Robust, Universal Tree Balance Indices

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Abstract

Balance indices that quantify the symmetry of branching events and the compactness of trees are widely used to compare evolutionary processes or tree-generating algorithms. Yet existing indices have notable shortcomings, including that they are unsuited to the tree types commonly used to describe the evolution of tumours, microbial populations, and cell lines. The contributions of this article are twofold. First, we define a new class of robust, universal tree balance indices. These indices take a form similar to Colless’ index but account for node sizes, are defined for trees with any degree distribution, and enable more meaningful comparison of trees with different numbers of leaves. Second, we show that for bifurcating and all other full m-ary cladograms (in which every internal node has the same out-degree), one such Colless-like index is equivalent to the normalised reciprocal of Sackin’s index. Hence we both unify and generalise the two most popular existing tree balance indices. Our indices are intrinsically normalised and can be computed in linear time. We conclude that these more widely applicable indices have potential to supersede those in current use.

Tree balance indices – most notably those credited to Sackin (1972) and Colless (1982) – are widely used to describe speciation processes, compare cladograms, and assert the correctness of tree reconstruction methods (Shao and Sokal, 1990; Mooers and Heard, 1997). These indices have recently been introduced to oncology (Scott et al., 2020; Chkhaidze et al., 2019) because methods for determining and classifying modes of tumour evolution have clinical value (Maley et al., 2017). A problem here is that the trees that best describe tumour evolution are clone trees in which node sizes are informative and which frequently contain linear sections; indeed, developing methods to distinguish linear from branching tumour evolution is an important area of ongoing research (Davis et al., 2017). Existing tree balance indices are unsuited to these topologies and take no account of node size. Moreover, even when applied only to bifurcating cladograms, existing indices are unreliable for comparing trees with different numbers of leaves.

Here we develop a new class of robust, universal tree balance indices. Our definitions not only extend the tree balance concept and open up new applications but also unify the two main approaches to quantifying balance as proposed by Sackin and Colless. We describe several general advantages of our indices compared to those in current use.

Materials and Methods

Rooted trees

We consider exclusively rooted trees in which all edges are oriented away from the root (which in our figures will be coloured red). This orientation defines a natural order on the tree, from top to bottom: all edges are assumed to extend from the root to the other internal nodes and finally to the terminal nodes or leaves. The out-degree of a node i, written $d^{+}(i)$, is the number of direct descendants, ignoring descendant branches in which all nodes have zero size. Internal nodes have out-degree at least one, whereas leaves have out-degree zero.
Figure 1: Trees on eight leaves. a: Caterpillar tree with $I_S = 35$, $I_{S,norm} = 1$, $I_C = 21$, $I_{C,norm} = 1$, $I_\Phi = 56$, $I_{\Phi,norm} = 1$. b: Fully symmetric bifurcating tree with $I_S = 24$, $I_{S,norm} \approx 0.59$, $I_C = I_{C,norm} = 0$, $I_\Phi = 16$, $I_{\Phi,norm} \approx 0.29$. c: Star tree with $I_S = 8$, $I_{S,norm} = 0$, $I_C$ and $I_{C,norm}$ undefined, $I_\Phi = I_{\Phi,norm} = 0$.

Figure 2: Clone tree of the lung tumour CRUK0065 in the TRACERx cohort (Jamal-Hanjani et al., 2017). Nodes represented by empty circles correspond to extinct clones. The diameters of other nodes are proportional to the corresponding clone population sizes.

Figure 1 shows some trees with specific names. A **caterpillar tree** (or comb) is a bifurcating tree in which each internal node has one leaf. A **fully symmetric** tree is such that every internal node with the same depth has the same degree or, equivalently, for each internal node $i$ all the subtrees rooted at $i$ are identical. A **star tree** is a tree whose leaves are all attached to the root, which is the only internal node.

**Cladograms, species trees and clone trees**

Cladograms are trees that represent relationships between extant biological taxa (leaves) via edges linking them to hypothetical extinct ancestors (internal nodes). A common conception is that only bifurcating cladograms can be considered fully resolved and linear parts are inadmissible. However, as shown by Podani (2013), linear sections in cladograms are appropriate for representing anagenesis (in which a descendant replaces its ancestor), while budding (in which an ancestor produces a descendant and remains extant) can give rise to cladogram nodes with out-degree greater than two. An extant ancestor is represented in a cladogram by a leaf stemming from the internal ancestor node (so the two nodes represent the same taxon).

An alternative way to represent extant ancestors is as internal nodes (like in a genealogy with overlapping generations). Such diagrams are known to organismal biologists as species trees and to oncologists as clone trees. In a clone tree, each node represents a clone (a set of cells that share...
Figure 3: Muller plots (left column), species or clone trees (middle column), and cladograms (right column) representing evolution by splitting only (a) and both splitting and budding (b). Nodes represented by empty circles correspond to extinct types.

alterations of interest due to common descent) and edges represent the chronology of alterations. Clone tree nodes can have any out-degree, including $d^+ = 1$, and each node – including internal nodes – can be associated with a non-negative size, related to the clone population size at the time of observation (as in Figure 2). The size of a tree or subtree can then be defined as the sum of its node sizes.

When nodes are associated with sizes, the addition or removal of even vanishingly small terminal branches can change leaves into internal nodes or vice versa and so substantially change the value of existing tree balance indices. This behaviour is unsatisfactory because these small branches typically represent either newly-created types that have yet to experience evolutionary forces or types on the verge of extinction, and in either case their relative sizes convey negligible information about the mode of evolution. Data sets may also omit rare types due to sampling error or because genetic sequencing methods have imperfect sensitivity (Turajlic et al., 2018).

The change due to the addition of terminal nodes is greater when the tree is a cladogram rather than a species or clone tree. For example, when a three-node, two-leaf tree (as in Figure 3a) is augmented by adding a node $j$ to a leaf $i$ (as in Figure 3b), the three original nodes retain their positions in the species or clone tree (middle column), but in the cladogram (right column) node $i$ becomes two nodes, the larger of which is now further from the root. As the size of the new node $j$ is continuously reduced to zero, the species or clone tree changes continuously, whereas the cladogram undergoes an abrupt change of topology when the size of node $j$ reaches zero. We conclude that the species or clone tree representation is more robust than the cladogram representation in the general case in which nodes are associated with sizes and ancestors can be extant. Also an index that accounts for non-zero internal node sizes can be made more robust than one that does not.

Existing tree balance indices

The most widely used tree balance indices are in fact imbalance indices, such that more balanced trees are assigned smaller values. These indices were introduced to study cladograms and take no account of node size. The most popular are Sackin’s index and Colless’ index.

Sackin’s index

Let $T$ be a tree with set of leaves $L(T)$. For a leaf $l \in L(T)$, let $\nu_l$ be the number of internal nodes between $l$ and the root, which is included in the count. Then the index credited to Sackin (1972) is

$$I_S(T) = \sum_{l \in L(T)} \nu_l.$$ 

For two bifurcating trees on the same number of leaves, a less balanced tree has higher values of $\nu$ as the tree is in a sense less compact (compare trees a and b in Figure 1).
Since the value tends to increase with the number of nodes, Shao and Sokal (1990) proposed normalising $I_S$ with respect to trees on $n > 2$ leaves by subtracting its minimum possible value for such trees and then dividing by the difference between the maximum and minimum possible values. The minimal $I_S$ is reached on the star tree, such as tree $c$ in Figure 1, and hence $\min_n(I_S) = n$. The maximum is attained on the caterpillar tree, such as tree $a$: 

$$\max_n(I_S) = n - 1 + \sum_{i=1}^{n-1} \nu = n - 1 + n(n - 1)/2 = (n - 1)(n + 2)/2.$$ 

The normalised index is then

$$I_{S, \text{norm}}(T) = \frac{I_S(T) - n}{(n + 2)(n - 1)/2 - n}. \tag{1}$$

This normalised index is not very satisfactory as a balance index because it fails to capture an intuitive notion of balance. For example, it is not obvious why fully symmetric tree $b$ should be considered less balanced than star tree $c$ in Figure 1, yet its $I_{S, \text{norm}}$ value is much larger.

Another critical shortcoming of $I_{S, \text{norm}}$ is that it is not defined for trees containing linear parts. This is because, whatever the number of leaves, $I_S$ can be made arbitrarily large by adding nodes that have out-degree one. Moreover, since $I_{S, \text{norm}}$ doesn’t account for node size, it is highly sensitive to the addition or removal of relatively tiny terminal branches. Hence Sackin’s index is neither universal nor robust.

**Colless’ index**

For an internal node $i$ of a bifurcating tree $T$, define $n_{i_1}$ as the number of leaves of the left branch of the subtree rooted at $i$, and $n_{i_2}$ as the number of leaves of the right branch. Then the index defined by Colless (1982) is

$$I_C(T) = \sum_{i \in \tilde{V}(T)} |n_{i_1} - n_{i_2}|,$$

where $\tilde{V}(T)$ is the set of internal nodes. The index can be normalised for the set of trees on $n > 2$ leaves by dividing by its maximal value, $\binom{n - 1}{2}$, which is reached on the caterpillar tree (as in Figure 1a).

To generalise Colless’ index to multifurcating trees, Mir et al. (2018) recently introduced a family of Colless-like balance indices, including $I_C$ as a special case. Each of these indices $C_{D,f}$ is determined by a weight function $f$, which assigns a size to each subtree as a function of its out-degree, and a dissimilarity function $D$. By definition of $D$, Colless-like indices are zero if and only if each internal node divides its descendants into subtrees of equal size according to $f$. But since these indices are normalised by dividing by the maximal value for trees on the same number of leaves, they are unreliable for comparing trees with different numbers of leaves. For example, $I_{C, \text{norm}} = 1$ for every caterpillar tree, even though long caterpillar trees are intuitively less balanced than short ones. In common with Sackin’s index, the total cophenetic index $I_S$ (Mir et al., 2013) (see Appendix), and other existing indices, the Colless-like indices so far defined are neither universal nor robust.

**Desirable properties of a universal, robust tree balance index**

Our aim is to derive a tree balance index $J$ that is useful for classifying and comparing rooted trees that can have any distributions of node degrees and node sizes. Here we specify five desirable properties that such an index should have. The first two axioms relate to extrema and universality, in the sense of an index being defined for trees with any degree distribution. The other three axioms are concerned with robustness and are relevant only when nodes can have arbitrary sizes.

Conventionally, a tree is considered maximally balanced only if every internal node splits its descendants into subtrees on the same number of leaves (Shao and Sokal, 1990). We generalise this concept by requiring that every internal node splits its descendants into at least two subtrees of equal size, as in Figure 4a, b and c. We term this the equal splits property. We then set necessary and sufficient conditions for maximal balance.
Figure 4: Trees with extremal $J$ values. Empty nodes have null population. 

**a**: A fully symmetric bifurcating tree on eight leaves, for which $J = 1$.  

**b**: A tree that is not fully symmetric, but in which each internal node splits its descendants into subtrees with the same number of equally-sized leaves, and hence $J = 1$.  

**c**: A tree in which each internal node splits its descendants into subtrees of equal size, and hence $J = 1$. This tree can be considered balanced only according to an index that accounts for node size.  

**d**: A linear tree, for which $J = 0$.

**Axiom 0.1** (Maximum value). $J(T) \leq 1$ for all trees $T$, and $J(T) = 1$ only if $T$ has equal splits. Furthermore, if $T$ has equal splits and every internal node of $T$ has null size (or equivalently represents an extinct taxon) then $J(T) = 1$.

Another convention is that narrow trees with relatively many internal nodes are considered highly imbalanced. Linear trees (that is, trees in which every node $i$ has $d^+(i) \leq 1$, as in Figure 4d) are even narrower than caterpillar trees. Also the most unequal binary split is one that assigns all descendants to one branch and none to the other. Hence our second desirable property:

**Axiom 0.2** (Minimum value). $J(T) \geq 0$ for all trees $T$, and $J(T) = 0$ if and only if $T$ is a linear tree.

Our third desirable property is that our index should be insensitive to the presence of uninformative terminal branches:

**Axiom 0.3** (Leaf limit). Let $T$ be a tree with finitely many nodes and $l$ be a leaf of $T$. Suppose we create a new tree $T'$ by adding to $T$ a subtree $T_l$ with finitely many nodes, rooted at $l$. As the size of $T_l$ excluding its root approaches zero, so $J(T') \to J(T)$.

Our fourth desirable property ensures that a linear section of a tree is regarded as a maximally unequal split:

**Axiom 0.4** (Linear limit). Let $j$ be a node of a tree $T$ with $d^+(j) = 1$. Suppose we create a new tree $T'$ by adding to $T$ a subtree with finitely many nodes, rooted at $j$. As the size of $T_j$ excluding its root approaches zero, so $J(T') \to J(T)$.

Lastly, we require continuity with respect to varying node size:

**Axiom 0.5** (Continuity). If the population of any node of any tree $T$ varies continuously in $\mathbb{R}^+$, then $J(T)$ varies continuously.

The wording of Axiom 0.1 raises an important question: Can trees with non-zero-sized internal nodes be considered maximally balanced? The following proposition provides the answer.

**Proposition 0.6.** Axioms 0.3 and 0.4 each imply that equal splits are not sufficient for maximal balance.
Proof. Suppose that equal splits are sufficient for maximal balance. First consider a one-node tree $T$. If we add a vanishingly small linear subtree to $T$ then the new tree $T'$ will have $J(T') = 0$. But if we instead add two vanishingly small subtrees of equal size to $T$ then we obtain $J(T') = 1$. This implies that whatever value we assign to $J(T)$, we cannot satisfy Axiom 0.3. Second, consider a linear tree $T$ in which the sum of the non-root node sizes is $\delta$. Then $J(T) = 0$. But if we add another subtree to the root, also of size $\delta$, then the new tree $T''$ will have $J(T'') = 1$, even as $\delta \to 0$. This contradicts Axiom 0.4.

We therefore face a choice: either weaken Axioms 0.3 and 0.4 or accept that equal splits are not sufficient for maximal balance. We choose the second option (and as a corollary obtain $J = 0$ for the single-node tree) because we want our indices to be not only universal but also highly robust when applied to real, imperfect data. We will further argue that this choice is appropriate from a biological viewpoint and is consistent with the ideas underlying previous tree balance indices.

Results

General definition of universal, robust tree balance indices

Before defining a new class of balance indices we need to introduce some more notation. For a tree $T$, we will use $V(T)$ to denote the set of all nodes of $T$, which we will abbreviate to $V$ when the identity of the tree is unambiguous. Let $f(v) \geq 0$ denote the size of node $v$ (not necessarily a function of the out-degree). Then $S_i$ denotes the size of the subtree $T_i$ rooted at $i$, and $S_i^*$ is the size of $T_i$ excluding its root:

$$S_i := \sum_{v \in V(T_i)} f(v); \quad S_i^* := \sum_{v \in V(T_i) \setminus \{i\}} f(v) = S_i - f(i).$$

We will use $\tilde{V}(T)$ or simply $\tilde{V}$ to denote the set of all internal nodes such that $\{i \in \tilde{V} : S_i^* > 0\}$.

We then introduce three continuous functions of subtree sizes:

• An importance factor $g : \mathbb{R}_{> 0} \to \mathbb{R}_{> 0}$ with $g(x) \to 0$ as $x \to 0$;

• A non-root dominance factor $h : \mathbb{R}_{> 0} \times \mathbb{R}_{> 0} \to (0, 1]$ with $h(x_1, x_2) \to 0$ as $x_1 \to 0$, and $h(x_1, x_2) = 1$ if and only if $x_1 = x_2$;

• A balance score $W$ that assigns $W_i \in [0, 1]$ to each internal node $i$ such that $W_i = 0$ if and only if $d^+(i) = 1$, and $W_i = 1$ if and only if $i$ splits its descendants into at least two equally sized subtrees.

To allow us to define $W$ more rigourously, let $\mathcal{S}$ denote the set of vectors with positive components that sum to unity:

$$\mathcal{S} := \bigcup_{k \geq 1} \{(x_1, \ldots, x_k) | x_1, \ldots, x_k > 0, x_1 + \ldots + x_k = 1\}.$$ 

Then $W : \mathcal{S} \to [0, 1]$ is such that, for all $(x_1, \ldots, x_k) \in \mathcal{S}$:

• For every permutation $\pi$, $W(x_1, \ldots, x_k) = W(x_{\pi(1)}, \ldots, x_{\pi(k)})$;

• $W(x_1, \ldots, x_k) = 1$ if and only if $k > 1$ and $x_1 = \ldots = x_k$;

• $W = 0$ if and only if $\max(x_1, \ldots, x_k) = 1$;

• $W$ is a continuous function with respect to each of its arguments.

We then define a balance index in terms of subtree sizes as

$$J := \frac{1}{\sum_{k \in V} g(S_k^*)} \sum_{i \in V} g(S_i^*) h(S_i^*, S_i) W_i,$$

where $W_i = W(S_{i_1}/S_i^*, \ldots, S_{i_p}/S_i^*)$ and $i_1, \ldots, i_p$ are the children of node $i$. A short proof that this type of index satisfies our five axioms for robustness and universality is presented in the Appendix.
Interpretation of factors $W$, $g$, and $h$

The balance score $W$ in our general definition (Equation 2) measures the extent to which an internal node splits its descendants into equally sized subtrees. The importance factor $g$ assigns more weight to nodes that are the roots of large subtrees. In biological terms, this means giving more weight to types that have more descendants. The continuous function $h$ quantifies the extent to which a node should be considered a leaf (which doesn’t contribute to determining tree balance in Colless-like indices) as opposed to an internal node (which does). From a biological point of view, nodes that are large relative to their descendants represent extant populations whose evolutionary fate remains largely undetermined.

Factors $g$ and $h$ together ensure that our indices consider a tree imbalanced unless there is strong evidence to the contrary. For example, the one-node tree provides no evidence and is considered maximally imbalanced. The inclusion of $h$ also means that a tree can achieve maximal balance only if all its internal nodes have zero size, which is equivalent to all ancestors being extinct, as in a cladogram. This requirement can be removed simply by omitting $h$ from the definition, but then, as per Proposition 0.6, our robustness Axioms 0.3 and 0.4 will not be satisfied.

Sackin’s and Colless’ indices similarly assign more weight to nodes that have more descendant leaves or are closer to the root. As Mooers and Heard (1997) have remarked, it is reasonable to put more weight on nodes deeper within the tree because “those nodes are the most informative, as the subclades they define are older and therefore sample longer periods of evolutionary time.”

A specific index based on the Shannon entropy

In defining a specific index, we start by opting for the simplest choices of importance and non-root dominance factors:

$$g(x) = x, \quad h(x_1, x_2) = x_1 / x_2.$$  

The role of the balance score function $W$ is to quantify the extent to which a set of objects (specifically subtrees) have equal size. A well-known index that satisfies the necessary conditions is the normalised Shannon entropy.

Assume a population is partitioned into $P \in \mathbb{N}$ types. Each type has a proportion $p_i$ for $i \in \{1, \ldots, P\}$ such that $\sum_{i=1}^{P} p_i = 1$. Then the Shannon entropy is defined as $^1H := - \sum_{i=1}^{P} p_i \ln p_i$. If all types have equal frequencies $p_i = 1/P$ then $^1H = \ln P$. If the types have unequal sizes then $^1H < \ln P$. And if the abundance is mostly concentrated on one type $j$, such that $p_j \to 1$, then $\ln p_j \to 0$ and $p_i \to 0$ for all $i \neq j$, and hence $^1H \to 0$.

Let $C(i)$ denote the set of children (immediate descendants) of a node $i$, and for $j \in C(i)$ let $p_{ij} := S_j / S_i^*$ denote the relative size of subtree $T_j$ compared to all subtrees attached to $i$. A balance score based on the normalised Shannon entropy is then

$$W_i^1 = \sum_{j \in C(i)} W_{ij}^1,$$

with

$$W_{ij}^1 = \begin{cases} -\frac{p_{ij} \ln p_{ij}}{\ln d^+(i)} & \text{if } p_{ij} > 0 \text{ and } d^+(i) \geq 2, \\ 0 & \text{otherwise}. \end{cases}$$  

From aforementioned properties of the Shannon entropy, it follows that $W_i^1 \in [0, 1]$, with $W_i^1 = 0$ if and only if $d^+(i) = 1$, and $W_i^1 = 1$ if and only if $i$ splits its descendants into at least two equally sized subtrees. Therefore the following specific balance index satisfies our robustness and universality axioms:

$$J^1 := \frac{1}{\sum_{k \in V} S_k} \sum_{i \in V} S_i^* S_i^* W_i^1.$$  

The definition simplifies when we restrict the domain to the set of multifurcating cladograms in which all leaves have equal size $f_0$ (corresponding to equally important extant types) and internal nodes have zero size (representing extinct ancestors). For all internal nodes $i$ in such trees, $S_i^* = S_i = f_0 n_i$, where $n_i$ is the number of leaves of the subtree rooted at $i$. The general definition of Equation 2 then becomes a weighted average of node balance scores:

$$J = \frac{1}{\sum_{k \in V} n_k} \sum_{i \in V} n_i W_i,$$  

(5)
Figure 5: Three multifurcating trees on three leaves. The root is labelled 1. The population of each node is written in blue.

and the specific definition of Equation 4 becomes

$$J^1 = -\frac{1}{\sum_{i \in V} n_i} \sum_{i \in V} \frac{1}{\ln d^+(i)} \sum_{j \in C(i)} n_j \ln \frac{n_j}{n_i}.$$  \hspace{1cm} (6)

There are of course many alternative options for $W$. In particular, since the Shannon entropy belongs to a family of generalised entropies (Chao et al., 2014) parameterised by $q \geq 0$, the above reasoning can be generalised to define a balance score $W^q$, and hence a robust, universal balance index $J^q$, for every $q > 0$ (see Appendix). We prefer $W^1$ mostly because, as we will show, it is the only function for which Equation 5 is a generalisation of the inverse Sackin index.

Numerical examples

To illustrate the definition, we compute $J^1$ for the three trees shown in Figure 5.

• Tree a has only one internal node (the root), so the importance factor is 1. The root splits the tree perfectly evenly, so its balance score is $W^1 = 1$. Hence

$$J^1 = \frac{S^1}{S_1} = \frac{30}{31} \approx 0.97.$$  

Tree a is considered highly balanced.

• In tree b, both the importance factor and the balance score are again equal to 1 but the non-root dominance factor is much smaller:

$$J^1 = \frac{S^1}{S_1} = \frac{3}{13} \approx 0.23.$$  

Tree b is considered less balanced than tree a because of its relatively larger root, which makes it more similar to the uninformative one-node tree.

• In tree c, nodes 2, 3 and 4 have null balance scores as their subtrees are linear. These nodes nevertheless contribute to the importance of the root, which has a balance score of 1 as it splits the tree evenly. Thus the balance index of tree c is

$$J^1 = \frac{1}{S^1 + S_2 + S_3 + S_4} S^1 = \frac{1}{\frac{S^1}{S_1}} = \frac{1}{\frac{1}{45}} \times \frac{30}{31} \approx 0.65.$$  

Tree c is considered less balanced than tree a mostly because of its linear branches.

Relationship with Colless’ index

Like Colless’ index and Colless-like indices as previously defined, our new family of tree balance indices is based on the intuitive idea of assigning a value to each internal node, summing these values, and then normalising the sum. A Colless-like index in the sense of Mir et al. (2018) depends
on a function \( f : \mathbb{N} \to \mathbb{R}_{\geq 0} \), which assigns node sizes, and a dissimilarity score \( D : \mathcal{R} \to \mathbb{R}_{\geq 0} \), where \( \mathcal{R} \) is the set of non-null real vectors. Before normalisation, such an index has the form

\[
\mathcal{C}_{D,f} = \sum_{i \in \mathcal{V}} D(\delta_f(T_i), \ldots, \delta_f(T_{ik})),
\]

where \( \{i_1, \ldots, i_k\} \) are the children of node \( i \). The function \( \delta_f \) assigns a size to each subtree by summing the node sizes: \( \delta_f(T) = \sum_{j \in \mathcal{V}(T)} f(d^+(j)) \). Neglecting the initial normalising factor, our general definition (Equation 2) has a similar form and can be considered Colless-like in only a slightly broader sense. Our definition nevertheless differs in three important ways.

First, whereas the unbounded dissimilarity index \( D \) measures both the relative imbalance of subtrees and their combined size, and is undefined for nodes with out-degree one, we split these two roles into a normalised balance score \( W \) and an unbounded importance factor \( g \) and – crucially – we assign a \( W \) value (specifically zero) to nodes with out-degree one. This difference enables us to extend the balance index definition to trees with any degree distribution. It also makes it easy to normalise our index for any tree, simply by dividing by the sum of the importance factors. Furthermore, our normalisation is universal, rather than being based on comparison with other trees with the same number of leaves. For example, our index judges long caterpillar trees less balanced than short ones (Figure 6), whereas previous indices consider them equally imbalanced.

Second, we multiply the balance score by an additional non-root dominance factor \( h \). This factor makes the balance index robust when internal nodes can have non-zero size, which blurs the distinction between internal nodes and leaves. Non-root dominance plays no role if all internal nodes have null size, as in cladograms (because then \( h \equiv 1 \)).

Third, instead of assigning a size to each node as a function of its out-degree, we associate a broader measure of subtree imbalance and an unbounded importance factor

\[
\sum_{i \in \mathcal{V}} \delta_f(T_i) = \sum_{i \in \mathcal{V}} \delta_f(T_{im}),
\]

where \( \sum_{k \in \mathcal{V}} n_k \) is just another way of expressing Sackin’s index (summing over internal nodes instead of leaves). Therefore \( J \) in Equation 5 is essentially a weighted Sackin index (with each term in the sum weighted by the balance score \( W \)) divided by the unweighted Sackin index. In the special, important case of full \( m \)-ary cladograms, the weighted sum in \( J^1 \) (Equation 6) simplifies yet further. Let \( \mathcal{T}_{n,m} \) denote the set of all trees on \( n \) leaves such that all internal nodes have the same out-degree \( m > 1 \), every internal node has null size, and all leaf sizes are equal. Then we obtain a remarkably simple relationship between \( J^1 \) and Sackin’s index:

**Proposition 0.7.** If \( T \) is a tree in \( \mathcal{T}_{n,m} \) then

\[
J^1(T) = \frac{\min_{n,m} I_S}{I_S(T)} = \frac{n \log_m n}{I_S(T)},
\]

where \( \min_{n,m} I_S \) is the minimum \( I_S \) value of trees in \( \mathcal{T}_{n,m} \).

The above result is somewhat surprising as it unifies our Colless-like index, which can be viewed as a weighted average of internal node balance scores, and Sackin’s index, which is the sum of all leaf depths. A short proof of Proposition 0.7 is presented in the Appendix. The converse result, which is also proved in the Appendix, justifies our choice of \( W^1 \) instead of alternative balance score functions:

**Proposition 0.8.** Let \( J \) be a tree balance index such that

\[
J(T) = \frac{1}{\sum_{k \in \mathcal{V}} n_k} \sum_{i \in \mathcal{V}} n_i W\left(\frac{n_{i_1}}{n_i}, \ldots, \frac{n_{i_p(i)}}{n_i}\right),
\]

where \( i_1, \ldots, i_{p(i)} \) are the children of node \( i \), and \( W \) is a balance score satisfying the conditions stated before Equation 2. Suppose that for all trees \( T \in \mathcal{T}_{n,m} \), \( J(T) = n \log_m n / I_S(T) \). Then \( W = W^1 \) and hence \( J = J^1 \).
The right-hand side of the equation in Proposition 0.7 incidentally provides an alternative way of normalising Sackin’s index on full \(m\)-ary cladograms, including the bifurcating cladograms on which the index was originally defined. This normalised inverse Sackin index, which we can define as \(J_S := n \log_m n / I_S\), provides a more satisfactory way of comparing trees with different numbers of leaves. Unlike the usual normalisation (Equation 1), \(J_S = 1\) if and only if the tree has minimal depth given \(m\), which is equivalent to being fully symmetric (because in this case \(\log_m n = \nu_l\) for every leaf \(l\)). Hence \(J_S\) is a sound tree balance index in the sense defined by Mir et al. (2018). For \(m > 1\), we have \(J_S > 0\) but \(\min J_S \to 0\) as \(n \to \infty\), which makes sense because trees with more leaves can be made less balanced. In particular, when \(T\) is a caterpillar tree on \(n \geq 2\) leaves,

\[
J_S(T) = \frac{\min_{n,2} I_S}{\max_{n,2} I_S} = \frac{2n \log_2 n}{(n-1)(n+2)},
\]

as illustrated in Figure 6. The definition of \(J_S\) can be naturally extended to the case \(m \leq 1\) by setting \(J_S(T) := 0\) if \(T\) is linear or has only one node. From this point of view, \(J^1\) (a Colless-like index) is a generalisation of \(J_S\) (the normalised reciprocal of Sackin’s index) to the domain of trees with arbitrary degree distributions and arbitrary node sizes.

Correlations with preexisting indices

When internal nodes have non-zero size, the non-root dominance factor \(h\) becomes influential and the relationship between \(J^1\) and previous indices is more difficult to deduce. Therefore, to compare \(J^1\) to normalised versions of Sackin’s index, a Colless-like index, and the total cophenetic index (defined in the Appendix) on a more diverse set of trees, we generated 200 random trees on 100 leaves using the R package CollessLike (Mir et al., 2018). These are multifurcating trees with constant node size and without linear parts.

As shown in Figure 7, our new balance index correlates negatively with the previously defined imbalance indices on this set of trees (Spearman’s \(\rho < -0.6\) in all three cases), indicating that it captures a similar notion of balance. The strongest correlation is between \(J^1\) and the total cophenetic index (Spearman’s \(\rho \approx -0.8\)). We also note that nearly all the random trees have balance values less than 0.25 according to the previous indices, whereas \(J^1\) values are more evenly distributed between zero and one, with mean and median between 0.5 and 0.6. The two-dimensional distribution of points suggests that \(J^1\) is approximately related to the reciprocals of the other indices.

Discontinuities

Although our indices are robust to the addition or removal of uninformative nodes, the addition of informative nodes – however small – can create a discontinuity. Consider a node that splits its descendants into several subtrees of similar size. The addition of a new, relatively small subtree...
Figure 7: Scatter plots of $J^1$ versus normalised Sackin’s, Colless-like, and total cophenetic indices for 200 random trees on 100 leaves. The histograms in the margins show the marginal distributions. Dashed reference curves are $y = 0.1/x$ and $y = 0.01/(x + 0.01)$. We use the Colless-like index with $f(n) = \ln(n + e)$ and $D$ the mean deviation from the median, as recommended by Mir et al. (2018).

Implementation and algorithmic complexity

Assuming the matrix describing the tree is sorted by node identity from root to leaves, all our new indices can be computed in linear time via depth-first search. The initialisation step required for efficient depth-first search fills several vectors of length $N$, where $N$ is the number of nodes, and hence takes $O(N)$ time. The search then visits each node exactly once. During the search process, the size of every subtree rooted at each internal node is also computed. The computation of the balance index takes at most $\sum_{i=1}^{N} |\text{Adj}[i]| = |E| = N - 1$ time, where $\text{Adj}[i]$ is the adjacency list of node $i$. Therefore the total complexity of the procedure is $O(N)$. If the tree matrix is not initially sorted then we need to add a sorting step, which increases the runtime to $O(N \log(N))$. Efficient R code for calculating $J^q$ is shared in an online repository (Noble and Lemant, 2021).

Discussion

Here we have defined a new class of tree balance index that unifies, generalises, and in various ways improves upon existing definitions.

Being applicable to a wider set of trees, our new indices enable important new applications. We recently demonstrated the usefulness of $J^1$ in comparing real and simulated tumour clone trees (Noble et al., submitted). In the same study, using a large simulated data set, we showed that, unlike preexisting indices, $J^1$ is insensitive to the addition or removal of rare types. This is an important property because whether or not a rare type is detected in practice depends on both
methodology and chance (Turajlic et al., 2018). Besides oncology, our indices are well suited to the study of microbial evolution and any other system in which population sizes are important or linear evolution can occur.

In generalising conventional indices we also obviate their shortcomings. Even when restricted to the tree types on which previous indices are defined, our indices enable more meaningful comparison of trees with different degree distributions or different numbers of leaves. This advantage might make our indices preferable to other options more generally. Because of its unique relationship with Sackin’s index, we especially recommend $J^1$ – a weighted average of the normalised entropies of the internal nodes – as defined in general by Equation 4 and more simply for cladograms by Equation 6.

Nevertheless, an impediment to using $J^1$ in place of Sackin’s index or Colless’ index is that we have not yet determined the moments of $J^1$ under tree generating processes such as the Yule and uniform models, nor the sensitivity of $J^1$ in detecting departures from such models. We will report progress in this direction in a forthcoming article.

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**Author contributions**

RN conceived the project. JL and RN developed the balance indices with helpful input from CLS. JL and RN obtained mathematical results with helpful input from VM. JL and RN wrote the paper, based on a chapter from JL’s ETH Zurich master’s thesis. All authors have read and approved this manuscript.

**References**


Appendix

Definition of the total cophenetic index

The cophenetic value \( \phi(l, m) \) of a pair of leaves \((l, m)\) is the depth of their lowest common ancestor. The total cophenetic index (Mir et al., 2013) is then the sum of the cophenetic values over all pairs of leaves:

\[
I_\phi(T) = \sum_{N-n+1 \leq l < m \leq n} \phi(l, m),
\]

where \( N \) is the number of nodes and \( n \) the number of leaves. As in Sackin’s index, the principle is that an unbalanced tree stretches more than a balanced tree. Being explicitly defined for all multifurcating
trees, the total cophenetic index permits meaningful comparison of any two multifurcating trees on the same number of leaves.

For trees on \( n > 2 \) leaves, the minimum of the total cophenetic index is reached on the star tree, with \( \min_n(I_\Phi) = 0 \). The maximum is attained on the caterpillar tree:

\[
\max_n(I_\Phi) = \sum_{l=2}^{n-1} \sum_{m=1}^{l-1} m = \sum_{l=2}^{n-1} \frac{l(l-1)}{2} = \frac{1}{2} \left( (n-1)n(2n-1) - n(n-1) \right) = \frac{n(n-1)(n-2)}{6} = \binom{n}{3}.
\]

Hence a normalised version of the total cophenetic index is

\[
I_{\Phi,\text{norm}}(T) = \frac{I_\Phi(T)}{\binom{n}{3}},
\]

and \( I_{\Phi,\text{norm}}(T) = 0 \) if and only if \( I_\Phi(T) = 0 \). This implies that, like Sackin’s index, the normalised total cophenetic index is not minimal for all fully symmetric trees. For example, the cophenetic value of the two leftmost leaves of fully symmetric tree \( b \) in Figure 1 is two, and so both the unnormalised and normalised cophenetic indices of tree \( b \) will be nonzero.

**Proof that the index of Equation 2 satisfies our five axioms**

*Proof*. Axiom 0.1 (Maximum value): We have \( J \leq 1 \) since \( h \) and \( W \) lie between zero and one by definition. Also if any internal node \( j \) of tree \( T \) doesn’t split its descendants into at least two equally sized subtrees then \( W_j < 1 \) by definition and so

\[
\sum_{i \in \tilde{V}} g(S_i^*) h(S_i^*, S_i) W_i < \sum_{k \in \tilde{V}} g(S_k^*) \quad \Rightarrow \quad J(T) < 1.
\]

Finally, let \( T \) be a tree such that every internal node splits its descendents into at least two equally sized subtrees. Then \( W_i = 1 \) for all \( i \in \tilde{V} \) by definition. And if every internal node has null population then \( S_i^* = S_i \), which implies \( h(S_i^*, S_i) = 1 \) for all \( i \in \tilde{V} \) by definition. Hence

\[
J(T) = \sum_{k \in \tilde{V}} g(S_k^*) \sum_{i \in \tilde{V}} g(S_i^*) = 1.
\]

Axiom 0.2 (Minimum value): We have \( J \geq 0 \) since \( g, h \) and \( W \) are always non-negative by definition. Also if \( T \) is a linear tree then \( W_i = 0 \) for all \( i \in \tilde{V} \) by definition, and hence \( J(T) = 0 \). Conversely, if some internal node \( j \) has \( d^+(i) > 1 \) then \( W_j > 0 \) by definition and, because \( g \) and \( h \) are always positive by definition, we must have \( J(T) > 0 \).

Axiom 0.3 (Leaf limit): Adding a subtree to a leaf \( l \) changes the tree balance value via the contributions of three sets of nodes: the newly added nodes, the former leaf \( l \), and all other internal nodes. First, for each internal node \( i \in V(T) \) with \( i \neq l \), as \( S_i^* \to 0 \) so also \( S_i^* \to 0 \) (because \( S_i^* \leq S_l^* \)), which implies \( g(S_i^*) \to 0 \) by definition, and hence the first contribution approaches zero. Next, as \( S_l^* \to 0 \), so \( h(S_l^*, S_l) \to 0 \) by definition, which implies that the second contribution approaches zero. Lastly, the third contribution approaches zero because \( g, h \) and \( W \) are continuous by definition.

Axiom 0.4 (Linear limit): Adding a subtree to a node \( j \), with previously \( d^+(j) = 1 \), changes the tree balance value via the contributions of the newly added nodes and of node \( j \). The first contribution approaches zero for the same reason as in the leaf limit proof. Now without loss of generality let \( j_1 \) denote the original child of \( j \), and \( j_2, \ldots, j_k \) denote the newly added children of \( j \). As \( S_{j_1} + \ldots + S_{j_k} \to 0 \) there are two possibilities. If we also have \( S_{j_1} \to 0 \) then \( S_j^* = S_{j_1} + S_{j_2} + \ldots + S_{j_k} \to 0 \), which implies \( h(S_j^*, S_j) \to 0 \) by definition. Otherwise, \( \max(S_{j_2}, \ldots, S_{j_k}) \to 0 \), which implies \( W(S_{j_1}, S_{j_2}, \ldots, S_{j_k}) \to 0 \) by definition. In either case the second contribution approaches zero.

Axiom 0.5 (Continuity): The continuity of \( J \) follows immediately from the continuity of \( g, h \) and \( W \). □
Other balance indices based on generalised entropies

As defined by Chao et al. (2014), generalised entropies for $q \geq 0, q \neq 1$ are

$$qH := \frac{1}{q-1} \left( 1 - \sum_{i=1}^{P} p_i^q \right).$$

Parameter $q$ determines the sensitivity to the type frequencies. $0H$ is simply the richness (minus 1) of the population, which corresponds to ignoring the frequencies and just counting the types. For $0 < q < 1$, rare types are given more weight than implied by their proportion, whereas for $q > 1$ abundant types matter more. $2H$ is the Gini-Simpson coefficient. In the limit $q \to 1$ we recover the Shannon entropy $1H$.

For $q > 0$, $qH$ attains its maximum value if and only if all types have equal frequency $p_i = \frac{1}{P}$:

$$\max(qH) = \frac{1}{q-1} \left( 1 - \frac{1}{P^{q-1}} \right) = \frac{P^{q-1} - 1}{P^{q-1}(q-1)}. $$

We can therefore define a normalised balance score $W_i^q$ for $q > 0, q \neq 1$ and $i \in \tilde{V}$:

$$W_i^q := \begin{cases} d^+(i)^{q-1} & \text{if } d^+(i) \geq 2 \\ 0 & \text{otherwise} \end{cases} \left( 1 - \sum_{j \in C(i)} p_i^q \right)$$

A balance index $J^q$ satisfying our axioms is then

$$J^q := \frac{1}{\sum_{k \in \tilde{V}} S_k} \sum_{i \in \tilde{V}} S_i \frac{S_i^*}{S_i} W_i^q,$$

for any $q > 0$. In the limit $q \to 1$, $J^q \to J^1$.

Proof of Proposition 0.7

We will prove a more general result of which Proposition 0.7 is a special case:

**Lemma .9.** Let $T$ be a tree on $n$ leaves with $d^+(i) = m > 1$ and $f(i) = 0$ for every internal node $i$. Then

$$J^1(T) = \frac{1}{\sum_{k \in \tilde{V}} S_k} \sum_{i \in L} f(i) \log_m \frac{f(i)}{S},$$

where $S = \sum_{i \in L} f(i)$ is the size of the entire tree.

**Proof.** By definition of $J^1$, for such a tree $T$,

$$J^1(T) = \frac{1}{\ln m} \sum_{k \in \tilde{V}} S_k \sum_{i \in L} S_j \ln \frac{S_j}{S_i}.$$

The number of distinct subtrees that contain a given leaf $l$ is equal to its number of ancestors, which is the same as $\nu_l$, the depth of $l$. So the sum of subtree sizes over the set of all internal nodes is equal to the sum of $\nu_l$ multiplied by leaf size over the set of all leaves:

$$\sum_{k \in \tilde{V}} S_k = \sum_{k \in L} \nu_k f(k).$$

Summing first over the internal nodes and then over their children gives the same result:

$$\sum_{i \in \tilde{V}} \sum_{j \in C(i)} S_j = \sum_{i \in \tilde{V}} S_i = \sum_{i \in L} \nu_i f(i) = \sum_{i \in L} f(i) \sum_{j=1}^{\nu_i} 1.$$
Let \(a(i, j)\) denote the ancestor of node \(i\) at distance \(j\), with \(a(i, 0) = i\) and \(a(i, \nu_i) = r\) (the root) for all \(i\). Then by extension,

\[
\sum_{i \in V} \sum_{j \in C(i)} S_j \theta(S_i, S_j) = \sum_{i \in L} f(i) \sum_{j=1}^{\nu_i} \theta(S_{a(i,j)}, S_{a(i,j-1)}),
\]

for any function \(\theta\). In particular, we have

\[
\sum_{i \in V} \sum_{j \in C(i)} S_j \ln S_j = \sum_{i \in L} f(i) \sum_{j=1}^{\nu_i} S_{a(i,j-1)}.
\]

Substituting this result into the expression for \(J^1\) we find

\[
J^1(T) = \frac{-1}{\ln m} \sum_{k \in V} S_k \sum_{i \in L} f(i) \ln S_{a(i,j-1)}
\]

\[
= \frac{-1}{\ln m} \sum_{k \in V} S_k \sum_{i \in L} f(i) \sum_{j=1}^{\nu_i} (\ln S_{a(i,j-1)} - \ln S_{a(i,j)})
\]

The right-hand sum is a telescoping series that collapses to give

\[
J^1(T) = \frac{-1}{\ln m} \sum_{k \in V} S_k \sum_{i \in L} f(i) (\ln S_{a(i,0)} - \ln S_{a(i,\nu_i)}).
\]

Now since \(i\) is a leaf, \(\ln S_{a(i,0)} = \ln S = \ln f(i)\). Also \(\ln S_{a(i,\nu_i)} = \ln S_r = \ln S\). Hence

\[
J^1(T) = \frac{-1}{\ln m} \sum_{k \in V} S_k \sum_{i \in L} f(i) (\ln f(i) - \ln S)
\]

\[
= \frac{-1}{\ln m} \sum_{k \in V} S_k \sum_{i \in L} f(i) \log_m \frac{f(i)}{S}.
\]

Substituting \(f(i) = f_0\) for all leaves \(i\), we immediately obtain the result of Proposition 0.7 (because then \(S = n f_0\) and \(\sum_{k \in V} S_k = f_0 I_S\)).

Proof of Proposition 0.8

Proof. Since \(\sum_{k \in V} n_k = I_S(T)\), the conditions are equivalent to

\[
I_S(T)J(T) = \sum_{i \in V} n_i W_i = n \log_m n, \quad \text{with} \quad W_i = W\left(\frac{n_{i_1}}{n_i}, \ldots, \frac{n_{\nu_i(i)}}{n_i}\right),
\]

where \(n_{i_1}, \ldots, n_{\nu_i(i)}\) are the children of \(i\). Let \(T\) be a tree in \(\mathcal{T}_{n,m}\) and \(i\) be an internal node of \(T\). Then \(T_i \in \mathcal{T}_{n_i,m}\) and \(T_j \in \mathcal{T}_{n_j,m}\) for every child \(j\) of \(i\). Therefore

\[
I_S(T_i)J(T_i) = n_i W_i + \sum_{j \in C(i)} J(T_j) = n_i W_i + \sum_{j \in C(i)} n_j \log_m n_j.
\]

Also, \(I_S(T_i)J(T_i) = n_i \log_m n_i\), so we have

\[
n_i W_i + \sum_{j \in C(i)} n_j \log_m n_j = n_i \log_m n_i
\]

\[
\implies W_i = \log_m n_i - \sum_{j \in C(i)} \frac{n_j}{n_i} \log_m n_j.
\]

Since \(\sum_{j \in C(i)} n_j = n_i\), this implies

\[
W_i = \sum_{k \in C(i)} \frac{n_k}{n_i} \log_m n_i - \sum_{j \in C(i)} \frac{n_j}{n_i} \log_m n_j = \frac{-1}{\ln m} \sum_{j \in C(i)} \frac{n_j}{n_i} \log_m n_j = W_i^1.
\]