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Does clustering of DNA barcodes agree with botanical classification directly at high taxonomic levels? Trees in French Guiana as a case study

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Running title: Clustering plants' barcodes at order level

2 Abstract

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Characterising biodiversity is one of the main challenges for the coming decades. Most diversity has not been morphologically described and barcoding is now complementing morphological-based taxonomy to further develop inventories. Both approaches have been cross-validated at the level of species and OTUs. However, many known species are not listed in reference databases. One path to speed up inventories using barcoding is to directly identify individuals at coarser taxonomic levels. We therefore studied in barcoding of plants whether morphological-based and molecularbased approaches are in agreement at genus, family and order levels. We used Agglomerative Hierarchical Clustering (with Ward, Complete and Single Linkage) and Stochastic Block Models (SBM), with two dissimilarity measures (Smith-Waterman scores, kmers). The agreement between morphological-based and molecular-based classifications ranges in most of the cases from good to very good at taxonomic levels above species, even though it decreases when taxonomic levels increase, or when using the tetramer-based distance. Agreement is correlated with the entropy of morphological-based classification and with the ratio of the mean within- and mean between-groups dissimilarities. The Ward method globally leads to the best agreement whereas Single Linkage can show poor behaviours. SBM provides a useful tool to test whether or not the dissimilarities are structured by the botanical groups. These results suggest that automatic clustering and group identification at taxonomic levels above species are possible in barcoding.

Keywords: taxonomy; barcoding; clustering; Stochastic Block Model; Ward method;

24 French Guianan Trees

$_{5}$ 1 Introduction

- Numerical taxonomy and hierarchical clustering have coevolved since the 1960s' (Cole,
- ²⁷ 1969; Sneath and Sokal, 1973). Both approaches rely on the assumption that the diversity

of life for taxonomy, or patterns in distances between some items in clustering, are organized as a nested hierarchy, modelled as a tree. This approach has survived the revolution of 29 molecular-based taxonomy (Hillis et al., 1996) and molecular phylogenies (Felsenstein, 2004; 30 Yang, 2006), with a current revival due to barcoding (Floyd et al., 2002; Hebert et al., 2003), 31 and metabarcoding (López-García et al., 2001; Sogin et al., 2006; Hajibabaei et al., 2011; 32 Taberlet et al., 2012; Kermarrec et al., 2013). As far as morphological-based taxonomy is 33 concerned, most of the diversity in many clades of organisms is still unknown. Leray and Knowlton (2015) point out that between 33% and 91% of all marine biodiversity has never been named. Currently many effort are devoted to speeding up the process of producing large inventories with metabarcoding by bypassing identified obstacles (Bik et al., 2012). 37 The notion of OTU (Operational Taxonomic Unit) has been coined (Floyd et al., 2002; 38 Blaxter et al., 2005). Such units are produced by clustering sets of barcodes by aggregation 39 at a level assumed to be similar to the level of species in morphological-based classifications. 40 The authors in Blaxter et al. (2005) emphasize that they are "agnostic" as to whether OTU 41 are species or not. Identifying OTUs in an environmental sample and organising molecular 42 diversity as the frequency of OTUs make it possible to produce molecular-based inventories at previously unparalleled speed. A classical approach is therefore to build OTUs and to map them on reference databases 45 that contain reference barcodes. A standard tool for mapping is BLAST (Altschul et al., 46 1990), but other more sophisticated solutions exist (e.g., the use of Bayesian Phylogenetics, 47 Munch et al., 2008). When taxonomic expertise and references exist at the species level, 48 the agreement between molecular and morphological-based classification can be excellent 49 (Ji et al., 2013), even if sometimes like for plants, introgression may blur the distinction 50 between species (Petit and Excoffier, 2009). It may happen that such a comparison is not feasible when morphological-based taxonomy is unknown or when only partial references exist. Leray and Knowlton (2015) report in their study that less than 12\% of their OTUs matched with GenBank or BOLD. The same observation was made in White et al. (2010)

regarding intestinal microbial flora. Hence most inventories with supervised learning are made at a grain often much coarser than the genus/species level.

Trying to complete databases at the species level is highly time-consuming. Another 57 solution is to build groups larger than OTUs, e.g. at the scale of families or orders, by 58 clustering¹ the barcodes. Then each group could be annotated as a taxon at this higher taxonomic level by looking for a match for one or several sequences of the group, in the reference database. This is in line with the conclusion of the study by Meiklejohn et al. (2019), on the accuracy of BOLD and GenBank: the authors suggest that a solution to address concerns with incorrect species identifications observed in their experiments would be to report the taxonomy at a higher level. This raises the question of the agreement between morphological-based and molecular-based taxonomy when clusters of sequences are built at a level coarser than species, e.g., class or order. Comparing morphologicalbased classifications and OTUs produced by barcode clustering has been thoroughly studied 67 (see, e.g., White et al., 2010). Several methods have been recently designed and widely 68 used for delineating species on the basis of barcodes (Pons et al., 2006; Fontaneto et al., 69 2008; Puillandre et al., 2012; Talavera et al., 2013; Zhang et al., 2013). However, to our knowledge, the question has seldom been addressed directly at coarser taxonomic levels such as orders. 72

Our objective here is to study whether the clustering of barcodes in molecular-based taxonomy makes it possible to directly recover the taxa present in a sample, for a given taxonomic level coarser than species, and, if so, with which tool, accuracy and robustness. More precisely, we consider the clustering of the barcodes in a reduced number of groups compared to a clustering into species, and we ask the question whether the classification obtained is similar or not with the botanical classification at genus, family or order levels.

This comparison is performed without annotating the classes: we only aim at comparing

¹In this article, the term *clustering* makes reference to any numerical method for the unsupervised grouping of the individuals, while the term *classification* designates the method's output, i.e. the partition of individuals into classes.

the two partitions of the sequences, the botanical one and the molecular-base one.

We have selected for this study a dataset of barcodes of trees in "Piste de Sainte Elie" 81 research station in French Guiana. The corresponding plot has been inventoried botanically 82 for decades (Madelaine et al., 2007). The data set represents about one third of the diversity 83 of the French Guianan tree flora (1458 sequences, from 20 orders, 56 families, 182 genera and 428 species). We selected flowering plants because the botanical classification is well known, both morphologically (it is organised as a nested system of different taxonomic levels as a classification system) and molecularly with the Angiosperm Phylogeny Group initiative, 87 even if it is under continuous revision (The Angiosperm Phylogeny Group et al., 2016). The dataset itself is composed of some 1,500 trees from French Guiana that have been botanically identified and sequenced with chloroplastic marker trnH-psbA using Sanger technology which produces high quality sequences (Caron et al., 2019). By selecting a small data set and a long resolutive sequence (trnH-psbA is about 450 bp long, with high variability), we are not confronted to the computational burden of treatment of massive 93 data sets as in metabarcoding data, and we can therefore concentrate on the analysis of agreement. The question of the scaling to metabarcoding with massive data sets of shorter reads of the clustering methods will be the object of further studies.

It can be expected that there is not a clear answer to the degree of agreement between 97 the two types of classification (morphological-based or molecular-based). There may be 98 favourable situations where the agreement is strong, and others where the two classifica-99 tions are surprisingly quasi-independent of each other. Moreover this can depend on the 100 taxonomic level. To identify potential factors that may explain variations in agreement in 101 our study: (i) we varied the taxonomic level at which the clustering is performed (order, 102 family, genus, species), (ii) we used two definitions of dissimilarity between sequences; and, 103 finally, (iii) we considered four numerical methods for the clustering of the molecular data. 104 Altogether, this leads to 32 possible combinations 105

More specifically, we first worked with 30 non random sub-samples of the whole dataset,

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each sub-sample comprising either all the individuals of an order or of a family. In each case,
we compared the botanical classification of the individuals at the next finer taxonomic level
with the molecular-based classifications. In a second step, we studied whether the mean
behaviour observed from these replicates is recovered when the set of individuals to be
classified is larger and more diverse, by comparing the botanical classification of the whole
dataset into orders with the molecular-based classification for the same number of classes.
We also performed the comparison at the family, gender and species levels.

Dissimilarities between sequences have been computed as edit distances (Levenstein, 114 1966; Gusfield, 1997). The score of local pairwise alignment (Smith and Waterman, 1981) 115 has been preferred to global pairwise alignment (Needleman and Wunsch, 1970) to avoid the 116 cost of slight lengths variations due to technological reasons in Sanger sequencing (Gusfield, 117 1997). Even if this algorithm relies on dynamic programming, thus making it very efficient 118 (and exact), its complexity is in $\mathcal{O}(n^2\ell^2)$ if n is the number of barcodes or reads, and ℓ 119 their length. This becomes prohibitive for large datasets. A classical way to circumvent 120 this difficulty is to use kmer-based distances (Sun et al., 2009; Mahé et al., 2014), a priori 121 with a decrease in the quality of the estimation of the dissimilarity, but much faster to 122 compute. A comparison between Smith-Waterman scores and kmer-based distances can 123 be found in Sun et al. (2009). The question here is to explore whether the loss in quality 124 remains acceptable and does not lead to a decrease in agreement between the botanical and 125 the molecular-based classifications. This is a preliminary step for developing further studies 126 on metabarcoding which require investment in scaling and accelerating the computation of 127 distances. 128

If the morphological-based taxonomic classification is a priori unique, this is not true for a molecular-based classification. A diversity of softwares for implementing hierarchical clustering has been proposed for more than a decade in metabarcoding with the objective of efficient scaling with respect to the growing size of environmental datasets. This includes Uclust (Edgar, 2010), ESPRIT (Sun et al., 2009) and SWARM (Mahé et al., 2014,

2015, 2019), which make it possible to cluster millions of barcodes on a laptop. Nearly all of the hierarchical clustering algorithms mentioned above rely at one step or another 135 on heuristics (like computing kmer-based distances, considering short distances only i.e. 136 working with sparse distance matrices) to make computation feasible within a reasonable 137 time with reasonable memory. SWARM uses kmers only as a first step to filter out pairs 138 of sequences which are distant and cannot belong to a same compact community. In this 139 study, we focus on understanding the agreement (or not) between molecular-based clas-140 sification from clustering and botanical classification, without computational constraints. We therefore consider Aggregative Hierarchical Clustering (AHC), whose above-mentioned algorithms can be seen as heuristic versions for scaling up, with three different aggrega-143 tion methods: Single Linkage, Complete Linkage, Ward (Murtagh, 1983; Müllner, 2013). 144 Statistical models like Bayesian classifiers with mixture models have also been considered 145 in the literature to cluster sequences (Hao et al., 2011). To extend the scope of statistical 146 modeling in molecular-based taxonomy, we explore here the potential interest of a model-147 based clustering method, the Stochastic Block Model (SBM, Holland et al., 1983; Daudin 148 et al., 2008; Lee and Wilkinson, 2019) as an alternative to AHC. SBMs are already widely 149 applied with success in domains like the social sciences (Barbillon et al., 2017), the anal-150 vsis of ecological interaction networks (Miele and Matias, 2017) and neurology (Faskowitz 151 et al., 2018). They rely on a more flexible definition of a cluster than AHC (searching for 152 general groups and not just communities), and we hypothesised that SBM and AHC could 153 be complementary in their capacity to distinguish meaningful groups of individuals in an 154 inventory. 155 In the following section, we provide a brief description of the dataset. We also de-156

156 In the following section, we provide a brief description of the dataset. We also de-157 scribe the method. Results on the quality of the agreement between molecular-based and 158 morphological-based classifications obtained on replicates are presented in Section 3.2, the 159 results obtained on the whole dataset are presented in Section 3.3.

¹⁶⁰ 2 Materials and methods

2.1 Dataset and computation of dissimilarities

This study relies on a dataset built from a collection of some 1,500 trees located in the 162 "Piste de Saint-Elie" experimental plot in French Guiana, mainly composed of lowland 163 tropical rainforest (Sabatier et al., 1997). The data used here are part of a dataset gath-164 ered for the study published in Caron et al. (2019), which focused on agreement or not 165 between botanical-based and molecular-based classification at the species level over a wide 166 range of diversity along the angiosperms tree. The main result in Caron et al. (2019) is 167 that molecular-based clustering is highly consistent with species delineation in a majority of 168 cases, and that introgression or incomplete lineage sorting are the most likely explanations 169 in the case of non-agreement. We focus here on a similar question but at the level of genera, 170 families and orders. The main elements for the material are recalled here, and the reader 171 can refer to Caron et al. (2019) for details. Among this dataset, 1,458 individual trees 172 were selected for this study. For each tree, we used the botanical name as given by field 173 botanists working with the Cayenne Herbarium of the French National Research Institute 174 for Sustainable Development, and a sequence of chloroplastic marker trnH-psbA, which is highly resolutive, despite the fact that it is variable in length. This drawback is mitigated because no multiple alignment is done: we work with pairwise distances only, computed ei-177 ther by local alignment or comparison of histogram of tetramer histograms. trnH-psbA has 178 been used is several studies or benchmarks in plant metabarcoding (Hollingsworth et al., 179 2009, 2011; Pang et al., 2012). These trees encompass 20 orders, 56 families, 182 genera 180 and 428 species. 181

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Three 1458×1458 matrices of pairwise distances or dissimilarities between sequences were built, a first one using the Smith-Waterman algorithm for local sequence alignment (Smith and Waterman, 1981), and two other ones for the distance between kmers distributions

(k = 4 and k = 6). The local alignment score is the most precise quantification of genetic 186 dissimilarities between sequences, but it is time consuming. Several methods for building 187 OTUs therefore rely on alternatives to local alignment scores. A classical way to circumvent 188 this computational burden is to build kmer counts for each sequence, and then compute 189 the distance between the normalised counts. A kmer is a contiguous sub-sequence of length 190 k in a given sequence. We selected short kmers here with k=4: there are $4^4=64$ dif-191 ferent tetramers which is a good compromise between longer ones with more resolution, 192 but too sparse histograms of counts, or smaller ones with coarse resolution and less empty 193 categories. If k = 6, there are $4^6 = 4096$ different hexamers. The length of the sequences 194 is about 450 bp, which means that at least 9 hexamers out of 10 have 0 count. For k = 8, 195 this increases up to 993 out of 1000. Moreover, for short sequences with bases labelled 196 N, there may be no hexamer without a N (met once in the dataset, this sequence has 197 been eliminated). We designed an efficient algorithm that counts the frequency of each 198 kmer in each sequence, and a short program that computes a distance between any pair 199 of frequency distributions as the ℓ^1 norm, i.e., the sum of absolute values of difference of 200 frequencies per kmer. We give here as an illustration the computation times on a standard 201 laptop. For Smith-Waterman scores, exact local alignment with dynamical programming, 202 programmed in C language: 5 hours, 39 minutes and 34 seconds. For kmer distances, with 203 k=4, programmed in C language as well: 13 minutes and 4 seconds. The time for k=6204 is 32 minutes and 6 seconds. 205

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The dataset used in the rest of this paper is composed of three files (see Frigerio et al., 2021):

- a csv file of botanical names for each sequence for order, family, genus and species;
- a csv file of pairwise dissimilarities computed with the Smith-Waterman algorithm;
 - a csv file of pairwise distances based on the comparison of tetramer and hexamer

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13 2.2 Visualisation of the whole dataset using MDS and t-SNE

A preliminary step is to propose a global picture of the dataset based on the dissimilarity 214 matrices, without a classification objective. Multidimensional Scaling (MDS) is a method 215 that, once a dissimilarity matrix between items is given, builds a point cloud in a Euclidean 216 space of prescribed dimension where each point corresponds to an item (here a sequence), 217 and such that the Euclidean distance between any two points is as close as possible to 218 the dissimilarity given in the matrix (Cox and Cox, 2001; Izenman, 2008). In our case, 219 we selected the so-called Classical Scaling, as proposed initially in Torgerson (1952). It is 220 expected that the projection of the point cloud on the first axis encompasses much of the 221 information about the structure of the point cloud. MDS was run with the dissimilarity 222 matrices built with the Smith-Waterman algorithm and as distances between tetramer 223 histograms. We also applied the t-SNE algorithm (van der Maaten and Hinton, 2008) to 224 obtain a complementary 2D representation of the point cloud. The t-SNE algorithm is 225 another technique for reduction dimension. It is based on the minimisation of a divergence 226 between a distribution probability on points' neighbours in the original space and in the 227 visualisation space. While MDS approximates at best the global structure of the distance 228 array, t-SNE gives a better summary of local structures (van der Maaten and Hinton, 2008). 229 MDS and t-SNE have been run on the whole data set (1458 sequences).

2.3 General approach

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Depending on the specific question addressed, we selected a different sample of the whole dataset. However, in all cases, the general approach for comparing two classifications was the same and can be broken down into four steps.

In step 1, we selected the sub-sample: either the whole dataset with filters, or only

the individuals of a particular order, or of a particular family. We then extracted the sub-matrix corresponding to the n individuals in the sample, from the global dissimilarity matrix based on the Smith-Waterman score. We also extracted the sub-matrix of the global kmer-based dissimilarity matrix, for k = 4 and k = 6. The next steps were applied for each sub-matrix.

In step 2, we built the classifications corresponding to AHC with the three aggregation methods, Ward, Complete Linkage (CL) and Single Linkage (SL), and to SBM (see SI for a description of these methods). The number of clusters K was provided by the botanical classification of the individuals of the sub-sample. For instance if we wanted to study agreement between the classification into families and the molecular-based ones, we cut the AHC hierarchy of classifications at K equal to the number of families in the sample, and we ran SBM for the same value. At the end of step 2, we had five different classifications of the individuals in the sub-sample.

In step 3, we compared the classifications, two by two, for each possible pair of classifi-249 cations (10 pairs in total). We used a visual tool for preliminary analysis of the agreement 250 between two classifications: Sankey plots. A Sankey plot is a flow chart in which the 251 width of an arrow is proportional to the flow. For instance, if there are $n_{kk'}$ sequences that 252 are in class k for the botanical classification and k' for a molecular-based classification, 253 there is a flow of width proportional to $n_{kk'}$ between those two clusters. We computed 254 an index as well, to quantify the agreement. Classification comparison is equivalent to 255 the comparison of two partitions of the same set, a dynamic research area with several 256 surveys (Pfitzner et al., 2009). Several indices were proposed and we chose the Normalised 257 Mutual Information (NMI1 in Pfitzner et al., 2009, see the Supplementary Information 258 for a formal definition). It is not empirical and has a sound basis in information theory, 259 as opposed to indices based on counting pairs of elements that may be non-symmetric or non-bounded or even be dependent on K or n, making comparison difficult. The Nor-261 malised Mutual Information is normalised and, as such, bounded by 0 and 1, facilitating interpretation and comparison of indices. A Normalised Mutual Information of 0 indicates independence between the two classifications, while a Normalised Mutual Information of 1 indicates a perfect agreement. For an easier interpretation, we also defined threshold on the Normalised Mutual Information values, to define domains of very good, good, average, poor and very poor agreement between two classifications. The method to compute the thresholds is based on simulated partition. It is presented in the Supplementary Material, together with the thresholds values (section 4 and Figure 1.

Finally, when replicates of the experiment are performed like in Section 2.4, in a fourth step, we analysed the distributions of the Normalised Mutual Information for a given pair of classifications in order to study trends in the agreement using histograms and boxplot representations.

2.4 Comparison of botanical and molecular-based classification at the family and genus levels, on replicates

In order to have information on the variability of the results, we created 10 sub-samples 276 of the whole dataset each of them corresponding to the individuals of a particular order, 277 and 20 sub-samples each of them corresponding to the individuals of a particular family. 278 We selected only orders (respectively families) composed of at least 15 individuals, and structured into more than one family (respectively genus). The number of individuals in 280 the sub-samples at order level varies between 15 and 321. For the sub-samples at family level 281 it varied between 17 and 127. Then, for the samples at the order level, we performed the 282 four molecular-based clustering with K equals to the number of families in that order. For 283 the samples at the family level, we chose K equals to the number of genera. The orders are 284 Malpighiales, Ericales, Sapindales, Laurales, Myrtales, Magnoliales, Gentianales, Rosales, 285 Oxalidales and Malvales. 286

We structured the empirical analysis of the Normalised Mutual Information obtained

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 $(30 \times 10 \text{ values})$ into four different analyses addressing the following questions: (i) What is 288 the level of agreement between the botanical classification and each of the four molecular-289 based ones? (ii) Are the classifications provided by the four molecular-based clusterings 290 similar? (iii) Can we identify elements of the dissimilarity matrix that explain the vari-291 ability observed in the answer to question (i) and that would be indicators of the agree-292 ment/independence between the botanical classification and the molecular-based ones? (iv) 293 How does the agreement change between the botanical classification and the molecular-294 based ones when substituting kmer-based distances for Smith-Waterman dissimilarities? 295 In practice, for question (i), we only considered the Normalised Mutual Information in-296 volving the botanical classification and any of the four molecular-based ones, whereas for 297 question (ii), we only considered the Normalised Mutual Information between any pairs of 298 the molecular-based classifications. For question (iii), we studied three factors: the taxo-299 nomic level of the groups, the entropy of the botanical classification (defined as the entropy 300 of the normalised vector of the groups sizes), and the structure of the dissimilarity matrix. 301 The latter was measured by three different ratios between the within-group dissimilarities 302 and the between-group dissimilarities (see Supplementary Information). We only present 303 here the one for which we observed a relationship with the Normalised Mutual Informa-304 tion values on the data: r_{mean} , defined as the mean of the larger within-class dissimilarity 305 over the mean of the smaller between-class dissimilarity. Intuitively when the dissimilarity 306 matrix is well structured into several groups each with a small within-class dissimilarity, 307 then r_{mean} will be lower than 1. On contrary, when there are no clearly delimited groups 308 of similar individuals then r_{mean} will be larger than 1. This is illustrated of Figure 2 in the 309 Supplementary Information. 310

2.5 Comparison of botanical and molecular-based classification on the whole data set

We looked at whether or not clustering on the whole dataset could directly retrieve botanical classification at levels higher than species (genera, families, orders). In addition, the same comparison was performed for species as well, as a benchmark. Since several taxa are singletons, regardless of the level, or have a very small number of sequences (e.g. Apiales are represented by three sequences only in the whole sample), we built one sub-sample for each taxonomic level by filtering out taxa with less sequences than a given threshold. The size of those sub-samples are given in Table 1, with the number of sequences and of different taxa per level, and the threshold selected for filtering sequences.

For a given taxonomic level, we ran SBM and AHC with Ward, CL and SL, on the 321 sub-matrix of the associated sub-sample and for K equal to the number of taxa present 322 in this sub-sample. This was done both on the matrix of dissimilarities between scores 323 of the Smith-Waterman algorithm and on distances between tetramer frequencies. 324 compared each of these four classifications with the botanical one using Normalised Mutual 325 Information. Note that a good Normalised Mutual Information at a low taxonomic level 326 does not automatically imply a good Normalised Mutual Information at a higher level. If 327 there are K_s species and K_g genera, the SBM classification into K_g classes is build without 328 using the SBM classification into K_s classes. By construction the AHC classification into 329 K_s classes is embedded into the one into K_g but depending on the structure of the dis-330 similarity matrix, the successive merges can make the AHC move away from the botanical 331 classification. 332

As for the study of the replicates, we also computed the entropy and the r_{mean} ratio of the botanical orders, families, genera and species classifications. For each of the taxonomic levels, we produced a visual graphical analysis by generating Sankey plots.

336 Results

$_{\scriptscriptstyle 37}$ 3.1 Visualisation of the whole dataset using MDS and t-SNE

We represented the shape of the point cloud on the first two axes built with MDS on 338 the dissimilarity matrix, with points coloured according to the order that they belong to 339 (see Figure 1, left). For Smith-Waterman-based dissimilarities, axis 1 clearly distinguishes 340 Ericales (in purple) and Sapindales (dark green), and axis 2, Malpighiales (in light green). 341 Axis 3 distinguishes Fabales (blue), and the set of Laurales and Magnoliales (red and 342 orange), which are primitive Eudicots. When using t-SNE (see Figure 1, right), clusters of 343 sequences appears more clearly, with less overlapping than with MDS. These clusters are in 344 general composed of sequences of the same order. However an order can be split into several 345 clusters. This phenomenon is reduced for families (see Figure 3, right, in Supplementary 346 Information), which indicated a stronger link between dissimilarities and families, than 347 between dissimilarities and orders. 348

The organisation of the point cloud is different for tetramer-based dissimilarities (see 349 Figure 4 in Supplementary Information). For MDS, the point cloud is more compact. Axis 350 1 distinguishes the same set of Laurales and Magnoliales, and axis 2 distinguishes Fabales. 351 With t-SNE also, the separation between groups is less obvious when using tetramer-based 352 dissimilarities. Clearly, the shape of the point cloud based on Smith-Waterman distances is 353 more closely related to the organisation of specimens in botanical orders. Such a connection 354 is blurred for tetramer-based distances. This allowed us to predict that agreement between 355 the botanical classification and the molecular-based ones will be lower when using tetramer-356 based distances. 357

358 3.2 Comparison of botanical and molecular-based classification at the family and genus levels on replicates

We present first the results obtained with Smith-Waterman scores for points (i) to (iii) raised in Section 2.4. We then show how results change when working with kmer-based distances (point (iv)).

- (i) Level of agreement between the botanical classification and the molecular-363 based ones. For SBM, Ward, and CL, the shape of the histogram of the 30 Normalised Mutual Information is the same (see Figure 5 of Supplementary Material). The mode is 365 observed at large values and 50 % of the values correspond to good to very good agree-366 ment, according to our definition of Normalised Mutual Information categories (see Figure 367 2). Then, intermediate values of the Normalised Mutual Information (corresponding to an 368 average agreement according to our thresholds) are not often observed. In the case of the 369 Normalised Mutual Information between SL and the botanical classification, the mode is 370 also observed at values corresponding to very good agreement, however the second mode 371 is for values of very poor agreement. So globally we observe a range of values that cor-372 respond to good to very good agreement between the botanical and the molecular-based 373 classification, with better performance for the Ward method. 374
- methods. There is a good agreement between the three AHC methods (see Figure 3). We observed larger Normalised Mutual Information between Ward and CL than between Ward and SL or CL and SL, but the median values are all in the categories good or very good. The SBM classification is globally in good agreement with Ward, in average agreement with CL and in poor agreement with SL, if we consider the median value of the Normalised Mutual Information.
- (iii) Factors explaining variability in the results. We observed no clear difference in the distribution of the Normalised Mutual Information (between the botanical classifi-

cation and the molecular-based ones) when computed on replicates whose groups are at the family level or those at the genus level or when pooling the replicates (see Table 2).

We observed a trend towards an increase in agreement between botanical and molecularbased classifications when the entropy of the sub-sample increases (Figure 4 left). We also observed a clear decrease of the agreement when the ratio r_{mean} increases (see Figure 4 right). When a dissimilarity matrix is associated with a ratio larger than 1, it can be the case that several sequences exist that are closer to sequences belonging to a different genus or family than to sequences in their own genus or family. This can lead to low Normalised Mutual Information.

(iv) Influence of the choice of dissimilarity. We observed a decrease of the Nor-393 malised Mutual Information when substituting the Smith-Waterman dissimilarity with the 394 tetramer-based or 6mer-based distances (Table 2). For k = 4, this decrease ranged between 395 6~% to 39~% depending on the taxonomic level of the groups and the molecular-based clus-396 tering method. For k=6 it is lower and ranged between 0 % and 28 %. As with the 397 Smith-Waterman dissimilarity, the agreement with the botanical classification is the high-398 est for the Ward-based classification, and we still observed the influence of the entropy of 399 the botanical classification and of r_{mean} on the agreement (Figures 6 and 7 in Supplemen-400 tary Material). From now on, we present only results for the Smith-Waterman dissimilarity 401 and for tetramer-based distances, to illustrate the best and the worst case. 402

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In conclusion, agreement between the botanical classification and molecular-based ones can be good to very good. However, there are also situations were the agreement is low. We have identified several factors that can influence the level of agreement: the choice of the clustering method, with Ward leading to the greatest agreement; the choice of the dissimilarity, with a greater agreement for Smith-Waterman dissimilarities than for kmer-based distances; the entropy of the botanical classification, with greater agreement for larger entropies; r_{mean} , with greater agreement for lower ratios.

3.3 Comparison of botanical and molecular-based classification on the whole data set

The results presented here extend the results on the replicates with four new experiments:
we compared, on the one hand, the botanical classifications of the whole dataset partitioned
into 11 orders, 20 families and 36 genera, as well as 55 species as a benchmark, (see Table 1)
and, on the other hand, the molecular-based classifications obtained for the same number
of classes.

(i) Level of agreement between the botanical classification and the molecular-418 based ones. On Figure 5 one curve is associated to one numerical method and gives the 419 value of the Normalised Mutual Information for the taxonomic levels ordered from the finer 420 to the coarser: species, genera, families and orders. All curves, regardless of the molecular-421 based clustering method and the dissimilarity, display a decrease from species to orders. 422 All of the methods are excellent for identifying species (Normalised Mutual Information are 423 in categories good or very good), and decreases depend on the method: a slight decrease 424 for the Ward method, a sharp decrease for the SL method, and an intermediate decrease 425 for CL or SBM. When groups are at orders or even families levels, SL seems to lead to 426 the lower indices, regardless of the dissimilarity used. This result illustrates that it is not 427 granted that the aggregation from fine to coarse level follows the same path in botanics 428 and in the dendrogram of the AHC. The cut of the dendogram at K_s groups, K_s being the 429 number of species, can be in good agreement with the botanical classification into species, 430 but the next merging steps of AHC may not be consistent with families and orders. 431

The correspondence between botanical, Ward and SBM classifications obtained with Smith-Waterman dissimilarities are graphically visualised in Figure 6 for orders and Figure 7 for families, with Sankey plots. We can note two types of behaviour: a botanical group is split into several groups in Ward or SBM classifications or, on the contrary a Ward or SBM group is composed of individuals from several botanical groups. The latter is more

- problematic when interpreting molecular-based classifications. On Figure 8, we can observe that the low Normalised Mutual Information for SL at the order level is due to the creation of a giant cluster formed by almost all of the orders present in the dataset.
- 440 (iii) Factors explaining variability in the results. The fact that agreement be-441 tween the molecular-based and the botanical-based classifications decreases when the tax-442 onomic level of the groups searched increases is in agreement with the influences of the 443 entropy and of the r_{mean} observed on the replicates. Indeed entropy here decreases and 444 r_{mean} increases when moving from the classification into species and genera towards families 445 and orders (see Table 3).
- (iv) Influence of the choice of dissimilarity. Regardless of the clustering method, 446 when groups are species or genera, the Normalised Mutual Information is equivalent for 447 Smith-Waterman-based dissimilarities and for kmer-based distances (the variation is at 448 most of 6%). When the groups are families or orders there is a decrease in the Normalised 449 Mutual Information when performing HAC with tetramer-based distances: Normalised 450 Mutual Information varied between 2% and 60% with the larger decrease observed for 451 SL. On contrary, for SBM, we observe a larger Normalised Mutual Information with the 452 tetramer-based distance, when groups are families or orders. 453
- Note that for AHC, the running times varied between 1 and 3 seconds, whatever the subset of sequences considered and the level of the groups searched. For clustering with SBM on tetramer distances, we used the Gaussian distribution and the running time was about 5 minutes for clustering the whole data set into orders and about 1 hour for clustering the whole data set into families. Running time was multiplied by two when using SBM on the Smith-Waterman dissimilarity with the Poisson distribution.

4 Discussion

In this study, several numerical methods were compared on a dataset of approximately 1,500 specimens of trees in a French Guianese forest for the purpose of quantifying the agreement between, on the one hand, botanical classification and, on the other hand, molecular-based classification on an array of genetic distances, on deep taxonomic levels of the classification. We discuss here the results obtained.

466 4.1 Agreement between botanical and molecular-based classifi467 cations

There is one pattern common to the study based on the clusterings of the 30 replicates and 468 the clusterings performed on the whole dataset: regardless of the combination between tax-469 onomic level and dissimilarity, AHC with the Ward aggregation criterion provides the best 470 agreement. Other methods rank differently depending on these combinations. Agreement 471 can be high (good or very good values of Normalised Mutual Information), in particular 472 when the molecular-based clustering is based on the Smith-Waterman dissimilarity. How-473 ever, we also occasionally observed low agreement, and we will discuss the reasons for this. 474 When interpreting Normalised Mutual Information values, it is important to have in mind 475 that Normalised Mutual Information is conservative in the sense that a strong agreement 476 is required to obtain a large Normalised Mutual Information value. The strength of the 477 agreement could be higher with another choice of index, but we selected Normalised Mutual 478 Information partly for this conservative behaviour. 479

A strong assumption in our study is that the number of groups K in the botanical classification is known when performing the molecular-based clustering. This is obviously not the case in real situations, like in metabarcoding of environmental samples. When K is not available, the Integrated Classification Likelihood criterion (Biernacki et al., 2000) for model selection can be used to estimate a number of groups that lead to a trade-off

between a good explanation of the dissimilarity matrix and parsimony. This criterion has the advantage to take into account the objective of clustering when comparing two 486 models (i.e. two values for K). A version for selecting K in a SBM has been proposed 487 in Daudin et al. (2008). For AHC, choosing K amounts to choosing where to cut the 488 dendogram, and heuristics have been proposed (Husson et al., 2010; Zumel and Mount, 489 2014) However, these approaches do not include a goal of agreement with the botanical classification. In White et al. (2010), to compare molecular-based clustering at the OTU 491 level and the taxonomic classification, the authors used partial assignment of the sequences 492 and the VI-cut algorithm (Navlakha et al., 2010) to automatically determine the number 493 of OTUs that optimally matches this partial knowledge. The method relies on the Value of 494 Information to compare two classifications, which we did not select for our study because it 495 is not normalised. However, the VI-cut method could easily be extended to the Normalised 496 Mutual Information and therefore provide a way to estimate the number of groups, driven 497 by the partial taxonomic knowledge that is available on some sequences of the inventory. 498 Although neighbor-joining (Saitou and Nei, 1987) is one of the reference methods in 499 phylogenies, and based on distances, we have not retained it in our study for two rea-500 sons. First, the agreement between orders and clades² (monophyly of orders) in the tree 501 is not excellent (see section 5 and Figure 8 in Supplementary Information), and second, 502 neighbor-joining is not a clustering method (Limpiti et al., 2014): the outcome cannot be 503 automatically organized as a partition into clusters. 504

4.2 Agreement of botanical classification and the AHC classifications

In our result, a variability of agreement is observed according to the linkage method. If the dataset is organised as a set of isolated clusters, all linkage methods will find them

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²A clade here is an internal node with its descent.

and provide the same classification. If not, different linkage methods will yield different classifications. Not surprisingly, we recover these behaviours in our experiments on molecular-based clustering of the tree specimens.

Ward method: The Ward method nearly always led to the best agreement with botanical classification, regardless of the measurement of distance (Smith-Waterman or kmers) and the taxonomic level of the groups (Sections 3.2 and 3.3).

Complete linkage Method The CL method generally led to the second-best agreement with the botanical classification. It provided classifications very similar to those obtained with the Ward method (see Table 3).

Single linkage method: In contrast, agreement between the classification provided by the SL method and the botanical classification was highly variable and could be either very 519 good or very poor. The agreement was very poor with the classification into orders of the 520 whole dataset (the Normalised Mutual Information is equal to 0.06 for Smith-Waterman dis-521 similarities and to 0.02 for tetramer-based distances, which is very close to independence), 522 better but still low for the classification into families (the Normalised Mutual Information 523 is equal to 0.44 for Smith-Waterman dissimilarities, and to 0.34 for tetramer-based dis-524 tances). As we explained, reason for that can be seen on Figure 8: the SL classification is 525 composed of a huge cluster, containing sequences from all orders, and a set of much smaller 526 clusters, each containing one, seldom two, orders. The creation of the huge cluster may 527 be due to low dissimilarities existing between the orders. By nature, the SL criterion will 528 link these orders by the well known "chaining effect" which produces long and thin clusters 529 which are not compact (Ros and Guillaume, 2019). 530

4.3 Interest of SBM models for molecular-based classification

Even if the SBM clustering and the botanical one are in very good agreement in some of 532 the experiments, globally, the Normalised Mutual Information values for SBM are lower 533 than the Normalised Mutual Information for the best AHC method (see Table 2 and Figure 534 5). When agreement with the botanical classification is good, then the SBM classification 535 resembles the one obtained with the Ward method. This is the case when the dissimilarity 536 matrix is well structured into communities, and all clustering methods will perform well. 537 When agreement is low, our interpretation is the following. The main difference between AHC and SBM clustering is that AHC looks for groups with small within-group dissimilarity (communities), while SBM does not impose such a constraint on the groups. It seeks for groups such that (i) all individual in group k share the same pattern of connections with the other groups, and (ii) members of group k are almost at the same distance to 542 each others. However, this distance is not necessarily small, meaning that SBM groups should not be systematically interpreted as communities. When the matrix of the pairwise 544 dissimilarities is not clearly structured according to the botanical groups, SBM clustering 545 can create groups with individuals that are far from each other. This is what we observed 546 on the SBM classification of the whole dataset into orders (both for the Smith-Waterman 547 and the kmer-based clustering). For several SBM groups, the estimated parameter char-548 acterising the mean within group distance was larger than the lower mean distance with 549 the other groups. In these situations, the Normalised Mutual Information between the 550 botanical and the SBM classification is obviously low, and the ratio r_{mean} is large. A SBM 551 classification with groups of large within-group mean distances should be a warning that 552 the matrix of dissimilarities is not entirely structured according to the botanical classes. 553 For similar reasons, SBM is also able to detect outlier individuals by gathering them into a 554 group, while methods looking for communities will force them to enter a community. For 555 these reasons, we think SBM should be considered as a valuable tool for (meta)barcoding.

$_{557}$ 4.4 Factors explaining the variations in the agreement.

One of the two main factors influencing the quality of agreement between the botanical 558 and the molecular-based classifications is the relative difference between the dissimilarities 559 within and between groups in botanical classification. This notion was well captured by 560 the r_{mean} ratio and, we obtained a clear tendency for Normalised Mutual Information 561 to decrease when the ratio increases on the 30 replicates (Figure 4). When considering 562 the clustering of the whole dataset, the same tendency was observed. The other factor 563 influencing the quality of agreement is the value of the entropy of the distribution of the group sizes in the botanical classification. We observed a tendency for an increase in 565 agreement when entropy increases, both on the 30 replicates and when clustering the whole dataset at different taxonomic levels. 567

In the latter experiments we obtained a clear decrease of agreement for high taxonomic levels, whereas in the experiments on the 30 replicates, agreement was better at the family level than at the genus level. These apparent contradictory results are actually explained by the fact that they correspond to two different protocols. On the 30 replicates the targeted set of sequences to cluster is different for each replicate: we did not search for families and genera among the same set of individuals. We instead searched for families (respectively genera) of sequences of the same order (respectively family).

The negative influence of the r_{mean} ratio and the positive influence of the entropy are global tendencies. We also observed variations around these main tendencies, which means that they are probably not the only factors explaining the Normalised Mutual Information values. Still, they are strong signals.

579 4.5 Biological interpretation

There may be several approaches to analyse the reasons for agreement/disagreement between botanical and molecular-based numerical classification. We first examine possible reasons arising from the structure of the molecular data, and second we propose some interpretations arising from the literature in plant molecular phylogenies.

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In our study on the whole dataset, the agreement between the molecular-based and the botanical-based classification is better when groups are at a low taxonomic level, hence more numerous, regardless of the method and the distance (see Figure 5). As discussed in Section 4.4, the r_{mean} ratio, involving distances within a group over distances between groups, is smaller at the family level than at the order level. This suggests that families are better delineated than orders by pairwise distances. The results shown in Figure 5 extend this observation to species over genera, and show that molecular-based delineation of taxa is more accurate at fine taxonomic levels than at coarse ones.

This is consistent with the evolution of plant classification system, where confidence 593 about delineation of higher groups, like orders, is lower than for lower groups, like gen-594 era. APG (Angiosperm Phylogeny Group) regularly updates phylogenetic classification of 595 plants, focusing on families and orders. Initial goal in APG has been to classify families 596 in orders, and later revisions have focused on delineations. In the first proposal, in 1998, 597 there were 40 orders and 462 families. In the fourth one, called APG IV (The Angiosperm 598 Phylogeny Group et al., 2016), there were respectively 64 orders and 416 families. This 599 is consistent with a stabilisation of family delineations, while there is still ongoing work 600 for stabilising orders. This might be an explanation for the drop in agreement for orders, 601 whereas the quality of agreement for families is similar to the one for genera and species 602 for some methods (see figure 5). 603

The commonly accepted notion for molecular-based classification is monophyly in molecular phylogenies (Hillis et al., 1996). The evolutionary distance between two species is the age of their Most Recent Common Ancestor. It is related to genetic distance as measured here by Smith-Waterman score, provided that the molecular clock hypothesis is accepted (see Yang (2006), chapter 7, for an overview). Even if the marker selected here is neutral

(intergenic spacer), it is highly likely that evolution rates over tens of millions of years across 609 all lineages have not remained constant. Main clades of angiosperms have radiated quickly 610 in Late Cretaceous (this is Darwin's "abominable mystery", see Friedman (2009) for a his-611 torical perspective), whereas they have diverged earlier in late Jurassic / Early Cretaceous. 612 Diversification occurred with heterogeneities in space and time (Ramirez-Barahona et al., 613 2020). It is highly likely that such heterogeneities have been reflected even partially in 614 evolution rates of markers, which may in turn lead to heterogeneities in agreement between 615 molecular based and botanical classifications at the level of orders. As a consequence, assuming that botanical classification reflects monophyletic clades can lead to a decrease of 617 agreement between botanical and molecular- based classification for higher taxa, especially 618 at the order level. 619

Our interpretation is that uncertainties on classification of plants (e.g. APG system)
are currently higher at high levels of taxonomy (orders and higher), and that this is shared
by clustering of barcodes (our numerical result).

623 4.6 Comparison between Smith-Waterman and kmer-based dis-624 similarities

Computing Smith-Waterman dissimilarity between two sequences is the most precise way 625 to compare them. However, it is time-consuming. Computing kmer-based distances is 626 much quicker, but at the cost of approximations. The histograms of Smith-Waterman dis-627 similarities and kmer-based (k = 4 and 6) distances are provided in Figures 9 and 10 of 628 Supplementary Information. A coarse correlation can be observed between both quantifi-629 cations of dissimilarities (see Figure 11 in the Supplementary Information), stronger for 630 small dissimilarities. However, a significant number of pairs of sequences exists with a 631 very low Smith-Waterman dissimilarity and a significant tetramer-based distance. This 632 is due to the high variability in length of the trnH-psbA marker. For instance, a small 633

Smith-Waterman dissimilarity means that the smallest sequence is nearly identical to a contiguous sub-sequence of the larger one. However, due to the dissimilarity in length of 635 the two sequences, the kmer histograms cannot be similar, and the kmer-based distance is 636 large. Therefore, some small values of the Smith-Waterman dissimilarity can be associated 637 with median values of the kmer-based distance. Since the AHC classification (regardless 638 of the linkage) builds groups of individuals with small within group distances, it can be 639 expected that the Smith-Waterman-based and the kmer-based classifications will be different. In practice, as expected, we observed that agreement decreases when substituting kmers for Smith-Waterman regardless of the combination between the taxonomic level and the clustering method (but for SBM sometimes). However, substituting kmer-based distances for Smith-Waterman dissimilarities did not strongly modify the agreement between the molecular-based classifications and the botanical one.

⁶⁴⁶ 4.7 Perspectives for metabarcoding

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The dataset is sufficiently small for all calculations to be run on a laptop in a reasonable time, making it possible to focus on the comparison of the methods. Some methods are clearly more accurate than others to retrieve orders or families in our dataset. The expectation is that those methods are those that will permit inventories or clustering at higher taxonomic levels such as families, orders or phyla in metabarcoding studies. However, we underline two issues.

We have worked with trnH-psbA which is highly resolutive but longer than markers currently used in metabarcoding of environmental samples or with degraded DNA. It is an issue to study whether the results found here can be extended either to other groups than plants in barcoding or with shorter markers for metabarcoding. A second issue is the scaling of the clustering methods used in the study, to data sets with hundreds of thousands of reads.

We recommend first using AHC with the Ward method for clustering regardless of the

taxonomic level, and not using AHC with Single Linkage which may produce poor results,
despite the observation that current softwares scaling up with NGS massive data sets make
it possible to use it (like MOTHUR) or yield results very close to it (like SWARM). It
can be observed that SWARM has a step for preventing the formation of giant clusters
by irrelevant aggregation between two clusters from different seeds, see Mahé et al. (2014,
2015).

Second we recommend using SBM classification to detect, via the estimated distribution of within cluster distances, situations where the molecular-based classifications may be poorly related to the morphological-based one (because the dissimilarity matrix is not clearly structured into communities).

These results and observations lead us to recommend the pursuit of methodological 670 efforts to analyse metabarcoding data for building inventories at the coarse level (i.e., 671 between phyla and orders). Inventories at the coarse level are a first step towards the 672 global exploration of diversity of unknown groups. This can be done in two ways. First, 673 nearly all surveys about clustering emphasize that there is no method that is perfect and 674 better than some others for all evaluation criteria (see, e.g., Fahad et al., 2014). Therefore, 675 it may be useful to produce classifications by several numerical methods and to extract 676 the shared elements. These are the ones in which the user can be more confident that 677 they actually reflect an actual structure in the data. Second, scaling-up methods that 678 have proven themselves to properly perform on well-known datasets, like AHC with Ward 679 linkage or SBM-based clustering, is a key issue. A very efficient method for clustering may 680 be to "divide and conquer": first, dividing the problem by building connected components 681 in a graph built from pairwise distances and, second, conquering by implementing AHC 682 Ward or SBM in each connected component. More globally, connecting these efforts with 683 studies on wider classes of methods under development for clustering for big-data (Fahad 684 et al., 2014) is a challenging issue for metabarcoding.

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696 6 Author contributions

A.F. and N.P. designed the study. M.A.A., A.F. and N.P. performed the research and analyzed the data. The paper was drafted by M.A.A., A.F. and N.P. and written by A.F. and N.P. All authors commented on and approved the final manuscript.

700 7 Data accessibility

Sequences used to compute distances are in NCBI under accession number KX247940–KX249593.

A fasta file with these sequences, a file with taxonomical assignation for each tree, as well

₇₀₃ as pairwise Smith-Waterman and kmer distances are publicly available at

https://doi.org/10.15454/XSJ079 in Inrae Data Portal.

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869 8 Tables

Taxonomic level	Number of sequences	Number of taxa	Minimal size of a taxon
Species	313	55	5
Genera	845	36	10
Families	1349	30	10
Orders	1357	11	15

Table 1: Characteristics of the four subsamples of sequences, one per taxonomic level. The number of sequences in the sample is lower for low taxonomic levels because we selected only taxa composed of a minimal number of sequences, and there are less sequences of a given species than of a given genera, etc. Each subsample is used for a comparison between the molecular-based clustering methods and the botanical classification.

		Families			Genera		Pooled			
Method		SW	4mer	6mer	SW	4mer	$6 \mathrm{mer}$	SW	4mer	$6 \mathrm{mer}$
AHC	Ward	1	0.61	0.72	0.83	0.73	0.74	0.87	0.71	0.74
	SL	0.51	0.48	0.65	0.75	0.59	0.72	0.70	0.58	0.68
	CL	0.85	0.63	0.66	0.75	0.71	0.75	0.75	0.67	0.68
SBM		0.67	0.52	0.51	0.82	0.62	0.71	0.73	0.61	0.68

Table 2: Normalised Mutual Information between the botanical classification (into families or into genera) and the four molecular-based classifications (row) for two dissimilarities (column). SW stands for Smith-Waterman, 4mer for kmer-based distances computed with kmers of length k=4 and 6mer for kmer-based distances computed with kmers of length k=6

). Results for families are median values over 10 samples and results for genera are median values over 20 samples. A sample is the set of sequences of an order (10 of them) or a family (20 of them).

	Orders	Families	Genus	Species
Entropy	2.15	3.01	3.39	3.98
r_{mean} with SW	2.22	1.3	0.60	0.30
r_{mean} with kmer	1.89	1.29	0.77	0.14

Table 3: Entropy and r_{mean} ratio (describing the ratio between mean larger within-group and mean smaller between-group dissimilarities) for the botanical classifications of the dataset into orders, families, genera and species. SW stands for Smith-Waterman and kmer for kmer-based distances computed with kmers of length k=4. Samples (one per taxonomic level) are those which have been built with the filters presented in Table 1.

9 Figures

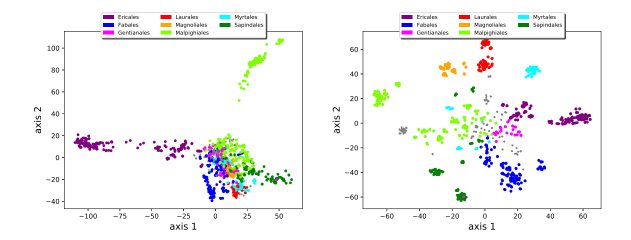


Figure 1: Visualisation of the sequences of the whole data set, as a point cloud. One dot is one sequence. The points of the eight more numerous orders are coloured, while the others are in grey. Dissimilarities are computed with the Smith-Waterman algorithm. Left: MDS, projected on axis 1 and 2. Right, t-SNE.

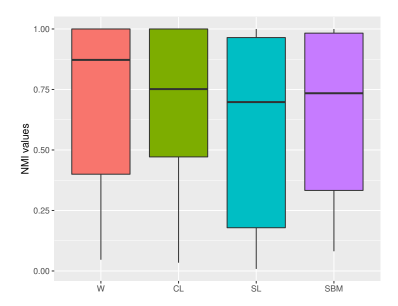


Figure 2: Boxplots on the distribution of the Normalised Mutual Information computed between each molecular-based classification and the botanical one. The data are the Normalised Mutual Information obtained on 30 replicates (10 classifications into families and 20 into genera). A replicate is the set of sequences of an order (10 of them) or a family (20 of them). Results obtained using the Smith-Waterman dissimilarity.

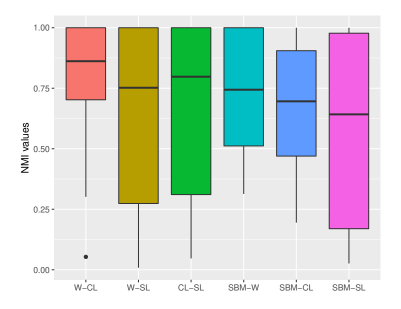


Figure 3: Boxplots on the distribution of the Normalised Mutual Information computed between each pair of molecular-based classifications. The data are the Normalised Mutual Information obtained on 30 replicates (10 classifications into families and 20 into genera). A replicate is the set of sequences of an order (10 of them) or a family (20 of them). Results obtained using the Smith-Waterman dissimilarity.

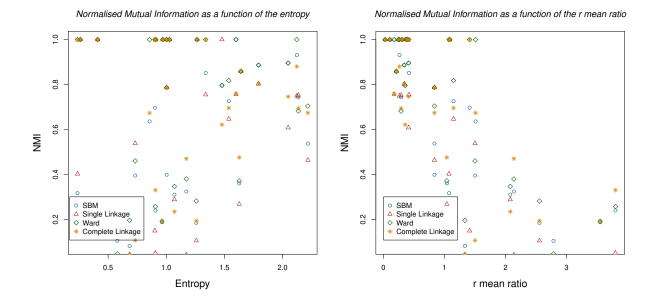


Figure 4: Normalised Mutual Information as a function of the entropy (left) and the ratio r_{mean} (right) computed on the botanical classification. Each point corresponds to one of the four molecular-based clustering method applied to one of the 30 replicates. The x-axis is the value of the entropy or ratio r_{mean} computed on the botanical classification, the y-axis is the Normalised Mutual Information between the botanical classification and the molecular-based one. The Clustering is made using the Smith-Waterman dissimilarity.

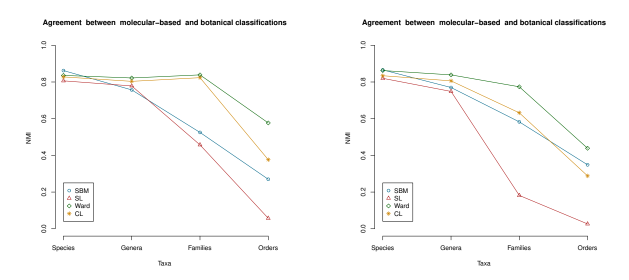


Figure 5: Agreement between molecular-based classifications and botanical classification from low to higher taxonomic levels. x axis: taxonomic levels, y axis: Normalised Mutual Information between molecular-based classification and botanical classification. One curve corresponds to one molecular-based classification. The Normalised Mutual Information are computed for classifications obtained on the samples (one per taxonomic level) presented in Table 1. Left: Smith-Waterman dissimilarities. Right: tetramer-based distances.

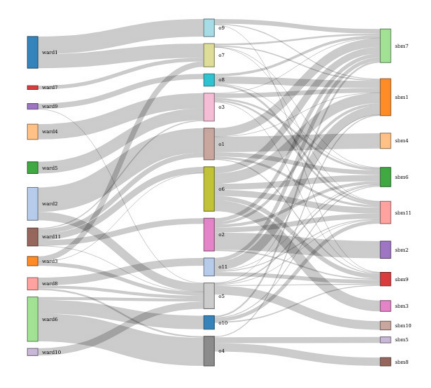


Figure 6: Sankey plot of correspondences between AHC with Ward (left column), botanical (central column) and SBM classification (right column) at the order level. The width of a flow between two classes is proportional to the number of sequences belonging to the two classes.

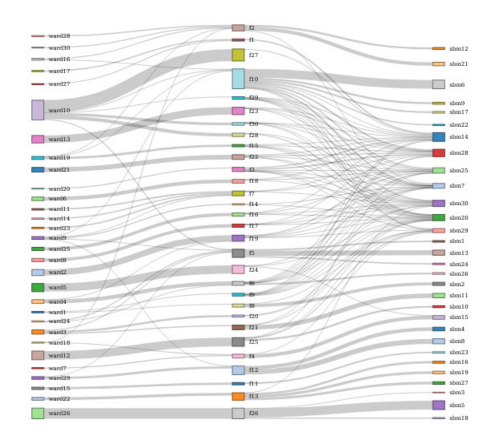


Figure 7: Sankey plot of correspondences between AHC with Ward (left column), botanical (central column) and SBM classification (right column) at the family level. The width of a flow between two classes is proportional to the number of sequences belonging to the two classes.

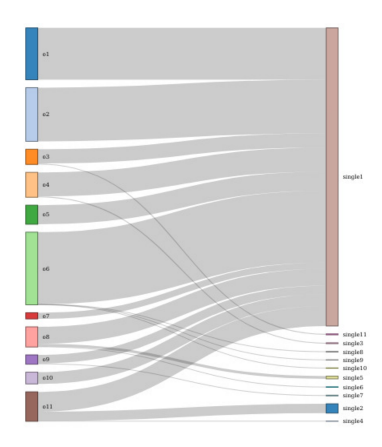


Figure 8: Sankey plot of correspondences between botanical classification (left column) and AHC with Single Linkage (right column), at the order level. The width of a flow between two classes is proportional to the number of sequences belonging to the two classes.