Generalized Langevin Equation

Stochastic Differential Equations

Sankarasubramanian Ragunathan

389851

RWTH Aachen University



Agenda

- 1 Scope of the Project
- 2 Introduction to Generalized Langevin Equation Introduction Mathematical Model Model Complications and Solutions Extended Variable GLE
- 3 Numerical Schemes for Solving GLE Numerical Schemes Comparison of Numerical Schemes Harmonic Potential Well
- 4 Sensitivity Analysis Need for Sensitivity Analysis Introduction Methods for calculating sensitivity Efficient Finite Difference Estimators Senitivity Analysis - Results
- 6 Conclusion
- **6** References

Scope of the Project

- What? Generalized Langevin Dynamics is a modeling technique that can be used to model anomalous diffusive phenomena observed in viscoelastic fluid flow and in biological systems.
- Why? Anomalous diffusion problems: Langevin model fails to capture sub-diffusive and super-diffusive behavior which the GLE succeeds in capturing. But GLE is Non-Markovian i.e. memory kernel depends on the history of velocity. This issue is overcome by using Extended Variable GLE that considers a finite dimensional subspace for the memory kernel.
- Where? Applications of GLE include but are not restricted to micro-rheology, biological systems, nuclear quantum effects and systems in which anomalous diffusion arise.
 - How? Study Extended Variable GLE using Prony series approximation. Accuracy of Implicit/Explicit Euler and Splitting Numerical schemes are also tested to find out the optimal scheme. Study the sensitivity of the solution to the changes in the parameters of the extended variable GLE.

Introduction

What? Langevin Dynamics: Large particles in a bath of small particles, motion of large particles directly integrated while the dynamics of small particles are "averaged out".

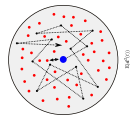
Why? *Molecular Dynamics* simulations involving all particles is computationally expensive. Langevin Equation model is computationally cheaper.

Drawback

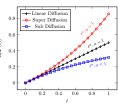
Anomalous diffusion problems arising due to *Power Law* behavior of solute-solvent systems cannot be solved.

Solution

Generalized Langevin Equation (GLE)







(b) Sub-diffusive and Super-diffusive behavior of solute-solvent systems

Mathematical Model

The velocity term of GLE is based on *Ornstein-Uhlenbeck* process.

GLE Equations

$$dX(t) = V(t)dt (1)$$

$$\boldsymbol{M}d\boldsymbol{V}(t) = \underbrace{\boldsymbol{F}^{c}(\boldsymbol{X}(t))dt}_{\text{Conservative}} - \underbrace{\int_{0}^{t} \boldsymbol{\Gamma}(t-s)\boldsymbol{V}(s)dsdt}_{\text{Temporally}} + \underbrace{\boldsymbol{F}^{r}(t)dt}_{\text{Random Correlated}}$$
Force due to Non-Local Drag (F^{d}) Force given by Potential Force FDT
$$\boldsymbol{X}(0) = \boldsymbol{X}_{0}, \quad \boldsymbol{V}(0) = \boldsymbol{V}_{0} \quad \text{(Initial Conditions)}$$
 (3)

Note:

 $oldsymbol{F}^r$ and $oldsymbol{F}^d$ are characterized by the memory kernel consistent with FDT.

Mathematical Model

Theorem

FDT (Fluctuation Dissipation Theorem) states that the equilibration to a temperature, T, requires that the two-time correlation of $\mathbf{F}^r(t)$ and $\Gamma(t)$ be related as:

$$\langle \boldsymbol{F}_{i}^{r}(t+s), \boldsymbol{F}_{j}^{r}(t) \rangle = k_{B}T\Gamma(s)\delta_{ij}, \quad s \geq 0$$
 (4)

where k_B is the Boltzmann's Constant and δ_{ij} is the Kronecker Delta.

Note:

- ${m F}^d(t)$ depends on the velocity history unlike in Langevin Equation where it depends on the velocity at that instant.
- The random forces are not just delta correlated but are correlated by the memory kernel. Memory Kernel choice and approximation important based on the problem to be studied.

Model Complications and Solutions

Complications

- Storage of subset of the time history of V(t).
- 2 Sequence of $\boldsymbol{F}^r(t)$ given by FDT.
- Numerical SDE solution should converge in distribution.

Solution

Using extended variable Prony Series for Memory Kernel.

$$\Gamma(t) pprox \sum_{k=1}^{N_k} rac{c_k}{ au_k} \exp\left[-rac{t}{ au_k}
ight], \qquad t \geq 0$$
 (5)

where N_k is the number of terms used in approximating the memory kernel.

Using a suitable integration scheme for the numerical method.

Model Complications and Solutions

Why use extended variable Prony Series?

- Approximation of memory kernel to map Non-Markovian GLE to Markovian system of N_k variables.
- Typically used for modelling Power Law based decay/growth as observed in sub/super diffusive systems.

Importance of choice of integration scheme?

- Conservation of moments of variables of interest such as displacement and velocity (usual variables of interest for MD simulations)
- Convergence of **GLE** to Langevin equation in the limit of small τ_k as observed in theory.

Extended Variable GLE

Main Extended Variable GLE Equations

$$m_i dV_i(t) = F_i^c(\mathbf{X}(t))dt + \sum_{k=1}^{N_k} S_{i,k} dt$$
 (6)

$$dX_i(t) = V_i(t)dt (7)$$

$$dS_{i,k}(t) = -\frac{1}{\tau_k}S_{i,k}(t)dt - \frac{c_k}{\tau_k}V_i(t)dt + \frac{1}{\tau_k}\sqrt{2\mathsf{k_B}Tc_k}dW_{i,k}(t) \tag{8}$$

Auxiliary Extended Variable GLE Equations

$$S_{i,k}(t) = Z_{i,k}(t) + F_{i,k}(t)$$
(9)

$$dZ_{i,k}(t) = -\frac{1}{\tau_k}Z_{i,k}(t)dt - \frac{c_k}{\tau_k}V_i(t)dt \qquad \qquad Z_{i,k}(t) = -\int_0^t \frac{c_k}{\tau_k} \exp\left[-\frac{(t-s)}{\tau_k}\right]V_i(s)ds \ \ (10)$$

$$dF_{i,k}(t) = -\frac{1}{\tau_k}F_{i,k}(t)dt + \frac{1}{\tau_k}\sqrt{2k_BTc_k}\,dW_{i,k}(t) \qquad \left\langle F_{i,k}(t+s),F_{i,k}(t)\right\rangle = \mathsf{k_B}T\frac{c_k}{\tau_k}\exp\left[-\frac{s}{\tau_k}\right] \tag{11}$$

$$F_i^T(t) = \sum_{k=1}^{N_k} F_{i,k}(t)$$
 (12)

Numerical Schemes

Numerical Schemes:

- Explicit Euler Scheme.
- Splitting Scheme.

Which numerical scheme to choose for solving the problem?

Scheme that is able to conserve the first and second moments of quantities of interest i.e. ${m V}(t)$ and ${m X}(t)$.

Reason: General observables/physical quantities that are measured to understand the behavior of a system. For example, *Mean Square Displacement*(MSD) in microrheology simulations.

Implementation Details:

- Uniform time-step size, Δt , where $N_t \Delta t = T_{\text{tot}}$ (T_{tot} represents the total time of the simulation and N_t represents the # of time-steps.)
- All N_p particles are seeded with the same constant X(0) and V(0) (Note: We could also seed the initial conditions based on the p.d.f if known.)
- The composite variable $S_{i,k}(t)$ is assumed to be zero initially.

Numerical Schemes

Explicit Euler Scheme

Input: X(0).V(0).S(0)

Output: X(t), V(t)

1: for n=0 to N_t do

2:
$$V_i^{n+1} = V_i^n + \frac{\Delta t}{m_i} F_i^c(\mathbf{X}^n) + \frac{\Delta t}{m_i} \sum_{k=1}^{N_k} S_{i,k}^n$$

3:
$$X_i^{n+1} = X_i^n + \Delta t V_i^n$$

$$4: \qquad S_{i,k}^{n+1} = \left(1 - \frac{\Delta t}{\tau_k}\right) S_{i,k}^n - \frac{c_k \Delta t}{\tau_k} V_i^n + \frac{1}{\tau_k} \sqrt{2 \mathsf{k}_\mathsf{B} T c_k} \Delta W_{i,k} \qquad \qquad \triangleright \mathsf{Advance} \ S(t) \ \mathsf{by a full step}$$

5 end for

Splitting Scheme

Input: X(0).V(0).S(0)

Output: X(t), V(t)

1: for
$$n=0$$
 to N_t do

2:
$$V_i^{n+1/2} = V_i^n + \frac{\Delta t}{2m_i} F_i^c (\mathbf{X}^n) + \frac{\Delta t}{2m_i} \sum_{k=1}^{N_k} S_{i,k}^n$$

$$hd Advance \ oldsymbol{V}(t)$$
 by a half step

3:
$$X_i^{n+1} = X_i^n + \Delta t V_i^{n+1/2}$$

$$\triangleright$$
 Advance $\boldsymbol{X}(t)$ by a full step

$$S_{i,k}^{n+1} = \theta_k S_{i,k}^n - (1 - \theta_k) c_k V_i^{n+1/2} + \alpha_k \sqrt{2 k_B T c_k} \Delta W_{i,k}$$

$$hd$$
 Advance $oldsymbol{S}(t)$ by a full step

5:
$$V_i^{n+1} = V_i^{n+1/2} + \frac{\Delta t}{2m_i} F_i^c \left(\mathbf{X}^{n+1} \right) + \frac{\Delta t}{2m_i} \sum_{k=1}^{N_k} S_{i,k}^{n+1}$$

$$riangleright$$
 Advance $oldsymbol{V}(t)$ by a half step

6: end for

Comparison of Numerical Schemes

Case Study:

- One dimensional problem, d=1
- Single mode in the Prony series approximation, $N_k=1$
- ullet Zero conservative force acting on the particles, $oldsymbol{F}^{c}\left(oldsymbol{X}(t)
 ight)=0$

The case study is simulated using both numerical schemes, *Explicit Euler* and *Splitting Method*, for three different τ and c values in the Prony Series approximation (Table 1)

Type of System	\boldsymbol{c}	au
Under-damped	1	1
Critically-damped	0.5	0.5
Over-damped	0.25	0.25

Table: c and τ values used for the case study

Comparison of Numerical Schemes - Results

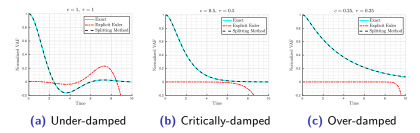


Figure: Comparison of different numerical schemes w.r.t. Normalized VAF

Normalized VAF (Velocity Autocorrelation Function)

$$\frac{\langle \boldsymbol{V}(t), \boldsymbol{V}(0) \rangle}{\langle \boldsymbol{V}(0), \boldsymbol{V}(0) \rangle} = \begin{cases} \exp\left[-\frac{t}{2\tau}\right] \left(\cos\left(\Omega t\right) + \frac{1}{2\tau\Omega}\sin\left(\Omega t\right)\right) & \text{for } \Omega \neq 0\\ \exp\left[-\frac{t}{2\tau}\right] \left(1 + \frac{t}{2\tau}\right) & \text{for } \Omega = 0 \end{cases}$$
(13)

$$\Omega = \sqrt{c/\tau - 1/4\tau^2}$$

Note: VAF is for one mode Prony series approx. without any conservative force terms.

Comparison of Numerical Schemes - Results

Observation: Explicit Euler scheme produces wrong results for the Normalized VAF as when compared to the Splitting scheme (*Reason:* Independent updates for $\boldsymbol{X}(t)$ and $\boldsymbol{V}(t)$ when using Explicit Euler scheme.)

Solution: Using the Splitting scheme as the preferred integration scheme for all proceeding cases.

Harmonic Potential Well

Harmonic Potential Well GLE

$$d\boldsymbol{V}(t) = \underbrace{-\omega_0^2 \boldsymbol{X}(t)}_{\text{Conservative Force}} dt - \underbrace{\int_0^t \frac{\gamma_\lambda}{\Gamma_0 \left(1-\lambda\right)} \left(t-s\right)^{-\lambda}}_{\text{Power Law decay}} \boldsymbol{V}(s) ds dt + \boldsymbol{M}^{-1} \boldsymbol{F}^r(t) dt$$

$$\underbrace{-\sum_{\text{Conservative Force arising from Harmonic Potential}}_{\text{monic Potential}} dt - \underbrace{\sum_{\text{Conservative Force from East Power Law decay}}_{\text{power Law decay function}} \boldsymbol{V}(s) ds dt + \boldsymbol{M}^{-1} \boldsymbol{F}^r(t) dt$$

Question?

Approximation of the memory kernel in Equation (14) using Prony series

Answer:

log-spaced values for τ_k from $\Delta t/10$ to $10N_t\Delta t$ and then linearly fitting c_k using least squares regression i.e.

$$\min_{\boldsymbol{x}} ||\boldsymbol{A} \cdot \boldsymbol{x} - \boldsymbol{b}||^2, \quad \boldsymbol{x} = \{c_k\} \quad \forall k = 1, \cdots, N_k$$

Harmonic Potential Well - Parameter Fitting

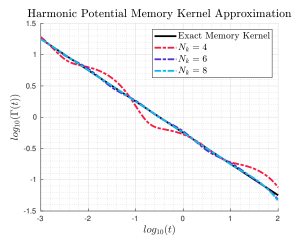
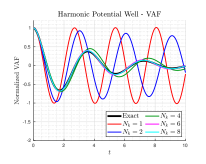
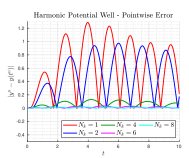


Figure: A Prony series fit of the *power law* memory kernel in eq. (14) for $\gamma_{\lambda}=1,\ \lambda=0.5$ for different N_k

Harmonic Potential Well - Normalized VAF & Pointwise Error





- (a) Normalized VAF at different times for (b) Pointwise error in VAF at different different no. of modes used in Prony series approximation
 - times for different no. of modes used in Prony series approximation

Figure: Approximation of the Harmonic Potential Well problem for $\gamma_{\lambda} = 1$, $\lambda = 0.5$ and $\omega_0 = 1.4$ using Prony series approximation

Harmonic Potential Well - Normalized VAF & Pointwise Error

Normalized VAF - Exact Solution

$$C_{V}(t) = \sum_{k=0}^{\infty} \frac{(-1)^{k}}{k!} \left(\omega_{0} t\right)^{2k} \mathcal{E}_{2-\lambda,1+\lambda k}^{(k)} \left(-\gamma_{\lambda} t^{2-\lambda}\right)$$

where $\mathscr{E}^{(k)}_{\alpha,\beta}(y)$ represents the kth derivative of the *Generalized Mittag-Leffler* function given by:

$$\mathscr{E}_{\alpha,\beta}^{(k)}\left(y\right) = \sum_{j=0}^{\infty} \frac{\left(j+k\right)! \; y^{j}}{j! \; \Gamma_{0}\left(\alpha(j+k)+\beta\right)}$$

Harmonic Potential Well - Normalized MSD

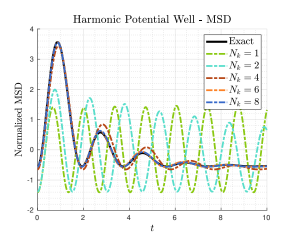


Figure: Normalized MSD at different times for different no. of modes used in Prony series approximation. ($\gamma_{\lambda}=1,\ \lambda=0.5$ and $\omega_{0}=1.4$)

Harmonic Potential Well - Normalized MSD

Normalized MSD - Exact Solution

$$\begin{split} \underbrace{\left\langle \left[\boldsymbol{X}(t+\tau) - \boldsymbol{X}(t) \right]^2 \right\rangle}_{\text{Normalized MSD}} &= \frac{2 \mathsf{k_B} T}{m} I(\tau) - 2 x_0 v_0 \omega_0^2 \left[G(t+\tau) - G(t) \right] \left[I(t+\tau) - I(t) \right] \\ &\quad + \left(v_0^2 - \frac{\mathsf{k_B} T}{m} \right) \left[G(t+\tau) - G(t) \right]^2 \\ &\quad + \omega_0^2 \left(x_0^2 \omega_0^2 - \frac{\mathsf{k_B} T}{m} \right) \left[I(t+\tau) - I(t) \right]^2 \\ &\quad I(t) &= \sum_{k=0}^{\infty} \frac{(-1)^k}{k!} \omega_0^{2k} t^{2(k+1)} \mathscr{E}_{2-\lambda, 3+\lambda k}^{(k)} \left(-\gamma_{\lambda} t^{2-\lambda} \right) & \text{(Kernel integral)} \\ &\quad G(t) &= \sum_{k=0}^{\infty} \frac{(-1)^k}{k!} \omega_0^{2k} t^{2k+1} \mathscr{E}_{2-\lambda, 2+\lambda k}^k \left(-\gamma_{\lambda} t^{2-\lambda} \right) & \text{(Relaxation function)} \end{split}$$

where v_0, x_0 represent the velocity and position of the particles at time t=0.

Sensitivity Analysis

Inference:

- Devised an integration scheme that is able to conserve first and second moments of $\boldsymbol{V}(t)$ and $\boldsymbol{X}(t)$ (Observables for understanding the behavior of physical systems.)
- Adequately able to approximate exact memory kernel using finite no. of modes of Prony series.

Question?

Need for performing sensitivity analysis

Answer:

- Extended variable GLE fitted by matching MSD or VAF obtained from experimental data \Rightarrow Prony series parameters c_k, τ_k, N_k sensitive to experimental data i.e. to errors in measurement or lack of data.
- *Ergodicity breaking:* Time averaged statistics such as MSD do not converge to ensemble averages.

Sensitivity Analysis

Sensitivity Analysis

Let $\mathcal{S}\left(t,\theta;f\right)$ denote the sensitivity of the stochastic process $\boldsymbol{X}_{t}\left(\theta\right)$ where θ is a parameter that affects the stochastic process and f is the given observable. We are interested in calculating:

$$\mathcal{S}(t, \theta; f) = \frac{\partial \mathbb{E}\left[f\left(\boldsymbol{X}_{t}\left(\theta\right)\right)\right]}{\partial \theta}$$

Methods of Calculation:

- Finite Difference Stencils: Approximating derivative by a finite difference stencil and obtain the required moments by Monte Carlo.
- Likelihood Ratio: Expressing sensitivity as an expectation of f under a change of measure.
- Malliavin Calculus: Extension of calculus of variations to stochastic processes.

Methods for calculating sensitivity

Finite Difference Stencil

$$S_{\varepsilon}(t, \theta; f) = \frac{\mathbb{E}\left[f\left(\boldsymbol{X}_{t}\left(\theta + \varepsilon\right)\right)\right] - \mathbb{E}\left[f\left(\boldsymbol{X}_{t}\left(\theta\right)\right)\right]}{\varepsilon}$$

Likelihood Estimator

$$S_{LR}(t,\theta;f) = \frac{\mathbb{E}\left[f\left(\boldsymbol{X}_{t}\left(\theta\right)\right)\right]}{\partial\theta} = \int f\left(x_{t}\right)\left[\partial_{\theta}\log g\left(\theta,x_{t}\right)\right]g\left(\theta,x_{t}\right)dx_{t}$$
$$= \mathbb{E}\left[f\left(\boldsymbol{X}_{t}\left(\theta\right)\right)\partial_{\theta}\log g\left(\theta,\boldsymbol{X}_{t}\right)\right]$$

where g represents the change in measure by a p.d.f. g is unknown \Rightarrow difficulty in calculating sensitivity using likelihood estimator

Malliavin Calculus

$$S_{M}\left(t, \theta; f\right) = \mathbb{E}\left[f\left(\boldsymbol{X}_{T}\right) h\left(\left\{\boldsymbol{X}_{s}\right\}_{0 \leq s \leq T}\right)\right]$$

where $h\left(.\right)$ represents the Malliavin weights. h(.) is computationally expensive to calculate.

Sensitivity Estimators using Finite Difference Stencils

Objective: To calculate the sensitivity of the stochastic process efficiently using finite difference stencils ⇒ right sampling strategy to reduce variance.

Methodology: Let $\hat{\phi}(\theta)$ represent the expected value of the observable $f(X_t(\theta))$. Then we can write,

$$\hat{\phi}(\theta) = M^{-1} \sum_{i=1}^{M} f(X_{i,t}(\theta)) \Rightarrow S_{\varepsilon}\left(t,\theta;\hat{\phi}\right) \approx \Delta_{c}(M,\varepsilon) = \frac{\hat{\phi}(\theta+\varepsilon) - \hat{\phi}(\theta-\varepsilon)}{2\varepsilon}$$

where each random variable is sampled independently of each other.

$$\begin{split} \Rightarrow \operatorname{Var}\left[\Delta_{c}\right] &= \varepsilon^{-2} \operatorname{Var}\left[\hat{\phi}\left(\theta + \varepsilon\right) - \hat{\phi}\left(\theta - \varepsilon\right)\right] \\ &= \varepsilon^{-2} M^{-1} \operatorname{Var}\left[\underbrace{f\left(\boldsymbol{X}_{t}\left(\theta + \varepsilon\right)\right)}_{R_{1}} - \underbrace{f\left(\boldsymbol{X}_{t}\left(\theta - \varepsilon\right)\right)}_{R_{2}}\right] \end{split}$$

where the variance can be rewritten as

$$Var[\Delta_c] = \varepsilon^{-2} M^{-1} (Var[R_1] + Var[R_2] - 2Cov[R_1, R_2])$$

Sensitivity Estimators using Finite Difference Stencils

Statistical Error is given by $\epsilon_M=\frac{C_{\alpha}\sigma}{\sqrt{M}}$ where $\sigma=\sqrt{{\rm Var}\left[\Delta_c\right]}$ and C_{α} is a constant based on the confidence level of the solution

Coupled vs. Decoupled Noise:

- If R_1 and R_2 are not correlated, then ${\sf Var}\left[\Delta_c\right]=\mathcal{O}\left(arepsilon^{-2}M^{-1}
 ight)$
- ullet If R_1 and R_2 are positively correlated, then ${\sf Var}\left[\Delta_c
 ight]=\mathcal{O}\left(M^{-1}
 ight)$

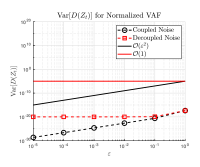
Observation:

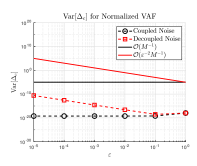
Let $\varepsilon = 0.1$. In order to reduce the error by a factor of 10.

- Coupled Noise: $M = 10^4$ samples
- Decoupled Noise: $M=10^2$ samples

Hence the statistical error becomes independent of the perturbation parameter on using a common random path coupling i.e. R_1 and R_2 use the same Wiener process dW.

Sensitivity Analysis - Results





(a)
$$\operatorname{Var}\left[D\left(Z_{t}\right)\right]$$
 for Normalized VAF where $D\left(Z_{t}\right)=\left(\hat{f}\left(\theta+\varepsilon\right)-\hat{f}\left(\theta-\varepsilon\right)\right)$

where
$$egin{aligned} \left(\mathbf{b}\right) & \mathrm{Var}\left[\Delta_c\right] & \mathrm{for} & \mathrm{Normalized} & \mathrm{VAF} & \mathrm{where} \\ \Delta_c &= \left(\hat{\phi}\left(\theta + \varepsilon\right) - \hat{\phi}\left(\theta - \varepsilon\right)\right)/2\varepsilon \end{aligned}$$

Figure: Order of variance reduction when using Coupled vs. Decoupled noise term on perturbing c_1 . ($M=200, \delta t=0.1, N_p=1000$)

Sensitivity Analysis - Results

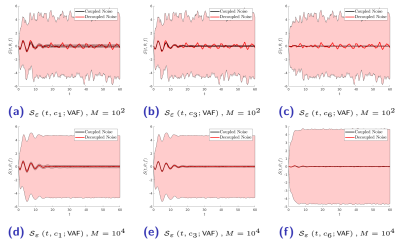


Figure: Sensitivity of solution due to perturbation of c ($\varepsilon = 0.01, \omega_0 = \frac{8}{9}, k_BT = 10^{-5}$)

Sensitivity Analysis - Results

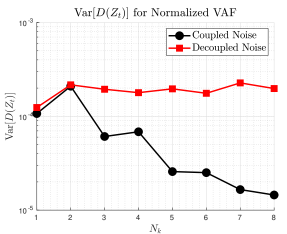


Figure: Var $[D\left(Z_{t}\right)]$ where the difference in solution, $D\left(Z_{t}\right)$, is between the nominal model with $N_{k}=n$ and the perturbed model with $N_{k}=n+1$ where $n=1,\cdots,8$ ($\omega_{0}=8/9,k_{B}T=10^{-5},t=100$)

Conclusion

Extended Variable GLE:

- Non-Markovian GLE problem converted to Markovian extended variable GLE problem using Prony series approximation.
- Explicit Euler scheme does not conserve 1^{st} and 2^{nd} moments of $\boldsymbol{X}(t)$ and $\boldsymbol{V}(t)$ unlike Splitting scheme.
- Number of modes used in Prony series approximation affects the memory kernel fit.

Sensitivity Analysis:

- Significant reduction in variance of the calculated sensitivity on using a common coupled random path for the diffusion term of GLE.
- Order of variance reduction for $D\left(Z_{t}\right)$ was observed to be $\mathcal{O}\left(\varepsilon^{2}\right)$ for coupled noise and $\mathcal{O}\left(1\right)$ for decoupled noise.
- Order of variance reduction for Δ_c was observed to be $\mathcal{O}\left(M^{-1}\right)$ for coupled noise and $\mathcal{O}\left(\varepsilon^{-2}M^{-2}\right)$ for decoupled noise.

References I



Andrew D. Baczewski and Stephen D. Bond.

Numerical integration of the extended variable generalized Langevin equation with a positive Prony representable memory kernel.

Journal of Chemical Physics, 139(4), 2013.



Bruce J. Berne, Jean Pierre Boon, and Stuart A. Rice.

On the calculation of autocorrelation functions of dynamical variables.

The Journal of Chemical Physics, 45(4):1086–1096, 1966.



M. A. Despósito and A. D. Viñales.

Subdiffusive behavior in a trapping potential: Mean square displacement and velocity autocorrelation function.

Physical Review E - Statistical, Nonlinear, and Soft Matter Physics, 80(2):1–7, 2009.



Eric J. Hall, Markos A. Katsoulakis, and Luc Rey-Bellet.

Uncertainty quantification for generalized Langevin dynamics.

Journal of Chemical Physics, 145(22), 2016.

References II



D. J. Higham.

An algorithmic introduction to numerical simulation of stochastic differential equations.

SIAM Review, 43(3):525-546, 2001.



K. Kawasaki.

Simple derivations of generalized linear and nonlinear Langevin equations.

Journal of Physics A: General Physics, 6(9):1289–1295, 1973.



Ben Leimkuhler and Charles Matthews.

Molecular Dynamics.

Springer International Publishing, 2015.



Martin Lysy, Natesh S. Pillai, David B. Hill, M. Gregory Forest, John W.R. Mellnik, Paula A. Vasquez, and Scott A. McKinley.

Model Comparison and Assessment for Single Particle Tracking in Biological Fluids.

Journal of the American Statistical Association, 111(516):1413–1426, 2016.

References III



A. D. Viñales and M. A. Despósito.

Anomalous diffusion: Exact solution of the generalized Langevin equation for harmonically bounded particle.

Physical Review E - Statistical, Nonlinear, and Soft Matter Physics, 73(1):5–8, 2006.