

ATLAS IMPLEMENTATION: USER GUIDE

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1. ATLAS OVERVIEW

The ATLAS algorithm was created by Miles Crosskey and Mauro Maggioni in the publication [1]. The overview below closely following sections 2 and 3 of [1], and for more details it is recommended to review those sections.

The ATLAS algorithm takes as input a simulator for a Markovian stochastic system whose dynamics are assumed to be concentrated near a low-dimensional manifold. The algorithm constructs the manifold from a sampling of points and approximate global dynamics from the local dynamics at patches of the manifold. The output is a reduced simulator in lower

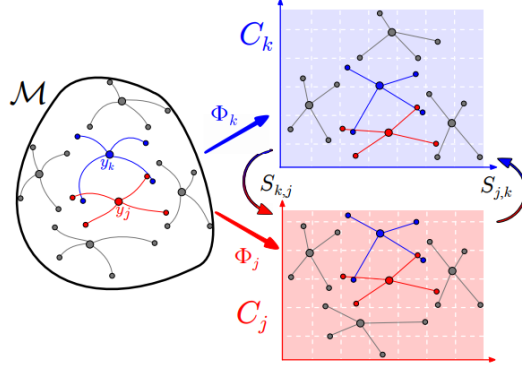


FIGURE 1. Learning the ATLAS: larger circles are net points, smaller circles are path endpoints [1]

dimensions with a coarser time scale and which approximates long term statistics of the original simulator.

The input stochastic system of interest $(X_t)_{t \geq 0}$ assumed to have state space in \mathbb{R}^D but concentrate near a manifold (the *effective state space*) \mathcal{M} of dimension $d \ll D$. The main inputs for the algorithm are the simulator \mathcal{S} for $(X_t)_{t \geq 0}$, a sampling of the manifold \mathcal{M} , the effective state space dimension d , and spatial granularity parameter δ .

The algorithm has three main steps:

- (i) **Net Construction:** Find set of points $\Gamma = \{y_k\}$ on \mathcal{M} which are well-distributed on \mathcal{M} and which are at least δ away from each other
- (ii) **Learning the ATLAS** (Figure 1):
 - (a) For each y_k , learn mapping Φ_k from neighborhood of y_k to local coordinate *chart* $C_k \subseteq \mathbb{R}^d$. The mapping Φ_k is learned via the Landmark Multi-Dimensional Scaling Algorithm (LMDS)
 - (b) Learn pairwise *transition maps* for changing coordinates between overlapping charts
- (iii) **Learning the Simulator**
 - (a) For each y_k , run ensemble of paths from y_k , map paths to coordinate chart $C_k \subseteq \mathbb{R}^d$
 - (b) Use mapped paths to learn a constant-coefficient SDE on the chart in \mathbb{R}^d , and use transition maps to stitch together the SDEs

2. NET CONSTRUCTION

A δ -net is a set of points $\{y_k\}_{k \in \Gamma} \subseteq \mathcal{M}$ such that with $d(y_i, y_j) > \delta$ for each i, j and such that for each $x \in \mathcal{M}$ there exists $y \in \{y_k\}_{k \in \Gamma}$ such that $d(x, y) < 2\delta$. Two net points y_i, y_j are connected neighbors if $d(y_i, y_j) < 2\delta$, and we denote this by $i \sim j$.

Many algorithms exist for constructing covering nets in this fashion, though a simple brute force algorithm for finding the net is downsampling a sufficiently dense sampling of \mathcal{M} . Hence, construction of the δ -net does assume the ability to sample \mathcal{M} .

3. LEARNING THE COORDINATE CHARTS

Given net point y_k , a mapping Φ_k is constructed from y_k and the connected neighbors $\{y_j\}_{j \sim k}$ to \mathbb{R}^d . The chart C_k is the image of the mapping $\Phi_k(B_{2\delta}(y_k)) \rightarrow \mathbb{R}^d$. For ATLAS, the Φ_k map is from the Landmark Multidimensional Scaling algorithm (LMDS) [2]. This is an extension of the Multidimensional scaling algorithm (MDS) [3].

Given y_k in the δ -net, the landmark points $\mathcal{A}_k \subseteq \mathcal{M}$ for y_k should be roughly δ away from y_k and well-spread in d directions. The landmarks are obtained from taking $m \geq d$ paths from the original simulator started at y_k . Then, all neighboring landmark points are collected in $L_k = \bigcup_{j \sim k} \mathcal{A}_j$, and the chart C_k is the image of $B_{2\delta}(y_k)$ under LMDS applied with landmarks L .

4. LEARNING THE SIMULATORS

For each y_k , the associated SDE for C_k , is learned from running p sample paths from the original simulator S up to time t_0 (on the order of δ). Each chart C_k has an associated simulator modeling an Ito diffusion SDE with constant coefficients:

$$(1) \quad d\bar{X}_t = \bar{b}_k dt + \bar{\sigma}_k dW_t \quad \bar{b}_k \in \mathbb{R}^d, \bar{\sigma}_k \in \mathbb{R}^{d \times d}$$

The solution to (1) is a Gaussian with mean $\bar{b}_k t_0$, covariance $\bar{\sigma}_k \bar{\sigma}_k^T t_0$. Thus, b_k and σ_k are estimated by taking sample mean \bar{b}_k and covariance $\bar{\sigma}_k$ of the endpoints of the p paths simulated.

After iterating through all net points y_k , the simulators $(\hat{S}_k)_{k \in \Gamma}$ with parameters $(\bar{b}_k, \bar{\sigma}_k)_{k \in \Gamma}$ are obtained.

4.1. Learning the Transition Maps and Simulating the SDE. In order to combine the above simulators, transition maps are needed between different charts, in particular near landmarks which overlap, which are denoted as $L_{k,j} = A_k \cup A_j$ for $k \sim j$. The transition map

T_{kj} from C_k to L_j is defined as $X^\dagger Y$, namely the solution to the least squares minimization $\|XT - Y\|_2$, where $X = \Phi_k(L_{k,j})$ and $Y = \Phi_j(L_{k,j})$. For $x \in C_k$, coordinates are changed through the transition

$$S_{k,j}(x) = (x - \mu_{k,j})T_{k,j} + \mu_{j,k},$$

where $\mu_{k,j}$ is the mean vector of X and $\mu_{j,k}$ is the mean vector of y .

Finally, the simulations are extended to a global SDE which advances in time by the following steps:

- Given simulation at time t , at $x \in \mathbb{C}_i \subseteq \mathbb{R}^d$
- Find closest coordinate chart $C_{i'}$ to x
- Compute a forward Euler step for simulator \hat{S}_i :

$$x \leftarrow x + \hat{S}_i \bar{b}_{i'} + \eta \hat{S}_i \bar{\sigma}_{i'} \sqrt{\delta t} \quad \eta \sim \mathcal{N}(0, I_d)$$

5. ATLAS IMPLEMENTATION

The ATLAS algorithm assumes an input stochastic ODE simulator S which can be started at a given initial condition and run for a given period of time. The simulator coded (`simulator.m`) uses the Euler-Maruyama method to simulate the SDE

$$dX_t = f(x)dt + dW_t, \quad X(0) = X_0,$$

where the function $f(x)$ can be specified by the user. To implement this simulator in ATLAS, we wrote a function for the simulator to be run by $S(X_0, m, T)$, which simulates m paths starting at $X(0) = X_0$ for time T and stores the endpoints of these paths.

TO DO: Adjust section based on current code

The ATLAS code is organized around the main construction steps outlined in the algorithm (see Figure 1):

- Constructing the δ -net $\{y_k\}_{n=1}^N$ (`delta_net.m`)
- Creating landmarks \mathcal{A} for each y_k in the δ -net and finding neighbors (`construction.m`)
- Constructing charts C_k for each y_k in the δ -net (`construction.m`)
- Learning the simulator associated to each chart C_k (`construct_local_sde()` in `construction.m`)
- Constructing a timestep function for the new approximate simulator (function `learned_simulator_step.m`)

The δ -net construction follows a simple brute force method: given an initial dense sampling of \mathcal{M} , downsample until the δ -net coarseness and covering requirement are met. The δ -net is stored an array $net = [y_1 \ y_2 \dots \ y_k]$, and a graph is created, where two net points

input : Simulator \mathcal{S} , Manifold sampling $\{x_j\}$, distance ρ

output: Reduced Simulator $\hat{\mathcal{S}}$

$\{y_k\}_{k \in \Gamma} = \delta - \text{net}(\{x_j\});$

for $k \in \Gamma$ **do**

Create landmarks for LMDS around y_k ;

$\{a_{k,l}\}_{l=1}^m = \mathcal{S}(y_k, m, t_0);$

$A_k = y_k \cup \{a_{k,l}\};$

end

Create charts and local simulators;

for $k \in \Gamma$ **do**

Simulate paths for creating local simulator;

$\{x_{k,l}\}_{l=1}^p = \mathcal{S}(y_k, p, t_0);$

Compute charts and chart centers;

$L_k = \bigcup_{i \sim k} A_i;$

$[L'_k, \{x'_{k,l}\}_{l=1}^p] = \text{LMDS}(L_k, \{x_{k,l}\}_{l=1}^p);$

$\{\mathcal{S}.c_{k,j}\} = \bigcup_{j \sim k} \{y'_j\}$ in L'_k ;

Center chart C_k and compute diffusion, drift;

$C_k = C_k - c_{k,k};$

$\hat{\mathcal{S}}.\bar{b}_k = \sum_l x'_{k,l}/pt_0;$

$\hat{\mathcal{S}}.\bar{\sigma}_k = (\text{Cov}(x'_{k,l})/t_0)^2;$

Compute transition maps T between charts;

for $j \sim k, j < k$ **do**

$L_{k,j} = A_k \cup A_j;$

$L'_{k,j} = L_{k,j}$ in chart C_k coordinates;

$L'_{j,k} = L_{j,k}$ in chart C_j coordinates;

$\hat{\mathcal{S}}.\mu_{k,j} = \mathbb{E}[L'_{k,j}];$

$\hat{\mathcal{S}}.\mu_{j,k} = \mathbb{E}[L'_{j,k}];$

$\hat{\mathcal{S}}.T_{k,j} = (L'_{k,j} - \mu_{k,j})^\dagger (L'_{j,k} - \mu_{j,k})^\dagger;$

end

end

Algorithm 1: Pseudocode for construction of the ATLAS, closely following [1] and implemented in `construction.m` function

are neighbors if $d(y_i, y_j) < 2\delta$. The edges are stored in `edges` and the neighbors in an array `neighbors` where each row is the length of the maximum degree, and row i has first $\text{degree}(i)$ entries as the neighbors of y_i , with zeros for the rest of the row.

All landmarks are stored in a three-dimensional array `L` where `L(:, :, n)` lists all landmarks associated to the net point y_n . The paths sample paths $\{x_{n,l}\}_{l=1}^p$ associated to y_n

are similarly stored in $\mathbf{X}(:, :, \mathbf{n})$ where \mathbf{X} is a three dimensional array storing sample paths from the net points.

For constructing charts, LMDS and MDS are implemented following the algorithm outlined in [2] and outlined in the ATLAS description earlier.

Further, the notation in 1 uses $\hat{S}.b_k$, $\hat{S}.\sigma_k$, and $\hat{S}.c_k$ to denote the drift vectors, diffusion coefficients and chart centers for chart C_k . Following this notation, we store these variables in a struct array *newS* where *newS*(*n*).*b* denotes the drift vector for chart y_k , and likewise for the other properties.

5.1. **Examples.** TO DO: Use language for a User Guide, instead of a report

(2)
$$dX_t = -\nabla V(x)dt + dW_t$$

with potential

$$V(x) = 16x^2(x - 1)^2 + \frac{1}{6} \cos(100\pi x),$$

also tested in [1].

A great visual example for the ATLAS algorithm is to apply to potential-driven problems, and to reconstruct potentials by the effective potentials obtained from the ATLAS algorithm. For 1D problems, *LMDS* is trivial and undoing it just requires multiplying by ± 1 . Then, the effective potential in a neighborhood of net point y_k is found by undoing MDS for b_k associated to the simulator for chart C_k (equation (1)) and then integrating. Unfortunately, a 1D problem is an extremely special case and in general we cannot construct the inverse for Φ_k .

5.1.1. *Two-Well Rough Potential.* This example is a simple test case to see if the ATLAS algorithm can take a system with microscale interactions and output a homogenized system with dynamics approximating macroscale interactions of the original. The high-frequency component in $V(x)$ from (2) emulate such microscale interactions, and homogenization from ATLAS should ideally return a system with smooth potential $U(x) = 16x^2(x - 1)^2$.

Following [1], the implementation for equation (2) has $\delta = 0.1$, $dt = 0.00005$ for the original simulator. The initial δ -net was created from subsampling points from the set of linear spaced points with distance 0.01 in $[-.3, 1.3]$.

As shown in our Figure 2(comparable to Figure 13 in [1]), the effective potential $\hat{V}(x)$ from ATLAS is a homogenized version of the rough potential $V(x)$. The ATLAS simulator obtained in this example has timestep roughly 100 times the original, so long simulations from the ATLAS will run about 100 times faster than the original simulator.

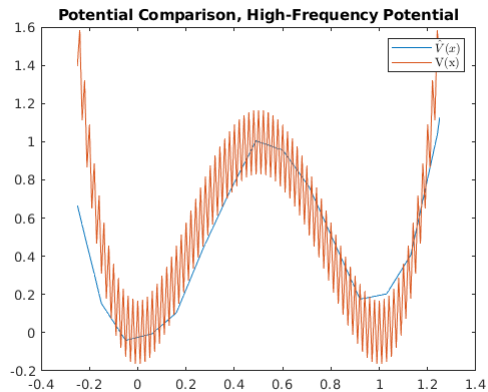


FIGURE 2. Effective potential $\hat{V}(x)$ from ATLAS plotted against true potential $V(x) = U(x) + \frac{1}{6} \cos(100\pi x)$

REFERENCES

- [1] M. Crosskey and M. Maggioni. Atlas: a geometric approach to learning high-dimensional stochastic systems near manifolds. *Multiscale Modeling & Simulation*, 15(1):110–156, 2017.
- [2] V. De Silva and J. B. Tenenbaum. Sparse multidimensional scaling using landmark points. Technical report, Technical report, Stanford University, 2004.
- [3] W. S. Torgerson. Multidimensional scaling: I. theory and method. *Psychometrika*, 17(4):401–419, Dec 1952.