# Maximum likelihood Phylogenetic model

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#### 1 Model and notation

A methylome is a sequence of CpG sites (or bins of CpG sites) with each site displaying a methylation state from the set  $\{0,1\}$ . During evolution, two processes jointly govern the methylation state inheritance across generations, and the correlation between neighboring sites within each generation.

The inheritance process is described with a continuous-time Markov process over the state space  $\{0,1\}$ . Let the initial distribution be  $\pi=(\pi_0,\pi_i)$ . Let the transition rate matrix matrix be

$$Q = \begin{bmatrix} -\lambda & \lambda \\ \eta & -\eta \end{bmatrix}.$$

In continuous-time Markov process, the transition probability between two time points separated by time t is determined by two terms  $(t(\lambda + \eta), \lambda/\eta)$ . Therefore, we let  $\eta = 1 - \lambda$ ,  $\lambda \in (0, 1)$ . The transition probability matrix becomes

$$P(t) = exp(Qt) = \begin{bmatrix} 1 - \lambda T & \lambda T \\ (1 - \lambda)T & 1 - (1 - \lambda)T \end{bmatrix},$$

where  $T = 1 - exp(-t) \in (0, 1)$  for t > 0.

The autocorrelation within a methylome is described with a discrete-time (corresponding to discrete CpG sites) Markov chain over the state space  $\{0,1\}$ . The initial distribution is  $\pi$  (only relevant in the root species). The transition probability matrix,

$$G = \begin{bmatrix} g_0 & 1 - g_0 \\ 1 - g_1 & g_1 \end{bmatrix}$$

is assumed to be homogeneous in all species (this assumption can be relaxed).

For a given set of extant species and their phylogenetic relationship, we use  $\tau = \{V, E\}$  to denote the phylogenetic tree, including the set of nodes, and the set of edges E. We assume that the tree structure is given, and the branch lengths are unknown parameters.

The model parameter space is thus

$$\Theta = \{E, \lambda, g_0, g_1, \pi_0\},\$$

where E is the collection of branch lengths, and  $\lambda$  is the transition rate from 0 to 1 in the continuous-time Markov process.  $\pi_0$  is the hypomethylation probability in the root node.

The two Markov processes are combined in the following way. The direction of inheritance process is obvious. For the discrete Markov process, let the first CpG site from the 5' end of the '+' strand to be the start position of a chain. The root methylome has no ancestor, so it is modeled only with the discrete time Markov chain described by  $\{\pi,G\}$ . For CpG sites at start position of a DNA fragment, the evolution of methylation states on this site is modeled only with the inheritance process described with  $\{\pi,Q,\tau\}$ . For CpG sites from internal nodes and leaf nodes that are not the start of a DNA fragment, we will combine the two processes to model its evolution. Let one such site be position j in node v. Let i=j-1 be the previous site in the DNA fragment. Let node u be v's parent in the phylogenetic tree. Use the notation  $u_j$  to denote the methylation state of node u at position j,  $u_j \in \{0,1\}$ . Let  $t_v = e(u,v)$  be the branch length between nodes u and v. We define the following transition probability:

$$\Pr(v_j|v_i, u_j) = \frac{G_{v_i v_j} P(t_v)_{u_j v_j}}{\sum_{s=0.1} G_{v_i s} P(t_v)_{u_j s}}$$
(1)

Assume hypomethylation probabilities at N sites are observed in the methylome at each leaf node of a phylogenetic tree. Let  $O_i$  be the observed hypomethylation probabilities at location i in all extant species,  $i=1,\ldots,N$ . Let  $O=O_1O_2\cdots O_N$  be the total observed methylation states at leaf nodes. Let  $S=S_1\cdots S_N$  be the unobserved methylation states at internal nodes.

## 2 Data Likelihood

The above description defines a hidden Markov model over the space

$$H = \{h \in \{0,1\}^{|V|} : \text{all possible combination of methylation states over the phylogenetic tree} \}.$$

Each state specifies the history of the methylation evolution at CpG site, and thus we call it History of Methylation Evolution (HME). The transition probability between two HMEs can be calculated using transition probabilities in the previous section. Let  $h_i$ ,  $h_j$  be the HMEs at two neighboring sites in the genome. Let the root node be r. Let v be an internal or leaf node in the phylogenetic tree, and let its parent node be u. Let  $r_i$  be the methylation state at node r as specified by HME  $h_i$ . Let the HME transition probability matrix be  $A = \{a_{ij}\}$ , where

$$a_{ij} = \Pr(h_j | h_i) = G_{r_i r_j} \prod_{(u,v) \in E} P(v_j | v_i, u_j).$$

The data likelihood calculation can be written as

$$L = P(O|\Theta) = \sum_{H_1, \dots, H_N \in H} \Pr(H_1) \prod_{i=1}^{N} \Pr(H_i|H_{i-1}, \Theta),$$
 (2)

where  $H_t$  is a HME random variable at position t. H is the HME space.

## Forward algorithm

Alternatively, we can use the forward algorithm for likelihood calculation. Define the forward variables as below:

$$\alpha_1(i) = \Pr(O_1, H_1 = h_i | \Theta) = \Pr(h_i | \Theta) \Pr(O_1 | h_i)$$

$$\alpha_{n+1}(j) = \sum_{i=1,\dots,|H|} \alpha_n(i) a_{ij} \cdot \Pr(O_{n+1} | h_i)$$
(3)

The data likelihood is thus

$$L = P(O|\Theta) = \sum_{i=1,\dots,|H|} \alpha_N(i).$$

And the log-likelihood is

$$l = \log L = \log \sum_{i=1,\dots,|H|} \alpha_N(i).$$

## 3 First order derivative

Our parameter space is  $\Theta = \{E, \lambda, g_0, g_1, \pi_0\}$ . Let  $E' = \{e' = 1 - exp(-e) : \forall e \in E\}$ . There is a 1-1 mapping between E and E', so the model parameter space is equivalently  $\Theta = \{E', \lambda, g_0, g_1, \pi_0\}$ . For simplicity, we use the same notation  $\Theta$ . Let x be a parameter of interest.

$$\frac{\partial l}{\partial x} = \frac{1}{L} \sum_{i=1,\dots,|H|} \frac{\partial}{\partial x} \alpha_N(i)$$

## Recursion by forward algorithm

Using the recursive definition of forward variables,

$$\frac{\partial}{\partial x}\alpha_{n+1}(j) = \sum_{i=1,\dots,|H|} \left[ \frac{\partial}{\partial x}\alpha_n(i) \cdot a_{ij} + \alpha_n(i) \cdot \frac{\partial}{\partial x}a_{ij} \right] \cdot \Pr(O_{n+1}|h_j),$$

the first order derivative computation comes down to computing derivatives for HME transition probabilities  $a_{ij}$ , and the base case forward variables  $\alpha_1(i)$ , where  $i, j \in \{1, \dots, |H|\}$ .

Some Preparation The continuous-time Markov process' transition probability matrix corresponding to a given branch (u, v) contains two parameters:  $\lambda$ , and  $T_u = 1 - exp(-t_u)$ . The derivatives with respect to these two parameters are

$$\frac{\partial}{\partial \lambda} P(t) = T \cdot \begin{bmatrix} -1 & 1 \\ -1 & 1 \end{bmatrix}$$

$$\frac{\partial}{\partial T} P(t) = Q$$
(4)

where T = 1 - exp(-t).

The discrete-time Markov chain involves two parameters,  $g_0$  and  $g_1$ .

$$\frac{\partial}{\partial g_0} G_{ij} = \delta_{\{i=0\}} \cdot (2 * \delta_{\{j=0\}} - 1) 
\frac{\partial}{\partial g_1} G_{ij} = \delta_{\{i=1\}} \cdot (2 * \delta_{\{j=1\}} - 1)$$
(5)

The combined transition probability  $Pr(v_i|v_i,u_j)$  contains parameter  $T_v,g_{v_i}$ , and  $\lambda$ .

$$\frac{\partial}{\partial T_{v}} \operatorname{Pr}(v_{j}|v_{i}, u_{j}) = \frac{G_{v_{i}v_{j}} \cdot \frac{\partial}{\partial T_{v}} P(t_{v})_{u_{j}v_{j}}}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}v_{j}}\right]} - \frac{G_{v_{i}v_{j}} P(t_{v})_{u_{j}v_{j}} \cdot \left[\sum_{s=0,1} G_{v_{i}s} \frac{\partial}{\partial T_{v}} P(t_{v})_{u_{j}s}\right]}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}s}\right]^{2}}$$

$$\frac{\partial}{\partial g_{v_{i}}} \operatorname{Pr}(v_{j}|v_{i}, u_{j}) = \frac{\frac{\partial}{\partial g_{v_{i}}} G_{v_{i}v_{j}} \cdot P(t_{v})_{u_{j}v_{j}}}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}v_{j}}\right]} - \frac{G_{v_{i}v_{j}} P(t_{v})_{u_{j}v_{j}} \cdot \left[\sum_{s=0,1} \frac{\partial}{\partial g_{v_{i}}} G_{v_{i}s} P(t_{v})_{u_{j}s}\right]}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}s}\right]^{2}}$$

$$\frac{\partial}{\partial \lambda} \operatorname{Pr}(v_{j}|v_{i}, u_{j}) = \frac{G_{v_{i}v_{j}} \cdot \frac{\partial}{\partial \lambda} P(t_{v})_{u_{j}v_{j}}}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}s}\right]} - \frac{G_{v_{i}v_{j}} P(t_{v})_{u_{j}v_{j}} \cdot \left[\sum_{s=0,1} G_{v_{i}s} \frac{\partial}{\partial \lambda} P(t_{v})_{u_{j}s}\right]}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}s}\right]^{2}}$$

$$= \frac{G_{v_{i}v_{j}} T_{v} \cdot (2\delta_{\{v_{j}=1\}} - 1)}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}v_{j}} T_{v} \left[\sum_{s=0,1} G_{v_{i}s} (2\delta_{\{s=1\}} - 1)\right]}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}s}\right]^{2}}$$

$$= \frac{G_{v_{i}v_{j}} T_{v} \cdot (2\delta_{\{v_{j}=1\}} - 1)}{\left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}v_{j}} T_{v} \left[\sum_{s=0,1} G_{v_{i}s} P(t_{v})_{u_{j}s}\right]^{2}}$$

Derivative of  $\lambda$ 

$$\frac{\partial}{\partial \lambda} \alpha_1(i) = \alpha_1(i) \times \Big[ \sum_{(u,v) \in E} \frac{T_v \cdot (2\delta_{\{v_i=1\}} - 1)}{P(t_v)_{u_i v_i}} \Big]$$
$$\frac{\partial}{\partial \lambda} a_{ij} = a_{ij} \times \Big[ \sum_{(u,v) \in E} \frac{\frac{\partial}{\partial \lambda} P(v_j | v_i, u_j)}{P(v_j | v_i, u_j)} \Big]$$

Derivative of branch length  $T_v$ 

$$\frac{\partial}{\partial T_v} \alpha_1(i) = \alpha_1(i) \times \frac{Q_{u_i v_i}}{P(t_v)_{u_i v_i}}$$
$$\frac{\partial}{\partial T_v} a_{ij} = a_{ij} \times \frac{\frac{\partial}{\partial T_v} \Pr(v_j | v_i, u_j)}{\Pr(v_i | v_i, u_j)}$$

Derivative of transition probability  $q_s$ 

$$\frac{\partial}{\partial g_s} a_{ij} = a_{ij} \times \left[ \delta_{\{r_i = s\}} \times \frac{(2\delta_{\{v_j = s\}} - 1)}{G_{r_i r_j}} + \sum_{(u,v) \in E} \delta_{\{s = v_i\}} \times \frac{\frac{\partial}{\partial g_{v_i}} P(v_j | v_i, u_j)}{P(v_j | v_i, u_j)} \right]$$

Derivative of initial distribution  $\pi_0$ 

$$\frac{\partial}{\partial \pi_0} \alpha_1(i) = \alpha_1(i) \times \frac{2\delta_{\{r_i = 0\}} - 1}{\pi_{r_i}}$$

All other partial derivatives of  $\alpha_n(i)$  and  $a_{ij}$  that are not specified above have value 0.

## 4 Reduce transition complexity

The total number of possible transitions between HMEs is  $2^{2|V|}$ . We introduce constraints on transition between HMEs to reduce the computational complexity.

**Neighbor HME** Let  $N(h) = \{h' \in H : P(H_{n+1} = h' | H_n = h) > 0\}$ , be the neighbor set of an HME h. We always allow self-transition, i.e.  $h \in N(h), \forall h \in H$ . For any two HMEs, if they only differ by methylation state at one node, then they are neighbors. In addition, if two HMEs differ at root node, and differ at only one other node, they are neighbors.

The neighbor relationship defined above is symmetric, *i.e.* if  $h \in N(h')$  then  $h' \in N(h)$ . Each HME has 2|V| neighbors, including itself. Now, the total number of possible transitions between HMEs is  $2^{|V|} \times 2|V|$ . The new transition probabilities are defined as below:

$$a'_{ij} = 1_{h_j \in N(h_i)} \frac{a_{ij}}{\sum_{k \in N(i)} a_{ik}}$$

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