Simulate binary-state epigenome evolution

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Assume the epigenome is a sequence of auto-correlated sequence of binary-state random variables. The auto-correlation reflects the organization of the epigenome as alternating domains bearing a certain modification or not. Let the epigenomic sequence be $S = s_1 s_2 \dots s_N$. As a graphical model, neighboring sites are connected with undirected edges. S evolves over time. Assume the stationary distribution for the epigenome has a Gibbs distribution that factorize over pairs of neighboring sites:

$$\Pr(S) = \frac{1}{Z} \exp\left\{\phi(s_1) + \sum_{n=1}^{N-1} \phi(s_n, s_{n+1}) + \phi(s_N)\right\}$$
(1)

The evolution of an individual site is context-dependent. The instantaneous mutation rate from state s to the alternative state \bar{s} is $\gamma(l,s,r)$, where l,s and r are the states of three consecutive sites. For a time interval [0,t), given that the states of l and r are not changed, then the states of site s follows a continouse-time Markov chain, and the holding time thus follows an exponential distribution. An observation of the path can be summarized with

$$L = \{s(0), k, \{t_i\}_{i=1}^k, t\},\$$

where s(0) is the state at time 0, k is the total number of jumps, t_i is the time when the i-th jump occurred, and t is the total length of the time interval.

Suppose L_l and L_r are given paths of two neighboring sites, then the union of their jumping times

$$\{0,t\} \cup \{t_{li}\}_{i=1}^{k_l} \cup \{t_{ri}\}_{i=1}^{k_r}$$

defines time intervals, within each of which the states of l and r stayed constant.

1 Stationary Gibbs measure as Markov chain

The stationary distribution in (1) is equivalent to the distribution of a Markov chain. We can derive the relationship between the factors in (1) and the transition probabilities of the Markov chain. The pair-wise potentials are $Q(a,b) = \exp(\phi(a,b))$, where $a,b \in \{0,1\}$ are binary states. The largest eigen value of Q is

$$q = \frac{1}{2} \{Q_{00} + Q_{11} + \sqrt{\Delta}\}, \text{ where } \Delta = (Q_{00} - Q_{11})^2 + 4Q_{01}Q_{10}.$$

Let r be a right eigenvector of Q corresponding to q, then we have $\frac{r_0}{r_1} = \frac{Q_{01}}{q - Q_{00}} = \frac{q - Q_{11}}{Q_{10}}$. Then the Markov chain transition matrix is

$$T(a,b) = \frac{Q(a,b)r(b)}{qr(a)}$$
, where $a, b \in \{0,1\}$.

To be more specific,

$$T(1,1) = \frac{2Q_{11}}{Q_{00} + Q_{11} + \sqrt{\Delta}},$$

$$T(0,0) = \frac{2Q_{00}}{Q_{00} + Q_{11} + \sqrt{\Delta}},$$

$$T(0,1) = \frac{4Q_{01}Q_{10}}{(Q_{00} + \sqrt{\Delta})^2 - Q_{11}^2},$$

$$T(1,0) = \frac{4Q_{01}Q_{10}}{(Q_{11} + \sqrt{\Delta})^2 - Q_{00}^2}.$$

$$(2)$$

The expected methylation level is thus $1-\frac{2Q_{01}Q_{10}}{(Q_{00}-Q_{11})^2+4Q_{01}Q_{10}+(Q_{11}-Q_{00})\sqrt{\Delta}}$

2 Relationship between mutation rates and stationary distribution

What transition rate function γ can lead to a stationary distribution determined by ϕ ? The Proposition 1 of Jensen & Pedersen (2000) gives a sufficient condition:

$$\frac{\gamma(l,s,r)}{\gamma(l,\bar{s},r)} = \frac{\exp(\phi(l,\bar{s}) + \phi(\bar{s},r))}{\exp(\phi(l,s) + \phi(s,r))},\tag{3}$$

which is derived from the reversibility property of the stationary distribution.

The proposition 2 gives a way of specifying γ from ϕ : mutation densities γ satisfy the relation (4) if and only if we can write

$$\log(\gamma(l,s,r)) = -\psi(l,s,r) + \ell(l,r),\tag{4}$$

The ℓ function in the original proposition is written as $\ell(s,t;l,r)$, which is symmetric in (s,t). The states in our problem setting are binary, thus ℓ is only a function of the two neighboring states (l,r). Moreover, we can directly verify that if we define mutation rates as follows,

$$\log(\gamma(l,s,r)) = \ell(l,r) + (\phi(l,\bar{s}) + \phi(\bar{s},r)),\tag{5}$$

then the rates satisfy the condition (4).

We can organzie the mutaion rates into a 8×8 matrix as follows:

$$\Gamma = \begin{pmatrix} 000 & 010 & 001 & 011 & 100 & 110 & 101 & 111 \\ 000 & 0 & a & 0 & 0 & 0 & 0 & 0 \\ 001 & b & . & 0 & 0 & 0 & 0 & 0 \\ 001 & 0 & 0 & . & c & 0 & 0 & 0 & 0 \\ 0 & 0 & d & . & 0 & 0 & 0 & 0 \\ 110 & 0 & 0 & 0 & 0 & . & c & 0 & 0 \\ 101 & 0 & 0 & 0 & 0 & 0 & . & e \\ 111 & 0 & 0 & 0 & 0 & 0 & 0 & f & . \end{pmatrix}$$

$$(6)$$

If a methylome sequence with mutation rates Γ has stationary distribution (1), then the condition in (4) holds, which leads to the following constraints on the mutation rates:

$$ad^2e = bc^2f. (7)$$

Given the mutation rates a, b, c, d, it is sufficient to derive the relationship between the potentials:

$$\phi(0,0) = \phi(0,1) + \frac{1}{2}\log(\frac{b}{a}), \text{ and } \phi(1,1) = \phi(0,1) + \frac{1}{2}\log(\frac{bc^2}{ad^2}). \tag{8}$$

In summary, the mutation rate matrix can have 5 free parameters. If we add a constraint on the expected number of changes per unit time, then there will be only 4 free parameters. The two ratios $\frac{b}{a}$ and $\frac{bc^2}{ad^2}$ can uniquely determine the stationary distribution (1) through equation (8).

3 Simulation scheme

We model the evolution of the entire sequence that with a continuous time Makov chain that allows instantaneous jump from one sequence to another only if they differ at one position. The jumping rate at each position is dependent on its state and the states of its neighboring sites. We assume that the states of the first and last sites are fixed throughout evolution.

Consider the $2^N \times 2^N$ transition rate matrix M, for any methylome a and methylome b that have a single difference at a position where a has state j and b has state \bar{j} , and the neighboring positions have states i and k in both methylomes, the rate of such a jump is $\lambda_{ijk} = \gamma(i,j,k) = \Gamma_{ijk,i\bar{j}k}$.

Given the current methylome a, the holding time

$$X_a \sim Exp(-M_{aa})$$

is an exponential variable. Its rate parameter $-M_{aa}$ is the sum of all instantaneous rates for jumps from a to a methylome that only differs with a at one position:

$$-M_{aa} = \sum_{i,j,k} c_{ijk}(a)\lambda_{ijk},$$

where $c_{ijk}(a) = \sum_{n=1}^{N-2} I(a_n = i, a_{n+1} = j, a_{n+2} = k)$ is the total number of the tripplet pattern ijk in methylome a.

Given that the first jump happened, the probability that the jump occurred in the context of ijk is proportional to $c_{ijk}(a)\lambda_{ijk}$. Given that a jump happend in context ijk, the jump is equally likely among positions with this context. The expected number of changes per site per unit time is $\sum_{ijk} \pi_{ijk}\lambda_{ijk}$, where π_{ijk} is the stationary probability of pattern ijk in the methylome.

In summary, we have the following simulation procedure for the evolution process for a methylome with N sites over time interval [0, T], given that the initial methylome is a(0):

- 1. Let $t \leftarrow 0$, and initialize all paths $L_n = \{a_n(0), k_n = 0, T_n = \emptyset, t\}$, for $n = 1, \dots, N$.
- 2. While t < T:

(a) Generate
$$x \sim \text{Exp}(-M_{a(t)a(t)})$$
, where $-M_{a(t)a(t)} = \sum_{i,j,k} c_{ijk}(a(t))\lambda_{ijk}$.

If t + x < T

– Choose pattern ijk from $\{ijk: i, j, k \in \{0, 1\}\}$ with probability proportional to $c_{ijk}(a(t))\lambda_{ijk}$.

- Scan methylome a(t), uniformaly choose one position n out of the c_{ijk} positions all with the pattern ijk in a(t).
- Set $a(t+x) \leftarrow a(t)_{1...n-1} \overline{a(t)_n} a(t)_{n+1...N}$.
- Add jump time to the path of position n:

$$k_n \leftarrow k_n + 1, \ T_n \leftarrow T_n \cup \{t + x\}.$$

Else
$$a(T) \leftarrow a(t)$$
.
(b) $t \leftarrow t + x$

4 Parameter inference

When we're given the complete epigenome evolution path from time 0 to time t, can we effectively recover the initial distribution and mutation parameters and evolutionary time? We are interested in the parameters describing the evolutionary process, which are the mutation rates $\{\lambda_{ijk}\}$. These parameters will be inferred from the state changes in the evolutionary path of the entire epigenome. Meanwhile, we do not require the process to be stationary, so we are also interested in the epigenome properties at time 0, which are characterized by the Markov chain transition probabilities T_0 . These transition probabilities are easy to infer given the complete observations at the time-0 epigenome.

Let
$$c_{ij} = \sum_{n=1}^{N-1} I\{s_n(0) = i, s_{n+1}(0) = j\}$$
. Then

$$\hat{T}(0,0) = \frac{c_{00}}{\sum_{n=1}^{N-1} I\{s_n(0) = 0\}}, \ \hat{T}(1,1) = \frac{c_{11}}{\sum_{n=1}^{N-1} I\{s_n(0) = 1\}}.$$

Let's first assume that the time span of this complete evolutionary history is known, i.e. the value of t is give.

Recall that $L_n = \{s_n(0), K, \{t_k\}_{k=1}^K, t\}$ is a full path at position n in the epigenome. We can pool all the jumping times at all positions in to a sorted sequence $J = \{(t_m, \mathsf{pos}_m, \mathsf{context}_m)\}_{m=1}^M$, where pos_m is the position of the m-th jump in the entire evolutionary history of the epigenome, $\mathsf{context}_m$ is the 3-tuple context of the mutation.

Let $\Delta_m = t_m - t_{m-1}$ be the holding time before the m-th jump. Then Δ_m is an exponential variable

$$\Delta_m \sim \operatorname{Exp}(\lambda_m), \text{ where } \lambda_m = \sum_{i,j,k} c_{ijk}^{(t_m - \epsilon)} \lambda_{ijk},$$

where constant $\epsilon \in (0, \min_{1 \leq m \leq M} \{\Delta_m\})$ so that $c_{ijk}^{(t_m - \epsilon)}$ is the sequence context distribution between the (m-1)th jump and the m-th jump.

The likelihood function for parameters $\{\lambda_{ijk}\}$ is thus

$$L = \prod_{m=1}^{M} \lambda_m \exp(-\lambda_m \Delta_m) \times \frac{\lambda_{\text{context}_m}}{\lambda_m} \times \frac{1}{c_{\text{context}_m}^{(t_m - \epsilon)}}$$

$$= \prod_{m=1}^{M} \frac{\lambda_{\text{context}_m}}{c_{\text{context}_m}^{(t_m - \epsilon)}} \exp(-\lambda_m \Delta_m)$$
(9)

The log-likelihood function is

$$l = \sum_{ijk} \left(\sum_{m=1}^{M} (\log \lambda_{ijk} - \log c_{ijk}^{(t_m - \epsilon)}) \times I_{\{\text{context}_m = ijk\}} - c_{ijk}^{(t_m - \epsilon)} \lambda_{ijk} \Delta_m \right)$$

$$= \sum_{ijk} \left(J_{ijk} (\log \lambda_{ijk} - \log c_{ijk}^{(t_m - \epsilon)}) - D_{ijk} \lambda_{ijk} \right)$$
(10)

where $J_{ijk} = \sum_{m=1}^{M} I_{\{\text{context}_m = ijk}\}$, and $D_{ijk} = \sum_{m=1}^{M} c_{ijk} (t_m - \epsilon) \Delta_m$. Constraints on the mutation rates $\{\lambda_{ijk}\}$ as indicated in equations (6,7) are

$$\begin{cases}
\lambda_{001} = \lambda_{100} \\
\lambda_{011} = \lambda_{110} \\
\lambda_{000} \lambda_{110}^2 \lambda_{101} = \lambda_{010} \lambda_{100}^2 \lambda_{111}
\end{cases}$$
(11)

Then the log-likelihood function (10) becomes

$$l = J_{000} \log \lambda_{000} - D_{000} \lambda_{000} + J_{010} \log \lambda_{010} - D_{010} \lambda_{010} + J_{101} \log \lambda_{101} - D_{101} \lambda_{101} + (J_{100} + J_{001}) \log \lambda_{001} - (D_{100} + D_{001}) \lambda_{001} + (J_{011} + J_{110}) \log \lambda_{011} - (D_{011} + D_{110}) \lambda_{011} + J_{111} \log(\frac{\lambda_{000} \lambda_{011}^2 \lambda_{101}}{\lambda_{010} \lambda_{001}^2}) - D_{111} \frac{\lambda_{000} \lambda_{011}^2 \lambda_{101}}{\lambda_{010} \lambda_{001}^2}.$$
(12)

When the time span of this complete evolutionary history is unknown, the value of t is also a model parameter to be estimated. We assume that all the jumping times are expressed as a fraction of t. Then the problem of estimating the mutation rates and evolutionary time becomes identifiable. We need an extra constraint – the unit branch length corresponds to 1 expected mutation per site. This is a common constraint in phylogenetic studies.

Here we explain how to formulate this constraint on the $\{\lambda_{ijk}\}$ parameters. Given $\{\lambda_{ijk}\}$, according to (8) $\frac{\lambda_{010}}{\lambda_{000}}$ and $\frac{\lambda_{001}}{\lambda_{011}}$ can uniquely determine the stationary distribution of the epigenome that is described with a Gibbs measure of form (1). The Gibbs measure for the epigenomic sequence, in turn, is equivalent to a Markov chain (2). Given the Markov chain formulation, we can easily compute the expected abundance of triplet patterns $p_{ijk} = \frac{1}{N} \mathbb{E}(c_{ijk})$, where c_{ijk} is the frequency of the triplet pattern in an epigenomic sequence of length N sampled from the Gibbs distribution. Then, we can compute the expected number of changes per position per unit time as $\sum_{ijk} p_{ijk} \lambda_{ijk}$. When the evolutionary time is also unknown, we add the following constraint:

$$\sum_{ijk} p_{ijk} \lambda_{ijk} = 1, \tag{13}$$

where $\{p_{ijk}\}$ as explained above are functions of $\{\lambda_{ijk}\}$. We maximize the log-likelihood (10) over $\{\lambda_{ijk}\}\cup\{t\}$ under the constraints (11) and (13), where the unknown parameter t is buried within $\{D_{ijk}\}$ in (10).

Posterior distribution of a path given two neighboring paths We assume that the model parameters $\{\lambda_{ijk}\}$ and total evolutionary time t are known, the starting state of the epigenome are known except for position n, and that the jumping times $J = \{(t_m, \mathsf{pos}_m, \mathsf{context}_m)\}_{m=1}^M$ are known for all positions except position n. How can we estimate the posterior distribution of path L_n (a sequence of jumping times)?

$$\Pr(L_n|L_{-n}, \{\lambda_{ijk}\}) \propto \Pr(L_n \cup L_{-n})$$

We propose to use MCMC to sample from the posterior distribution of L_n .

Method 1. First, sample a starting state s_0 from a Bernoulli distribution with probabilities (π_0, π_1) . Then, propose a number K of jumps from a Poisson distribution with rate parameter $\lambda = \frac{1}{8} \sum \lambda_{ijk}$, which is chosen to approximate average mutation rate among different contexts. Given K, sample jumping times uniformly on the time interval (0,t), *i.e.* from the Dirichlet distribution with concentration parameters all equal to 1. Therefore, the probability (density) of proposing a specific path $L' = \{s_0, K, \{t_k\}_{k=1}^K, t\}$ is

$$q(L') = \pi_{s_0} \frac{\lambda^K \exp(-\lambda)}{K!} \frac{1}{(K-1)!}.$$

This is an *independence sampler*. Suppose the current guess of path at position n is L_n , we accept the move to the proposed path L' with probability

$$\alpha(L') = \min\{\frac{\pi(L', L_{-n})/q(L')}{\pi(L_n, L_{-n})/q(L_n)}, 1\},\tag{14}$$

where $\pi()$ is the complete data likelihood function (9).

Method 2. The previous method proposes paths from an approximately uniform distribution. This may be a very inefficient (high rejection rate) sampling process, although Jensen & Pedersen (2000) used a sampling procedure in the same spirit. If we use more information from the paths at neighboring sites, we may be able to improve the efficiency of sampling. The neighboring paths L_{n-1} and L_{n+1} fragments the evolutionary time interval (0,t) into smaller segments, within each of which the states at positions n-1 and n+1 stay unchanged.

- Collect the time intervals from site n-1 and site n+1, so that within each of the intervals the states of the neighboring sites are unchanged. Let the intervals be represented by a sorted array $\{t_0, t_1, \dots, t_M\}$.
- For m = 1, ..., M:
 - Let X be a random variable from an exponential distribution, representing the jumping time of the middle site given that the states of its two neighbors are constant. For a time interval $[t_{m-1},t_m]$ during which the neighboring sites' states are unchanged, suppose $X \sim \operatorname{Exp}(\lambda_{ijk})$, where i and k are the states of the two neighboring sites in this time interval, and j is the starting state of position n.
 - Let $t = t_{m-1}$, and a sample value of X = x.
 - While $t + x < t_m$:
 - (1) Add 1 to the number of jumps $k_n \leftarrow k_n + 1$.

- (2) Update $T_n \leftarrow T_n \cup \{t+x\}$.
- (3) Update $t \leftarrow t + x$,
- (4) Sample another value of X = x from $\text{Exp}(\lambda_{ijk})$.

The proposal probability density q() is thus the product of the appropriate Exponential distribution probability densities for the holding times at position n. The proposal distribution is independent of any current guess of the path. Therefore, this is also an *independence sampler*. The rejection rule stays the same, as in (14).

Additional note: A jump at time τ in the path L_n will affect the probability of all jumps that occur after time τ regardless of their genomic position, because it affects $\{c_{ijk}(t)\}$ for any $t > \tau$. Therefore, we can't cancel out factors in the likelihood function. However, we may try to make things easy by only considering the evolutionary paths at positions n-2, n-1, n+1, and n+2.

5 Posterior distribution of a path given two neighboring paths in a tree structure with fixed leaf data

Suppose we are given a tree structure representing the evolutionary relationship between ancestral and extant species. Let (u, v), (v, a) (v, b) be three branches in the phylogenetic tree. Suppose we are given the evolutionary paths on the entire tree for all positions, except for the paths along these 3 branches at position n. In other words, the states at position n of species u, u, u are known, but how the state at position u evolved from u and diverge into states at u and u through their last common ancestor u is unknown. How do we sample from the posterior distribution of u which is the path at position u restricted to branches with one end as node u and the other end with known states.

Hobolth & Stone (2009) reviewed and compared three approaches to sample paths of discrete-state continuous-time Markov chain conditional on end-point states, namely the rejection sampling, direct sampling and uniformization. Method 1 in the previous section is similar to uniformization, since the neighboring sites may have state changes, and the rate of jumps depends on the contemporary states of the neighboring sites. We proposed to use an averaged rate to for the Poisson distribution, and let the acceptance probability to do most of the work of correcting the proposal distribution towards the posterior distribution. Method 2 is forward sampling, which is the basis of rejection sampling.

Rejection sampling Do forward sampling as described in Method 2 for position n along the branch (u,v) and (v,a). Reject the path unless the end state agree with the given state at node a at position n. Then do forward sampling along the branch (v,b), with the accepted simulated state of node v. If the end state does not agree with the given state at node b at position a, start over the sampling from the beginning, a i.e. start over from node a. Because we are not taking site a and a and a into consideration, we are not directly sampling from the posterior distribution of the paths, therefore the rejection sampling procedure is a way to propose viable paths, then we use the MCMC acceptance rule (14) to reshape the proposal distribution to the posterior distribution.

A couple of performance statistics should be measured for this proposed sampling method. These include (1) the success rate for generating valid paths that satisfy the end conditions; (2) the acceptance rate of proposed valid paths in MCMC sampling.

References

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