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RESOURCE ARTICLE



Optimal sequence similarity thresholds for clustering of molecular operational taxonomic units in DNA metabarcoding studies (1)

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Abstract

Clustering approaches are pivotal to handle the many sequence variants obtained in DNA metabarcoding data sets, and therefore they have become a key step of metabarcoding analysis pipelines. Clustering often relies on a sequence similarity threshold to gather sequences into molecular operational taxonomic units (MOTUs), each of which ideally represents a homogeneous taxonomic entity (e.g., a species or a genus). However, the choice of the clustering threshold is rarely justified, and its impact on MOTU over-splitting or over-merging even less tested. Here, we evaluated clustering threshold values for several metabarcoding markers under different criteria: limitation of MOTU over-merging, limitation of MOTU over-splitting, and trade-off between over-merging and over-splitting. We extracted sequences from a public database for nine markers, ranging from generalist markers targeting Bacteria or Eukaryota, to more specific markers targeting a class or a subclass (e.g., Insecta, Oligochaeta). Based on the distributions of pairwise sequence similarities within species and within genera, and on the rates of over-splitting and over-merging across different clustering thresholds, we were able to propose threshold values minimizing the risk of oversplitting, that of over-merging, or offering a trade-off between the two risks. For generalist markers, high similarity thresholds (0.96-0.99) are generally appropriate, while more specific markers require lower values (0.85-0.96). These results do not support the use of a fixed clustering threshold. Instead, we advocate careful examination of the most appropriate threshold based on the research objectives, the potential costs of over-splitting and over-merging, and the features of the studied markers.

KEYWORDS

alpha diversity, COI, metabarcoding marker, MOTU over-merging, MOTU over-splitting, sequence variant

1 | INTRODUCTION

DNA metabarcoding studies are typically based on a succession of experimental steps governed by important methodological choices (Zinger et al., 2019). These include (i) the definition of sampling design and the selection of sampling sites (Dickie et al., 2018), (ii) the approach used for the preservation of the starting material

(Guerrieri et al., 2021; Tatangelo et al., 2014), (iii) the protocol used for DNA extraction (Capo et al., 2021; Eichmiller et al., 2016; Lear et al., 2018; Taberlet et al., 2012; Zinger et al., 2016), (iv) the selection of appropriate primers to amplify a taxonomically informative genomic region (Elbrecht et al., 2016; Fahner et al., 2016; Ficetola et al., 2021), (v) the strategy adopted for DNA amplification and highthroughput sequencing of amplicons (Bohmann et al., 2022; Nichols

et al., 2018; Taberlet et al., 2018), (vi) the pipeline selected for bioinformatics analyses (Boyer et al., 2016; Calderón-Sanou et al., 2020; Capo et al., 2021; Couton et al., 2021; Macher et al., 2021; Mächler et al., 2021) and (vii) the statistical approach used to translate metabarcoding data into ecological information (Chen & Ficetola, 2020; Paliy & Shankar, 2016). Each of these methodological choices can heavily influence the reliability and interpretation of results (Alberdi et al., 2018; Zinger et al., 2019), and there is thus a critical need for development, proper assessment and optimization of methods specially dedicated to DNA metabarcoding.

When analysing metabarcoding data, bioinformatic pipelines generally produce a list of detected sequences that can be assigned to a given taxon with a more or less precise taxonomic resolution. However, the number of unique sequences obtained after bioinformatic treatment is generally much higher than the number of taxa actually present in the sample (Calderón-Sanou et al., 2020; Mächler et al., 2021). This stems from multiple reasons including genuine intraspecific diversity of the selected markers and errors occurring during the amplification or sequencing steps. Consequently, sequence clustering approaches are often used to collapse very similar sequences into one single molecular operational taxonomic unit (MOTU), which does not necessarily correspond to a species in the traditional sense (Antich et al., 2021; Bhat et al., 2019; Froslev et al., 2017; Kopylova et al., 2016). Sequence clustering can be performed using similarity thresholds, using Bayesian approaches or through single-linkage (Antich et al., 2021). Approaches based on similarity thresholds can have excellent performance and they display several advantages such as flexibility and easy implementation (Kopylova et al., 2016; Wei et al., 2021). However, when performing clustering based on sequence similarity, two key parameters have to be determined a priori. The first is the sequence to be selected as representative of the cluster. In the case of metabarcoding studies, keeping the most abundant sequence of the cluster as the cluster representative is a convenient way of merging sequence variants generated during the PCR or sequencing steps with the original sequence they derive from (Mercier et al., 2013). The second parameter is the similarity threshold (clustering threshold) used to build MOTUs (Calderón-Sanou et al., 2020; Clare et al., 2016; Wei et al., 2021). The choice of this threshold is delicate without prior knowledge of the marker and its intrinsic level of diversity. A too low threshold can collapse different taxa into the same MOTU (over-merging), while a too high threshold can create too many MOTUs (over-splitting) compared to the actual level of diversity (Clare et al., 2016; Roy et al., 2019; Schloss, 2021).

Some works suggest that the ecological interpretation of metabarcoding data can be relatively robust to the threshold selected for sequence clustering. For instance, Botnen et al. (2018) used thresholds of sequence similarity ranging from 0.87 to 0.99 to analyse multiple microbial communities, and obtained community structures highly coherent across thresholds. Nevertheless, levels of alpha diversity can be heavily impacted by the threshold selection. Ideally, the threshold used for clustering would depend on a trade-off between MOTU over-splitting and MOTU over-merging. A growing number of markers are currently being used in metabarcoding studies (Taberlet et al., 2018), with some allowing broad-scale

biodiversity assessment but having limited taxonomic resolution (e.g., 18S rDNA primers amplifying all eukaryotes; Guardiola et al., 2015) and others being highly specific to one single class or even family (e.g., Baamrane et al., 2012; Ficetola et al., 2021). Biodiversity surveys generally aim to generate a set of MOTUs that are each associated with a unique taxon, all taxa being ideally situated at the same level in the taxonomic tree, in order to facilitate comparisons. In these conditions, optimal clustering thresholds probably differ strongly across markers. One can, for example, expect high values for highly conserved markers, and lower values for markers showing high variability (Brown et al., 2015; Kunin et al., 2010). However, there is limited quantitative assessment of how optimal clustering thresholds vary across markers (but see Alberdi et al., 2018).

In this study, we analysed sequences from a public database (EMBL-European Molecular Biology Laboratory) to identify clustering thresholds for different markers and under different criteria. We considered nine metabarcoding markers (Table 1), ranging from generalist markers (i.e., targeting Bacteria or Eukaryota) to more specific markers (e.g., targeting Oligochaeta [earthworms], Insecta [insects] or Collembola [springtails]), and amplifying fragments situated either in protein-coding (e.g., cytochrome c oxidase subunit 1 [COI] mitochondrial gene) or nonprotein-coding (e.g., rDNA genes) genomic regions. We evaluated how clustering thresholds can change for each marker and taxonomic group, depending on the criterion adopted to set the threshold. We used two alternative strategies to identify thresholds, each time with different objectives in mind. First, following a procedure similar to the one adopted in barcoding studies (Machida et al., 2009; Meyer & Paulay, 2005), we compared the distribution probabilities of sequence similarities among different individuals of the same species and among different species of the same genus to identify values: (i) minimizing the risk that different sequences of the same species are split in different MOTUs (i.e., risk of over-splitting); (ii) minimizing the risk that distinct but related species are clustered in the same MOTU (i.e., risk of over-merging); and (iii) balancing the risk of over-splitting and over-merging (Figure 1a). Second, we calculated the over-splitting and over-merging rates of the studied markers for a range of clustering thresholds, to identify values that minimize the two error rates (Figure 1b). We expect that, if researchers want to minimize over-splitting, they should select lower clustering thresholds than if they want to minimize over-merging. Furthermore, we expect higher clustering thresholds for generalist markers compared to markers targeting one class or more restricted taxonomic groups, because of the lower taxonomic resolution and slower evolutionary rate of the former.

METHODS 2

Markers examined and construction of 2.1 sequence data sets

We focused on a set of nine DNA metabarcoding markers (Bact02, Euka02, Fung02, Sper01, Arth02, COI-BF1/BR2, Coll01, Inse01, Olig01) targeting different taxonomic groups and different genomic regions (Table 1). Four of these markers can be considered as

				Taxonom	Taxonomic resolution (%) ^a)a		
Marker	Target gene	Target group	Taxonomic level	Species level	Genus level	Genus level Family level	Order level	Reference(s)
Bact02	V4 region of the 16S rDNA gene	Bacteria	Superkingdom	19.6	55.7	55.1	60.2	Taberlet et al. (2018)
Euka02	V7 region of the 18S rDNA gene	Eukaryota	Superkingdom	47.0	59.5	68.3	67.1	Guardiola et al. (2015)
Fung02	ITS1 nuclear rDNA gene	Fungi	Kingdom	72.5	90.2	87.7	85.5	Epp et al. (2012); Taberlet et al. (2018)
Sper01	P6 loop of the intron of the chloroplastic <i>trnL</i> gene	Spermatophyta	Clade < kingdom	21.5	36.9	77.4	89.6	Taberlet et al. (2007)
Arth02	16S mitochondrial rDNA gene	Arthropoda	Phylum	68.6	89.6	97.5	100.0	Taberlet et al. (2018)
COI-BF1/BR2	Cytochrome c oxidase subunit 1 mitochondrial gene	Arthropoda	Phylum	85.6	97.0	95.1	93.5	Elbrecht and Leese (2017)
Coll01	16S mitochondrial rDNA gene	Collembola	Class	80.5	87.2	75.0	NA	Janssen et al. (2018)
Inse01	16S mitochondrial rDNA gene	Insecta	Class	87.8	96.8	95.4	79.3	Taberlet et al. (2018)
Olig01	16S mitochondrial rDNA gene	Oligochaeta	Subclass	89.3	95.7	100.0	100.0	Bienert et al. (2012); Taberlet et al. (2018)
^a Percentage of di	scriminated taxa among taxa amplified in si	lico, as calculated by	<pre> the ecotaxspecificit) </pre>	y program i	from the obitool	.s. Reported fron	n Taberlet et al	^a Percentage of discriminated taxa among taxa amplified in silico, as calculated by the ecotaxspecificity program from the oBITOOLS. Reported from Taberlet et al. (2018) for all markers, except for COI-BF1/

for which these values were determined using the sequences amplified in silico from EMBL r140.

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generalist (i.e., targeting entire superkingdoms or kingdoms): Bact02 targeting Bacteria, Euka02 targeting Eukaryota, Fung02 targeting Fungi and Sper01 targeting Spermatophyta (vascular plants). Two markers were intermediate (Arth02 and COI-BF1/BR2, both targeting arthropods, i.e., the most species-rich phylum on Earth). Finally, three markers were more specific (i.e., targeting groups from classes to subclasses): CollO1 targeting Collembola (springtails), InseO1 targeting Insecta and Olig01 targeting Oligochaeta (earthworms). Eight of these markers are situated in nonprotein-coding genes (Bact02, Arth02, Coll01, Inse01 and Olig01: 16 rDNA gene; Euka02: 18S rDNA gene; Fung02: ITS1 nuclear rDNA gene; Sper01: P6 loop of the intron of the chloroplastic trnL gene). The last marker, COI-BF1/ BR2, is situated in the COI mitochondrial gene (Table 1).

For each of these markers, a sequence database was built from EMBL release 140 (final sequence databases available at https:// doi.org/10.5061/dryad.crjdfn353) as follows. An in silico PCR was first carried out by running the program ECOPCR (Ficetola et al., 2010) using the corresponding primers (Table S1). Three mismatches per primer were allowed (-e option), and amplicon length (without primers) was restricted (-I and -L options) to the expected length interval (Table S1). The amplified sequences were further filtered by keeping only those belonging to the target taxonomic group, showing a taxonomic assignment (i.e., taxid) at the species and genus levels and having no ambiguous nucleotides. This allowed assembling a working data set, from which we extracted two subdata sets. The "withinspecies" data set was built by keeping only species for which at least two sequences (identical or not) were available; if more than two sequences were available for a given species, we randomly selected two sequences for that species using the obiselect command of the OBITOOLS. The "within-genus" data set was built by keeping only genera for which at least two sequences were available; if more than two sequences were available for a given genus, we randomly selected two sequences for that genus using the obiselect command. For some markers (Bact02, Euka02, Fung02, Inse01, Sper01), the within-species data set and sometimes the within-genus data set still contained a very large number of sequences (>10,000). To limit computation time for these markers, we randomly selected a subset of 5000 different taxa, to reach a final number of sequences equal to 10,000. An example of data set preparation is provided in Script1_ Arth02 DatasetsPreparation.sh (Appendix S1), and Table S2 summarizes the number of sequences in the different data sets.

Calculation of sequence similarities and 2.2 probability distributions

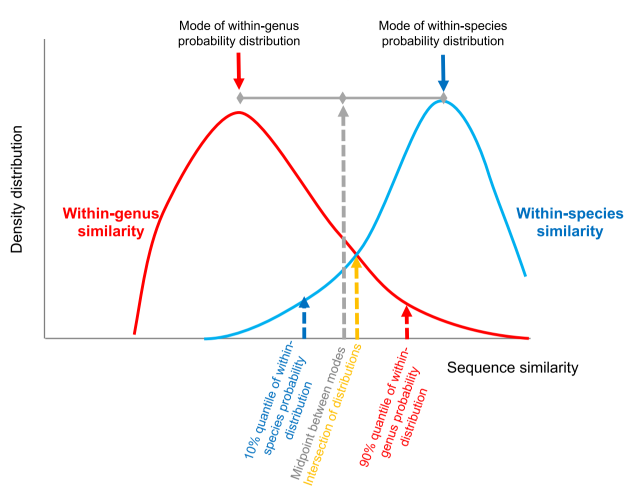
As a measure of sequence similarity, we computed the pairwise LCS (longest common subsequence) scores between pairs of sequences in the within-species and within-genus data sets using the SUMATRA program (Mercier et al., 2013; see Script2A_ Arth02_PairwiseSimilarities_Sumatra.sh from the Appendix S1). Methodological comparisons showed that this algorithm provides an excellent balance between performance and computation efficiency

Characteristics of the nine studied markers

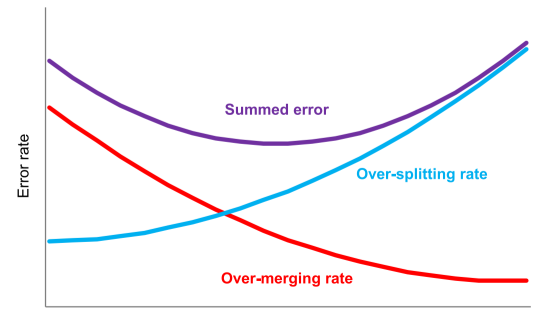
TABLE 1

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(a) Approach based on within-species and within-genus sequence similarities



(b) Approach based on over-splitting and over-merging rates



Clustering threshold

FIGURE 1 Different approaches to identify the most appropriate clustering thresholds. (a) Approach based on similarities between sequences belonging to different individuals from the same species (blue curve), and similarities between sequences belonging to different species from the same genus (red curve). One can choose to minimize the risk that different sequences from the same species are split into different MOTUs (over-splitting risk; e.g., 10% quantile of the distribution of within-species similarities), the risk that sequences from different species belonging to the same genus are clustered in the same MOTU (over-merging risk; e.g., 90% quantile of within-genus similarities), or one can try to find a balance between the risks of over-splitting and over-merging (e.g., with the intersection between probability distributions, or the midpoint between the modes of both distributions). (b) Approach based on rates of over-splitting and over-merging. One can compare the over-splitting (blue) and the over-merging (red) rates, and/or one can identify the thresholds minimizing the sum of these rates (violet).

(Bhat et al., 2019; Jackson et al., 2016; Kopylova et al., 2016). As SUMATRA provides pairwise scores for all possible pairs of sequences, the similarity scores resulting from the within-species data set were filtered in R (R Core Team, 2020) to keep only those representing similarities between sequences of the same species. Similarly, the scores resulting from the within-genus data set were filtered to keep only those representing similarities between different species of the same genus (see first part of Script2B_Arth02_DensityPlots.Rmd from the Appendix S1).

2.3 | Approach to identify clustering thresholds on the basis of within-species and within-genus sequence similarities

We first examined within-species and within-genus sequence similarities to evaluate four different strategies (Figure 1a) and determine the similarity value that: (i) avoids over-splitting; (ii) avoids over-merging; or (iii) finds a balance between over-splitting and overmerging, with two distinct procedures based on the intersection (iii-a) or on modes (iii-b) of the density probability distributions (see Script2B_Arth02_DensityPlots.Rmd from the Appendix S1). These strategies are analogous to those adopted in traditional barcoding studies to set the limit between intra- and interspecific diversity (Meyer & Paulay, 2005).

2.3.1 | Avoid over-splitting

In this case, the aim is to avoid distributing different sequences belonging to the same species in different clusters (i.e., to limit the probability of generating additional spurious MOTUs). For this purpose, we selected as clustering threshold the 10% quantile of the distribution of similarities between sequences from the same species (within-species data set). With this approach, the sequences belonging to the same species according to EMBL are gathered in the same cluster in 90% of the cases.

2.3.2 | Avoid over-merging

In this case, the aim is to avoid gathering sequences attributed to different species of the same genus in the same cluster (i.e., to limit the probability of merging related species in the same MOTU). For this purpose, we selected as clustering threshold the 90% quantile of the distribution of similarities between different species belonging to the same genus. With this approach, the sequences attributed to different species belonging to the same genus are assigned to different clusters in 90% of the cases.

2.3.3 | Find a balance between over-splitting and over-merging

In this case, the aim was to minimize both over-splitting and overmerging. We considered two distinct approaches. First, we obtained the probability distribution of within-species and within-genus sequence pairwise similarities using the *density* function from R, with biased cross-validation (bw = "bcv") as smoothing bandwidth selector and a Gaussian smoothing kernel (kernel = "gaussian"; Venables & Ripley, 2002). We tested other possible smoothing bandwidth selectors, but biased cross-validation was the approach best fitting the score histograms for all markers and all data sets (Figures S1– S9). The balance threshold iii-a was then identified as the intersection between the probability distributions of the within-species and within-genus similarities. As an alternative approach to balance over-merging and over-splitting (iii-b), we calculated the midpoint between the modes of the within-species and within-genus probability distributions.

2.4 | Rates of over-merging and over-splitting

For each marker, over-merging and over-splitting rates were evaluated at different clustering thresholds using the within-species data set described in the section "Markers examined and construction of sequences data sets." This data set contains two sequences at random, identical or not, for a number of species belonging to the taxonomic group of interest.

For each within-species data set, clustering was performed using the sumactust program (Mercier et al., 2013; see Script3A_Arth02_ Clustering.sh from the Appendix S1) with the *-n* option (normalization by alignment length) based on the sequence similarities first calculated using the sumatra program (see above; Mercier et al., 2013). Threshold values (*-t* option) ranging from 0.90 to 1 at steps of 0.01 were tested for all markers except Coll01 and Olig01 for which wider ranges ([0.70–1] and [0.80–1], respectively) were selected based on the within-genus and within-species sequence similarity probability distributions determined previously (see Figure 2). Clustered data sets were then explored to calculate five different variables at each clustering threshold (see Script3B_Arth02_Oversplitting_ Overmerging.Rmd from the Appendix S1): (1) the number of clusters; (2) the percentage of MOTUs containing one single species; (3) the percentage of MOTUs containing one single genus; (4) the percentage of species gathered in one single MOTU; and (5) the percentage of genera gathered in one single MOTU among genera represented by several sequences. Variables 2 and 3 are indicative of appropriate MOTU merging of sequences at the species and genus levels, respectively, while variables 4 and 5 are indicative of appropriate MOTU splitting at the species and genus levels, respectively.

These values were also used to calculate three measures of error. We defined the over-merging rate as (100-the percentage of MOTUs containing one single species)/100; and the oversplitting rate as (100-the percentage of species gathered in one single MOTU)/100. These two values belong to a [0,1] interval. The summed error rate was then calculated as the sum of the overmerging and over-splitting rates. For this estimate, we assigned the same weight to over-splitting and over-merging.

3 | RESULTS

Our in-silico PCRs amplified between 101,955 (Arth02) and 3,202,507 (Bact02) sequences per marker (Table S2). After data filtering, we retained between 510 (Coll01) and 707,874 (Bact02) sequences per marker. The within-species data set comprised between 118 (Coll01) and 10,000 (Bact02, Euka02, Fung02, Sper01, COI-BF1/BR2, Inse01) sequences, while the within-genus data set comprised between 74 (Coll01) and 10,000 (Euka02 and Sper01) sequences per marker.

3.1 | Clustering thresholds determined from probability distributions of within-species and withingenus sequence similarities

The probability distributions of within-species and within-genus sequence similarities showed very contrasting patterns between the generalist and the specific markers (Figure 2). For Arth02 and most of the markers targeting broad taxonomic groups (Bact02, Euka02 and Sper01), the distributions of within-species and within-genus similarities were rather similar, both showing a mode at very high similarity values (Figure 2). Fung02 showed a slightly different pattern, as the within-genus similarities had a very broad distribution. Conversely, for COI-BF1/BR2 and the more specific markers (Coll01, Inse01 and Olig01), the distributions of sequence similarities were very different, with two clearly distinct peaks. Within-species similarities remained very high (mostly above 0.95), while within-genus similarities generally showed lower values (mode around 0.88–0.90 for COI-BF1/BR2 and Inse01, and below 0.80 for Olig01 and Coll01).

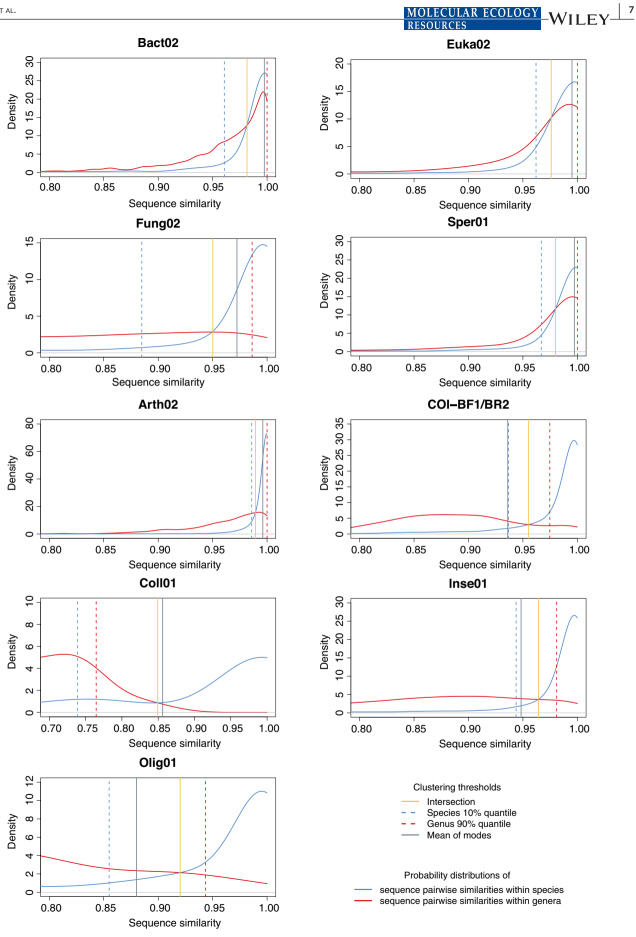
For all markers, criterion i (avoid over-splitting) yielded the lowest thresholds (Table 2), with very low values for Coll01 and Olig01. Conversely, criterion ii (avoid over-merging) yielded extremely high values, except for Coll01. For all generalist markers and Arth02, limiting over-merging would require setting clustering thresholds at 0.99 or higher. The same objective would entail a slightly lower threshold for COI-BF1/BR2 and Inse01 (0.98) and down to 0.94 for Olig01. For Coll01, criterion ii resulted in a very low threshold (0.77), because many within-genus comparisons showed very low similarity values.

Criteria iii-a and iii-b, searching a balance between over-merging and over-splitting, yielded somehow contrasting results across markers. For COI-BF1/BR2 and the three specific markers (Coll01, Inse01 and Olig01), the within-genus and within-species similarities showed clearly distinct peaks (Figure 2). As a consequence, the intersection between the two curves could effectively represent the point minimizing both over-merging and over-splitting (see Discussion), and the midpoint between the modes also identified rather similar threshold values. By contrast, for the generalist markers and Arth02, the within-species and within-genus similarities showed very high overlap and similar modes, and the density distributions actually intersected at values lower than both modes. The midpoint between the modes continued to identify threshold values intermediate between the peaks of within-species and within-genus similarities.

3.2 | Rates of over-splitting and over-merging

For all markers, irrespective of the clustering threshold examined (values ≥ 0.70 for Coll01, ≥ 0.80 for Olig01 and ≥ 0.90 for the other markers), the percentage of MOTUs containing one single species was higher than 50%, and that of MOTUs containing one single genus was higher or close to 70% (Figure 3). Overall, for the generalist and intermediate markers, these two percentages showed a regular increase with the clustering threshold. For the specific markers as well as Fung02 and COI-BF1/BR2, they reached values close to 100% for high thresholds. Unsurprisingly, the two percentages tended to be lower for the generalist markers than for the specific markers at a given threshold, indicating that the former are more sensitive to over-merging. Fung02 was a notable exception, since about 87% and 97% of MOTUs contained one single species and one single genus, respectively, at the 0.97 threshold, which is

FIGURE 2 Density probability distributions of sequence pairwise similarities within species (blue lines) and within genera (red lines) for the nine studied markers. For each marker, vertical dotted lines represent the 10% quantile of the within-species probability distribution (blue; threshold limiting over-splitting) and the 90% quantile of the within-genus probability distribution (red; threshold limiting over-merging). Vertical full lines represent the intersection of the within-species and within-genus probability distributions (yellow, balance-a) and the midpoint between modes (grey, balance-b).



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Citerion I: Avoid over-splitting (10% quantile of within) 0.961 0.885 0.964 0.855 species probability distribution) 0.001 0.000 0.986 0.976 0.975 0.975 0.976 0.974 0.855 Criterion II: Avoid over-merging (90% quantile of within-genus probability distribution) 1.000 1.000 0.986 1.000 0.976 0.947 0.944 0.944 Criterion III:-S Balance-a (intersection of within-species 0.982 0.949 0.976 0.949 0.976 0.949 0.949 0.944 0.944 Criterion III-S Balance-a (intersection of within-species 0.982 0.949 0.949 0.946 0.944 Criterion III-S Balance-b (intersection of within-species 0.976 0.949 0.976 0.976 0.976 0.976 0.949 0.964 0.920 Criterion III-S Balance-b (intersection of within-species 0.976 0.976 0.976 0.976 0.976 0.976 0.976 0.976 0.976 0.976 0.976 0.976 0.976 0.976 0.964 0.926 <th></th> <th>Bact02</th> <th>Euka02</th> <th>Fung02</th> <th>Sper01</th> <th>Arth02</th> <th>COI-BF1/BR2</th> <th>Coll01</th> <th>Inse01</th> <th>CESO 10gilo</th>		Bact02	Euka02	Fung02	Sper01	Arth02	COI-BF1/BR2	Coll01	Inse01	CESO 10gilo
1.000 1.000 0.986 1.000 1.000 0.975 0.981 cies 0.982 0.949 0.980 0.989 0.955 0.849 0.964 0.997 0.995 0.997 0.996 0.936 0.948 0.948	id over-splitting (10% quantile of within- bability distribution)	0.961	0.962	0.885	0.967	0.986	0.937	0.739	0.944	URCES 558.0
ies 0.982 0.976 0.949 0.980 0.989 0.955 0.849 0.964 0.97 0.995 0.972 0.996 0.936 0.936 0.948	oid over-merging (90% quantile of us probability distribution)	1.000	1.000	0.986	1.000	1.000	0.975	0.765	0.981	0.944
0.997 0.995 0.972 0.997 0.996 0.936 0.856 0.948	Criterion iii-a: Balance-a (intersection of within-species and within-genus probability distributions)	0.982	0.976	0.949	0.980	0.989	0.955	0.849	0.964	0.920
	Balance-b (midpoint between modes)	0.997	0.995	0.972	0.997	0.996	0.936	0.856	0.948	0.880

Values of the different thresholds estimated for the nine studied markers on the basis of within-species and within-genus sequence similarities

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frequently adopted as the clustering threshold for fungal ITS sequences. These values were comparable to those observed for COI-BF1/BR2 and the specific markers, for which >85% and >98% of MOTUs contained one single species or one single genus, respectively, for thresholds \geq 0.95.

The percentages of species and genera gathered in one single MOTU decrease at a similar rate with the clustering threshold, with generally a sharp drop at high thresholds (≥0.98; Figure 3). However, the pattern of MOTU splitting was less characteristic of generalist vs. specific markers. For some markers (Euka02, Sper01, Arth02, Inse01), the percentage of species or genera gathered in a single MOTU remained higher or close to 50% up to high thresholds (0.98). By contrast, for Bact02, Fung02, COI-BF1/BR2, Coll01 and Olig01, these percentages dropped quickly when the clustering threshold increased, indicating that these markers are susceptible to over-splitting.

For all markers, the number of clusters generally increased regularly with the clustering threshold up to 0.97–0.98 (Figure 3), followed by a sharp rise up to 1 (but which was less obvious for Euka02 and Olig01). For example, for Bact02, the number of clusters more than doubled between 0.97 (2862 clusters) and 1 (6461 clusters).

Our results showed clear patterns for over-merging and oversplitting rates, with over-splitting quickly increasing and overmerging quickly decreasing at high clustering thresholds (Figure 4). For several markers, the summed error showed a relatively clear minimum at specific clustering thresholds (Figure 4): 0.96–0.99 for Bact02, 0.97–0.99 for Euka02 and Arth02, 0.96–0.98 for Sper01, 0.93–0.96 for COI-BF1/BR2 and 0.94–0.97 for Inse01. The minimum was much less evident for Fung02, Coll01 and Oligo01, these markers showing relatively similar summed error rates over a broad range of clustering thresholds (Fung02: 0.91–0.98; Coll01: 0.89–0.97, with multiple minima; Oligo01: 0.84–0.96, with multiple minima).

4 | DISCUSSION

Sequence clustering approaches are routinely used for the identification of MOTUs in metabarcoding studies, and they often resort to methods based on similarity values. Still, selecting a clustering threshold for a given marker more than often relies on common practices and rules of thumb rather than on proper scientific argument. By analysing extensive sequence data deposited in public databases for a range of generalist and specialist markers, we showed that different thresholds can be selected depending on the marker and on the criterion favoured by researchers. All studied markers except one (COI-BF1/BR2) are situated in nonprotein-coding genes (Table 1), and this has an influence on levels of sequence diversity. More variability might be expected in protein-coding genes due to the redundancy of the genetic code. Yet, for all markers including COI-BF1-BR2, the 10% quantile of the within-species similarity probability distribution was almost always lower than the 0.97 clustering threshold traditionally used in barcoding for markers targeting protein-coding genes such as COI (Hebert et al., 2003), or for

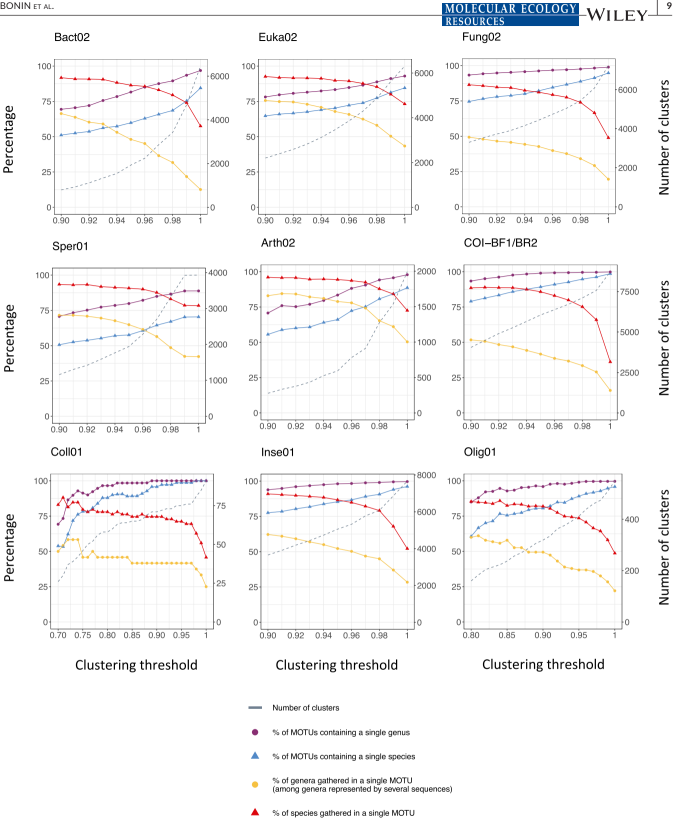


FIGURE 3 Evolution of over-splitting and over-merging rates for a range of clustering thresholds, for the nine studied markers.

microbial MOTU delimitation (Bálint et al., 2016). This indicates that some level of over-splitting can occur when using this threshold.

COI-BF1/BR2 is the only marker amplifying a fragment of a protein-coding gene, and it would have been logical to observe singular patterns for this marker. However, this was not the case, and COI-BF1/BR2, although designed to target arthropods (Elbrecht & Leese, 2017) like Arth02, actually showed a behaviour very similar to the more specific Inse01 targeting insects. The similarity

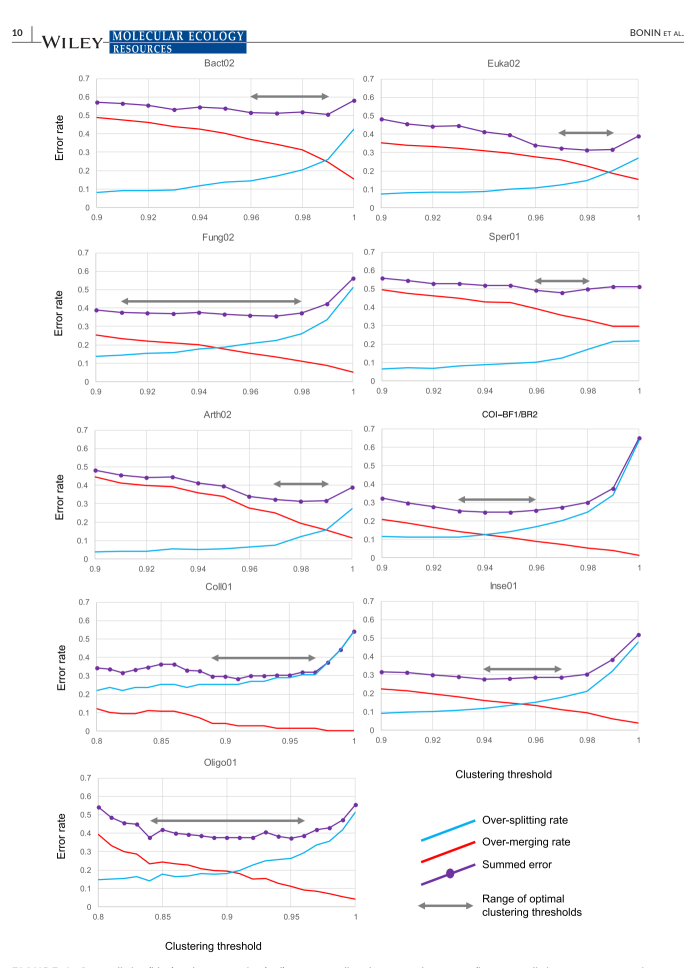


FIGURE 4 Over-splitting (blue) and over-merging (red) rates, as well as the summed error rate (i.e., over-splitting rate + over-merging rate; violet), for the nine studied markers across a range of clustering thresholds. Horizontal grey arrows indicate the range for which the summed error rate is minimal.

between COI-BF1/BR2 and the more specific markers might be related to their high resolution, which allows the successful distinction of closely related species even on the basis of relatively short sequences (Elbrecht & Leese, 2017; Ficetola et al., 2021). Furthermore, at 0.94, which is a suitable clustering threshold for COI-BF1/BR2, about 88% of the MOTUs contain a single species, and about 88% of the species are gathered in a single MOTU (Figure 3), indicating that MOTU richness at this threshold is a reasonably good proxy for the number of species detected with this marker. This is corroborated by the number of clusters observed at this threshold (5659), which is comparable to the expected number of species (5000; Table S2) in the within-species data set used to obtain Figure 3. Several COI markers are routinely used in metabarcoding, and COI-BF1/BR2 shows a large overlap with many of them (Elbrecht & Leese, 2017). We can thus expect that optimal clustering thresholds for COI-BF1/ BR2 can also be rightfully applied to markers targeting a slightly different COI region.

Although the within-genus similarity values were generally lower than the within-species similarities for all the markers, the overlap between the two distributions was dependent on the generalist vs. specific nature of the marker. For some specific markers (e.g., Coll01 and Olig01), distinct peaks were visible for the two similarity metrics (Figure 2). Within-species similarities generally were >0.90, while within-genus values were < 0.80. Such a pattern is expected for markers with an excellent taxonomic resolution and designed to identify taxa at the species level. Conversely, for the generalist markers, within-species and within-genus similarity probability distributions largely overlapped and the differences between the peaks were minimal. Nevertheless, even for these markers, the density of the within-species similarity distribution was consistently higher than that of the within-genus similarity distribution at high similarity values. This suggests that the probability of observing the corresponding sequence similarity is higher within species than within genera. In other words, at high sequence similarities, an MOTU is more likely to represent a species than a genus. This result is confirmed by the fact that the percentage of MOTUs containing a single species is always higher than 50%, whatever the clustering threshold or the marker considered (Figure 3).

The sequences used as a primary source of information in this study were downloaded from the EMBL public database, and therefore our results are probably highly dependent on the quality of the data deposited. Even though broad-scale analyses suggest that sequence data from public databases are generally reliable (Leray et al., 2019), errors in the sequence itself (e.g., wrong nucleotide, or more complex errors such as insertions, deletions, inversions, duplications or pseudogene sequences) and taxonomic mislabelling can occur. Organisms that are difficult to identify based on morphology are particular susceptible to wrong taxonomic information (Bidartondo, 2008; Bridge et al., 2003; Mioduchowska et al., 2018; Valkiūnas et al., 2008). While errors in the sequence will affect within-species sequence similarity negatively, the effect of taxonomic mislabelling is more diffuse. For example, in a group such as springtails where species delimitation is tricky (Porco et al., 2012), MOLECULAR ECOLOGY RESOURCES -WILEY

the existence of cryptic species will decrease within-species sequence similarity while increasing over-splitting rates. In a group such as Bacteria, type strains are sometimes entered at the species level in the NCBI (EMBL) taxonomy (Federhen, 2015), leading to an inflation of within-genus similarity and over-merging rates. Regardless, database errors will make within-species and within-genus similarity distributions more difficult to distinguish and clustering thresholds trickier to identify, and thus the over-splitting or over-merging rates reported here could be artificially higher than in reality.

In this work, we came up with a global measure of the error associated with a given clustering threshold, which we called the "summed error." We calculated it by summing over-splitting and over-merging rates, assuming both have the same cost for biodiversity studies. However, it is possible to assign a differential weight to over-splitting and over-merging. For instance, if the aim is to reach conservative estimates of alpha diversity (i.e., avoid over-splitting), more weight can be assigned to over-splitting rate. Conversely, if the aim is to tease apart closely related species, which differ in their sensitivity to environmental stressors or in threat levels, one may prefer to avoid over-merging, particularly when extensive reference databases are available (Lopes et al., 2021; Roy et al., 2019).

For most of the markers we examined, the summed error approach provided relatively clear results and identified a range of threshold values that minimized the summed error. For instance, for Euka02, the summed error was relatively low at thresholds between 0.96 and 0.99 (Figure 4), indicating a good trade-off between overmerging and over-splitting. Interestingly, this range of values was also highlighted by the analysis of probability distributions (Table 2). Indeed, 0.96 is the threshold minimizing over-splitting for Euka02 while 0.99 is the balance (midpoint) threshold. The consistency of values obtained with very different approaches supports the robustness of our conclusions.

However, for a few markers, the threshold values minimizing summed error yielded somewhat less clear patterns. For Fung02, the summed error rate was rather constant (36%-37%) at all the thresholds between 0.91 and 0.98, while it guickly increased for higher clustering thresholds. For Coll01 and Oligo01, the summed error rate showed multiple minima, some of which were at very low clustering thresholds (Figure 4). In principle, increasing the threshold value should determine a monotone decrease of overmerging, and a monotone increase of over-splitting (Figure 1b). However, at low similarity values this was not always the case (Figure 4). This probably occurs because for these markers a large proportion of sequences have pairwise similarities of 0.80-0.85 (Figure 2), and this might affect the identification of clusters, with some sequences clustering together, for example, at 0.85 but not at 0.86 similarity values. We also note that these similarity values match those corresponding to the intersection between the within-genus and within-species similarities for these markers (Table 2). It is also possible that, at this level of sequence similarity, there is strong uncertainty between MOTUs representing different hierarchical levels of taxonomy.

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Our results provide quantitative data that can help researchers set their optimal clustering thresholds and understand the consequences of choosing low or high threshold values. If a clear minimum exists for the summed error rate, it probably represents an excellent trade-off between over-merging and over-splitting. In this sense, a threshold value ranging from 0.97 to 0.99 is probably appropriate for both Bact02 and Euka02, while Arth02 should accommodate a slightly higher range (0.98-0.99) and a threshold of 0.97 seems to be more suitable for Sper01. For Inse01 and COI-BF1/BR2, lower threshold values (0.94–0.97 and 0.93–0.96, respectively) are more judicious. All these values match with those obtained on the basis of within-species and within-genus similarities (Table 2). However, for Coll01, Oligo01 and Fung02, the summed error rate does not provide clear indications, and within-species and within-genus similarity distributions (e.g., midpoint between modes) might be more informative to set the clustering threshold (Figure 2 and Table 2).

The selection of clustering thresholds can have a strong effect in the estimates of MOTU richness (Figure 3), yet it is important to remember that it often does not have a tremendous effect on the ecological message conveyed by metabarcoding data. For instance, Clare et al. (2016) examined different clustering thresholds to analyse dietary overlap between skinks and shrews in Mauritius. Although high clustering thresholds yielded a larger number of MOTUs, ecological conclusions remained rather consistent overall. Therefore, provided that appropriate parameters are considered (e.g., alpha diversity measured using Hill's numbers with q > 0 instead of richness, beta diversity estimates), the interpretation of data can be relatively robust (Calderón-Sanou et al., 2020; Clare et al., 2016; Mächler et al., 2021; Roy et al., 2019). Nevertheless, we discourage the blind application of one single clustering threshold such as the classical 0.97, as it can have very different meaning across markers, and can inflate MOTU richness for fast-evolving markers. Instead, we advocate the ad-hoc definition of the most appropriate thresholds, depending on the research aims, the potential costs of over-splitting and over-merging, and the features of the studied markers.

AUTHOR CONTRIBUTIONS

All authors conceived the idea for the manuscript, A.B. and G.F.F. designed the study, A.B. performed the analyses, A.B. and G.F.F. generated the figures and drafted the manuscript, and all authors contributed with discussions and edits.

CONFLICT OF INTEREST

The authors declare that they have no conflicts of interest.

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OPEN RESEARCH BADGES

This article has earned an Open Data badge for making publicly available the digitally-shareable data necessary to reproduce the reported results. The data is available at [https://doi.org/10.5061/dryad.crjdfn353].

DATA AVAILABILITY STATEMENT

Raw data obtained from EMBL r140 (*ecopcr* files) and example scripts to prepare the datasets and perform the analyses are available on Dryad: https://doi.org/10.5061/dryad.crjdfn353.

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SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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