance of our spline latent space representation in reconstructing detailed simulation outputs. Although the figure presents the reconstruction of slip-rate time series along one observation point (195 km along fault), the reconstruction PCC score is calculated over all observation points within  $W$ . For a lengthy 33-cycle SSE simulation, whose original outputs  $(\hat{\mathbf{t}}, \hat{\Phi}, \hat{\mathbf{S}}, \hat{\Psi}, \hat{\mathbf{S}})$  comprise  $1.8 \times 10^8$  floating-point numbers, a clear relationship is observed between reconstruction fidelity and the number of spline knots utilized. With the ratio of knots fixed at  $K_1 = 0.1K_0$ , which was found to be adequate based on trial and error, a representation with  $K_0 = 1000$  spline knots per observation point (illustrated by the blue curve in Figure 7(a)) reproduces the general characteristics of the slip-rate waveform at a specific observation point, yet it fails to capture crucial details such as slip-rate peaks and performs poorly during the inter-seismic period.

The target reconstruction accuracy, defined by a PCC score exceeding  $1 - \epsilon$  (where  $\epsilon = 0.001$ ), is achieved with  $K_0 = 12000$  spline knots (red curve, Figure 7(a)). This results in a latent representation size of  $l \approx 4 \times 10^6$  floating-point numbers, corresponding to a 97.5% data compression. The performance of this latent space representation was also evaluated for individual SSE cycles (Figure 7(c)-(j)), which are fundamental to our ROM scheme (Section 3.1.2), across a range of model parameters ( $\xi_1$  through  $\xi_4$ ). The original output data for these single-cycle simulations range from  $4 \times 10^6$  to  $7 \times 10^6$  floating-point numbers. For these single-cycle cases, the target PCC threshold is met when the number of spline knots  $K_0$  equals 250, 300, 450, and 350 per observation point for simulations using  $\xi_1 = (30.5 \text{ km}, 4 \text{ MPa})$ ,  $\xi_2 = (38 \text{ km}, 6 \text{ MPa})$ ,  $\xi_3 = (48 \text{ km}, 2 \text{ MPa})$  and  $\xi_4 = (68 \text{ km}, 3 \text{ MPa})$ , respectively. This level of reconstruction yields data com-



**Figure 6.** Comparison of spline interpolation accuracy when the slip-rate ( $\dot{s}$ ) is represented in time-domain and phase-space. The plot demonstrates the number of spline knots required to achieve a PCC exceeding 0.999 (red horizontal line) when reconstructing the slip-rate. The phase-space based spline (gray lines) computed via  $\mathcal{B}^{-1}(\mathbf{k}_\phi, \mathbf{c}_s, \tilde{\phi}(t))$  (see Section 3.1.1, Eq. (13)) consistently requires considerably fewer spline knots than time-domain representation (black lines, computed via  $\mathcal{B}^{-1}(\mathbf{k}_t, \mathbf{c}_s, t)$ ), demonstrating the efficiency of phase-space representation of the SSE FOM simulation output. Panel (a) shows the PCC for spline calculated with number of spline knots  $< 700$ , while panel (b) illustrates the performance difference for spline calculated with number of spline knots  $> 5000$ . In the case of the gray lines, the number of knots corresponds to  $K_1$ .

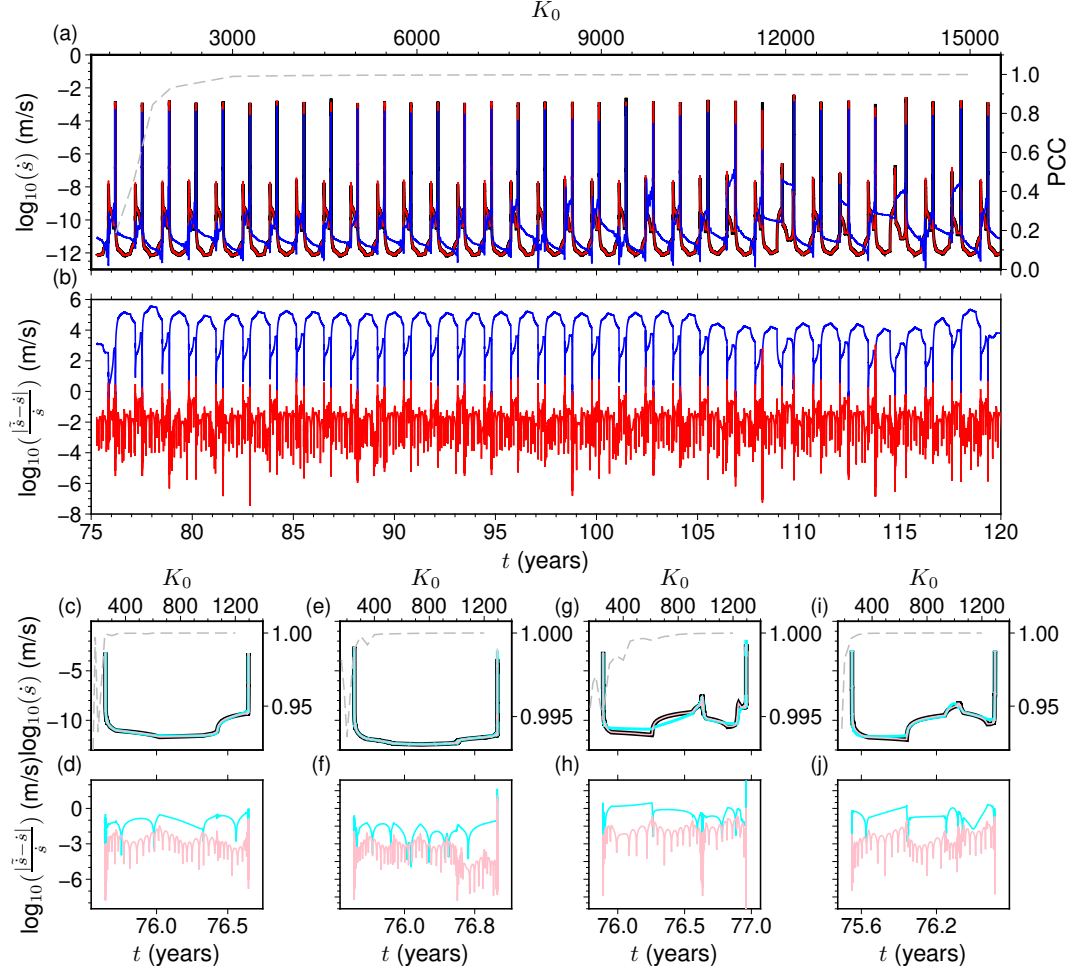
pression ratios between 97.3% and 97.8%, comparable to the multi-cycle simulation reconstruction.

Throughout the remainder of this work, we construct ROMs using  $K_0 = 800$  and  $K_1 = 80$  spline knots for the time to phase-progression ( $\mathbf{k}_t$ ) and the phase to physical variable ( $\mathbf{k}_\phi$ ) mapping respectively. This choice ensures that reconstruction fidelity comfortably exceeds the desired threshold while still achieving an approximate 95% reduction in data size compared to the original FOM output. With  $m = 346$  fault observation points, this results in a total latent vector length of  $l = 388,320$  for each SSE cycle simulation (Eq. (16)).

### 4.3 ROM results

The objective of developing the ROM is to accelerate the inference of SSE cycles over the parameter space defined by the width of the low effective normal stress zone ( $W$ ) and its magnitude ( $\sigma_W$ ). This section details the performance and accuracy of the constructed ROM.

The development of the ROM involves an initial offline phase, which includes running the 76 FOM simulations to generate the training dataset, transforming these simulations into their spline latent space representations, performing POD on the matrix of latent vectors  $\mathbf{D}$  for each SSE cycle, and constructing the RBF interpolators for the POD coefficients (Figure 2 - offline computations). The generation of the 76 FOM simulations incurred a cumulative cost of approximately 233,565 CPU hours (Section 4.1). The subsequent offline computations for the ROM construction, of calculating spline coefficients in order to transform the FOM to their latent space require an additional  $\sim 360$  CPU hours all other offline steps have negligible run time compare to the first two.



**Figure 7.** Accuracy of the spline latent space reconstruction of the FOM slip-rate  $\dot{s}(t)$ . Panels (a) and (b) analyze the spline accuracy when applied to an entire time series consisting of 33-cycle SSEs simulation with  $\xi = (50.5 \text{ km}, 4 \text{ MPa})$ . Panel (a): Slip-rate at an along-dip of 195 km for the FOM (black line); spline reconstruction using  $K_0 = 1000$  spline knots (blue line) and  $K_0 = 12000$  spline knots (red line). The gray dashed line shows the PCC (right y-axis) as a function of the number of spline knots (top x-axis). Note that the PCC is calculated for all observation points within  $W$ . Panel (b): Reconstruction errors,  $|\tilde{s}(t) - \dot{s}(t)|/\dot{s}(t)$  when using  $K_0 = 1000$  (blue) and  $K_0 = 12000$  (red) splines knots. Panels (c), (e), (g), (i): Spline reconstruction for slip-rate when applied to a single SSE cycle using  $K_0 = 200$  spline knots (cyan) and  $K_0 = 800$  spline knots (pink) with  $\xi$  given by (30.5 km, 4 MPa), (38 km, 6 MPa), (48 km, 2 MPa) and (68 km, 3 MPa) respectively. Panels (d), (f), (h), (j): Spline reconstruction errors associated with (c), (e), (g), (i) respectively.

Once all the offline tasks are complete, evaluating the ROM for a new parameter vector  $\xi^*$  (online task) is very fast. A single ROM evaluation, which involves interpolating the POD coefficients using the RBFs (Eq. (22)), reconstructing the latent vector  $q^*$  (Eq. (26)), and then transforming it back to the physical space using the inverse spline transformation  $G^{-1}$  (Eq. (27)) to obtain the time series for slip-rate, state variable, and cumulative slip takes approximately 30 seconds with one CPU. This represents a speedup of  $3.68 \times 10^5$  compared to the average FOM runtime of  $\sim 3000$  CPUh.

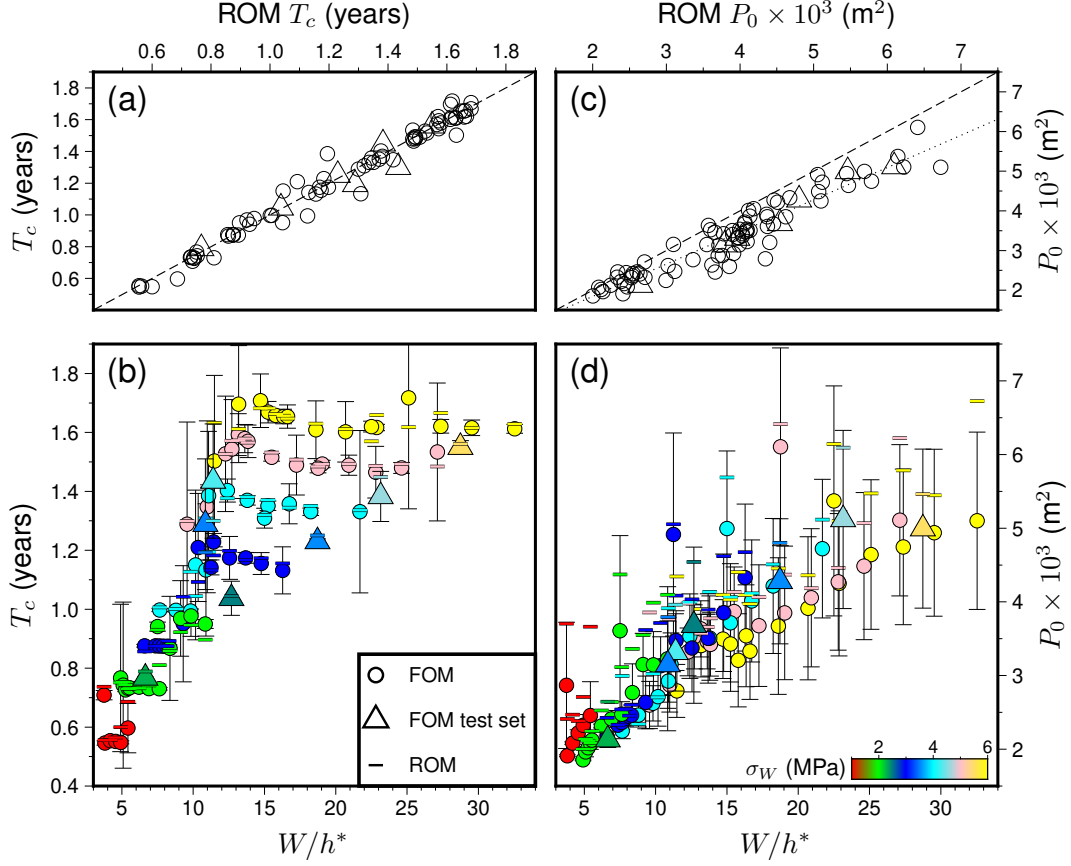
The accuracy of the ROM was systematically assessed using the LOOCV procedure detailed in Section 3.2. In this process, for each of the 76 FOM simulations, the ROM was trained on the remaining 75 simulations, and its predictions for the held-out parameter set were compared against the actual FOM results. Figure 8 illustrates the ROM's capability in reproducing key physical characteristics of SSEs, namely the recurrence time  $T_c$  and potency  $P_0$ . For recurrence time, the ROM's predictions largely fall within the FOM's cycle-to-cycle standard deviation, with only one exception among the 76 cases. The mean absolute difference between FOM and ROM  $T_c$  was 11.56 days, relative to an average SSE recurrence time of 440 days for the dataset. Consequently, the  $R^2$  score comparing ROM-predicted  $T_c$  to FOM  $T_c$  is 0.98, indicating a strong linear relationship and a near one-to-one correspondence.

For the potency, the majority of ROM predictions also align with the FOM results within the FOM's own standard deviation, which, as shown in Figure 8(d), can exhibit considerable cycle-to-cycle variability. However, notable discrepancies between FOM and ROM potency were observed for a few parameter vectors, specifically  $\xi = (68 \text{ km}, 6 \text{ MPa})$ ,  $\xi = (60.5 \text{ km}, 6 \text{ MPa})$ ,  $\xi = (30.5 \text{ km}, 4 \text{ MPa})$ , and  $\xi = (30.5 \text{ km}, 6 \text{ MPa})$ . These cases generally lie at the periphery of our sampled parameter space, often near corners where the interpolation scheme has limited surrounding data. The ROM appears to exhibit a systematic overestimation of potency. While the  $R^2$  score for ROM predicted  $P_0$  versus FOM  $P_0$  is 0.59, on the other hand the PCC between the two datasets is substantially higher at 0.946, suggesting a strong underlying correlation despite the offset. This systematic overestimation, further discussed in Section 5.5, though not ideal, might be addressable through post-processing corrections.

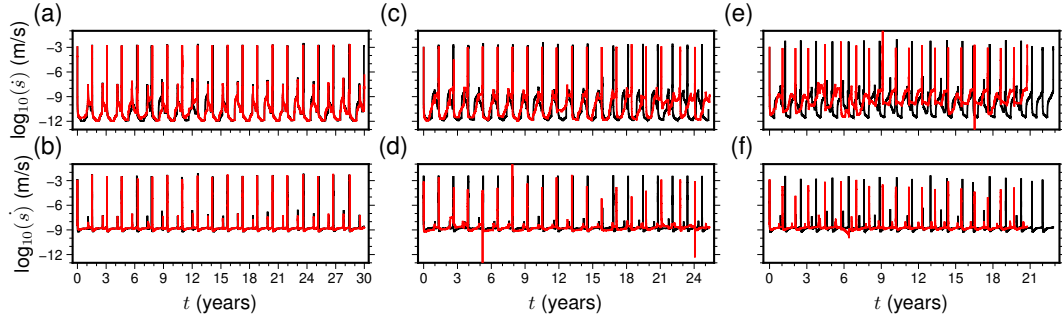
To further illustrate the ROM's predictive capabilities, Figures 9, ??, ?? show direct comparisons of the slip-rate, cumulative slip, and state variable evolution at two specific fault observation points: 195 km and 220 km along dip. These locations correspond to a VW region within  $W$  and the stability transition zone ( $a - b = 0$ ), respectively. These predictive capabilities are presented for three distinct parameter sets:  $\xi_1 = (W = 37.375 \text{ km}, \sigma_W = 5 \text{ MPa})$ , which yielded among the best prediction results in the LOOCV;  $\xi_2 = (W = 60.5 \text{ km}, \sigma_W = 4 \text{ MPa})$ , which showed median prediction accuracy; and  $\xi_3 = (W = 43.0 \text{ km}, \sigma_W = 3 \text{ MPa})$ , which represents the cases with the largest LOOCV discrepancies.

For both  $\xi_1$  and  $\xi_2$ , the ROM mostly captures the overall SSEs timeseries characteristics successfully, including peak slip-rates and recurrence times. For  $\xi_3$ , the recurrence time prediction is less accurate; although the mean recurrence time predicted by the ROM remains within one standard deviation of the FOM's mean recurrence time, the difference between the means is approximately two months, which is comparable to the cycle-to-cycle standard deviation observed in the FOM for this parameter set. Nevertheless, the prediction of event potency for  $\xi_3$  remains reasonable, with the FOM yielding an average potency of  $3153 \text{ m}^2$  and the ROM predicting  $3097 \text{ m}^2$ .

In machine learning applications, it is common practice to employ a validation set for hyperparameter tuning and model construction, and a separate test set for an unbiased evaluation of the finalized model (Kohavi, 1995). While we utilized an LOOCV scheme for validation (Section 3.2), we also established an independent test dataset to further assess our ROM. For this purpose, an additional seven FOM simulations were



**Figure 8.** Results of the LOOCV test assessing ROM performance. (a) Comparison of FOM results versus ROM-predicted recurrence time ( $T_c$ ). The dashed line indicates a 1:1 ratio. (b) Characteristic recurrence time ( $T_c$ ) as a function of normalized fault width ( $W/h^*$ ). Vertical bars represent the standard deviation derived of the FOM results, and colored horizontal bars show the corresponding ROM-calculated values. (c) Comparison of FOM results versus ROM-predicted potency ( $P_0$ ). The dashed line indicates a 1:1 ratio and the dotted line shows the best linear fit between the ROM and FOM results. (d) Potency ( $P_0$ ) as a function of normalized fault width ( $W/h^*$ ). Vertical bars represent the standard deviation of the FOM results, and colored horizontal bars indicate the ROM-calculated values.



**Figure 9.** LOOCV comparison of slip-rate predictions from the ROM (red curves) versus those from the FOM (black curves). Results are displayed for three different parameter sets at two along-dip observation points: 195 km (panels (a), (c), (e)) and 220 km (panels (b), (d), (f)). The parameters ( $W, \sigma_W$ ) used were:  $\xi_1 = (37.375 \text{ km}, 5 \text{ MPa})$  ((a)-(b));  $\xi_2 = (60.5 \text{ km}, 4 \text{ MPa})$  ((c)-(d));  $\xi_3 = (43.0 \text{ km}, 3 \text{ MPa})$  ((e)-(f)).

run, distributed across our parametric domain. These simulations were not used during the ROM construction process.

The test dataset yielded prediction results for  $T_c$  and  $P_0$  that were similar to those obtained from the LOOCV procedure (Figure 8), achieving  $R^2$  scores of 0.92 and 0.58, respectively. The test dataset results for  $P_0$  also showed the systematic overestimation previously observed in the LOOCV analysis, although the predicted potency maintained a strong linear correlation with the simulated potency, evidenced by a PCC of 0.98. The availability of this test dataset also facilitated the evaluation of a post-processing correction for the potency overestimation. The corrected potency,  $P_0^c$ , is calculated as:

$$P_0^c = (m - 1)P_0 - n, \quad (31)$$

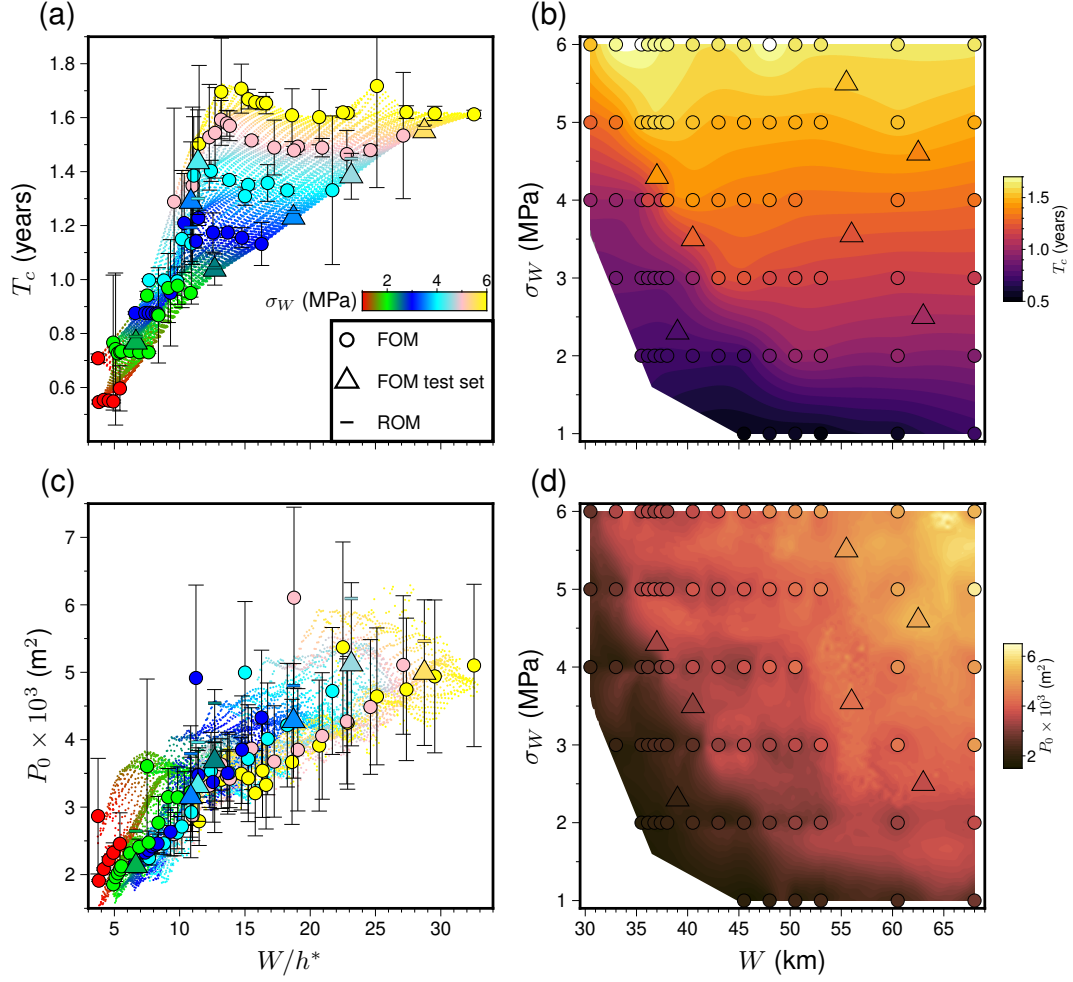
where  $m$  and  $n$  are the coefficients of the first-degree polynomial that best fits the potency predictions from the LOOCV analysis. Applying this correction to the potency values in the test set improved the  $R^2$  score to 0.95 (Figure ??).

The computational efficiency and demonstrated accuracy of the ROM facilitate a detailed exploration of the parameter space. Figure 10 presents the ROM predictions for  $T_c$  and the corrected  $P_0$  on a finely sampled grid of  $W$  and  $\sigma_W$ . We initially sampled the parametric space uniformly with 10000  $P^*$  pairs, subsequently removing pairs that fell outside the convex hull of the FOM training parameters (illustrated as white space in the bottom left corner of Figure 10), which resulted in a total of 9116 ROM evaluations. The ROM smoothly interpolates between the FOM training points, revealing complex dependencies that might be overlooked with coarser parameter space sampling. For instance, analyzing the FOM results solely as a function of  $W/h^*$  might suggest that for normalized fault lengths in the range of 4–15, SSE recurrence times increase linearly, and then gradually approach a constant for  $W/h^* > 15$ . In contrast, the ROM’s dense parametric view of  $T_c$  as a function of both  $W$  and  $\sigma_W$  (Figure 10(b)) shows a more complex picture. A diagonal band, roughly delineated by the points ( $W = 53$  km,  $\sigma_W = 1$  MPa) and ( $W = 30.5$  km,  $\sigma_W = 6$  MPa), exhibits a high gradient, indicating rapid changes in recurrence time over short parametric distances. To the upper-right and lower-left of this band, smaller gradients are observed, with  $T_c$  variations appearing to be predominantly influenced by changes in effective normal stress. This detailed mapping of SSE characteristics as a function of fault properties is critical for understanding the underlying physics and for constraining these parameters against geodetic observations. The construction of these high-resolution parameter maps using the ROM required only 75 CPU hours, in stark contrast to the estimated  $3 \times 10^7$  CPU hours that would have been necessary if one exclusively used a FOM.

#### 4.4 Uncertainty quantification of width and amplitude of low effective normal stress regions governing slow slip events

The extensive parameter space exploration detailed in Section 4.3, while showcasing the ROM’s efficiency, could theoretically be achieved with FOMs, if sufficient parallel computing resources were available, although at a vastly greater cost. However, certain tasks in model-based inference, such as global optimization or Bayesian parameter estimation via Markov Chain Monte Carlo (MCMC) methods, are inherently sequential or have limited parallelizability. For such methodologies, the computational cost of FOMs renders them practically infeasible. The rapid evaluation capabilities of our ROM framework, on the other hand, opens the door to utilize these techniques, allowing for robust uncertainty quantification of model parameters based on observations.

Here, we describe how the ROM can be employed within an MCMC framework to invert for the uncertainties in the fault parameters  $W$  (width of the low effective normal stress zone) and  $\sigma_W$  (magnitude of low effective normal stress), constrained by observed characteristics of Cascadia SSEs. We will show that in our models the recurrence



**Figure 10.** ROM predictions for characteristic properties of SSEs. (a) Recurrence time ( $T_c$ ) as a function of the normalized fault width ( $W/h^*$ ). (b) Recurrence time ( $T_c$ ) as a function of the width ( $W$ , x-axis) and magnitude ( $\sigma_W$ , y-axis) of the low effective normal stress zone. This panel highlights complex dependencies, such as a diagonal band of high  $T_c$  gradient (c) Corrected potency ( $P_0$ ) as a function of the normalized fault width ( $W/h^*$ ). (d) Corrected potency ( $P_0$ ) as a function of  $W$  (x-axis) and  $\sigma_W$  (y-axis). The uncorrected potency results are presented in Figure ??.

interval of SSEs is mainly controlled by the magnitude of effective normal stress, whereas their magnitudes primarily depend on the width of the low-stress region.

We employ the Metropolis-Hastings MCMC algorithm to sample the posterior probability distributions for  $W$  and  $\sigma_W$ . The observational constraints derived from Cascadia SSE studies are:

1. The mean recurrence interval  $\mu_{T_c}^{obs} = 1.17$  years (14 months) with a standard deviation  $\sigma_{T_c}^{obs} = 0.17$  years (2 months), assuming a normal distribution (Schmidt & Gao, 2010; Gombert et al., 2016).
2. The seismic moment  $M_0$  is assumed to follow a normal distribution with mean  $\mu_{M_0}^{obs} = 7.28 \times 10^{18}$  Nm (corresponding to  $M_w \approx 6.45$ ) and standard deviation  $\sigma_{M_0}^{obs} = 4.22 \times 10^{18}$  Nm, encompassing the typical range of  $M_w \sim 6.2 - 6.7$  for Cascadia SSEs (Behr & Bürgmann, 2021; Schmidt & Gao, 2010).

For given parameter vector  $\xi = (W, \sigma_W)$ , the ROM can be used to compute  $T_c^{ROM}$  and  $P_0^{ROM}$  (in units of  $m^2$ ). The ROM potency is corrected by the best fitted linear relation to 1 to 1 relation correction (Figure 8) and then converted to seismic moment using  $M_0^{ROM} = \mu W_s P_0^{ROM}$ , where  $\mu$  is the shear modulus (Table 1) and  $W_s$  is the assumed along-strike width of 60 km.

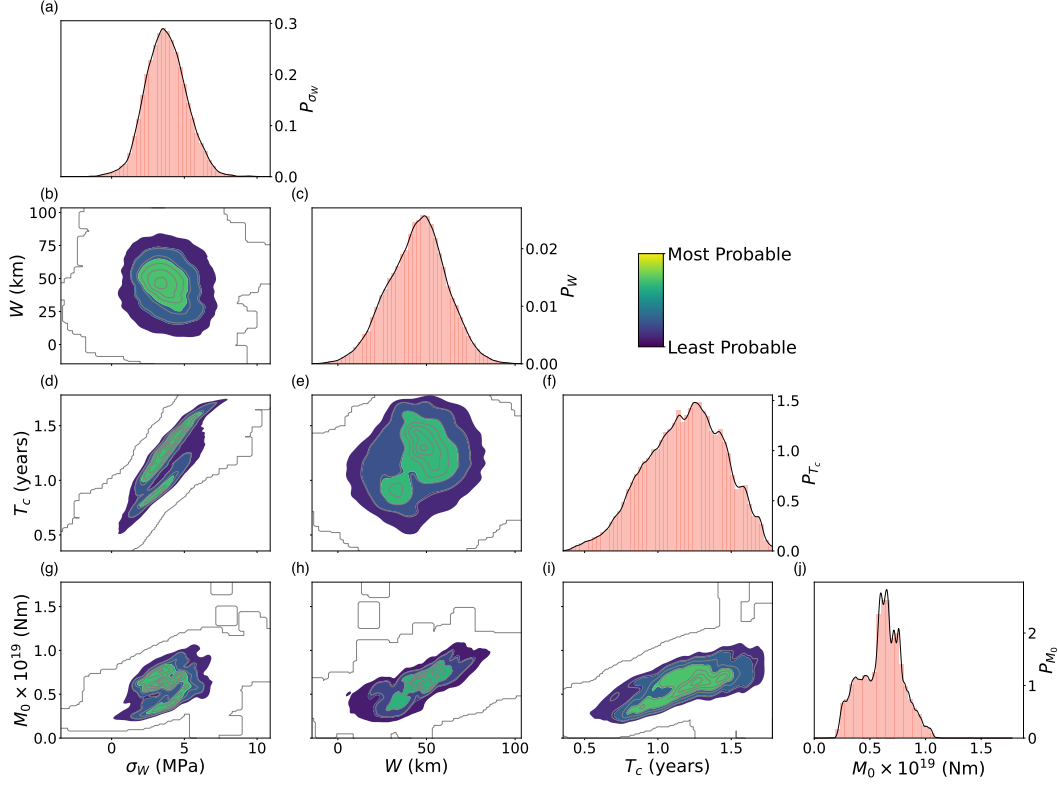
The likelihood function  $\mathcal{L}(\mathbf{D}^{obs} | (W, \sigma_W))$  for a proposed parameter vector, given the observed data  $\mathbf{D}^{obs} = (T_c^{obs}, \log_{10}(M_0^{obs}))$ , is:

$$\mathcal{L} \propto \exp \left( -\frac{1}{2} \left[ \left( \frac{T_c^{ROM} - \mu_{T_c}^{obs}}{\sigma_{T_c}^{obs}} \right)^2 + \left( \frac{\log_{10}(M_0^{ROM}) - \mu_{\log_{10}(M_0)}^{obs}}{\sigma_{\log_{10}(M_0)}^{obs}} \right)^2 \right] \right). \quad (32)$$

We initiate the chains with a uniform prior distributions to  $W$  and  $\sigma_W$  over the ranges  $W \in [30.5, 68]$  km and  $\sigma_W \in [1, 6]$  MPa. Then the chains are propagated with the Metropolis-Hastings algorithm (Hastings, 1970) which iteratively proposes new parameter vector  $(W_{i+1}, \sigma_{W,i+1})$  from the current state  $(W_i, \sigma_{W,i})$ . The ROM is evaluated at the proposed state, the likelihood  $\mathcal{L}_{i+1}$  is computed, and the proposal is accepted with probability  $\alpha = \min \left( 1, \frac{\mathcal{L}_{i+1} \cdot \text{Prior}_{i+1}}{\mathcal{L}_i \cdot \text{Prior}_i} \right)$ .

To sample the posterior distributions, we ran 10 parallel chains, each for 6000 iterations (including a 600 burn-in iteration) requiring approximately 50 hours of computation. This extent of MCMC analysis is computationally tractable only due to the ROM's efficiency. Relying on the FOM would be prohibitive; the sequential nature of each chain, requiring 6000 model evaluations, would translate to an estimated 50 years of computation time per chain, in addition to the general immense total computational effort required for all  $10 \times 6000$  FOM evaluations.

The convergence of the MCMC inversion was monitored using diagnostics presented in Figure ???. We employed the potential scale reduction factor,  $\hat{R}$ , which compares variance between chains to variance within each chain, values approaching 1 indicate convergence to a common target distribution (Vehtari et al., 2021). We also calculated the Effective Sample Size (ESS) to quantify the number of independent samples in the correlated MCMC output, crucial for reliable posterior inference (Gelman et al., 1995). Figure ??? indicates that  $\hat{R}$  values for model parameters generally fell below 1.01 after approximately 4000 total MCMC evaluations (summed across all 10 chains, not including their burn-in periods). The ESS for key parameters typically surpassed 200, a level often considered sufficient for robust estimation of posterior means and standard deviations (Gelman et al., 1995), after approximately 8000 total MCMC evaluations. Attaining these levels of convergence and sample independence, corresponding to thousands of individual model evaluations, is babyhood what is feasible with FOM simulations.



**Figure 11.** Posterior probability distributions and correlations for fault parameters characteristic properties of SSEs. The parameters shown are: the magnitude of low effective normal stress ( $\sigma_W$ ), the width of the low effective normal stress zone ( $W$ ), the recurrence interval ( $T_c$ ), and the seismic moment ( $M_0$ ). Diagonal plots (panels (a), (c), (f), (j)) display the marginal posterior probability distributions for each parameter. The y-axis of each marginal plot indicates the probability density. Off-diagonal plots (panels (b), (d), (e), (g), (h), (i)) illustrate the bivariate joint posterior distributions for pairs of parameters. For example, panel (b) shows the joint posterior of  $\sigma_W$  and  $W$ .

The MCMC analysis, informed by the observed Cascadia SSE recurrence intervals and seismic moments, yields posterior probability distributions for the fault parameters  $W$  and  $\sigma_W$ , visualized in Figure 11. The posterior distribution for the width of the low effective normal stress zone,  $W$ , is characterized by a mean of 44.7 km and a standard deviation of 16.2 km. For the magnitude of the low effective normal stress,  $\sigma_W$ , the inferred posterior has a mean of 3.8 MPa and a standard deviation of 1.4 MPa. These values represent the constrained estimates and associated uncertainties for these parameters, conditional on the observational data and the physics assumed by our model.

Analysis of the relationships within the posterior samples (Figure 11) further illuminates the control of these parameters on SSE characteristics. The SSE recurrence interval ( $T_c$ ) exhibits a strong dependence on  $\sigma_W$ , with a PCC between their posterior samples of 0.86, whereas its correlation with  $W$  is considerably weaker (PCC = 0.2). Conversely, the seismic moment ( $M_0$ ) is primarily correlated with  $W$  (PCC = 0.75), and shows a more moderate correlation with  $\sigma_W$  (PCC = 0.45). These findings suggest that, within our model framework for Cascadia-like SSEs, the magnitude of effective normal stress predominantly governs the timing of SSEs, while the spatial extent of this low-stress region is the primary factor controlling their magnitude.

## 5 Discussion

### 5.1 Implications for the state of stress and pore fluid pressure in Cascadia

Our findings support the critical role of low effective normal stress, e.g., maintained by high pore fluid pressure (e.g.  $\beta$ ,  $\beta$ ) (Behr, 2021), in enabling and modulating SSEs at the CSZ. Our MCMC inversion, constrained by observed Cascadia SSE characteristics, suggests that these events are consistent with low effective normal stress of  $3.8 \pm 1.4$  MPa. This aligns well with conditions hypothesized for SSE generation, which necessitate near-lithostatic pore fluid pressures at the depths where Cascadia SSEs typically occur. Such low effective normal stress is consistent with the findings of Audet and Kim (2016), who highlighted near-lithostatic pore-fluid pressure as a dominant control in SSE environments, often evidenced by seismic observations like high  $V_p/V_s$  ratios in Cascadia's episodic tremor and slow slip zone. Based on 3D dynamic rupture simulations (Madden et al., 2022) proposed that pore fluid pressure likely averages near 97% of lithostatic pressure in the Sumatra megathrust. Our results give even higher pore fluid pressure of  $99.6 \pm 0.17\%$  of lithostatic pressure assuming constant crust density of  $2.8 \times 10^3$  kg/m<sup>3</sup>. We are also in quantitative agreement with numerical models by Perez-Silva et al. (2023), which require effective normal stresses in the range of 1-5 MPa to generate SSEs on rate-strengthening faults, a scenario compatible with the transitional stability regime investigated in our study.

The inferred upper depth limit of the SSE generation zone from our MCMC inversion,  $30.44 \pm 2.8$  km, or  $175.3 \pm 16.2$  km along-dip provides quantitative constraints on the transition from locked seismogenic behavior to aseismic creep along the Cascadia megathrust. This depth is consistent with observations and models suggesting that SSEs in Cascadia initiate down-dip of the primary locked seismogenic zone. For instance, Audet and Kim (2016) noted that deep non-volcanic tremors, which are often correlated with SSEs, generally occur at depths of 30 to 45 km, some distance down-dip of the main seismogenic zone. Michel et al. (2019), who inverted geodetic observations using secular linear motion for interseismic locking and for SSE slip distribution, also describe the zone of SSEs and tremors in their Cascadia models as lying inland from the coastline, clearly disconnected from and down-dip of the locked portion of the megathrust by a shallow creeping section. Their modeling indicates this transition zone, characterized by stationary fault creep, spans between approximately 100 km and 150 km away from the trench. The down-dip limit of this shallow creeping section at around 150 km along dip, as identified by Michel et al. (2019), falls within the uncertainty bounds of our MCMC inversion result for the updip limit of the SSE zone, although our specific model setup does not explicitly impose a creeping zone between the locked and SSE-prone sections.

### 5.2 Secondary controls on the recurrence time of SSEs

Our parameter exploration, enabled by the ROM, allows us to investigate parameter regimes beyond those typically accessible via computationally intensive direct FOM studies, expanding upon the linear trends reported by Liu and Rice (2009) and highlighting more complex, second-order dependencies on both  $W$  and  $\sigma_W$  (Figs 5, 10, 11). Our MCMC inversion (Figure 11) uncovers the posterior distributions of  $W$  and  $\sigma_W$ , constrained by characteristic SSE observations from the northern CSZ.

Given that the extent of our parameter exploration allows  $W/h^*$  to vary by a factor of 6 due to changes in  $\sigma_W$  (holding  $W$  constant), versus a factor of 1.6 due to changes in  $W$  (holding  $\sigma_W$  constant), the general gradient of  $T_c$  across the sampled parameter space appears predominantly aligned with the  $\sigma_W$  axis. This observation is supported by the MCMC inversion, where  $T_c$  exhibits a PCC of 0.83 with  $\sigma_W$ , compared to only 0.2 with  $W$ . Notably, some regions exhibit high gradients in recurrence time with a strong dependence on  $W$ . For instance, at  $\sigma_W = 2.75$  MPa,  $T_c$  jumps from 0.85 years to 1.1

years as  $W$  increases from 38 km to 43 km. This represents a 30% increase in recurrence time but only a 7% increase in the normalized fault width  $W/h^*$ .

### 5.3 Forward modeling method

A crucial aspect underpinning our ROM development is the verified accuracy of the FOMs that generated the training data. Our FOM simulations, performed using the volumetric discontinuous Galerkin finite element code tandem (Section 2, Uphoff et al., 2022), successfully reproduce the key relationships between normalized fault width ( $W/h^*$ ), SSE recurrence interval, and mean SSE slip previously established by Liu and Rice (2009) using a boundary element method (BEM) (Figure 5). This agreement verifies our physical model setup prior to the development of our ROM.

While BEM offers computational advantages for fault-dominated problems by reducing dimensionality compared to our tandem-based FOMs, for more complex model setups BEM simulations may still be computationally expensive. For example, Tainpakdipat et al. (2025) report that simulations for their SSEs models demand between 20 and 250 CPU hours. Therefore, BEM may be similarly limited as volumetric codes when calculating more than 10,000 model inferences, as done with our ROM for exploring the parameter spaces and estimating uncertainties. Furthermore, classical BEM techniques typically assume a homogeneous or layered elastic medium and simplified model and fault geometries (Rice & Gu, 1983; Lapusta et al., 2000; Liu & Rice, 2005; Lapusta & Liu, 2009; Segall & Bradley, 2012; Li & Liu, 2016b, 2017; Barbot, 2019). Despite recent developments to incorporate more complex models (e.g., Mallick et al., 2022; Mallick & Sathiakumar, 2024), efficiently handling widespread and complex off-fault material variations, or complex subsurface geometries and topography, remains a challenge for BEM compared to volumetric codes.

It is important to note that the ROM framework presented in this study is agnostic to the specific numerical method of the FOM. Consequently, our ROM scheme could be readily applied to training data generated from BEM-based simulations or from volumetric simulations that accommodate complex off-fault material properties.

Nie and Barbot (2021) explore SSEs in a 2D anti-plane strain setting, varying both the Dieterich-Ruina-Rice number ( $R_u$ ), which is proportional to  $W/h^*$  by a constant factor close to unity and  $R_b = (b - a)/b$ , a parameter that controls the ratio of dynamic to static stress drops (Gabriel et al., 2012). In our model, a larger width of the low effective normal stress zone incorporates a greater portion of the VW section of the fault (Figure 1), thereby increasing the apparent  $R_b$  of the SSE-producing zone. Although Nie and Barbot (2021) did not directly investigate the change in recurrence time as a function of  $R_b$ , their observed data shows rapid changes in the peak slip-rate and the system limit cycle style which can consequently alter the recurrence times of events. The picture emerging from our parametric space exploration reveals three distinct dependencies of recurrence time on the normalized fault length: for  $R_b < 0.175$ , recurrence time increases steeply over a relatively short interval of  $W/h^*$  (from 7.5 to 12.5); in contrast, for  $R_b > 0.21$ , the recurrence interval becomes linearly dependent on the normalized fault length, and a transition zone between these behaviors is identified for  $R_b$  values in the range  $0.175 \leq R_b \leq 0.21$  (Figure ??). These detailed dependencies of recurrence time on  $W/h^*$  agrees with Nie and Barbot (2021) in that SSEs characteristics have at least secondary dependence on  $R_b$ .

### 5.4 Comparison with previous work

The application of ROM techniques to accelerate computationally intensive simulations is gaining traction in earthquake science (Rekoske et al., 2023; Kaveh et al., 2024; Rekoske et al., 2025; Ragu Ramalingam et al., 2025; Hobson & May, 2025a) and beyond

(e.g., Degen et al., 2023; Quiaro et al., 2025; Hobson & May, 2025b). Our ROM methodology, which combines a spline-based latent space representation with RBF interpolation for POD coefficients builds a ROM for each individual slow slip event expanding the approach by Rekoske et al. (2025). A key difference in our approach is the two-step nature of the model order reduction. The first step involves the transformation of the complex SSE cycle data into an efficient, low-dimensional spline-based latent representation. Key advantage of this latent space is the representation of simulations as fixed-length vectors, which addresses the challenge of variable timestep outputs and differing durations across simulations. This is a prerequisite for the following matrix-based ROM analyses and could also be utilized by other machine learning methods, such as many neural networks, which require uniform-length inputs (Lecun et al., 1998). Another aspect of our ROM framework the per-cycle ROM construction helps effectively managing and reducing the dimensionality of the complex time-history data that are characteristic of SSE cycle simulations.

Kaveh et al. (2024) employ a POD-based ROM to forecast extreme events in a rate-and-state friction fault model that produces SSEs. They focus on identifying precursor states to SSEs by building their ROM from simulation snapshots of the inter-event periods only, and over one simulation settings (no change in initial parametrization), thereby capturing the system’s characteristics while excluding the SSEs themselves. This allows them to define an optimization problem within the reduced-order space to find extreme events precursors. In contrast, our ROM is designed to efficiently simulate the entire SSE cycle, including the SSEs.

Physics-Informed Neural Networks (PINNs, Fukushima et al., 2023; Okazaki et al., 2022) represent a class of deep learning models that are trained to solve PDEs by directly incorporating the equations, along with initial and boundary conditions, into the neural network’s loss function. Recently, PINNs have emerged as a promising candidate for a data-driven approach to solving and inverting fault and rate-and-state friction equations in both laboratory (Borate et al., 2024) and numerical settings (Rucker & Erickson, 2024; Fukushima et al., 2025). Fukushima et al. (2025) employed PINNs for the direct inversion of spatially distributed frictional parameters ( $a, a-b, L$ ) from geodetic observations. Parameters are determined by minimizing a composite loss that includes both data misfit and PDEs residuals, the underlying physics thus serves as an inherent regularization constraint during the learning process itself. In such an intrusive framework, the learning phase is coupled with the inversion for specific fault frictional properties. Conversely, our ROM functions as a non-intrusive, data-driven surrogate for the underlying physics, i.e., is agnostic toward the kind of forward model used. The ROM approximates the complex input-output relationships of the system from a dataset of FOM simulations, without requiring explicit knowledge or direct utilization of the governing equations during its construction phase. This process yields an efficient forward model, which can subsequently be integrated into established inversion frameworks. As a result, our ROM approach is well-suited for uncertainty quantification, as showcased by our MCMC analysis. This is a capability not as directly featured in a PINNs parameter estimation framework.

## 5.5 Limitations

A primary consideration for the scalability of our ROM approach is the offline cost associated with generating the FOM simulations required for training. In this study, with a two-dimensional parameter space ( $\dim(\mathcal{P}) = 2$ , for  $W$  and  $\sigma_W$ ), the 76 FOM simulations, though computationally intensive, provide a sufficient basis for constructing an accurate ROM. However, the number of FOM evaluations needed to adequately sample the parameter space and train a robust ROM can, in theory, increase exponentially, with the number of parameters being surrogated in the parameter space  $\mathcal{P}$ . While the speedup achieved during the online phase is substantial, the initial investment in FOM

simulations for higher dimensional parameter spaces could become a limiting factor. Future work might explore more adaptive or sparse sampling strategies to mitigate the challenge of FOM generation in higher-dimensional parameter spaces for ROMs (Bui-Thanh et al., 2008).

Another limitation observed in our results is the systematic overestimation of potency by the ROM (Figure 8(c),(d)). Although we demonstrated that a simple linear post-processing correction, derived from the LOOCV results, can effectively mitigate this bias for the test set (Figure ??), this is not an ideal solution. The source of this systematic discrepancy may lie in the RBFs interpolation of POD coefficients, in the accumulation of minor errors through the multi-step ROM construction process or potentially in the ROM lacking the ability to perfectly capture the transition between consecutive SSE cycles when they are concatenated in time. Ideally, the ROM should predict potency accurately without requiring such a correction. This suggests an area for future refinement, perhaps through the exploration of alternative interpolation schemes for the POD coefficients, or by investigating strategies like introducing small overlaps or more sophisticated blending techniques between individual SSE cycle ROMs when reconstructing longer time series.

Furthermore, as with most data-driven surrogate models, at its core, our ROM is an interpolatory method. Its accuracy can be trusted only within the convex hull of the training parameter sets ( $\xi$ ) and it should not be used for extrapolation beyond the sampled parameter range. The quality of the ROM predictions is also contingent on the density and distribution of the FOM training samples. While our iterative refinement strategy aimed to address regions of high error, ensuring comprehensive coverage for complex, high-gradient parameter responses remains a challenge.

Finally, the physical limitations inherent in the FOMs themselves will propagate to the ROM. Our current FOMs, for instance, are 2D and do not capture 3D effects such as along-strike variations in fault properties (Brudzinski & Allen, 2007; Li & Liu, 2017) or in SSE kinematics (Takagi et al., 2019; Li & Gabriel, 2024). Similarly, more complex rheologies (Gao & Wang, 2017) or fluid-transport mechanisms (Perez-Silva et al., 2023; Ozawa et al., 2024), if not included in the FOMs, cannot be represented by the ROM. The ROM's are ultimately bounded by the underlying full-order model, the ROM cannot learn, or capture time-dependence and or physics which is not present in the FOM.

## 5.6 Future work

A natural future extension of our SEAS ROM approach involves applying it to more complex simulations, such as those incorporating fast earthquakes in addition to SSEs, more extensive frictional parameter variations, or to extend the forward simulations to 3D SSE models, all requiring to approximate more parameters. For example, a transition from 2D to 3D domains (and consequently from a 1D to a 2D fault) would increase the number of coupled rate-and-state friction equations. These added complexities could potentially disrupt the well-behaved nature of the phase-space limit cycle trajectories, possibly leading to chaotic behavior (Barbot, 2019; Cattania, 2019), which would be more challenging to capture with a ROM. Wang (2024) studied the effect on cycle simulations of a 1D fault embedded in a 2D domain and found that an increase in the number of interacting rate-and-state friction points in space can lead to more complicated time-dependence behavior, potentially resulting in more complex phase-space trajectories. Distinguishing between quasi-periodic and truly chaotic behavior is not straightforward, and the former can be misinterpreted as the latter (Wang, 2024).

The stability of these limit cycles and their potential transition to chaotic behavior can be linked to the friction parameters explored. Viesca (2016b) demonstrated that as the ratio of rate-and-state friction parameters  $a/b$  approaches 1, the system tends towards instability and can exhibit chaotic characteristics. Conversely, smaller  $a/b$  values

are associated with stable, periodic limit cycles. However, Viesca (2016b) also notes that in most seismic cycle models  $a/b \leq 0.8$ , in this regime, slip instability does not develop chaotically but rather in a universal manner. In the context of our Cascadia-like model, the  $a - b$  parameters vary along the fault, transitioning from VW to VS behavior. In the VW sections, our chosen  $b$  value of 0.0045 and  $a$  values (Figure 1(c)) result in  $a/b$  ratios that are generally sufficiently less than 1. This suggests that the SSEs generated in our 2D FOMs should, and do, exhibit stable limit cycles. However, extending the ROM to exploring parameter regimes where  $a/b$  is closer to unity would necessitate careful consideration of these potential transitions to more complex, possibly chaotic or quasi-periodic, behaviors. The adaptability of the spline latent space representation and the POD-RBFs framework, which handles the simulation cycle-by-cycle, makes it a promising candidate for future work aiming at capturing such behavior, provided the training FOMs adequately sample these complex regimes of the parameter space.

## 6 Summary

We present a two-step, scientific machine learning reduced-order modeling (ROM) framework that accelerates rate-and-state friction simulations of the slow slip cycle by  $3.6 \times 10^5$  compared with full-order sequences of earthquakes and aseismic slip (SEAS) models. First, each simulated slow slip event (SSE) is recast into a compact spline-based latent space using a phase-space representation of slip rate and state. Second, proper orthogonal decomposition (POD) combined with radial-basis-function (RBF) interpolation emulates the simulations with varying initial conditions. Our ROMs are validated with leave-one-out cross-validation and comparison to earlier, independent SSE simulations. We use the ROMs to explore complex, non-linear dependencies of northern Cascadia-like SSE characteristics on the width  $W$  and magnitude  $\sigma_W$  of a deep low effective normal stress zone. We perform a Bayesian Markov-chain Monte-Carlo inversion, constraining these parameters and their uncertainties to  $W = 44.7 \pm 16.2$  km and  $\sigma_W = 3.8 \pm 1.44$  MPa. These values imply near-lithostatic pore fluid pressure ( $99.6 \pm 0.17\%$  lithostatic) and place the upper SSE source boundary (i.e., the frictional transition zone) at  $30.44 \pm 2.8$  km depth, which is consistent with geophysical observations. Because the method is non-intrusive and agnostic to the underlying forward model, future work may extend our ROMs to even higher-dimensional parameter spaces, mixed seismic-aseismic cycles, and fully 3-D geometries, offering a practical route to systematic uncertainty quantification throughout the earthquake cycle. By systematically linking megathrust properties to rate-and-state governed slow slip cycle characteristics, our study helps to constrain first- and second-order controls on how plate boundaries slip, providing input for seismic hazard assessment and future 3-D modeling.

## Appendix A Mathematical symbols and definitions

Table A1: Table of mathematical symbols and definitions used throughout the study

Symbol	Definition	Dimension
$\tau$	Fault shear stress	$\mathbb{R}^{D-1}$
$\dot{\mathbf{s}}$	Slip-rate	$\mathbb{R}^{D-1}$
$\psi$	State variable in rate-and-state friction	Scalar
$a, b$	Empirical friction parameters	Scalar
$\dot{s}_0$	Reference slip-rate	Scalar
$L$	Characteristic slip distance	Scalar
$f_0$	Reference friction coefficient	Scalar

*Continued on next page*

Symbol	Definition	Dimension
$\theta$	State variable in standard aging law	Scalar
$\dot{s}_p$	Plate convergence velocity (11.7 cm/year)	$\mathbb{R}^{D-1}$
$h^*$	Characteristic nucleation size	Scalar
$\Lambda$	Process zone size	Scalar
$d_{\text{poly}}$	Polynomial degree of basis functions	Scalar
$\sigma_W$	Effective normal stress	Scalar
$W_l$	Up-dip extent of the low effective normal stress region	Scalar
$W_r$	Down-dip extent of the low effective normal stress region	Scalar
$W$	Width of low effective normal stress seismogenic region	Scalar
$\xi$	Parameter pair	$\mathbb{R}^k$
$\mathcal{P}$	2D parameter space	$\mathbb{R}^{N \times b}$
$\mathbf{Q}$	Data set of simulation outputs	$\mathbb{R}^n \times \mathbb{R}^{m \times n} \times \mathbb{R}^{m \times n} \times \mathbb{R}^{m \times n}$
$\mathbf{t}$	FOM simulation time steps vector	$\mathbb{R}^n$
$\mathbf{\dot{S}}$	FOM simulation slip-rate outputs	$\mathbb{R}^{m \times n}$
$\mathbf{\Psi}$	FOM simulation state variable outputs	$\mathbb{R}^{m \times n}$
$\mathbf{S}$	FOM simulation cumulative slip outputs	$\mathbb{R}^{m \times n}$
$\text{ROM}_i(\cdot)$	Reduced-order model for cycle $i$	Function
$\xi^*$	Parameter pair input for the ROM ( $\xi^* \notin \mathcal{P}$ )	$\mathbb{R}^k$
$\mathbf{Q}^*$	ROM simulation output set	$\mathbb{R}^n \times \mathbb{R}^{m \times n} \times \mathbb{R}^{m \times n} \times \mathbb{R}^{m \times n}$
$\mathbf{q}$	Latent space representation of $\mathbf{Q}$	$\mathbb{R}^l$
$G(\cdot)$	Simulation to spline latent space projector	Function
$G^{-1}(\cdot, \cdot)$	Spline latent space to simulation reconstruction	Function
$H(\cdot)$	Trajectory of the phase-space parametric curve $\log_{10}(\dot{s}(t), \psi(t))$	Function
$\phi(t)$	Progression variable along trajectory in phase-space	Scalar
$\mathcal{B}(\cdot, \cdot)$	B-spline transformation	Function
$\mathcal{B}^{-1}(\cdot, \cdot, \cdot)$	Inverse B-spline transformation	Function
$\mathbf{k}_x$	B-spline knot vector placed along $\mathbf{x}$	$\mathbb{R}^M$
$\mathbf{c}_f$	B-spline coefficient vector evaluated along $f(x)$	$\mathbb{R}^M$
$\mathbf{D}$	Matrix of latent vector representations	$\mathbb{R}^{N \times l}$
$\mathbf{u}_k$	Basis vector of the image of $\mathbf{D}$	$\mathbb{R}^l$
$\alpha_k^j$	POD coefficient	Scalar
$\varphi(\cdot)$	RBF kernel	Function
$T_c$	SSEs recurrence interval	Scalar
$P_0$	SSEs potency	Scalar
$W_s$	along strike fault length	Scalar
$N$	number of FOM evaluation	Scalar
$i$	FOM parameter index (e.g. $\xi_i$ )	Scalar
$n$	length of time steps vector	Scalar
$m$	number of fault observation points	Scalar
$j$	index for specific observation point (e.g. $\mathbf{Q}_j$ )	Scalar

*Continued on next page*

Symbol	Definition	Dimension
$p$	number of SSEs cycles	Scalar
$k$	index for specific SSE cycle (e.g. $Q^k$ )	Scalar

## Appendix B B-spline knot placement strategy

The accuracy and efficiency of our spline-based latent space representation depends on the strategic placement of B-spline knots. An optimal knot distribution allocates more knots to regions of high functional complexity, increases representation fidelity with a minimal number of spline knots. Given the multi-scale nature of the SSE cycle simulations data, we developed a two-step, adaptive knot placement strategy for our two distinct spline mappings: (1) mapping the phase progression  $\phi$  from the simulation time  $t$  using  $\mathbf{k}_t \in \mathbb{R}^{K_0}$  knots, and (2) mapping the physical variables ( $\dot{s}$ ,  $s$ ,  $\psi$ ) from the phase progression variable  $\phi$  using  $\mathbf{k}_\phi \in \mathbb{R}^{K_1}$  knots.

For the first mapping, which connects simulation time  $t$  to the phase progression  $\phi$ , the primary challenge is the highly non-uniform distribution of time steps from the full-order model’s adaptive time-stepping scheme. As shown in Figure B1, the relationship  $\phi(t)$  is characterized by a long, low-gradient inter-event period, preceded and followed by an abrupt change during the SSEs. To model this, we use a combination of quantile-based and uniform knot placement along time. The adaptive time stepping used in the FOM generates a high density of time steps during the rapid slip of an SSE, placing knots according to the quantiles of the time steps vector ( $\mathbf{t}$ ) allocates more knots to the SSE period itself. On the other hand, uniformly placed knots ensure coverage in the inter-event period, where time steps can be sparse. This combined approach is crucial for accurately resolving the sharp onset and evolution of the slow slip event while efficiently representing the long, quasi-static interseismic period with fewer knots. Based on trial and error for the best reconstruction fidelity, a ratio of 0.7 is chosen between quantile and uniform placement, which gives  $0.7K_0$  quantile knots and  $0.3K_0$  uniform knots along  $t$ .

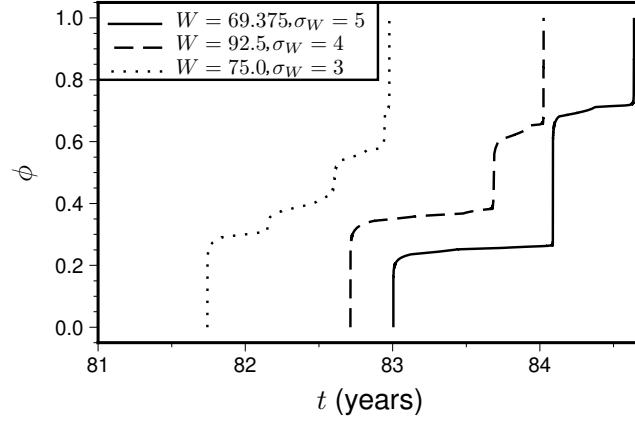
For the second mapping, which represents the trajectory in phase space, a similar approach is taken for placing knots along the  $\phi$ . However, the mapping from  $\phi$  to the fault variables is much smoother and lacks sharp, abrupt gradients (Figure 3, Figure 6, and Section 4.2). Consequently, a majority of the knots are placed uniformly. A ratio of 0.2 is chosen between quantile and uniform placement, which gives  $0.2K_1$  quantile knots and  $0.8K_1$  uniform knots along  $\phi$ .

## Open Research Section

All codes and data products supporting this study are openly available. The input files required to reproduce the tandem simulations, a static version of the tandem code, and the ROM analysis code are publicly hosted on Zenodo (Magen et al., 2025). The complete raw outputs from all simulations are archived on the National Data Platform S3 bucket. The Zenodo repository provides the direct link and instructions for accessing these raw data.

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**Figure B1.** The relationship between simulation time  $t$  and phase-space progression  $\phi$ . The plot illustrates the evolution of the phase-space progression variable  $\phi$  as a function of time  $t$  for a single representative SSE cycle from three different simulations.

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