Markov invariants for phylogenetic rate matrices
derived from embedded submodels

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Abstract

We consider novel phylogenetic models with rate matrices that arise via the embedding of a progenitor model on a small number of character states, into a target model on a larger number of character states. Adapting representation-theoretic results from recent investigations of Markov invariants for the general rate matrix model, we give a prescription for identifying and counting Markov invariants for such 'symmetric embedded' models, and we provide enumerations of these for low-dimensional cases. The simplest example is a target model on 3 states, constructed from a general 2 state model: the '2→3' embedding. We show that for 2 taxa, there exist two invariants of quadratic degree, that can be used to directly infer pairwise distances from observed sequences under this model. A simple simulation study verifies their theoretical expected values, and suggests that, given the appropriateness of the model class, they have greater statistical power than the standard (log) Det invariant (which is of cubic degree for this case).

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1 Introduction

Phylogenetic inference based on molecular sequence data typically involves the selection of one or more specific models for state substitutions. There is a well-known hierarchy of classes of $4 \times 4$ rate matrices, with varying complexity and numbers of free parameters (Posada & Crandall, 1998). However, for a given data set it is not always (if ever) clear which substitution model is most appropriate or “best”. For example, the software package ModelTest (Posada & Crandall, 1998) selects a model of nucleotide substitution that best fits a given data set under a likelihood, information theoretic or Bayesian framework. While it is useful to be able compare results derived from different models, given the dangers of over-parametrisation the question of which datasets best conform to which class of models is difficult to resolve. Elaborations such as allowing rate variation across sites and invariant sites are also standard ingredients which allow for more flexibility in data-fitting. More extreme measures, which relinquish the conventional picture of species evolution via Markov models on trees, lead to generalisations such as mixtures (Pagel & Meade, 2004) (sites which have probabilities for following different models), “mosaics” (Woodhams et al., 2009) (edge classes or subtrees which have different weights for different models), or ultimately network models (Bandelt & Dress, 1992; Holland & Moulton, 2004), for example using acyclic directed graphs.

In recent work we have introduced so-called “Markov invariants” (Sumner et al., 2008; Sumner & Jarvis, 2009), which are polynomial quantities built up of the phylogenetic pattern frequencies or divergence arrays. Markov invariants are distinct from “phylogenetic invariants” (Cavender & Felsenstein, 1987; Lake, 1987) in that they are defined to behave as a (one-dimensional) “representation” of continuous unfolding of the Markov process. This means that as the Markov process proceeds in time by an amount $\tau$, the expectation value of the Markov invariants simply scales with a product of the multiplicative constants $\det(\text{tr}(Q_e)\tau)$, where $m_e = e^{Q_e\tau}$ is the transition matrix associated with the edge $e$. With this understanding, the “invariance” property of the Markov invariants is captured by the simple time-dependence

$$\tau \rightarrow \tau + \tau' \iff \text{tr}(Q_e)\tau \rightarrow \text{tr}(Q_e)\tau e^{\text{tr}(Q_e)\tau'} = e^{\text{tr}(Q_e)(\tau + \tau')}.$$ 

It is important to note that the definition of phylogenetic invariants stipulates no such constraint.

Markov invariants are phylogenetically informative for the most general Markov model, giving some information of both model parameters and tree topology, and can be implemented without the need for explicit parameter estimation (ie. via optimization of a likelihood function). These invariants generalise the “log Det” distance measure which has precisely this feature: pairwise distances can be directly estimated whose expected value turns out to be the sums of rate parameters multiplied by time. We have identified Markov invariants for diverse combinations of numbers of taxa and numbers of characters. For example there are three so-called “squangle” invariants (Sumner et al., 2008; Sumner & Jarvis, 2009) for quartets of taxa and four character states, whose values directly resolve and distinguish the three unrooted trees 12|34, 13|24, 14|23 for the most general Markov model, without the need for parameter estimation. Indeed, there are consistency arguments to suggest that the Markov invariants, as defined simply as functions of the phylogenetic pattern data, are in fact identical to their maximum likelihood estimators (or, more technically, belong to the ideal generated by the solutions of the likelihood equations). This is trivially true for the log Det estimator (Allman & Rhodes, 2009) and its proof in the general case is a subject of future work.

Notwithstanding these promising developments for the general Markov model, it is still of great interest to have at hand tools for exploring the full range of models available for phylogenetic inference. In this vein, Huelsenbeck et al. (2004) identified up to 203 submodels of the general time reversible model (GTR) for four characters, the number being based simply on the combinatorial
problem of counting compositions and refinements as parameters are turned on and off at various positions in the rate matrix.

In view of the discussion above concerning model classes, a natural criterion for model and sub-model selection is that of the multiplicative closure of the edge transition matrices, or semi-group property. Multiplicative closure is sometimes held out to be required for establishing the unrootedness of phylogenetic trees [Isaev, 2004; Semple & Steel, 2003], but for a given tree it seems not to be necessary. Its importance arises more directly from the methodology and interpretation of phylogenetic reconstruction. In doing tree searches for example, a swing of a leaf edge from one subtree to another entails a cut-and-rejoin operation: the incoming and outgoing Markov edge matrices from the source node of the originating leaf edge must be multiplied, while the Markov matrix from the target edge where the leaf is rejoined, must be expressed in turn as the product of Markov matrices for two new edges. Again, the possibility of extinctions along some edges, or of incomplete taxon sampling, suggests that, to allow for correct marginalisation, multiplicativity is mandatory for a consistent interpretation. It is clear that scarcely any of the GTR matrices identified by [Huelsenbeck et al., 2004] will be multiplicative (indeed, only some well-known models comply, for example the symmetric models such as the Kimura model, as well as Felsenstein’s TR, non-symmetrical model). Indeed it is easy to show that the GTR model itself is not multiplicative, and this poses serious interpretive questions if the GTR class is used in generalized models where more than a single rate matrix is implemented in the analysis.

In this paper we introduce, and explore through Markov invariants, new classes of submodels for a given number of characters, generated from general models in smaller numbers of characters. These we term ‘symmetric embeddings’. Clearly, in essence, these submodels contain similar information to the originating model in lower dimensions (with a smaller number of characters, and fewer parameters), and it is the manner in which this intuition is realised in technical detail, which we wish to elaborate here. In contrast to the naïve identification of submodels by the mere presence or absence of additional parameters however, our embedded models are by construction multiplicatively closed. Moreover we are able to adapt the technical setting of Markov invariants to this new situation, and so derive new invariants of different structure and polynomial degree from the standard ones, which play an equivalent role to them. Again, these new invariants fulfill the expectation (because their underlying models have fewer parameters) that they should be of lower degree than the standard ones.

In §2 below we introduce the symmetrically embedded models. These are given in a general setting, but we concentrate in detail on the general 2 state model embedded into 3 character models, called the ‘2 → 3’ case. A variety of Markov invariants, for diverse degrees and numbers of characters and taxa, is enumerated in §3 after adapting the group representation method for identifying Markov invariants [Sumner et al., 2008; Sumner & Jarvis, 2009] to the present setting. The simplest case is again that of ‘2 → 3’, with two taxa, where it is shown that, apart from the (degree 3) determinant function (guaranteed to exist for the general Markov model and any number of characters, and well known as the log Det measure), there are two additional quadratic degree invariants called $I_{3,1}$ and $I_{2,2}$. These are constructed explicitly and their properties are explored.

Finally in §4 the paper is summarised, and the conclusions supported by some simple simulated data analysed for comparison using “Det” invariant, or $I_{3,3}$ in our notation, as well as $I_{3,1}$ and $I_{2,2}$ invariants. As expected, the invariants of lower degree (and “weight”, see below), are apparently statistically better behaved, at least from this preliminary numerical test. The paper ends with some concluding remarks and prospects for further work. An appendix, §A, gives an adaptation of a technical representation-theoretic result from [Sumner et al., 2008] enabling the Markov invariants for embedded submodels to be enumerated and constructed as presented in §3 for low-dimensional cases.
2 Symmetrically embedded character substitution models

In this section we introduce the concept of symmetrically embedded models, concentrating initially on the two state case, and developing the analysis to be able to present the $2 \leftrightarrow 3$, $2 \leftrightarrow 4$ and $2 \leftrightarrow K$ rate matrices in detail. The discussion finishes with an overview of the general case.

Consider the rate matrix for the general Markov model on two characters,

$$Q = \begin{pmatrix} -\alpha & \beta \\ \alpha & -\beta \end{pmatrix} = \alpha \begin{pmatrix} -1 & 0 \\ 1 & 0 \end{pmatrix} + \beta \begin{pmatrix} 0 & 1 \\ 0 & -1 \end{pmatrix} = \alpha L_\alpha + \beta L_\beta$$

(1)

where as usual in the Markov matrix $m(t) := \exp(Qt)$, the matrix elements $m_{ab}$ have the interpretation

$$m_{ab} = \mathbb{P}[X(t) = a \mid X(0) = b]$$

for the random variable $X(t)$ in character space[1] describing the probability of change (along each edge after time $t$),

$$p_a(t) = \sum_{b=1}^{2} m_{ab} p_b(0),$$

(2)

or $p(t) = m(t) \cdot p(0)$, where we have $K = 2$ characters and the edge probability distribution is $p_a(t) = \mathbb{P}(X(t) = a)$.

We have chosen to write $Q$ in terms of the natural basis of column-sum zero ‘stochastic generator matrices’ \{\(L_\alpha, L_\beta\)\} of the group $GL_1(2)$; the subgroup of the general linear group $GL(2)$ of invertible $2 \times 2$ matrices together with the probabilistic constraint of unit-column sums [Johnson, 1983; Mourad, 2004]. This is relevant for considerations of multiplicative closure of models, which might arise in applications where different rate matrices are allowed on different parts of a phylogenetic tree; where potentially missing taxa may need to be inserted into edges; or where re-evaluations of phylogeny may require edge rearrangements. In this case of a general Markov rate model, in continuous time, closure of the product $M_1M_2 = \exp Q_1 \exp Q_2$ is guaranteed by the so-called BCH formula which requires closure of the commutator brackets $[Q_1, Q_2] := Q_1Q_2 - Q_2Q_1$ of the $Q$’s:

$$\exp Q_1 \exp Q_2 = \exp(Q_1 + Q_2 + \frac{1}{2}[Q_1, Q_2] + \frac{1}{12}[Q_1, [Q_1, Q_2]] - \frac{1}{12}[Q_2, [Q_1, Q_2]] + \cdots).$$

(3)

Referring to the BCH formula (3), it is immediately clear that closure is assured if the rate matrices form a Lie algebra, which in this case follows as we have chosen the most general two-state model. Specifically,

$$[L_\alpha, L_\beta] = L_\alpha - L_\beta = -[L_\beta, L_\alpha]$$

with of course $[L_\alpha, L_\alpha] = 0 = [L_\beta, L_\beta]$.

How can we use (1) to infer rate matrices for ‘target’ models on different numbers of characters (larger than 2)? A natural observation from the linear $m$-action on the array (vector) $p_a$ is that a similar linear action can be obtained not only on the components of $p$, but also on any homogeneous polynomials in the components. Specifically we can regard the $k+1$ distinct monomials in components of the $p$ at fixed degree $k$, as the formal components of a new, $k + 1$-dimensional array.

Consider for instance the case $k = 2$, $k + 1 = 3$, and the monomials $p_1^2$, $p_1p_2 = p_2p_1$, and $p_2^2$. We write four terms to emphasize that the new character probabilities, say $P_1 = p_1^2$,

\[1\]In our notation the the random change process is implemented by the left matrix action, so that the column sum of $m$ is unity (the column sum of $Q$ vanishes).
$P_2 = 2p_1p_2$, $P_3 = p_2^2$ are really components of a symmetric array (tensor) $p_{ab} := p_ap_b$, with of course $p_{ab} = p_{ba}$, and to motivate the choice of scaling; we use the binomial expansion $(p_1 + p_2)^2 = p_1^2 + 2p_1p_2 + p_2^2 = 1$. If we take the differential form of the change rule (2), $dp/dt = Qp$, and write the induced transformation on $P$ (considered as a three component vector) as $dP/dt = Q^{(3)}P$, it is then easy to infer $Q^{(3)}$ by considering $dp_{ab}/dt$ and referring to (1). Following this through we find that

$$Q^{(3)} = \begin{pmatrix}
-2\alpha & \beta & 0 \\
2\alpha & -\alpha - \beta & 2\beta \\
0 & \alpha & -2\beta
\end{pmatrix} = \alpha \begin{pmatrix}
-2 & 0 & 0 \\
2 & -1 & 0 \\
0 & 1 & 0
\end{pmatrix} + \beta \begin{pmatrix}
0 & 1 & 0 \\
0 & -1 & 2 \\
0 & 0 & -2
\end{pmatrix} \equiv \alpha L^{(3)}_{\alpha} + \beta L^{(3)}_{\beta}. \quad (4)$$

By construction, the new generator matrices $L^{(3)}_{\alpha}$ and $L^{(3)}_{\beta}$ form a subalgebra of the Lie algebra of $GL_1(3)$ and satisfy the same commutation relations as their $2 \times 2$ progenitors, namely $[L^{(3)}_{\alpha}, L^{(3)}_{\beta}] = L^{(3)}_{\alpha} - L^{(3)}_{\beta}$. Thus technically we have an embedding of the Lie algebra of $GL_1(2)$ into that of $GL_1(3)$, which we shall denote $2 \rightarrow 3$ (and of course multiplicative closure for this class of $3 \times 3$ model is guaranteed).

The generalisation to the $2 \rightarrow 4$, or $2 \rightarrow K$, character case is immediate. For $K = 4$ we have

$$Q^{(4)} = \begin{pmatrix}
-3\alpha & \beta & 0 & 0 \\
3\alpha & -2\alpha - \beta & 2\beta & 0 \\
0 & 2\alpha & -\alpha - 2\beta & 3\beta \\
0 & 0 & \alpha & -3\beta
\end{pmatrix} = \alpha \begin{pmatrix}
-3 & 0 & 0 & 0 \\
3 & -2 & 0 & 0 \\
0 & 2 & -1 & 0 \\
0 & 0 & 1 & 0
\end{pmatrix} + \beta \begin{pmatrix}
0 & 1 & 0 & 0 \\
0 & -1 & 2 & 0 \\
0 & 0 & -2 & 3 \\
0 & 0 & 0 & -3
\end{pmatrix} \equiv \alpha L^{(4)}_{\alpha} + \beta L^{(4)}_{\beta},$$

based on a totally symmetric, rank three tensor $p_{abc} := p_ap_bp_c$ with binomial constants of proportionality derived, as above, and using the constraint $(p_1 + p_2)^3 = 1$ to form the vector $P$ with four components $P_1 = p_1^3$, $P_2 = 3p_1p_2^2$, $P_3 = 3p_1^2p_2$ and $P_4 = p_2^3$. In the $2 \rightarrow K$ case (corresponding to a rank $k = K - 1$ tensor array $p_{a_1 \cdots a_k}$), the rate generator matrices $L^{(K)}_{\alpha}$, $L^{(K)}_{\beta}$ have lower and upper diagonal entries $K-1$, $\cdots$, 2, 1 and 1, 2, $\cdots$, $K-1$, respectively, with the diagonals ensuring that the zero column sum condition is satisfied. Again these matrices satisfy the same commutator bracket relations (Lie algebra) as their $2 \times 2$ progenitors and hence generate phylogenetic models that are guaranteed to satisfy the closure property.

## 3 Markov invariants

In this section we adopt the background context of group actions on which we base our general theorems on Markov invariants. Details are provided in §3 where we restate our previous technical results. We then explore the counting of Markov invariants for symmetrically embedded models in diverse dimensions (number of characters of the generating model, embedding rank and hence number of characters of the derived model, number of taxa, and polynomial degree of the invariant) and tabulate several low-dimensional cases. Finally we give details of the quadratic degree invariants for the $2 \rightarrow 3$ case, and explicit constructions of them along with the cubic, determinant function for comparison.

As explained above and in systematic terms in §3, embedded submodels are associated with particular matrix group constructions, whereby the character probability distribution $p$ for a starting model on $K'$ characters, is regarded as a progenitor for a target model deriving from a composite tensor array. If the starting model has dimension $K'$ and the tensor $p$ is of rank $k$ and totally symmetric (the only case we consider), then $p$ has $K = K'(K'+1)(K'+2)\cdots(K'+k-1)/k!$
components, and so $K$ is number of characters of the the target model. Table 4 gives a list of several cases of interest for low-dimensional examples; of course $K' = 2$, $k = 2, 3, \ldots K-1$ are the $2 \leftrightarrow 3$, $2 \leftrightarrow 4$, and $2 \leftrightarrow K$ cases already identified.

Markov invariants (see Sumner et al. (2008); Sumner & Jarvis (2009), and §A) are formally polynomials in the components of a phylogenetic tensor $P$ with components $P_{a_1 a_2 \ldots a_L}$ representing the probability of observing the character state pattern $(a_1, a_2, \ldots, a_L)$ at the leaves of the tree. Hence for $L$ taxa $P$ is a tensor with $k^L$ components, indexed by $L$ sets of $k$ multi-indices. Markov invariants are constructed such that, under time evolution associated with the model on the pendant edges, they change at most by a multiplicative factor. For clarity, at the pendant edges of the tree let $m_i, i = 1, \ldots, L$, be the $K' \times K'$ transition matrices of the starting model, and denote the embedding into the target model\footnote{In the previous discussion the corresponding rate matrices were distinguished by a superscript, $^{(K)}$.} as $M_i \equiv M(m_i), i = 1, \ldots, L$. Then given the transformation rule, (A-1), for $P$ for pendant edge evolution under the model, a Markov invariant $I$ must satisfy (A-4), namely

$$I(P') = \det(m_1)^{w_1} \det(m_2)^{w_2} \cdots \det(m_L)^{w_L} I(P),$$

for some integers $w_i$. We note that for a continuous-time Markov model with rate matrices $Q_i$, we have $m_i = e^{Q_i \tau}$ and $\det(m_i) = e^{\nu(Q_i) \tau}$, as in the introduction.

Recall that a partition $\mu$ of an non-negative integer $m$ is a set of non-negative integers $\lambda_1, \lambda_2, \ldots, \lambda_r$ such that $\lambda_1 + \lambda_2 + \ldots + \lambda_r = m$. It is usual to write $\mu = (\lambda_1, \lambda_2, \ldots, \lambda_r)$ with $\lambda_1 \geq \lambda_2 \geq \ldots \geq \lambda_r$ and to use exponents for repeated parts. For example we write $\mu = (4, 3, 3, 3, 2, 2, 1) \equiv (4, 3^3, 2^2, 1)$, with $\mu$ being a partition of 18. Markov invariants $I$ of degree $D$ are then identified by associating them with certain partitions of special shape, which in turn label certain irreducible group representation characters of the general linear group.

Whether admissible $\mu$ arise at each $D$ and number of leaves $L$, and how many occurrences thereof, must be answered for each case by evaluating a certain representation-theoretic branching rule (see §A Theorems 1 & 2, for details). Instances of such invariants are enumerated in Table 2 for the cases identified in Table 1. They are listed by $K'$, $k$, $K$ (to define the embedding type), by $L$ for small numbers of leaves, and then by degree $D$ up to 4. From the tables it is evident that there exists a plethora of Markov invariants for embedded submodels. Further information on the independent invariants for phylogenetic tensors constructed under the Markov model can be accessed by studying the isotropy subgroup of leaf permutations on a tree, as in Sumner & Jarvis (2009). We defer discussion on the general results, including commentary on cases of possible biological interest, to the conclusions.

For now we resume consideration of the lowest dimensional situation which motivated the present study, $2 \leftrightarrow 3$, and the lowest-degree (quadratic) invariants for the simplest situation of two leaves ($L = 2$), namely $K' = 2$, $k = 2$, $K = 3$, $L = 2$, $D = 2$. Here we outline briefly the manner in which these objects are constructed by standard tensor symmetrisation techniques. The end result will be the explicit forms (5) and (6) below.

Recall that we handle the 3 state embedded model via a rank two phylogenetic probability

| $K'$ | 2 | 2 | 2 | 3 | 3 | 4 | 4 |
|------|---|---|---|---|---|---|---|
| $k$  | 2 | 3 | 4 | 2 | 3 | 2 | 3 |
| $K$  | 3 | 4 | 5 | 6 | 10 | 10 | 20 |

Table 1: Identification of embedded submodels for low-dimensional cases of interest. ($K', k, K$) gives the number of characters of the progenitor model, the rank of the embedding tensor, and the number of characters of the target model, respectively, with $K = K'(K' + 1)(\cdots)(K' + k - 1)/k!$.  

For now we resume consideration of the lowest dimensional situation which motivated the present study, $2 \leftrightarrow 3$, and the lowest-degree (quadratic) invariants for the simplest situation of two leaves ($L = 2$), namely $K' = 2$, $k = 2$, $K = 3$, $L = 2$, $D = 2$. Here we outline briefly the manner in which these objects are constructed by standard tensor symmetrisation techniques. The end result will be the explicit forms (5) and (6) below.
Table 2: Enumeration of linearly independent candidate Markov invariants for the embedded models listed in Table 1 above, for small numbers of taxa $L$, and degrees $D$ up to 4. $(K', k, K)$ gives the number of characters of the progenitor model, the rank of the embedding tensor, and the number of characters of the target model, respectively. The linear invariants simply record overall probability conservation for each phylogenetic tensor. The invariants $I_{3,1}$ and $I_{2,2}$ studied in this paper are the two nonzero algebraically independent quadratic invariants from the count of 5 identified for the (2,2,3) model for $D = L = 2$. 

| $D$ | $L$ | $D$ | $L$ | $D$ | $L$ |
|-----|-----|-----|-----|-----|-----|
| (2,2,3) | 1   | 2   | 1   | (3,2,6) | 1   | 2   | 1   |
|      | 3   | 1   |     |      | 3   | 1   |     |
|      | 4   | 1   |     |      | 4   | 1   |     |
|      | 2   | 2   | 5   |     | 2   | 2   | 0   |
|      | 3   | 14  |     |     | 3   | 0   |     |
|      | 4   | 41  |     |     | 4   | 0   |     |
|      | 3   | 2   | 9   |     | 3   | 2   | 2   |
|      | 3   | 58  |     |     | 3   | 4   |     |
|      | 4   | 401 |     |     | 4   | 8   |     |
|      | 4   | 22  |     |     | 4   | 22  |     |
|      | 3   | 321 |     |     | 3   | 31  |     |
|      | 4   | 5597|     |     | 4   | 274 |     |
| (2,3,4) | 1   | 1   | 1   | (3,3,10) | 1   | 2   | 1   |
|      | 3   | 1   |     |      | 3   | 1   |     |
|      | 4   | 1   |     |      | 4   | 1   |     |
|      | 2   | 2   | 8   |     | 2   | 2   | 0   |
|      | 3   | 32  |     |     | 3   | 0   |     |
|      | 4   | 128 |     |     | 4   | 0   |     |
|      | 3   | 26  |     |     | 3   | 2   | 5   |
|      | 3   | 292 |     |     | 3   | 13  |     |
|      | 4   | 3464|     |     | 4   | 41  |     |
|      | 4   | 100 |     |     | 4   | 19  |     |
|      | 3   | 3688|     |     | 3   | 338 |     |
|      | 4   | 158384|   |     | 4   | 6532|     |
array $p_{01}p_{02} = p_{02}p_{01}$. Given the probability sum $p_{11} + p_{12} + p_{21} + p_{22} = 1$, the correct transcription between the two-state and three-state basis is a relabelling $p_{11} \rightarrow P_1$, $p_{12} \rightarrow \frac{1}{2}P_2$, $p_{21} \rightarrow \frac{1}{2}P_2$, $p_{22} \rightarrow P_3$ between $p_{01}p_{02}$ and a three component vector $P_1$. By the same token, for a model on 2 leaves, the phylogenetic tensor will be an object $p_{ab,\alpha\beta}$ built from edge transition matrices and a root probability in the usual way. Quantities at quadratic degree $p_{ab,\alpha\beta}P_{cd,\gamma\delta}$ therefore admit only certain compatible tensor symmetrisations between the index labels $a, b, c, d$ and $\alpha, \beta, \gamma, \delta$. Table 2 lists 5 invariants corresponding to symmetry types identified in the discussion in [A]. Here we take up the invariants $I_{3,1}$ and $I_{2,2}$, respectively.

Consider for example the $\lambda = (3, 1)$ form quadratic in the component,\(^4\) written down according to standard row and column Young symmetrisation and antisymmetrisation operations (on the sets $a, b, c, d$ and $\alpha, \beta, \gamma, \delta$ separately.\(^5\)

$$W_{abc,\alpha\beta\gamma\delta} = p(ab, \alpha\beta)p(cd, \gamma\delta) + p(ac, \alpha\beta)p(bd, \gamma\delta) - p(bd, \alpha\beta)p(ac, \gamma\delta) - p(cd, \alpha\beta)p(ab, \gamma\delta) + p(ab, \gamma\beta)p(cd, \alpha\delta) + p(ac, \gamma\beta)p(bd, \alpha\delta) - p(bd, \beta\gamma)p(ac, \alpha\delta) - p(cd, \beta\gamma)p(ab, \alpha\delta)$$

The next step is to identify the part of this array which provides the Markov invariant. This is most natural in a transformed basis for the character states in which the probability mass is treated according to standard row and column Young symmetrisation and antisymmetrisation operations (on the usual way. Quantities at quadratic degree $w = 2$, respectively, is how they transform under phylogenetic evolution. Namely, as $t \rightarrow t + \tau$, we have

$$I_{3,1} \rightarrow I'_{3,1} = \det(m_1) \det(m_2)I_{3,1}, \quad I_{2,2} \rightarrow I'_{2,2} = \det(m_1)^2 \det(m_2)^2I_{2,2},$$

where $m_i = e^{Q_i\tau}$.\(^6\)

Table 2 also lists several invariants at degree 3 (see [A]). One of these, $I_{3,3}$ in our notation, is nothing but the Markov invariant coming from the general Markov model on 3 states, and well

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\(^4\)Writing $p_{ab,\alpha\beta}$ as $p(ab, \alpha\beta)$ for clarity.

\(^5\)See [Sumner, 2008] for details.
known via its log (as used for distance and likelihood studies) as the log Det measure\(^5\), whose form (as a cubic polynomial) is obvious and will provide a standard by which the behaviour of \(I_{3,1}\) and \(I_{2,2}\) under simulation can be compared (see the concluding results and discussion below):

\[
\text{Det} \equiv I_{3,3} = P_{11}P_{22}P_{33} + P_{12}P_{23}P_{31} + P_{13}P_{32}P_{21} - P_{11}P_{23}P_{32} - P_{13}P_{22}P_{31} - P_{12}P_{21}P_{33}.
\]

Of course the Det is itself a Markov invariant of weight \(w = 3\) and satisfies the transformation rule

\[
\text{Det} \rightarrow \text{Det}' = \det(m_1)^3 \det(m_2)^3 \text{Det}.
\]

4 Results

In this section we present a simple simulation study that compares the performance of the Markov invariants \(\text{Det}, I_{22}\) and \(I_{31}\) as pairwise distance estimators for data generated under our symmetric embedded model. We do this by taking the theoretical probability distribution for a two leaf tree generated under the model and then sampling from the multinomial distribution for a range of sequence lengths. We discuss the derivation of unbiased estimators of the invariants and observe that the invariants \(I_{22}\) and \(I_{31}\) have superior statistical estimation power over that of the (log) Det. This adds credibility to the intuitive notion that it should be invariants of lower degree and lower weight that can be expected to perform best in practical contexts.

If we take a root probability distribution \(\pi_i, i = 1, 2, 3\), the “starting” (zero-edge length) probability distribution on a two leaf tree is \(P^0_{ii} = \pi_i, P^0_{ij} = 0\) if \(i \neq j\). For non-zero edge lengths, \(I_{3,1}\) and \(I_{2,2}\) are then determined by this \(P^0\) and the transformation rules \((7)\). Starting with the rate matrix \(Q\) as in \((1)\), we have the standard form

\[
m(t) = e^{tQ} = \begin{pmatrix} 1 - \alpha \lambda & \beta \lambda \\ \alpha \lambda & 1 - \beta \lambda \end{pmatrix},
\]

where \(\lambda = \frac{1}{\alpha + \beta}(1 - e^{-(\alpha + \beta)t})\), with determinant \(\det m = \exp(-(\alpha + \beta)t)\). Thus choosing as independent parameters \(\alpha\) and \(t\), with \(\alpha + \beta = 1\), if the edge distances are \(t_1\) and \(t_2\), the theoretical values for \(I_{3,1}\) and \(I_{2,2}\) become after evaluating them on \(P^0\),

\[
I_{3,1} = e^{-t_1}e^{-t_2}(4\pi_1\pi_3 + \pi_1\pi_2 + \pi_2\pi_3), \quad I_{2,2} = e^{-2t_1}e^{-2t_2}\frac{1}{8}(\pi_2^2 + 8\pi_1\pi_3).
\]

These are to be compared to the “Det” function, with theoretical value

\[
\text{Det} = e^{-3t_1}e^{-3t_2}(\pi_1\pi_2\pi_3).
\]

We take as data an alignment of two sequences consisting of three character states corresponding to \(i = 1, 2, 3\) reduced to the pattern frequencies

\[
F_{ij} := \{\text{number of occurrences of the pattern (ij)}\}.
\]

It is possible to use this data and the above invariant functions to obtain an estimator of the pairwise distance \(\Delta = t_1 + t_2\) under the symmetric embedded model \(2 \leftrightarrow 3\), as follows.

\(^5\)See Sumner et al. (2008) for comments on the relationship between log Det and other Markov invariants for the general model, and also Sumner & Jarvis (2003, 2006).
Consider the estimators of the invariants constructed by simply making the “obvious” replacement \( P_{ij} \rightarrow f_{ij} := \frac{1}{N} F_{ij} \) given by

\[
\tilde{I}_{3,1} := 4 (f_{33} + \frac{1}{2} (f_{23} + f_{32}) + \frac{1}{4} f_{22}) - 4 \left( \frac{1}{2} f_{12} + \frac{1}{2} f_{22} + \frac{1}{2} f_{32} + f_{13} + f_{23} + f_{33} \right) \cdot \left( \frac{1}{2} f_{21} + \frac{1}{2} f_{22} + \frac{1}{2} f_{23} + f_{31} + f_{32} + f_{33} \right),
\]

\[
\tilde{I}_{2,2} := f_{33} + 2 \left( f_{33} + \frac{1}{2} (f_{23} + f_{32}) + \frac{1}{4} f_{22} \right)^2 + (f_{13} + f_{23} + f_{33}) \cdot (f_{31} + f_{32} + f_{33})
- 2 \left( \frac{1}{2} f_{12} + \frac{1}{2} f_{22} + \frac{1}{2} f_{32} + f_{13} + f_{23} + f_{33} \right) \cdot \left( \frac{1}{2} f_{23} + f_{33} \right)
- 2 \left( \frac{1}{2} f_{21} + \frac{1}{2} f_{22} + \frac{1}{2} f_{23} + f_{31} + f_{32} + f_{33} \right) \cdot \left( \frac{1}{2} f_{32} + f_{33} \right),
\]

\[
\tilde{\Det} := f_{11} f_{22} f_{33} + f_{12} f_{23} f_{31} + f_{13} f_{32} f_{21} - f_{11} f_{23} f_{32} - f_{13} f_{22} f_{31} - f_{12} f_{21} f_{33}.
\]

We work under the assumption that frequency array \( \hat{F}_{ij} \) is generated by sampling \( N \) patterns with probability \( p_{ij} \) generated under our model on a two taxa tree. This means that the probability of observing a given frequency array \( \hat{F}_{ij} \) is given by the multinomial form

\[
\frac{N!}{F_{11}! F_{12}! \ldots F_{33}!} P_{11}^{F_{11}} P_{12}^{F_{12}} \ldots P_{33}^{F_{33}}.
\]

Under these conditions it is easy to show using generating function techniques that we have

\[
E \left[ \tilde{I}_{3,1} \right] = N(N - 1) I_{3,1}^{(2)} + N \left( I_{3,1}^{(1)} - (P_{22} + 2P_{32} + 2P_{23} + 4P_{33}) \right),
\]

where \( I_{3,1}^{(2)} \) and \( I_{3,1}^{(1)} \) are the quadratic and linear parts of \( I_{3,1} \) respectively (see Sumner et al. (2008) for example calculations of this kind). Thus we observe that \( E \left[ \tilde{I}_{3,1} \right] \neq I_{3,1} \), so that \( \tilde{I}_{3,1} \) is a biased estimator of \( I_{3,1} \).

This is easily rectified by defining the unbiased estimator:

\[
\tilde{I}_{3,1}^{(ub)} := \frac{1}{N(N - 1)} \left( \tilde{I}_{3,1}^{(2)} + (N - 1) \tilde{I}_{3,1}^{(1)} + F_{22} + 2F_{23} + 2F_{32} + 4F_{33} \right).
\]

This estimator then takes on the expectation value

\[
E \left[ \tilde{I}_{3,1}^{(ub)} \right] = I_{3,1} = e^{-(t_1 + t_2)} (4 \pi_1 \pi_3 + \pi_1 \pi_2 + \pi_2 \pi_3).
\]

A similar observation for \( I_{2,2} \) leads to its unbiased estimators

\[
\tilde{I}_{2,2}^{(ub)} := \frac{1}{N(N - 1)} \left( \tilde{I}_{2,2}^{(2)} + (N - 1) \tilde{I}_{2,2}^{(1)} + (F_{33} - \frac{1}{8} F_{22} + \frac{1}{2} F_{23} + \frac{1}{2} F_{32}) \right),
\]

\[
\tilde{\Det}^{(ub)} := \frac{1}{N(N - 1)(N - 2)} \tilde{\Det},
\]

with expectation values

\[
E \left[ \tilde{I}_{2,2}^{(ub)} \right] = I_{2,2} = e^{-2(t_1 + t_2)} \left( \frac{1}{8} (\pi_2^2 + 8 \pi_1 \pi_3) \right),
\]

\[
E \left[ \tilde{\Det}^{(ub)} \right] = \Det = e^{-3(t_1 + t_2)} (\pi_1 \pi_2 \pi_3).
\]

We now define the marginalizations \( F^{(1)}_i := \sum_j F_{ij} \) and \( F^{(2)}_j := \sum_i F_{ij} \) and, as is standard for the “log Det”, estimate the values \( \pi_1, \pi_2, \pi_3 \) by assuming the process was stationary and taking the harmonic means:

\[
\hat{\pi}_i = \sqrt{F^{(1)}_i F^{(2)}_i}.
\]

\footnote{More formally, this equates to taking the \( f_{ij} \) as our best estimate of the probabilities \( P_{ij} \) for this data.}
With these estimators in hand, we are now in a position to define six reasonable estimators of $\Delta = t_1 + t_2$ that can readily be evaluated directly from the pattern counts $F_{ij}$:

\[
\begin{align*}
\Delta_{3,1} &:= -\log (\hat{I}_{3,1}) + \log (4\hat{\pi}_1\hat{\pi}_3 + \hat{\pi}_1\hat{\pi}_2 + \hat{\pi}_2\hat{\pi}_3), \\
\Delta_{2,1} &:= -\log (\hat{I}_{2,1}) + \log (\hat{\pi}_1^2 + 8\hat{\pi}_1\hat{\pi}_3), \\
\Delta_{\text{Det}} &:= -\log (\hat{\text{Det}}) + \log (\hat{\pi}_1) + \log (\hat{\pi}_2) + \log (\hat{\pi}_3), \\
\Delta_{3,1}^{(ab)} &:= -\log (\hat{I}_{3,1}) + \log (4\hat{\pi}_1\hat{\pi}_3 + \hat{\pi}_1\hat{\pi}_2 + \hat{\pi}_2\hat{\pi}_3), \\
\Delta_{2,1}^{(ab)} &:= -\log (\hat{I}_{2,1}) + \log (\hat{\pi}_1^2 + 8\hat{\pi}_1\hat{\pi}_3), \\
\Delta_{\text{Det}}^{(ab)} &:= -\log (\hat{\text{Det}}^{(ab)}) + \log (\hat{\pi}_1) + \log (\hat{\pi}_2) + \log (\hat{\pi}_3).
\end{align*}
\]

To compare the performance of these estimators we performed a simulation study over a range of sequence lengths, from very long $N = 10^6$ to very short $N = 10$, with fixed rate parameters $\alpha = 0.45, \beta = 0.55$ and $t_1 + t_2 = 1$. The results are presented in Figure 1. Careful inspection of the results shows that it is consistently the lower degree invariants that have greater statistical power and that taking unbiased forms also provides a significant improvement.

5 Conclusion

In this paper we have described a novel approach to phylogenetic model construction using symmetric tensor embeddings that ensures multiplicative closure. Although this model construction is of interest in its own right, we went further to exploit the simple structure of the embedding to give examples of Markov invariants for these models. We also showed how these invariants can be exploited effectively as distance estimators with favourable statistical properties (as compared to the standard “log Det”).

We should emphasize that we do not make any claim that the symmetrically embedded models discussed have any particular direct appeal as biologically realistic rate matrices for molecular phylogenetics – this is indeed why we did not shy away from considering the 3-state case in detail. We do however argue strongly that the multiplicative closure that is present in our model is itself highly desirable from a biological perspective, and make the point that the general time reversible model (perhaps the most popular currently in use) does not satisfy the closure property.

The work presented in this paper thus serves as an elementary example that illustrates how one may go about constructing models with multiplicative closure. We are currently directing work into expanding our knowledge of such “closed” phylogenetic models and we expect that furthering the connection to Lie algebras will be vital in this regard.

A Appendix: Groups and representation theory in phylogenetic models

A.1 General results

The Markov model for phylogenetic branching sketched in §2 above can be given a formal setting for considering phylogenetic invariants and related constructions, and in particular, Markov invari-
Figure 1: **Comparison of performance of Markov invariants.** The barplots give the mean (dark shade) and variance (light shade) of $10^7$ runs with sequence length ranging from $N = 100$ to $N = 100,000$.

The starting point of our approach is to regard the model of stochastic change via Markov matrices, (2), in terms of linear actions of certain matrix groups $G$ affiliated with stochastic matrices (Johnson, 1985; Mourad, 2004). Most generally, consider a phylogenetic tensor $P$ of rank $L$ (the array of pattern frequencies for $L$ leaf edges of a presumed phylogenetic tree under the general Markov model). The equation equivalent to (2) describing the evolution along the leaf edges\footnote{Here we do not need to consider the full phylogenetic tree model, which involves labelling all edges by stochastic matrices and appropriate summations over characters. For remarks on such internal structure in the context of Markov invariants see (Sumner et al., 2008; Sumner & Jarvis, 2009).} is simply the action of an element $(M_1, M_2, \ldots, M_L)$ of the $L$-fold direct product group $G \times G \times \cdots \times G$.

$$P' = M_1 \otimes M_2 \ldots \otimes M_L \cdot P,$$

(A-1)
G. Where there is a rate matrix in continuous time models, or generically if the $M_i$ are nonsingular, then $G$ can simply be taken to be the general linear group $GL(K)$. Specifically however, we restrict consideration to probability-conserving, stochastic matrices fulfilling a unit column sum condition, so the relevant matrix group is $GL_1(K)$, which from (Johnson, 1982; Mourad, 2004) is isomorphic to the affine group $\mathbb{A} \cong GL(K-1) \ltimes \mathbb{C}^{K-1}$.

Including the symmetric embeddings construction, the full chain of edge subgroups becomes

$$GL(K) > GL_1(K) > GL_1(2) > GL(1). \quad (A-2)$$

The last step simply asserts that with a fixed rate matrix $Q$, a continuous-time Markov chain generically provide a $K \times K$ representation of the time evolution group $\mathbb{R}^+; \text{ the specialisation } GL_1(K) > GL_1(2) \text{ describes our present } '2 \rightarrow K' \text{ situation, and would be replaced for example, by } GL_1(K) > GL_1(K') \text{ in the } 'K' \rightarrow K' \text{ case, } K' < K. \text{ Formally then, the assertion } (A-1) \text{ is that of the full group branching rule for the reduction }$$

$$GL(K_L) > GL(K) \times GL(K) \times \cdots \times GL(K) \quad (A-3)$$

followed by the reduction $A-2$ on each edge. From the formal point of view, the problem of dealing with polynomial functions of $P$, say of degree $D$, is thus to implement the corresponding branching rule for the group representations arising. Specifically the Markov invariants $I(P)$, now for the embedded $GL_1(2)$ submodel, or $GL_1(K')$ in general, are in correspondence with the 1-dimensional representations occurring. Specifically, representing $M_i = M(m_i)$, with $m_i \in GL_1(K')$, $i = 1, \cdots, L$ as the embedded $K' \rightarrow K$ submodel, then under $A-1$ we have by definition

$$I(P') = \det(m_1)^{w_1} \det(m_2)^{w_2} \cdots \det(m_L)^{w_L} I(P) \quad (A-4)$$

for a Markov invariant of weight $(w_1, w_2, \cdots, w_L)$.

The two stages of the above representation-theoretic problem have been solved in Sumner et al. (2008), and for completeness we quote the relevant theorems. Firstly recall that the polynomial ring $\mathbb{C}[P]$ is isomorphic to the symmetric tensor algebra $\vee(P)$, that is, at each degree, symmetric tensor powers of the module corresponding to $P$.

**Theorem 1: Polynomial covariants for embedded models.**

Consider the embedding $GL(K_L) \supset \times^L GL(K')$ defined by the branching rule for the fundamental $K^L$-dimensional representation

$$\{1\} \rightarrow \{\lambda_1\} \otimes \{\lambda_2\} \otimes \cdots \otimes \{\lambda_L\}.$$  

The corresponding branching rule for the $D$'th symmetric tensor power is given by

$$\begin{align*}
\{D\} & \rightarrow \sum_{\sigma_1 \ast \cdots \ast \sigma_L \vdash D} (\{\lambda_1\} \boxtimes \{\sigma_1\}) \otimes (\{\lambda_2\} \otimes \{\sigma_2\}) \otimes \cdots \otimes (\{\lambda_L\} \otimes \{\sigma_L\}). \quad (A-5)
\end{align*}$$

Here standard partition notation $\lambda$ or $(\cdot)$ has been adopted for irreducible tensor representations, with $\{\cdot\}$ denoting the corresponding characters (symmetric functions, as in Littlewood, 1955). The operation $\ast$ of inner multiplication corresponds to the evaluation of tensor products of irreducible representations in the symmetric group $\mathfrak{S}_D$ for partitions $\sigma_i \vdash D$. The symbol $\otimes$ stands for the operation of plethysm on group characters. For a recent discussion of the calculus of plethysms see Fauser et al. (2006).

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8 Or unit row sum condition; however we use left multiplication for our actions.

9 We discuss groups and representations over $\mathbb{C}$ and regard real parametrisations as a separate issue.
The above result has been stated in full generality allowing for the possibility of heterogeneous edges (models with different numbers of characters). In practice we restrict attention to standard embeddings $K' \hookrightarrow K$ where $K$ is the dimension of a certain irreducible representation $\lambda$ of $GL(K')$. Moreover, for the cases of interest, $\lambda$ is a symmetric tensor of rank $k$; specifically for $2 \hookrightarrow K$, $K = k + 1$. With the above branching rule in hand, the occurrence of one dimensional representations (including multiplicities) can be read off using the following result:

**Theorem 2: Polynomial invariants for phylogenetic models and embedded models.**

Linearly independent polynomial invariants at degree $D$ of the groups (i) $\times^LGL(K')$, (ii) $\times^LGL_1(K')$ and (iii) $\times^LGL_{1,1}(K')$ for the general phylogenetic model are given by the one dimensional modules of these groups within polynomial representations of $\times^LGL(K')$ corresponding to the following partitions:

\[
\times^LGL(K') : \{r^K\} \otimes \{r^K\} \otimes \cdots \otimes \{r^K\}, \quad \text{where} \quad K'r = kD,
\]

\[
\times^LGL_1(K') : \{r_1 + s_1, r_1^{K-1}\} \otimes \{r_2 + s_2, r_2^{K-1}\} \otimes \cdots \otimes \{r_L + s_L, r_L^{K-1}\}, \quad (A-6)
\]

where $K'r_i + s_i = kD$,

\[
\times^LGL_{1,1}(K') : \{r_1 + s_1, r_1^{K-2}\}, t_1 \} \otimes \{r_2 + s_2, r_2^{K-2}\}, t_2 \} \otimes \cdots \otimes \{r_L + s_L, r_L^{K-2}\}, t_L \},
\]

where $(K'-1)t_i + s_i + t_i = kD$.

The number of admissible partitions of the given forms $\{\mu_1\} \otimes \{\mu_2\} \otimes \cdots \otimes \{\mu_L\}$ in each case (i), (ii), (iii), deriving from (A-5), is the number of times the inner product $\sigma_1 \ast \sigma_2 \ast \cdots \ast \sigma_L$ of irreducible representations of the symmetric group $\mathfrak{S}_D$ contains the one-dimensional irreducible representation $(D)$. This is also the number of linearly independent polynomial invariants in each case. \qed

Clearly the case $k = 1, K' = K$ corresponds to the standard situation treated in [Sumner et al., 2008], whereas $k \neq 1, K' < K$ covers embedded submodels. For the $K' = 2$ model (general phylogenetic model on 2 characters) and polynomial degree $D$, the Markov invariants (invariants of $GL(2)$, and a fortiori of $GL(2)$) occur as characters of the type $\{r, r\}$ with $2r = kD$ (one dimensional tensor representations of $GL(2)$), denoted $I_{rr}$, of weight $r$ or more generally are associated with characters of the type $\{r + s, r\}$ with $2r + s = kD$, denoted $I_{r+s,r}$ also of weight $r$.

### A.2 Enumeration of invariants for low-dimensional cases

Finally we turn to the enumeration of invariants for concrete, low-dimensional, cases. For the $2 \hookrightarrow 3$, $k = 2$, $K' = 2$, $K = 3$ case, admissible characters (partition shapes) $\{\mu_i\}$ according Theorem 2 are obviously $\{4\}$, $\{2, 2\}$ and $\{3, 1\}$ at degree $D = 2$, and $\{6\}, \{5, 1\}$ and $\{4, 2\}$ and $\{3, 3\}$ at degree $D = 3$. Their precise occurrence is determined by the evaluations of the appropriate plethysms. The following are well known and can be checked on dimensional grounds:

For $D = 2$:

\[
\{2\} \otimes \{2\} = \{4\} + \{2, 2\}, \quad \{2\} \otimes \{1^2\} = \{3, 1\};
\]

For $D = 3$:

\[
\{2\} \otimes \{3\} = \{6\} + \{4, 2\} + \{2^3\}, \quad \{2\} \otimes \{2, 1\} = \{5, 1\} + \{4, 2\} + \{3^2\}, \quad \{2\} \otimes \{1^3\} = \{41^2\} + \{3, 3\}.
\]

In turn, for $L = 2$ taxa, the possible invariants are associated with pairs $\{\mu_1\} \otimes \{\mu_2\}$ for such admissible characters, weighted by a combinatorial factor $2$ if $\{\mu_1\} \neq \{\mu_2\}$, and weighted by the (nonzero) multiplicity of the trivial representation $(D)$ of the symmetric group $\mathfrak{S}_D$ in the reduction.
of the inner product of the parent \{\sigma_1\} \ast \{\sigma_2\}. Given the rules

\[
D = 2 \text{ (in } \mathfrak{S}_2) : \quad \{2\} \ast \{2\} = \{2\}, \quad \{2\} \ast \{1^2\} = \{1^3\}, \quad \{1^2\} \ast \{1^2\} = \{2\}
\]

\[
D = 3 \text{ (in } \mathfrak{S}_3) : \quad \{3\} \ast \{3\} = \{3\}, \quad \{3\} \ast \{2,1\} = \{2,1\}, \quad \{2,1\} \ast \{2,1\} = \{3\} + \{1^3\} + \{2,1\},
\]

\[
\{2,1\} \ast \{1^3\} = \{2,1\}, \quad \{1^3\} \ast \{1^3\} = \{3\},
\]

it is clear that there are 5 = 2 + 1^2 linearly independent invariants at quadratic degree (namely \{4\} \otimes \{4\}, \{3,1\} \otimes \{4\}, \{4\} \otimes \{3,1\}, \{3,1\} \otimes \{3,1\}, and \{2,2\} \otimes \{2,2\}), and 9 = 2^2 + 2^2 + 1^2 linearly independent invariants at cubic degree. At the quadratic level, \{4\} \otimes \{4\} just represents probability conservation for \(P\), while \{3,1\} \otimes \{4\} and \{4\} \otimes \{3,1\} turn out to vanish identically for this case.\(^{10}\) These results verify that at quadratic degree, both \{2,2\} \otimes \{2,2\} and \{3,1\} \otimes \{3,1\} occur in the appropriate plethysms for admissible inner products, and lead to the invariants \(I_{2,2}\) and \(I_{3,1}\) constructed explicitly in \(^{3}\) Their properties are explored numerically in the final \(^{4}\) via some simple simulation studies. At cubic level it turns out that there are various invariants involving \{4,2\}, \{5,1\} and two invariants \{3,3\} \otimes \{3,3\} (again, cases such as \{6\} \otimes \{6\} are trivial by probability conservation).\(^{11}\) One of the latter, associated with the \((k = 1, K' = K = 3) GL(3)\) antisymmetric invariant, via \(\{1^3\} \ast \{1^3\} = \{3\}\), is in fact identical to the determinant function.

The above results for \(2 \leftarrow 3\) prove to be the simplest of a plethora of cases, some of which may be of interest for phylogenetic applications (see the concluding remarks, \(^{11}\)), but whose existence serves to illustrate our general philosophy. In \(^{3}\) we record invariants for symmetric embeddings for diverse (low-dimensional) cases of initial and target models \(K'\), \(k\) and \(K\) (see table \(^{12}\) for a tabulation of models and table \(^{12}\) for an enumeration of invariants for them).\(^{12}\) All manipulations with products, plethysms and group branching rules can be evaluated symbolically using an appropriate group theory package. The program \texttt{Schur} (Wybourne, \(2004\)) works with combinatorial algorithms based on manipulations of the group characters encoded as the celebrated Schur functions (symmetric polynomials in \(n\) indeterminates representing the eigenvalues of a \(n \times n\) matrix).

The general algorithm for identification of higher invariants follows the above pattern. Consider for example the following plethysms\(^{13}\) at degree 5 for a rank 2 embedding:

\[
\{2\} \otimes \{3,2\} = \{82\} + \{73\} + \{721\} + \{64\} + \{631\} + 2\{62^2\} + \{541\} + \{532\} + \\
\quad + \{531^2\} + \{52^21\} + \{4^22\} + \{4321\} + \{42^3\} + \{3^221^2\}.
\]

Thus admissible \(\{\mu\}\) for the 2 state model \(2 \leftarrow 3\) are \{82\}, \{73\}, and \{64\}; for the 3 state model\(^{14}\) \(3 \leftarrow 6\), we have \{6,2^2\} with multiplicity 2, and for the 4 state model \(4 \leftarrow 10\), the candidate\(^{15}\) \{42^3\}. Enumeration of invariants entails counting those products of admissible \(\{\mu\}\) whose parent inner product \(\prod_i \ast \{\sigma_i\}\), weighted by the correct multinomial factor, of symmetric group characters in \(\mathfrak{S}_D\) contains the trivial one-dimensional representation \(\langle D\rangle\) (weighted by multiplicity \(\geq 1\)).

\(^{10}\)They are examples of mixed weight invariants which in principle give different information for each edge (other examples of such mixed invariants were noted in Sumner et al. (2008)).

\(^{11}\)In relation to the remarks about algebraic independence below, note that the product \(I_4I_{2,2}\) of the linear invariant \(I_4\) and the quadratic \(I_{2,2}\) is also of cubic degree and has weight \(w = 2\).

\(^{12}\)At least up to linear independence: formally they form a ring, but the question of algebraic independence is beyond the scope of the present investigation (see Sumner et al. (2008)).

\(^{13}\)Using \texttt{Schur}.

\(^{14}\)According to Theorem 2, for choices of parameters giving a symmetric model on 3 states, there would be additional candidates \{721\}, \{631\}, \{541\}, \{532\}, and \{4^22\}.

\(^{15}\)Likewise for a starting symmetric 4 state model, there is the additional candidate \{52^21\}. 
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