Efficient k-mer based curation of raw sequence data: application in Drosophila suzukii

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Abstract

Several studies have highlighted the presence of contaminated entries in public sequence repositories, calling for special attention to the associated metadata. Here, we propose and evaluate a fast and efficient k-mer-based approach to assess the degree of mislabeling or contamination. We applied it to high-throughput whole-genome raw sequence data for 236 Ind-Seq and 22 Pool-Seq samples of the invasive species Drosophila suzukii. We first used Clark software to build a dictionary of species-discriminating k-mers from the curated assemblies of 29 target drosophilid species (including D. melanogaster, D. simulans, D. subpulchrella, or D. biarmipes) and 12 common drosophila pathogens and commensals (including Wolbachia). Counting the number of k-mers composing each query sample sequence that matched a discriminating k-mer from the dictionary provided a simple criterion for assignment to target species and evaluation of the entire sample. Analyses of a wide range of samples, representative of both target and other drosophilid species, demonstrated very good performance of the proposed approach, both in terms of run time and accuracy of sequence assignment. Of the 236 D. suzukii individuals, five were reassigned to D. simulans and eleven to D. subpulchrella. Another four showed moderate to substantial microbial contamination. Similarly, among the 22 Pool-Seq samples analyzed, two from the native range were found to be contaminated with 1 and 7 D. subpulchrella individuals, respectively (out of 50), and one from Europe was found to be contaminated with 5 to 6 D. immigrans individuals (out of 100). Overall, the present analysis allowed the definition of a large curated dataset consisting of > 60 population samples representative of the worldwide genetic diversity, which may be valuable for further population genetics studies on D. suzukii. More generally, while we advocate careful sample identification and verification prior to sequencing, the proposed framework is simple and computationally efficient enough to be included as a routine post-hoc quality check prior to any data analysis and prior to data submission to public repositories.
1. Introduction

With the democratization of sequencing technologies, the availability of genomic sequence in public repositories is increasing at an unprecedented rate. This is enabling the construction of large and highly informative combined datasets for an increasing number of model and non-model species, which in turn is refining the power and resolution of population genomics inference (e.g. Kapun et al., 2021). However, this increased availability of data comes at the cost of increased heterogeneity in the resulting combined dataset. For example, data sets may combine different sequencing library preparation protocols or technologies that are rapidly evolving with variable sequence quality or coverage. Similarly, for a given species, publicly available data may refer to original studies based on different sampling strategies consisting of either sequencing individuals (aka Ind-Seq) or pools of individuals (aka Pool-Seq) representative of some populations, the latter approach being quite popular due to its cost-effectiveness (Schlötterer et al., 2014). Nevertheless, such technical characteristics can be taken into account in downstream analyses if an appropriate statistical framework is used.

More problematically, several recent studies have highlighted the high level of contamination in public repositories, which requires special attention when relying on the associated metadata description files (Cornet and Baurain, 2022; Francois et al., 2020; Steinegger and Salzberg, 2020). For example, working with wild-caught samples of species that are difficult to distinguish from other closely related species sharing the same habitat may lead to taxonomic errors or biological contamination of the sample. Such potential problems have already been reported in population genetic studies of Drosophila melanogaster, where sample contamination with D. simulans individuals was not uncommon (Kapun et al., 2021; Machado et al., 2021). In addition to biological sources, contamination may be of experimental (e.g., sample contamination or mislabeling) and/or computational (e.g., during data processing) origin (Cornet and Baurain, 2022). It should also be noted that these contamination problems are obviously not specific to publicly available data and may be even more pronounced in newly generated data that have not yet been analyzed.

In recent years, several software packages have been developed to assess the level of contamination in genomic datasets, which has been greatly facilitated by the active field of metagenomics. As recently reviewed by Cornet and Baurain (2022), the available approaches can be classified into either database-free or reference-based methods. Database-free methods roughly
consist of partitioning sequences based on their DNA composition (e.g. GC content or frequencies in short DNA sequences of a few nt), but they are not well suited for the analysis of large amounts of samples as they require a case-by-case inspection of the results (Cornet and Baurain, 2022). Reference-based methods consist of aligning sequences to a set of tagged sequences representative of all or part (e.g., genes) of the genomes of candidate species. In practice, this may allow either negative and/or positive filtering (i.e., removal of contaminating sequences or identification of sequences from some species of interest) of the sequencing data (Cornet and Baurain, 2022). To accomplish this task, approaches based on the exact matching of k-mers (i.e., k nt long DNA words) constituting the query sequences to a dictionary of labeled k-mers (built from target species genomes) have proven highly efficient and are now very popular for sequence taxonomic classification in the metagenomic field (Ounit and Lonardi, 2016; Ounit et al., 2015; Wood et al., 2019; Wood and Salzberg, 2014).

Taking advantage of the high quality assemblies available for several dozen drosophilid genomes (Kim et al., 2021), the aim of this study was to rely on a k-mer-based approach to assess the level of contamination in public sequence data for the spotted wing Drosophila D. suzukii. D. suzukii originates from Asia and has recently invaded the entire European and American continents to become a major invasive insect pest causing dramatic losses in fruit production (Asplen et al., 2015; Cini et al., 2012). This species has thus become of great scientific interest, particularly to population geneticists, and several recent studies have provided informative samples for characterizing the structuring of its genetic diversity at global and whole-genome scales. Here, we focused on two recently published and publicly available Pool-Seq and Ind-Seq datasets, consisting of whole-genome sequences (WGS) for i) 22 pools of individual DNA (with n=50 to n=100 individuals per pool) representative of populations sampled both in the Asian native range (n=6) and in the European (n=8) and American invaded ranges (n=8) (Olazcuaga et al., 2020); and ii) 236 individuals collected mainly in North America but also at several sites in Europe, Brazil and Asia (Lewald et al., 2021). A combined analysis of these two datasets using standard descriptive approaches revealed anomalous behavior of some samples (not shown), thus motivating a systematic screening of all samples for putative contamination or (taxonomic) misidentification problems. Indeed, as highlighted by Piper et al. (2022), rapid morphological identification of D. suzukii on wild-caught specimens can be tricky. For example, the distinctive spots observed on the wing extremities are present only in (non-juvenile) males, and this feature is shared with two of its sister species D. biarmipes and D. subpulchrella, whose distributions overlap all or part of that of D. suzukii in its native Asian range (Ometto et al., 2013; Takamori et al., 2006). In addition, both D. suzukii and D. subpulchrella females possess a large and serrated ovipositor that allows them to penetrate under the skin of ripening fruits and lay eggs (Atallah et al., 2014), making the distinction between these two species even more difficult.

To assess contamination in publicly available D. suzukii raw sequencing data, we developed and evaluated a fast and efficient approach based on k-mer-based methods implemented in the software Clark (Ounit et al., 2015). We first build dictionaries of species-discriminating k-mers from the curated assemblies of 29 target drosophilidae species and 12 common drosophilidae pathogens and commensals. WGS data for individual samples representative of both the target and other drosophilidae species were then analyzed to evaluate the performance of the proposed approaches, both in terms of run time and accuracy of sequence assignment. Finally, we analyzed publicly available WGS data for the aforementioned 236 Ind-Seq (Lewald et al., 2021) and 22 Pool-Seq (Olazcuaga et al., 2020) samples of the invasive species Drosophilia suzukii, allowing us to identify unambiguously contaminated samples.

2. Material and Methods

2.1. Construction of the Clark and Clark-l target dictionaries of species-discriminating k-mers

Of the 136 reference genome assemblies available for species belonging to the genus Drosophila in the NCBI repository (https://www.ncbi.nlm.nih.gov/datasets/genomes/ accessed in February 2022), 29 were retained based on assembly quality criteria such as contiguity (evaluated with contig N50) and completeness (using BUSCO scores, Manni et al., 2021); but also and mostly
based on phylogenetic criteria (Figure 1). Our goal was to obtain a good representation of species closely related to *D. suzukii*, focusing on those belonging to the two subgenera *Sophophora* and *Drosophila* that are not unambiguously resolved (see Discussion). For subgroups or groups represented by multiple species (among those with good quality assemblies available), only one target species was selected, favoring the most cosmopolitan or temperate species (Jezovit et al., 2017), except for the species most closely related to (and likely to be confounded with) *D. suzukii* (e.g., *D. subpulchrella* and *D. biarmipes*). To further improve the representation of *D. suzukii* in the k-mer dictionary, the draft assembly of Ometto et al. (2013) was also downloaded from the ENA repository (https://www.ebi.ac.uk/ena/browser/home). Although this assembly was of lower quality than the reference (Paris et al., 2020), it was obtained from a different isofemale line and was based on short read sequences from a pool of females and males. Similarly, for *D. subpulchrella* (the sister species of *D. suzukii*), the assembly from (Kim et al., 2021) was considered in addition to the latest NCBI reference assembly (Durkin et al., 2021), because it is based on male individuals and therefore contain Y-linked contigs. The high quality *D. simulans de novo* assembly from Chang et al. (2022) was also included for similar reasons.

The resulting 32 assemblies, described in Table 1, were further screened for non-Drosophila contaminating sequences using the program Kraken2 v2.1.2 (Wood et al., 2019) by querying a database constructed from the NCBI non-redundant nucleotides (nt) released in February 2020. A contig or scaffold sequence was considered contaminating if it was assigned to a taxonomic identifier unrelated to any drosophilid species. Note that contigs assigned to Wolbachia endosymbionts were also flagged as contaminating, as we chose to consider Wolbachia specifically here (see below). Of the 90,071 sequences (i.e., contigs or scaffolds) from all 32 assemblies (5.96 Gb in total), 16,123 sequences (17.9%) were found to be contaminating. As detailed in Table S1, these contaminating sequences were mostly short, ranging from 110 bp to 1,478,327 bp.

![Figure 1](https://www.ncbi.nlm.nih.gov/taxonomy) - Relationship between the 29 target drosophilid species (adapted from https://www.ncbi.nlm.nih.gov/taxonomy). Species habitat was defined according to Jezovit et al. (2017), except for *D. miranda*. *D. suzukii* is highlighted in bold.
bp (median size of 1,522 bp), totaling only 102.7 Mb (i.e. 1.72% of all sequences). It should be noted that Wolbachia-related sequences represented only 6,173,139 bp of the contaminating sequences (6.01%), with the major contributor being the \textit{D. ananassae} assembly (6,078,940 bp), which may be explained by the widespread lateral gene transfer from Wolbachia described in this species (Klasson et al., 2014). The other Wolbachia contaminating sequences belonged to the assemblies for \textit{D. suzukii} (83,189 bp) of Ometto et al. (2013) and \textit{D. willistoni} (11,010 bp). Finally, out of the 32 assemblies, only three (for \textit{D. albomicans}, \textit{D. innubila} and \textit{D. melanogaster} species) were found to be free of any contaminant, the most contaminated assemblies being those for \textit{D. immigrans} and \textit{D. willistoni} with 10.6% and 8.1% of their total length contaminated, respectively (Table 1). As expected, the completeness of the assemblies (except for the draft assembly for \textit{D. suzukii} from Ometto et al., 2013 mentioned above) remained quite good after filtering out contaminating sequences, with more than 98% (resp. 95%) of the 3,285 BUSCO genes of the \textit{diptera_odb10} dataset (Manni et al., 2021) identified in 26 (31) of the assemblies (Table 1). Finally, in addition to the drosophilid species and following Kapun et al. (2021), 13 genome assemblies representing twelve different common drosophilid commensals and pathogens were included in the construction of the \textit{k-mer} dictionaries (Table 1). Note that the two reference assemblies for the Wolbachia endosymbiont of \textit{D. melanogaster} and \textit{D. simulans} were used to represent Wolbachia.

From the 45 reference assemblies representing the 29 drosophilid, commensal, and pathogen species, two different dictionaries of species-discriminating \textit{k-mer}s (i.e., \textit{k-mer}s that occur exclusively in the genome of a species represented by one or more assemblies) were then constructed using versions 1.2.6.1 of Clark (default \textit{k}=31) and Clark-l (default \textit{k}=27), respectively (Ounit et al., 2015). Clark-l is a variant of Clark designed for use when the amount of RAM is limited, with minimal impact on assignment accuracy. Both Clark and Clark-l were run in single-threaded mode on a computer cluster grid. Building the \textit{k-mer} dictionary (on a single thread of a cluster node equipped with a processor Intel® Xeon® CPU E5-2683 v4 @2.10GHz) took 2h46min with a peak RAM usage of 128G using Clark and 55s with a peak RAM usage of 2.65G using Clark-l. The resulting database consisted of 3,714,249,662 31-mers and 50,311,519 27-mers, respectively, and required 47.8 Gb and 1.97 Gb of RAM to load when computing the query sequence classification with Clark and Clark-l, respectively.

2.2. Query short-read sequencing data

A total of 301 short read WGS data sets were downloaded from the public SRA repository (https://www.ncbi.nlm.nih.gov/sra). These include 43 samples used for the empirical evaluation of \textit{k-mer}-based assignment accuracy, derived from the sequencing of laboratory strains representative of different drosophilid species (including data on 12 of the 29 target species available for the strains used to generate the corresponding assemblies) and the Wolbachia endosymbiont of \textit{D. melanogaster} (Table S2). As detailed in Table S2, all of these data were obtained from paired-end (PE) sequencing (2×150 nt) on an Illumina HiSeq 4000 instrument, except for ten samples sequenced on an Illumina i) GAIX in PE125 mode (n=1); ii) NextSeq550 in PE150 mode (n=4); iii) HiSeq 2000 in PE100 (n=2) and PE150 (n=1) modes; iv) MiSeq in PE300 mode (n=1); or v) HiSeq Ten X in PE150 (n=1). The second type of data corresponded to WGS data for 236 \textit{D. suzukii} individuals (Ind-Seq data) representative of 40 population samples (4-10 ind. per sample, mean=5.9) published by Lewald et al. (2021). These samples were mainly collected in the continental USA (n=31). The other regions represented are Brazil (n=1); Europe (n=2; Ireland and Italy) for two of them; China (n=2); South Korea (n=2), but also Japan (n=1) and Hawai’i (n=1), via two laboratory strains. These were all sequenced on an Illumina HiSeq4000 in PE150 (n=201) or PE100 (n=35) mode (Table S3). The last type of data corresponded to WGS data from 22 pools of \textit{D. suzukii} individuals (Pool-Seq data) representing 22 worldwide populations representative of the Asian native range (n=6) and the European (n=8) and American (n=8) invaded ranges, published by Olazcuaga et al. (2020). These were all sequenced on an Illumina HiSeq2500 in PE125 mode (Table S3).

Raw PE reads were filtered with \texttt{fastp} 0.23.1 (Chen et al., 2018) with the default options to remove contaminating adaptor sequences and trimmed for poor quality bases (i.e. with a phred
Table 1 – Description of the reference genome assemblies for the 29 drosophilid species (n=32 assemblies) and 12 common commensals and pathogens (n=13 assemblies) used to build the target \(k\)-mer dictionaries. All genome assemblies were downloaded from the NCBI repository (https://www.ncbi.nlm.nih.gov), except for the two additional assemblies for \(D\). simulans and \(D\). suzukii, which were downloaded from the Dryad (https://datadryad.org) and ENA (https://www.ebi.ac.uk/ena) repositories, respectively (with accession ID in italics in the third column). The size and N50 of all assemblies are given in the fourth and fifth columns. For drosophilid species, these correspond to the assemblies after filtering out the identified contaminant contigs (or scaffolds), the percentage of the original assembly retained is given in parentheses in the fourth column. Similarly, the BUSCO scores in parentheses correspond to the percentage of complete genes identified among the 3,285 genes of the diptera_odb10 dataset (Manni et al., 2021).

<table>
<thead>
<tr>
<th>ID</th>
<th>Species</th>
<th>Reference</th>
<th>Size in Mb (% init.)</th>
<th>N50 in Mb (BUSCO)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dalbo</td>
<td><em>Drosophila albomicans</em></td>
<td>GCA_009650485.1</td>
<td>165.85 (100.0)</td>
<td>33.43 (96.6)</td>
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<td>Danan</td>
<td><em>Drosophila ananassae</em></td>
<td>GCA_017639315.1</td>
<td>207.74 (97.16)</td>
<td>26.43 (99.1)</td>
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<td>Dath</td>
<td><em>Drosophila athabasca</em></td>
<td>GCA_008121215.1</td>
<td>191.06 (99.17)</td>
<td>52.10 (98.3)</td>
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<td>Dbiar</td>
<td><em>Drosophila biarmipes</em></td>
<td>GCA_018148935.1</td>
<td>183.51 (99.03)</td>
<td>23.38 (98.9)</td>
</tr>
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<td>Dbipe</td>
<td><em>Drosophila bipectinata</em></td>
<td>GCA_018153845.1</td>
<td>189.91 (98.71)</td>
<td>15.79 (99.1)</td>
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<td>Deleg</td>
<td><em>Drosophila elegans</em></td>
<td>GCA_018152505.1</td>
<td>177.57 (99.51)</td>
<td>21.93 (99.1)</td>
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<td>Derec</td>
<td><em>Drosophila erecta</em></td>
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<td>146.49 (99.97)</td>
<td>22.15 (99.2)</td>
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<td>Deugr</td>
<td><em>Drosophila eugracilis</em></td>
<td>GCA_018153835.1</td>
<td>158.76 (96.33)</td>
<td>2.299 (98.5)</td>
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<td>Dficu</td>
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<td>158.79 (94.61)</td>
<td>14.22 (98.7)</td>
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<td>5.150 (98.9)</td>
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<td><em>Drosophila immigrans</em></td>
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<td>163.77 (89.36)</td>
<td>11.45 (98.9)</td>
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<td>Dinnu</td>
<td><em>Drosophila innubila</em></td>
<td>GCA_004354385.1</td>
<td>166.28 (100.0)</td>
<td>29.57 (98.8)</td>
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<td>Dkikk</td>
<td><em>Drosophila kikkawai</em></td>
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<td>131.51 (99.89)</td>
<td>23.40 (99.0)</td>
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<td>Dmela</td>
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<td>25.29 (98.6)</td>
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<td>3.93 (98.4)</td>
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<td>24.81 (98.8)</td>
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<td>21.50 (99.0)</td>
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<td>Dsubp</td>
<td><em>Drosophila subobsccura</em></td>
<td>GCA_003285905.1</td>
<td>151.30 (98.41)</td>
<td>11.59 (98.7)</td>
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<td>11.59 (98.9)</td>
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<td>Dsyrp</td>
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<td>GCA_015803253.1</td>
<td>255.10 (98.87)</td>
<td>1.467 (96.8)</td>
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<td>2.610 (97.4)</td>
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<td>24.18 (98.7)</td>
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<td>162.96 (100.0)</td>
<td>24.81 (98.8)</td>
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<td>Dwill</td>
<td><em>Drosophila willistoni</em></td>
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<td>24.72 (98.7)</td>
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<td>Apomo</td>
<td><em>Acetobacter pomorum</em></td>
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<td>Cinte</td>
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<td>Efaec</td>
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<td>Palc</td>
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<td>NC_0002978.6</td>
<td>1.268 1.268</td>
<td>1.268 (98.7)</td>
</tr>
</tbody>
</table>

For drosophilid species, these correspond to the assemblies after filtering out the identified contaminant contigs (or scaffolds), the percentage of the original assembly retained is given in parentheses in the fourth column. Similarly, the BUSCO scores in parentheses correspond to the percentage of complete genes identified among the 3,285 genes of the diptera_odb10 dataset (Manni et al., 2021).
quality score <15). In addition, the --merge and --include_unmerged options were used to merge the detected overlapping PE reads into a single sequence. Finally, the --stdout option was enabled to generate an interleaved fastq output, which was converted to fasta format (losing quality and pairing information) with a simple awk one-liner for assignment analysis. As shown in Figure S1 and detailed in Tables S2 and S3, the quantity (and quality) of sequencing data was highly variable between samples, with the percentage of non-overlapping sequences ranging from 5.79 to 94.4 (median 35.0) as a consequence of different insert sizes; and the estimated percentage of duplicate reads ranging from 0.69 to 24.8 (median 4.44) (Figure S1B). Note that the sequencing data were not de-duplicated here, although this may be possible using the latest version of fastp (Chen et al., 2018).

2.3. Assignment of query sequences and contamination estimation

For each sample, the sequences contained in the filtered fasta files were matched to the target dictionaries of species-discriminating k–mers using Clark and Clark-l (Ounit et al., 2015). Briefly, analyzing a sequence consists of first decomposing it into its constituent k–mers (i.e., a L nt long sequence can be decomposed into \( L - k + 1 \) k–mers of length k nt) of length k = 31 and \( k = 27 \) for Clark and Clark-l, respectively. Each k–mer is then searched in the corresponding target dictionary and, if found, assigned to the underlying target species. Counting the number of k–mers assigned to the different species then provides a simple decision criterion for sequence classification. More specifically, for a given sequence, let \( t_1 \) and \( t_2 \) be the target species with the highest and second highest counts \( (k_q(t_1) \text{ and } k_q(t_2) \leq k_q(t_1)) \) of matching k–mers. If no species-discriminating k–mer was found in the sequence (i.e., \( k_q(t_1) = 0 \) for all target species \( t \)), the sequence is unassigned. If \( k_q(t_1) > 0 \), the sequence is assigned to species \( t_1 \) with a ‘confidence score’ defined as \( c_q(t_1) = \frac{k_q(t_1)}{k_q(t_1) + k_q(t_2)} \), noting that \( c_q(t_1) = 1 \) if all the matching k–mers are assigned to \( t_1 \) (i.e., \( k_q(t) = 0 \) for all \( t \neq t_1 \)). At the sample level, the origin and level of contamination can then be further assessed by counting the number of sequences assigned to the different target species. In practice, Clark was run with option \(-s 2 \) to load only half of the species-discriminating k–mers in the target dictionary, following the manual recommendation indicating that this value ‘represents a good trade-off between speed, accuracy and RAM usage’. Both Clark and Clark-l were run with the options \(-n 1 \) (i.e., on a single thread) and \(-m 0 \) (to compute the confidence score). The resulting csv files were parsed with a custom awk script to count for each sample i) the total number of sequences with no matching k–mer; ii) the total number of sequences with at least one matching k–mer; and iii) the proportion of sequences assigned to each target species. Four different criteria were considered for assigning sequences to their inferred species \( t \), taking into account both the minimum number \( n_{k\text{min}}(t) \) of matching k–mers and the confidence score \( c_q(t) \): i) \( n_{k\text{min}}(t) \geq 1 \) and \( c_q(t) > 0.9 \); ii) \( n_{k\text{min}}(t) \geq 1 \) and \( c_q(t) > 0.95 \); iii) \( n_{k\text{min}}(t) \geq 5 \) and \( c_q(t) > 0.9 \); and the most stringent iv) \( n_{k\text{min}}(t) \geq 5 \) and \( c_q(t) > 0.95 \). All subsequent analyses were performed using the R software (R Core Team, 2017).

3. Results

3.1. Clark and Clark-l run times

Publicly available short-read WGS data for 301 different samples derived from i) laboratory strains representing different drosophilid species (n=43); ii) 236 (putative) D. suzukii individuals representing 40 different populations; and iii) 22 pools of D. suzukii individuals representing 22 different populations were assigned to two different species-discriminating k–mer dictionaries built from the curated assemblies available for 29 drosophilid species (Figure 1) and 12 common drosophila commensals and pathogens (Table 1), using the k–mer-based approaches implemented in Clark and Clark-l (Ounit et al., 2015). Although this step is not required for assignment, the raw PE reads were filtered to limit the potential impact of varying sequence quality on the assessment of assignment efficiency and accuracy, particularly with respect to the observed proportion of unassigned sequences per query sample. After filtering, the total number of sequences per sample ranged from \( 1.61 \times 10^6 \) to \( 367 \times 10^6 \) (median of \( 18.5 \times 10^6 \)) for a total
Tables S2 and S3 show the total Clark and Clark-l run times $t_r$ for each sample, together with the time $t_l$ required to load the corresponding k–mer target dictionary and the time $t_a$ required to assign all sequences ($t_r = t_l + t_a$). As summarized in the Table 2, $t_l$ was a few seconds for Clark-l and a few minutes for Clark, the Clark-l target dictionary containing about 75 times less k–mer than Clark’s (see M&M). In addition, Clark-l required much less RAM than Clark (1.97 Gb vs 47.8 Gb), allowing it to run on a standard laptop. Note that Clark and Clark-l were run sequentially on each sample on a computer grid, but the samples were analyzed in parallel. Therefore, the run times between samples may be somewhat dependent on the characteristics of the underlying node, which explains the observed variation in dictionary loading times.

Given the size of the data sets, most of the analysis time was spent on sequence assignment which was almost linearly related to the number of sequences (Figure S2) as sequence length was similar across samples (Figure S1C). On average, the analysis of 1 million sequences (i.e., ∼0.56X of the D. suzukii genome with 150 nt reads) took 0.619 and 1.05 minutes with Clark-l and Clark, respectively (Table 1), making both approaches highly computationally efficient.

### 3.2. Proportion of assigned sequences

The percentage of sequences with no matching k–mer (i.e., not assignable) was similar between Clark (ranging from 2.29% to 85.5% with a median value of 20.1%) and Clark-l (ranging from 4.07% to 86.1% with a median value of 15.7%) (Figures 2A and 2B). Surprisingly, this percentage tended to be slightly lower for the D. suzukii sample (Ind-Seq or Pool-Seq) when analyzed with Clark-l, which may be related to the smaller k–mer size (k=27 for Clark-l and k=31 for Clark) leading to lower specificity. However, the proportion of sequences with no matching k–mer remained higher for Clark-l analyses for samples representative of the other species either represented or not represented in the target dictionaries (Figure 2B). As expected, and regardless of the program used, the highest percentages were observed for samples belonging to species not represented in the target dictionaries (up to 85.5% and 86.1% of sequences with no matching k–mer for the D. repleta sample analyzed with Clark and Clark-l, respectively), although the distribution was very wide and almost bimodal due to some samples being represented by closely related target species (see below). The sample representing target species had the lowest number of sequences with no matching k–mer, most of them (including D. suzukii) corresponding to short-read sequence data obtained from the same strains used to generate the reference assembly, with the notable exception of D. melanogaster, D. simulans, and the Wolbachia sample (see below), which were also outliers in the distributions of Figure 2B (see Table S4). Their values was actually similar to wild-caught D. suzukii samples (see below). The D. simulans sample was obtained from Madagascar individuals (Palmieri et al., 2014) thus distantly related to the two reference assembly strains, which may explain the observed pattern (see Discussion). Likewise, the analyzed D. melanogaster sample corresponded to a pool of 162 isogenic strains from the DGRP panel and may thus display higher genetic diversity (Zhu et al., 2012).

Consistent with a lower specificity of Clark-l (suggested by the unexpectedly slightly lower proportion of sequences with no matching k–mer in D. suzukii individuals), the percentages of
Figure 2 – Sequence assignment rate for the 301 samples analyzed with Clark and Clark-L. A) Percentage of sequences with no matching k-mer in the target dictionaries. Samples are colored according to their origin, i.e. i) dark blue if from species represented in the target dictionary (‘Target sp.’); ii) light blue if from drosophilid species not represented in the target dictionary (‘Other sp.’); iii) green for D. suzukii individuals from Lewald et al. (2021) (‘Ind-Seq’); and iv) red for pools of D. suzukii individuals from Olazcuaga et al. (2020) (‘Pool-Seq’). B) Violin plots showing the distribution of the percentage of sequences with no matching k-mer in the corresponding target dictionary with Clark (left panel) and Clark-L (right panel) analyses. For each analysis, four distributions are shown for the different sample origins (same color code as in A). C) Distribution of the percentage of assigned sequences (among those with at least one species-discriminating k-mer from the target dictionary) for four filtering criteria on i) the number nk of matching k-mers (nk ≥ 1 or nk ≥ 5); and ii) the assignment confidence score c as defined in the main text (c > 0.9 or c > 0.95).

assigned sequences among assignable sequences (i.e., containing at least one k-mer matching the dictionary of target species discriminating k-mers) were much lower with Clark-L than with Clark (Figure 2C). The percentages of assigned sequences always decrease with the stringency of the filtering criteria on the number nk of matching k-mers (nk ≥ 1 or nk ≥ 5) and the assignment confidence score c (as defined above in the M&M section), with the threshold on nk having the strongest effect. At the most stringent criterion (nk ≥ 5 and c > 0.95), which was chosen for
the remainder of this study, 84.8% and 26.4% of sequences with at least one matching \( k \)-mer were assigned to Clark and Clark-I, respectively, on average (see Tables S4 and S5 for details).

### 3.3. Assignment accuracy for samples representative of target and other species

![Figure 3](image-url)

**Figure 3** – Bubble plots summarizing assignment results obtained with Clark using the most stringent sequence assignment criterion (i.e., \( nk \geq 5 \) and \( c > 0.95 \), see the main text) for 13 samples (labeled in dark blue on the top of the y-axis) belonging to target species represented in the target \( k \)-mer dictionary and 30 other unrepresented drosophilid species.

To empirically evaluate the extent to which the proportion of assigned sequences from a sample provides an accurate proxy for species assignment, we focused on the results obtained for the 13 short-read datasets derived from strains representative of one of the target species (including Wolbachia), but also on 30 additional samples representative of unrepresented drosophilid species, considering our most stringent filtering threshold for sequence assignment (Figures 3 and S3 for Clark and Clark-I results, respectively). The results obtained were highly consistent for all 13 samples representing the target reference species. More precisely, with Clark, the percentage of sequences assigned to their species of origin was >99% or close to 99% (with 98.9% for both D. suzukii and D. subpulchrella) for 9 of these samples. The remaining four were those belonging to i) D. biarmipes (94.0%), due to yeast contamination with 5.76% of the sequences assigned to S. cerevisiae; ii) D. melanogaster (94.1%) with 3.78% of the analyzed sequences assigned to Wolbachia and 1.11% to D. simulans; iii) D. simulans (93.6%) with 3.14% of the analyzed sequences assigned to Wolbachia and 2.54% to D. melanogaster; and iv) Wolbachia with only 5.58% actually assigned to Wolbachia and 94.0% to D. melanogaster (Table S3). Note that
this latter Wolbachia sample was actually obtained from sequencing a *D. melanogaster* strain, and the observed level of contamination was in close agreement with the 5% of reads mapping to the Wolbachia wMel genome by the original authors (Newton and Sheehan, 2015). Similar results were obtained when scanning these 13 samples with Clark-l (Figure S3 and Table S5), with some notable differences. Indeed, the percentage of sequences assigned to their species of origin was also above 99% (including the *D. subpulchrella* one) or close to it (with 98.0% for *D. yakuba*) for 8 of the 9 samples that showed similarly high assignment rates with Clark. However, it was substantially lower for the *D. suzukii* sample (92.1%), with 7.22% of its sequences assigned to the *D. subpulchrella* sister species. Similarly, only 86.2% of the *D. melanogaster* sample sequences were assigned to *D. melanogaster*, with 6.86%, 2.62%, 1.65%, and 1.40% assigned to *D. suzukii*, *D. simulans*, *D. virilis*, and Wolbachia, respectively. Conversely, the percentage of correctly assigned sequences was higher with Clark-l than with Clark for the *D. biarmipes* (96.0%); *D. simulans* (98.1%) and Wolbachia (40.0% with 55.8% assigned to *D. melanogaster*) samples, the latter apparently being overestimated.

Of the 30 samples from non-target species, 16 had more than 96% of their reads assigned to a single target species by Clark (Figure 3). As expected, the corresponding species was generally the most closely related (Kim et al., 2021). More precisely, samples from i) *D. paulistorum* and *D. insularis* (*D. willistoni* subgroup) and *D. suinea* and *D. nebulosa* (*bocainensis* subgroup from the willistoni group) had 99.7%, 99.7%, 98.1%, and 97.9% of their sequences assigned to *D. willistoni*, respectively; ii) *D. parabipectinata*, *D. malerkotliana pallens*, *D. malerkotliana malerkotliana*, *D. pseudoananassae*, and *D. pseudoananassae nigrens*, all of which belong to the ananassae subgroup, had 99.2%, 99.1%, 99.0%, 96.5%, and 96.0% of their sequences assigned to *D. bipectinata* (ananassae subgroup), respectively; iii) *D. ambiguа* and *D. tristis* (*obscura* subgroup) had 98.7% and 97.3% of their sequences assigned to *D. obscura*, respectively; iv) *D. americana* and *D. littoralis* (*virilis* group) had 99.2% and 98.6% of their sequences assigned to *D. virilis*, respectively; and finally v) *D. carrolli*, *D. fuyamai*, and *D. kurseongensis* (*rhopaloa* subgroup) had 98.2%, 98.0%, and 97.7% of their sequences assigned to *D. rhopaloa*, respectively. As shown in Figure S4A, these 16 samples also had percentages of sequences with no matching k-mer in the range of those observed for samples from target species (Figure 2), i.e. <40% except for *D. sucinea* and *D. nebulosa*. For the other samples from the most distantly related species, both the highest observed assignment rate (to a target species) and the percentage of sequences with no matching k-mer clearly suggested that the target repository was not representative. At the extreme, the most represented target species capture less than 30% of the assigned sequences for the samples from *D. repleta*, *D. pruinosa*, *D. ohnishii*, and *D. bocqueti* (Figures 3 and S4A). Such species may therefore be considered unassignable with the current version of the k-mer dictionary. Despite a higher proportion of no matching species-discriminating k-mer, very similar results were obtained with Clark-l (Figure S3 and S4B).

### 3.4. Scanning 236 Ind-Seq and 22 Pool-Seq *D. suzukii* WGS data

As summarized in Figure 4 (see Table S4 for details), sequences from the 236 Ind-Seq (Lewald et al., 2021) and 22 Pool-Seq (Olaezuaga et al., 2020) *D. suzukii* were generally assigned to *D. suzukii* by Clark. More precisely, 215 of the 236 Ind-Seq and 17 of the 22 Pool-Seq showed >95% of their (assigned) sequences assigned to *D. suzukii*, with a median proportion of 97.5% over the 258 samples. It should be noted that these 215 individuals and 17 pools, which can be unambiguously considered as fully *D. suzukii*, all had a non-negligible fraction of their sequences assigned to *D. subpulchrella* with a median of 1.94% (ranging from 1.50% to 3.12%) and 2.20% (ranging from 1.96% to 2.64%), respectively. These proportions were higher than the one observed for the *D. suzukii* reference sample (0.433%) and may be related to the incomplete representation of genetic diversity within *D. suzukii* by the k-mer dictionary (see Discussion). Conversely, the results allowed 16 clearly mislabeled *D. suzukii* individuals to be identified as *D. simulans* (n=5) or *D. subpulchrella* (n=11). These consist of i) the 5 individuals (with US-Ca2 ID prefix, Table S2) sampled simultaneously in Watsonville (California, USA) with 92.1% to 96.9% of their sequences assigned to *D. simulans* (96.9% to 98.4% if Wolbachia is also included); ii) the 5 individuals (with Ko-Nam ID prefix, Table S2) sampled in Namwon (South Korea) with 97.9% to
Figure 4 – Barplots summarizing assignment results obtained with Clark using the most stringent sequence assignment criterion (i.e., \(nk \geq 5\) and \(c > 0.95\), see the main text) for the *D. suzukii* Ind-Seq (n=236) and Pool-Seq (n=22) samples. For each sample, the proportions of sequences assigned to the 7 target species that contribute at least 5% of the sequences of one of any of the 258 samples are shown using the color code indicated in the top-left legend. The proportions of sequences assigned to the 34 other target species are shown in gray.

98.7% of their sequences assigned to *D. subpulchrella*; iii) one of the 10 individuals (with Ko-San ID prefix, Table S2) sampled in Sancheong (South Korea) with 96.4% of its sequences assigned to *D. subpulchrella* (the other 9 individuals showing only 1.71% to 2.09% of their sequences assigned to *D. subpulchrella*; and iv) four of the five individuals (with CN-Kun ID prefix, Table S2) sampled in Kunming (Yunnan, China) with 97.3% to 97.6% of their sequences assigned to
D. subpulchrella. The last CN-Kun individual had a unique pattern with 88.1% of its sequences assigned to D. subpulchrella and 9.58% assigned to D. suzukii, which may be consistent with a recent hybrid origin (see Discussion). For the 10 individuals that can be unambiguously considered as fully D. subpulchrella (i.e. with >95% of their sequences assigned to D. subpulchrella), an assignment pattern opposite to that of the D. suzukii individuals was observed, as all of them had a non-negligible fraction of their sequences assigned to D. suzukii with a median value of 1.61% (ranging from 1.14% to 2.78%).

Among the 22 Pool-Seq samples, two to three pools were found to be likely contaminated with non-D. suzukii individuals. These are i) the DE-Jen pool of 100 individuals sampled in Jena (Germany), which contains 5.79% of sequences assigned to D. immigrans; ii) the CN-Nin pool of 50 individuals sampled in Ningbo (Zhejiang, China), which contains 15.0% of sequences assigned to D. subpulchrella (and 83.8% to D. suzukii); and iii) the JP-Tok pool of 50 individuals sampled in Tokyo (Japan), with 4.47% of sequences assigned to D. subpulchrella (and 94.9% to D. suzukii).

Assuming an equal contribution of pool individuals to the Pool-Seq sequences, the DE-Jen pool may actually contain up to 6 D. immigrans individuals (and 94 D. suzukii individuals). Furthermore, to estimate the number of D. subpulchrella individuals in contaminated pools while accounting for D. suzukii and D. subpulchrella cross-assignment of sequences, let \( \alpha = \frac{\text{Pool}_{\text{sub}}}{\text{Pool}_{\text{sub}} + \text{Pool}_{\text{suz}}} \) be the relative proportion of sequences assigned to D. subpulchrella. Based on the median proportions observed in the Ind-Seq samples, the following rough estimates were obtained: \( \hat{\alpha}_{\text{suz}} = 0.0194 \), \( \hat{\alpha}_{\text{sub}} = 0.0151 \), and \( \hat{\alpha}_{\text{sub}} = 0.985 \) for D. suzukii individuals and \( \hat{\alpha}_{\text{sub}} = 0.985 \) for D. subpulchrella individuals. The number of D. subpulchrella individuals \( n_{\text{sub}} \) in a contaminated pool of \( n \) individuals can then simply be derived from these estimates using their observed relative proportion \( \alpha_{\text{o}} \) as \( n_{\text{sub}} = \frac{n \cdot \alpha_{\text{o}} - \hat{\alpha}_{\text{suz}}}{\hat{\alpha}_{\text{sub}} - \hat{\alpha}_{\text{suz}}} \).

This leads to an estimated number of D. subpulchrella individuals of \( n_{\text{sub}} = 6.85 \) and \( n_{\text{sub}} = 1.32 \), i.e. probably 7 and 1 D. subpulchrella individuals within the CN-Nin and JP-Tok pools, respectively.

Overall, very low levels of Wolbachia contamination were detected within the Ind-Seq and Pool-Seq samples, with median proportions of assigned sequences of 3.80 × 10^-14% and 0.145%, respectively. However, 14 samples (Ind-Seq only) had more than 1% of their sequences assigned to Wolbachia. They consisted of i) the five US-Ca2 individuals mentioned above, which are actually D. simulans, with proportions ranging from 1.08% to 6.17%; ii) the four individuals with the CN-Dan ID prefix (Table S2), sampled in Dandeong (China), with proportions ranging from 1.07% to 8.82%; iii) three of the five individuals with the US-Ca1 ID prefix (Table S2) sampled in Davis (California, USA) with proportions ranging from 1.29% to 1.55%; iv) one of the five individuals with the BR-Pel ID prefix sampled in Pelotas (Brazil) with a proportion of 2.07%; and v) one of the five individuals with the IT-Tre ID prefix sampled in Trento (Italy) with a proportion of 1.92%.

Finally, a few Ind-Seq and Pool-Seq samples showed non-negligible to substantial contamination with five of the 11 other microbial species represented in the k-mer target dictionary. For example, more than 1% of the sequences were assigned to the L. plantarum bacterial gut symbiont for five samples corresponding to i) the four CN-Dan individuals (see above), with proportions ranging from 1.55% to 22.6%; and ii) the US-Sok pool of 50 individuals sampled in Dayton (Oregon, USA) with a proportion of 1.87%. Similarly, > 1% of the sequences were assigned to S. cerevisiae yeast for five samples corresponding to i) one of the four CN-Dan individuals with a proportion of 1.12%; ii) three individuals (with ID prefixes US-Nc2, US-Nc3, and US-Nc4, Table S2) sampled in different locations in North Carolina (USA) with proportions ranging from 1.33% to 9.58%; and iii) the US-Sdi pool of 50 individuals sampled in San-Diego (California, USA) with a proportion of 1.02%. At the margin, three other microbial species were also found to be represented by more than 1% of the sequences in at least one sample. These are i) the A. pomorum gut bacteria in two Chinese (CN-Dan) individuals (with proportions of 1.04% and 1.46%) and in the CN-Shi pools of 50 individuals sampled in Shiping (China) with 1.56%; ii) the L. brevis intestinal bacteria also found in two Chinese (CN-Dan) individuals with proportions of 1.12% and 4.19%; and iii) the E. faecalis pathogens in an Irish individual with proportion of 1.65%.

As expected from the assignment of D. suzukii and D. subpulchrella reference samples, Figure S5 (see Table S5 for details) suggested a worse performance of Clark-l. The proportions of D.
suzukii sequences appeared to be substantially underestimated, with a higher effect of cross-assignment with D. subpulchrella. In addition, C\textcopyright{}l\textcopyright{}ark-l did not allow to detect the presence of the microbial target species as detected by C\textcopyright{}l\textcopyright{}ark.

4. Discussion

The primary objective of this study was to propose and evaluate a computationally fast and accurate method for assessing contamination levels in publicly available WGS data for the D. suzukii species, which has been increasingly studied over the past decade. The availability of high quality genome assemblies for a wide range of drosophilid species (Kim et al., 2021) made it possible to rely on a \textit{k–mer}-based approach consisting of constructing and querying dictionaries of species-discriminating \textit{k–mers}. Such an approach has already proven to be quite valuable and benefits from the availability of optimized software, such as Kraken (Wood et al., 2019; Wood and Salzberg, 2014) or C\textcopyright{}l\textcopyright{}ark (Ounit and Lonardi, 2016; Ounit et al., 2015), which were primarily developed for metagenomics applications but have also been proposed for contaminant detection (Cornet and Baurain, 2022). As in the latter case, our primary goal here was to classify sequences at the level of predefined (target) species, and C\textcopyright{}l\textcopyright{}ark thus seemed particularly attractive due to its computationally efficient, tractable, and flexible way of both constructing and querying user-defined \textit{k–mer} dictionaries. Although Kraken may be able to further assign higher-level taxonomic labels by considering phylogenetic relationships among target species, this feature was not critical for our purpose. In fact, it may have made it more difficult in practice, since the phylogeny among Drosophilidae species is far from being fully and unambiguously resolved. In particular, Finet et al. (2021) recently provided evidence for a paraphyletic status of the subgenus Sophophora, to which most of the target species belong (Figure 1). However, as illustrated by the assignment of sequences from species closely related to one of the represented groups or subgroups (e.g., \textit{ananassae} or \textit{obscura}) but not included in the construction of the \textit{k–mer} dictionary, species-level assignment provided consistent results about their origin. Yet, assignment of samples to species belonging to groups or subgroups less well represented by the target species should be interpreted with caution, especially when the observed proportion of non-matching \textit{k–mers} is high (Figure S4). In such cases, analysis with a newly built \textit{k–mer} dictionary including more closely related species may be valuable. Indeed, our main focus was on the evaluation of D. suzukii samples. We therefore chose to deliberately overrepresent the suzukii subgroup in the \textit{k–mer} dictionary construction by including the high quality genome assemblies available for D. suzukii, D. subpulchrella, and D. biarmipes. The latter two species were in fact the most likely confounders in field-collected samples from the Asian range of D. suzukii (see Introduction). Interestingly, the inclusion of these closely related species seemed to have only a limited effect on the number of discriminating \textit{k–mers} in the resulting dictionary, with the percentage of sequences with no assigned \textit{k–mer} for their corresponding reference samples being in the range of that observed for reference samples from other target species (Figure 2).

Searching the resulting \textit{k–mer} dictionary of target species sequences with C\textcopyright{}l\textcopyright{}ark (Ounit et al., 2015) was highly efficient in terms of both run time and memory requirements. This makes analyses of common short-read sequencing data tractable on standard workstations or computer grids, and even on a standard laptop when using the lighter C\textcopyright{}l\textcopyright{}ark version (Ounit et al., 2015), although at some moderate cost in assignment accuracy. More specifically, it took only a few minutes and about 50 Gb of RAM to load the C\textcopyright{}l\textcopyright{}ark dictionary (<1 min and <2 Gb of RAM for the C\textcopyright{}l\textcopyright{}ark-l dictionary), and the mapping took about one minute per million of typical 150 nt short reads. Such assignment analyses could thus be performed routinely and may be worth including as a standard part of the quality control of sequencing data, at least for the D. suzukii sample. Note that here we have chosen to screen sequences after filtering raw PE reads with \texttt{fastp} (Chen et al., 2018), primarily to limit the potential impact of varying sequence quality across samples on the assessment of assignment accuracy (e.g., proportion of sequences assigned). Although this is not required in practice when trying to assign samples or assess their contamination levels, it seems to be a reasonable strategy when combined with other quality control procedures. Finally, for contamination assessment at the whole-sample level, \textit{k–mer}-based approaches represent an
attractive and efficient mapping-free alternative to competitive mapping methods that consist of mapping sequencing reads to hologenomes constructed from target species assemblies (e.g. Kapun et al., 2021). It also allows for easy interrogation of a wider range of target species, providing good quality genome assemblies are available. For sequence filtering purposes, however, such approaches must be used with caution because they rely on species-discriminating k-mer and thus may leave a substantial fraction of sequences unassigned. More advanced (and computationally expensive) methods may then be valuable, such as the one implemented in Clark-s (Ounit and Lonardi, 2016), which allows some mismatches in k-mer matching to improve the sensitivity of sequence assignment or even Kraken (Wood et al., 2019; Wood and Salzberg, 2014), which was used here to identify contaminating contigs in the assemblies of the target species. Indeed, this program can rely on k-mer shared by several species for sequence assignment, and not only species discriminating k-mers, since all the k-mers of the target dictionary (possibly built from very large databases such as the NCBI nt) are mapped to the nodes of a phylogenetic tree (species discriminating k-mers to terminal nodes and shared k-mers to internal nodes).

Overall, the results obtained from the analysis of WGS data for reference samples belonging to different target species and single or pools of D. suzukii individuals demonstrated the high accuracy of the k-mer-based approach. It also allowed the unambiguous identification of 16 mislabeled D. suzukii individuals among the 236 (i.e. 6.78%) from the Lewald et al. (2021) study. Five corresponded to D. simulans individuals collected at the same site in Watsonville (California, USA). It should be noted that Lewald et al. (2021) discarded these samples from their analysis because they displayed too low mapping rates like the Dandong (China) sample, which was found here to be substantially contaminated with microbial (and Wolbachia) sequences. The eleven other non-D. suzukii individuals from three different locations in Asia could all be assigned to D. subpulchrella individuals. These were also identified as D. subpulchrella by Lewald et al. (2021) and discarded from their analysis) using a phylogenetic analysis of the mitochondrial COX2 gene. Two of the 22 Pool-Seq samples of (Olazcuaga et al., 2020) collected in the Asian native area were also, and unexpectedly, found to be contaminated with D. subpulchrella individuals, namely CN-Nin with 7 D. subpulchrella individuals and to a lesser extent JP-Tok with 1 D. subpulchrella individual (both out of 50 individuals in total). More surprisingly, but confirming a gene-based analysis by D. Obbard (pers. comm.), the DE-Jen pool collected in Jena (Germany) was found to be contaminated with 5 to 6 D. immigrans individuals (out of 100). These observations may indicate that great care should be taken when analyzing sequencing data from wild-caught samples, and that more attention should probably be paid to species identification prior to sequencing. High-throughput metabarcoding and non-destructive approaches, such as those recently proposed by Piper et al. (2022), may represent valuable alternatives to sometimes difficult morphological identification by allowing rapid and efficient diagnosis of D. suzukii samples at any life stage. Such efforts may be even more critical for Pool-Seq experiments, since filtering out contaminated sequences (e.g., using competitive mapping) is far more challenging than discarding mislabeled Ind-Seq samples, especially when the sample is contaminated by individuals from very closely related species (such as D. subpulchrella for D. suzukii).

Although two different D. suzukii genome assemblies were used to build the species discriminating k-mer dictionary, all (pure) D. suzukii Ind-Seq and Pool-Seq samples showed a small but non-negligible fraction of their sequences (from 1.14% to 2.78%) assigned to D. subpulchrella by the most stringent criterion. Because i) the D. suzukii reference genome assemblies were derived from isofemale lines established from individuals sampled in the North American (Chiu et al., 2013) and European (Ometto et al., 2013) invaded areas; and ii) D. subpulchrella has not been yet described (to our knowledge) outside the Asian native range of D. suzukii; it is highly unlikely that this pattern is the result of pervasive gene flow between the two species, but rather can be explained by the close phylogenetic relationship between the two species. Indeed, some D. subpulchrella-discriminating k-mers may actually map to orthologous regions not represented in the D. suzukii reference assemblies and/or capture shared genetic variation between the two species due to incomplete lineage sorting (ILS). Including more reference assemblies (e.g., from different strains) for each target species may be considered as a valuable strategy to improve both the sensitivity (by ‘positive filtering’ of the discriminating k-mers that capture intraspecific
genetic variation) and specificity (by 'negative filtering' of the incompletely sorted \(k\)-mers). The optimal number of representative assemblies is thus likely to both depend on the relatedness of the selected target species and for each target species on their genetic diversity. Alternatively, the misassigned short read sequences found in the analyzed samples can be included in the construction of the \(k\)-mer dictionary, assuming that the considered samples are not contaminated and are 'pure' representatives of the corresponding target species. Such refined target dictionaries may even further allow providing (rough) estimates of the genome-wide level of interspecific gene flow, or at least the identification of highly admixed individuals. Hence, in the sample of identified \(D.\) \textit{subpulchrella} individuals, if about 2\% of the short-read sequences were assigned to \(D.\) \textit{suzukii} (in a similar but reversed pattern as observed for \(D.\) \textit{suzukii} individuals), one (presumably) \(D.\) \textit{subpulchrella} individual had nearly 10\% of its sequences assigned to \(D.\) \textit{suzukii}. The status of this sample may be of special interest for further study as it could represent a previously unreported case supporting some recent (i.e., only a few generations back) admixture events between \(D.\) \textit{suzukii} and \(D.\) \textit{subpulchrella}. As discussed by Lalyer et al. (2021), if no such recent events have been reported to date, several studies suggest that hybridization has occurred between these two sister species (Conner et al., 2017).

Overall, the present analysis allowed the definition of a large curated dataset consisting of > 60 population samples representative of global genetic diversity, which may be valuable for further \(D.\) \textit{suzukii} population genetics studies. Although constructed with the analysis of \(D.\) \textit{suzukii} samples in mind, the \(k\)-mer dictionary developed here may be directly relevant to the analysis of the level of contamination of samples from other target species such as \(D.\) \textit{simulans} or \(D.\) \textit{melanogaster}. Likewise, the current dictionary also allows for the rapid identification of Wolbachia-infected samples, which may be of interest for a first rapid screening of \textit{drosophilids} samples since the set of Wolbachia-discriminating \(k\)-mers was built by combining \(D.\) \textit{simulans} and \(D.\) \textit{melanogaster} Wolbachia assemblies. More generally, while we advocate careful sample identification and verification prior to sequencing, the proposed framework is straightforward and computationally efficient. It thus could be considered as a routine post-hoc quality check approach to be applied prior to any data analysis and prior to data submission to public repositories.

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Conflict of interest disclosure

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Data, script, code, and supplementary information availability

The Clark and \textit{Clark-l} \(k\)-mer databases and the (cleaned) assemblies used to build them have been made publicly available from the Data INRAE repository (Gautier, 2023b, https://doi.org/10.57745/HYTIBH). The compressed archive also contains scripts used to run Clark and \textit{Clark-l} analyses and parse the results. All sequencing data analyzed in this study are publicly available under the accession IDs reported in Tables 1, S2 and S3. Supplementary
Tables S1 to S5 and Supplementary Figures S1 to S5 are available online with the latest preprint version of the manuscript (Gautier, 2023a, https://doi.org/10.1101/2023.04.18.537389).

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