

1      **MitoChime: A Machine Learning Pipeline for**  
2      **Detecting PCR-Induced Chimeras in**  
3      **Mitochondrial Illumina Reads**

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13                         by

14                         Duranne Duran  
15                         Yvonne Lin  
16                         Daniella Pailden

17                         Adviser  
18                         Francis D. Dimzon, Ph.D.

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## Abstract

21 Next-generation sequencing (NGS) platforms have advanced research but re-  
22 main susceptible to artifacts such as PCR-induced chimeras that compromise  
23 mitochondrial genome assembly. These artificial hybrid sequences are prob-  
24 lematic for small, circular, and repetitive mitochondrial genomes, where they  
25 can generate fragmented contigs and false junctions. Existing detection tools,  
26 such as UCHIME, are optimized for amplicon-based microbial community ana-  
27 lysis and depend on reference databases or abundance assumptions unsuitable  
28 for organellar assembly. To address this gap, this study presents MitoChime,  
29 a machine learning pipeline for detecting PCR-induced chimeric reads in *Sar-*  
30 *dinella lemuru* Illumina paired-end data without relying on external reference  
31 databases.

32 Using simulated datasets containing clean and chimeric reads, a feature  
33 set was extracted, combining alignment-based metrics (e.g., supplementary  
34 alignments, soft-clipping) with sequence-derived statistics (e.g., k-mer com-  
35 position, microhomology). A comparative evaluation of supervised learning  
36 models identified tree-based ensembles CatBoost and Gradient Boosting as top  
37 performers, achieving an F1-score of 0.77 and an ROC-AUC of 0.84 on held-  
38 out test data. Feature importance analysis highlighted soft-clipping and k-mer  
39 compositional shifts as the strongest predictors of chimerism, whereas micro-  
40 homology contributed minimally. Integrating MitoChime as a pre-assembly  
41 step can aid in streamlining mitochondrial reconstruction pipelines.

42 **Keywords:** Chimera detection, Mitochondrial genome,  
Assembly, Machine learning

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<sup>136</sup> **Chapter 1**

<sup>137</sup> **Introduction**

<sup>138</sup> **1.1 Overview**

<sup>139</sup> The rapid advancement of next-generation sequencing (NGS) technologies has  
<sup>140</sup> transformed genomic research by enabling high-throughput and cost-effective  
<sup>141</sup> DNA analysis (Metzker, 2010). Among current platforms, Illumina sequencing  
<sup>142</sup> remains the most widely adopted, capable of producing millions of short reads  
<sup>143</sup> that can be assembled into reference genomes or analyzed for genetic variation  
<sup>144</sup> (Bentley et al., 2008; Glenn, 2011). Despite its high base-calling accuracy,  
<sup>145</sup> Illumina sequencing is prone to artifacts introduced during library preparation,  
<sup>146</sup> particularly polymerase chain reaction (PCR)-induced chimeras, which are ar-  
<sup>147</sup> tificial hybrid sequences that do not exist in the true genome (Judo, Wedel, &  
<sup>148</sup> Wilson, 1998).

<sup>149</sup> PCR chimeras form when incomplete extension products from one template

anneal to an unrelated DNA fragment and are extended, creating recombinant reads (Qiu et al., 2001). In mitochondrial genome assembly, such artifacts are especially problematic because the mitochondrial genome is small, circular, and often repetitive (Boore, 1999; Cameron, 2014). Even a small number of chimeric or misjoined reads can reduce assembly contiguity and introduce false junctions during organelle genome reconstruction (Dierckxsens, Mardulyn, & Smits, 2017; Hahn, Bachmann, & Chevreux, 2013; Jin et al., 2020). Existing assembly tools such as GetOrganelle and MITObim assume that input reads are largely free of such artifacts (Hahn et al., 2013; Jin et al., 2020). Consequently, undetected chimeras may produce fragmented assemblies or misidentified organellar boundaries. To ensure accurate reconstruction of mitochondrial genomes, a reliable method for detecting PCR-induced chimeras before assembly is essential.

This study focuses on mitochondrial sequencing data from the genus *Sardinella*, a group of small pelagic fishes widely distributed in Philippine waters. Among them, *Sardinella lemuru* (Bali sardinella) is one of the country's most abundant and economically important species, providing protein and livelihood to coastal communities (Labrador, Agmata, Palermo, Ravago-Gotanco, & Pante, 2021; Willette, Bognot, Mutia, & Santos, 2011). Accurate mitochondrial assemblies are critical for understanding its population genetics, stock structure, and evolutionary history. However, assembly pipelines often encounter errors or fail to complete due to undetected chimeric reads. To address this gap, this research introduces MitoChime, a machine learning pipeline designed to detect PCR-induced chimeric reads using both alignment-based and sequence-derived statistical features. The tool aims to provide bioinformatics laboratories, particularly the Philippine Genome Center Visayas (PGC Visayas), with an efficient

175 solution for improving mitochondrial genome reconstruction.

## 176 1.2 Problem Statement

177 Chimeric reads can distort assembly graphs and cause misassemblies, with par-  
178 ticularly severe effects in mitochondrial genomes (Boore, 1999; Cameron, 2014).  
179 Existing assembly pipelines such as GetOrganelle, MITObim, and NOVOPlasty  
180 assume that sequencing reads are free of such artifacts (Dierckxsens et al., 2017;  
181 Hahn et al., 2013; Jin et al., 2020). At PGC Visayas, several mitochondrial as-  
182 semblies have failed or yielded incomplete contigs despite sufficient coverage, sug-  
183 gesting that undetected chimeric reads compromise assembly reliability. Mean-  
184 while, existing chimera detection tools such as UCHIME and VSEARCH were  
185 developed primarily for amplicon-based community analysis and rely heavily on  
186 reference or taxonomic comparisons (Edgar, Haas, Clemente, Quince, & Knight,  
187 2011; Rognes, Flouri, Nichols, Quince, & Mahé, 2016). These approaches are un-  
188 suitable for single-species organellar data, where complete reference genomes are  
189 often unavailable.

## 190 1.3 Research Objectives

### 191 1.3.1 General Objective

192 This study aims to develop and evaluate a machine learning-based pipeline (Mi-  
193 toChime) that detects PCR-induced chimeric reads in *Sardinella lemuru* mito-

<sub>194</sub> chondrial sequencing data in order to improve the quality and reliability of down-  
<sub>195</sub> stream mitochondrial genome assemblies.

### <sub>196</sub> 1.3.2 Specific Objectives

<sub>197</sub> Specifically, the study aims to:

- <sub>198</sub> 1. construct simulated *Sardinella lemuru* Illumina paired-end datasets contain-  
<sub>199</sub> ing both clean and PCR-induced chimeric reads,
- <sub>200</sub> 2. extract alignment-based and sequence-based features such as k-mer compo-  
<sub>201</sub> sition, junction complexity, and split-alignment counts from both clean and  
<sub>202</sub> chimeric reads,
- <sub>203</sub> 3. train, validate, and compare supervised machine learning models for classi-  
<sub>204</sub> fying reads as clean or chimeric,
- <sub>205</sub> 4. determine feature importance and identify indicators of PCR-induced  
<sub>206</sub> chimerism,
- <sub>207</sub> 5. integrate the optimized classifier into a modular and interpretable pipeline  
<sub>208</sub> deployable on standard computing environments at PGC Visayas.

### <sub>209</sub> 1.4 Scope and Limitations of the Research

<sub>210</sub> This study focuses solely on PCR-induced chimeric reads in *Sardinella lemuru*  
<sub>211</sub> mitochondrial sequencing data, with the species choice guided by four consid-  
<sub>212</sub> erations: (1) to limit interspecific variation in mitochondrial genome size, GC

213 content, and repetitive regions so that differences in read patterns can be at-  
214 tributed more directly to PCR-induced chimerism, (2) to align the analysis with  
215 relevant *S. lemuru* sequencing projects at PGC Visayas, (3) to take advantage of  
216 the availability of *S. lemuru* mitochondrial assemblies and raw datasets in public  
217 repositories such as the National Center for Biotechnology Information (NCBI),  
218 which facilitates reference selection and benchmarking, and (4) to develop a tool  
219 that directly supports local studies on *S. lemuru* population structure and fisheries  
220 management.

221 The study emphasizes `wgsim`-based simulations and selected empirical mito-  
222 chondrial datasets from *S. lemuru*. It excludes naturally occurring chimeras, nu-  
223 clear mitochondrial pseudogenes (NUMTs), and large-scale assembly rearrange-  
224 ments in nuclear genomes. Feature extraction is restricted to low-dimensional  
225 alignment and sequence statistics, such as k-mer frequency profiles, GC con-  
226 tent, soft and hard clipping metrics, and split-alignment counts rather than high-  
227 dimensional deep learning embeddings. This design keeps model behaviour inter-  
228 pretable and ensures that the pipeline can be run on standard workstations at  
229 PGC Visayas. Testing on long-read platforms (e.g., Nanopore, PacBio) and other  
230 taxa is outside the scope of this project.

231 Other limitations in this study include the following: simulations with vary-  
232 ing error rates were not performed, so the effect of different sequencing errors on  
233 model performance remains unexplored; alternative parameter settings, including  
234 k-mer lengths and microhomology window sizes, were not systematically tested,  
235 which could affect the sensitivity of both k-mer and microhomology feature de-  
236 tection; and the machine learning models rely on supervised training with labeled  
237 examples, which may limit their ability to detect novel or unexpected chimeric

<sup>238</sup> patterns.

## <sup>239</sup> 1.5 Significance of the Research

<sup>240</sup> This research provides both methodological and practical contributions to mito-  
<sup>241</sup> chondrial genomics and bioinformatics. First, MitoChime detects PCR-induced  
<sup>242</sup> chimeric reads prior to genome assembly, with the goal of improving the con-  
<sup>243</sup> tiguity and correctness of *Sardinella lemuru* mitochondrial assemblies. Second,  
<sup>244</sup> it replaces informal manual curation with a documented workflow, improving au-  
<sup>245</sup> tomation and reproducibility. Third, the pipeline is designed to run on computing  
<sup>246</sup> infrastructures commonly available in regional laboratories, enabling routine use  
<sup>247</sup> at facilities such as PGC Visayas. Finally, more reliable mitochondrial assemblies  
<sup>248</sup> for *S. lemuru* provide a stronger basis for downstream applications in the field of  
<sup>249</sup> fisheries and genomics.

250 **Chapter 2**

251 **Review of Related Literature**

252 This chapter presents an overview of the literature relevant to the study. It  
253 discusses the biological and computational foundations underlying mitochondrial  
254 genome analysis and assembly, as well as existing tools, algorithms, and techniques  
255 related to chimera detection and genome quality assessment. The chapter aims to  
256 highlight the strengths, limitations, and research gaps in current approaches that  
257 motivate the development of the present study.

258 **2.1 The Mitochondrial Genome**

259 Mitochondrial genome (mtDNA) is a small, typically circular molecule found in  
260 most eukaryotes. It encodes essential genes involved in oxidative phosphorylation  
261 and energy metabolism. Because of its conserved structure, mtDNA has become  
262 a valuable genetic marker for studies in population genetics and phylogenetics  
263 (Anderson et al., 1981; Boore, 1999). In animal species, the mitochondrial genome

264 ranges from 15–20 kilobase and contains 13 protein-coding genes, 22 tRNAs, and  
265 two rRNAs arranged compactly without introns (Gray, 2012). In comparison to  
266 nuclear DNA, the ratio of the number of copies of mtDNA is higher and has  
267 simple organization which make it particularly suitable for genome sequencing  
268 and assembly studies (Dierckxsens et al., 2017).

### 269 **2.1.1 Mitochondrial Genome Assembly**

270 Mitochondrial genome assembly refers to the reconstruction of the complete mito-  
271 chondrial DNA (mtDNA) sequence from raw or fragmented sequencing reads. It is  
272 conducted to obtain high-quality, continuous representations of the mitochondrial  
273 genome that can be used for a wide range of analyses, including species identi-  
274 fication, phylogenetic reconstruction, evolutionary studies, and investigations of  
275 mitochondrial diseases. Because mtDNA evolves rapidly, its assembled sequence  
276 provides valuable insights into population structure, lineage divergence, and adap-  
277 tive evolution across taxa (Boore, 1999). Compared to nuclear genome assembly,  
278 assembling the mitochondrial genome is often considered more straightforward but  
279 still encounters technical challenges such as the formation of chimeric reads. Com-  
280 monly used tools for mitogenome assembly such as GetOrganelle and MITObim  
281 operate under the assumption of organelle genome circularity, and are vulnerable  
282 when chimeric reads disrupt this circular structure, resulting in assembly errors  
283 (Hahn et al., 2013; Jin et al., 2020).

## **284 2.2 PCR Amplification and Chimera Formation**

285 PCR plays an important role in NGS library preparation, as it amplifies target  
286 DNA fragments for downstream analysis. However as previously mentioned, the  
287 amplification process can also introduce chimeric reads which compromises the  
288 quality of the input reads supplied to sequencing or assembly workflows. Chimeras  
289 typically arise when incomplete extension occurs during a PCR cycle. This causes  
290 the DNA polymerase to switch from one template to another and generate hy-  
291 brid recombinant molecules (Judo et al., 1998). Artificial chimeras are produced  
292 through such amplification errors, whereas biological chimeras occur naturally  
293 through genomic rearrangements or transcriptional events.

294 In the context of amplicon-based sequencing, the presence of chimeras can in-  
295 flate estimates of genetic or microbial diversity and may cause misassemblies dur-  
296 ing genome reconstruction. Qin et al. (2023) has reported that chimeric sequences  
297 may account for more than 10% of raw reads in amplicon datasets. This artifact  
298 tends to be most prominent among rare operational taxonomic units (OTUs) or  
299 singletons, which are sometimes misinterpreted as novel diversity, further caus-  
300 ing the complication of microbial diversity analyses (Gonzalez, Zimmermann, &  
301 Saiz-Jimenez, 2004). As such, determining and minimizing PCR-induced chimera  
302 formation is vital for improving the quality of mitochondrial genome assemblies,  
303 and ensuring the reliability of amplicon sequencing data.

304    **2.3 Existing Traditional Approaches for Chimera**

305    **Detection**

306    Several computational tools have been developed to identify chimeric sequences in  
307    NGS datasets. These tools generally fall into two categories: reference-based and  
308    de novo approaches. Reference-based chimera detection, also known as database-  
309    dependent detection, is one of the earliest and most widely used computational  
310    strategies for identifying chimeric sequences in amplicon-based community studies.  
311    These methods rely on the comparison of each query sequence against a curated,  
312    high-quality database of known, non-chimeric reference sequences (Edgar et al.,  
313    2011).

314    On the other hand, the de novo chimera detection, also referred to as reference-  
315    free detection, represents an alternative computational paradigm that identifies  
316    chimeric sequences without reliance on external reference databases. This method  
317    infer chimeras based on internal relationships among the sequences present within  
318    the dataset itself, making it particularly advantageous in studies of under explored  
319    or taxonomically diverse communities where comprehensive reference databases  
320    are unavailable or incomplete (Edgar, 2016; Edgar et al., 2011). The underlying  
321    assumption on this method is that during PCR, true biological sequences are  
322    generally more abundant as they are amplified early and dominate the read pool,  
323    whereas chimeric sequences appear later and are generally less abundant. The  
324    de novo approach leverage this abundance hierarchy, treating the most abundant  
325    sequences as supposed parents and testing whether less abundant sequences can  
326    be reconstructed as mosaics of these templates. Compositional and structural  
327    similarity are also evaluated to check whether different regions of a candidate

328 sequence correspond to distinct high-abundance sequences.

329 In practice, many modern bioinformatics pipelines combine both paradigms  
330 sequentially: an initial de novo step identifies dataset-specific chimeras, followed  
331 by a reference-based pass that removes remaining artifacts relative to established  
332 databases (Edgar, 2016). These two methods of detection form the foundation of  
333 tools such as UCHIME and later UCHIME2.

### 334 2.3.1 UCHIME

335 UCHIME is one of the most widely used tools for detecting chimeric sequences in  
336 amplicon-based studies and remains a standard quality-control step in microbial  
337 community analysis. Its core strategy is to test whether a query sequence ( $Q$ ) can  
338 be explained as a mosaic of two parent sequences, ( $A$  and  $B$ ), and to score this  
339 relationship using a structured alignment model (Edgar et al., 2011).

340 In reference mode, UCHIME divides the query into several segments and maps  
341 them against a curated database of non-chimeric sequences. Candidate parents  
342 are identified, and a three-way alignment is constructed. The algorithm assigns  
343 “Yes” votes when different segments of the query match different parents and  
344 “No” votes when the alignment contradicts a chimeric pattern. The final score  
345 reflects the balance of these votes. In de novo mode, UCHIME operationalizes the  
346 abundance-skew principle described earlier: high-abundance sequences are treated  
347 as candidate parents, and lower-abundance sequences are evaluated as potential  
348 mosaics. This makes the method especially useful when no reliable reference  
349 database exists.

350        Although UCHIME is highly sensitive, it faces key constraints. Chimeras  
351      formed from parents with very low divergence (below 0.8%) are difficult to detect  
352      because they are nearly indistinguishable from sequencing errors. Accuracy in ref-  
353      erence mode depends strongly on database completeness, while de novo detection  
354      assumes that true parents are both present and sufficiently more abundant, such  
355      conditions are not always met.

356      **2.3.2    UCHIME2**

357      UCHIME2 extends the original algorithm with refinements tailored for high-  
358      resolution sequencing data. One of its major contributions is a re-evaluation  
359      of benchmarking practices. Edgar (2016) demonstrated that earlier accuracy es-  
360      timates for chimera detection were overly optimistic because they relied on un-  
361      realistic scenarios where all true parent sequences were assumed to be present.  
362      Using the more rigorous CHSIMA benchmark, UCHIME2 showed the prevalence  
363      of “fake models” or real biological sequences that can be perfectly reconstructed  
364      as apparent chimeras of other sequences, which suggests that perfect chimera de-  
365      tection is theoretically unattainable. UCHIME2 also introduces several preset  
366      modes (e.g., denoised, balanced, sensitive, specific, high-confidence) designed to  
367      tune sensitivity and specificity depending on dataset characteristics. These modes  
368      allow users to adjust the algorithm to the expected noise level or analytical goals.

369      Despite these improvements, UCHIME2 must be applied with caution. The  
370      website manual explicitly advises against using UCHIME2 as a standalone  
371      chimera-filtering step in OTU clustering or denoising workflows because doing so  
372      can inflate both false positives and false negatives (Edgar, n.d.).

### 373 2.3.3 CATch

374 As previously mentioned, UCHIME (Edgar et al., 2011) relied on alignment-based  
375 sequences in amplicon data. However, researchers soon observed that different al-  
376 gorithms often produced inconsistent predictions. A sequence might be identified  
377 as chimeric by one tool but classified as non-chimeric by another, resulting in  
378 unreliable filtering outcomes across studies.

379 To address these inconsistencies, Mysara, Saeys, Leys, Raes, and Monsieurs  
380 (2015) developed the Classifier for Amplicon Tool Chimeras (CATCh), which rep-  
381 resents the first ensemble machine learning system designed for chimera detection  
382 in 16S rRNA amplicon sequencing. Rather than depending on a single detec-  
383 tion strategy, CATCh integrates the outputs of several established tools, includ-  
384 ing UCHIME, ChimeraSlayer, DECIPHER, Pintail, and Perseus. The individual  
385 scores and binary decisions generated by these tools are used as input features for  
386 a supervised learning model. The algorithm employs a Support Vector Machine  
387 (SVM) with a Pearson VII Universal Kernel (PUK) to determine optimal weight-  
388 ings among the input features and to assign each sequence a probability of being  
389 chimeric.

390 Benchmarking in both reference-based and de novo modes demonstrated signif-  
391 icant performance improvements. CATCh achieved sensitivities of approximately  
392 85 percent in reference-based mode and 92 percent in de novo mode, with corre-  
393 sponding specificities of approximately 96 percent and 95 percent. These results  
394 indicate that CATCh detected 7 to 12 percent more chimeras than any individual  
395 algorithm while maintaining high precision.

396 **2.3.4 ChimPipe**

397 Among the available tools for chimera detection, ChimPipe is a pipeline developed  
398 to identify chimeric sequences such as biological chimeras. It uses both discordant  
399 paired-end reads and split-read alignments to improve the accuracy and sensitivity  
400 of detecting biological chimeras (Rodriguez-Martin et al., 2017). By combining  
401 these two sources of information, ChimPipe achieves better precision than meth-  
402 ods that depend on a single type of indicator.

403 The pipeline works with many eukaryotic species that have available genome  
404 and annotation data (Rodriguez-Martin et al., 2017). It can also predict multiple  
405 isoforms for each gene pair and identify breakpoint coordinates that are useful  
406 for reconstructing and verifying chimeric transcripts. Tests using both simulated  
407 and real datasets have shown that ChimPipe maintains high accuracy and reliable  
408 performance.

409 ChimPipe lets users adjust parameters to fit different sequencing protocols or  
410 organism characteristics. Experimental results have confirmed that many chimeric  
411 transcripts detected by the tool correspond to functional fusion proteins, demon-  
412 strating its utility for understanding chimera biology and its potential applications  
413 in disease research (Rodriguez-Martin et al., 2017).

## 414 2.4 Machine Learning Approaches for Chimera 415 and Sequence Quality Detection

416 Traditional chimera detection tools rely primarily on heuristic or alignment-based  
417 rules. Recent advances in machine learning (ML) have demonstrated that models  
418 trained on sequence-derived features can effectively capture compositional and  
419 structural patterns in biological sequences. Although most existing ML systems  
420 such as those used for antibiotic resistance prediction, taxonomic classification,  
421 or viral identification are not specifically designed for chimera detection, they  
422 highlight how data-driven models can outperform similarity-based heuristics by  
423 learning intrinsic sequence signatures. In principle, ML frameworks can integrate  
424 indicators such as k-mer frequencies, GC-content variation and split-alignment  
425 metrics to identify subtle anomalies that may indicate a chimeric origin (Arango  
426 et al., 2018; Liang, Bible, Liu, Zou, & Wei, 2020; Ren et al., 2020).

### 427 2.4.1 Feature-Based Representations of Genomic Se- 428 quences

429 Feature extraction converts DNA sequences into numerical representations suit-  
430 able for machine learning models. One approach is k-mer frequency analysis,  
431 which counts short nucleotide sequences within a read (Vervier, Mahé, Tournoud,  
432 Veyrieras, & Vert, 2015). High-frequency k-mers, including simple repeats such  
433 as “AAAAAA,” can highlight repetitive or unusual regions that may occur near  
434 chimeric junctions. Comparing k-mer patterns across adjacent parts of a read can  
435 help identify such regions, while GC content provides an additional descriptor of

436 local sequence composition (Ren et al., 2020).

437 Alignment-derived features further inform junction detection. Long-read tools  
438 such as Sniffles (Sedlazeck et al., 2018) use split alignments to locate breakpoints  
439 across extended sequences, whereas short-read aligners like Minimap2 (Li, 2018)  
440 report supplementary and secondary alignments that indicate local discontinu-  
441 ities. Split alignments, where parts of a read map to different regions, can reveal  
442 template-switching events. These features complement k-mer profiles and en-  
443 hance detection of potentially chimeric reads, even in datasets with incomplete  
444 references.

445 Microhomology, or short sequences shared between adjacent segments, is an-  
446 other biologically meaningful feature. Short microhomologies, typically 3–20 bp,  
447 are involved in template switching both in cellular repair pathways and during  
448 PCR, where they act as signatures of chimera formation (Peccoud et al., 2018;  
449 Sfeir & Symington, 2015). In PCR-induced chimeras, short identical sequences  
450 at junctions provide a clear signature of chimerism. Measuring the longest exact  
451 overlap at each breakpoint complements k-mer and alignment features and helps  
452 identify reads that are potentially chimeric.

## 453 2.5 Synthesis of Chimera Detection Approaches

454 To provide an integrated overview of the literature discussed in this chapter, Ta-  
455 ble 2.1 summarizes the major chimera detection studies, their methodological  
456 approaches, and their known limitations.

Table 2.1: Comparison of Chimera Detection Approaches and Tools

Method / Tool	Core Approach	Key Limitations
<b>Reference-based Detection</b>	Compares each query sequence against curated databases of verified, non-chimeric sequences; evaluates segment similarity to identify mosaic patterns.	Accuracy depends on database completeness; performs poorly for novel taxa or missing parents; limited sensitivity for low-divergence chimeras.
<b>De novo Detection</b>	Identifies chimeras using only internal dataset structure; leverages abundance hierarchy and compositional similarity to infer whether low-abundance sequences can be reconstructed from abundant parents.	Assumes true sequences are more abundant; fails when amplification bias distorts abundances; struggles when parental sequences are similarly abundant or highly similar.
<b>UCHIME</b>	Alignment-based model that partitions the query into segments, identifies parent candidates, and computes a chimera score via a three-way alignment; supports reference and de novo modes.	Reduced accuracy for very closely related parents (<0.8% divergence); sensitive to incomplete databases; de novo mode fails if parents are absent or not sufficiently more abundant.
<b>UCHIME2</b>	Updated UCHIME with improved benchmarking (CHSIMA) and multiple sensitivity/specificity presets; better handles incomplete references and dataset variability.	“Fake models” limit theoretical accuracy; genuine variants may mimic chimeras; not recommended as a standalone step in OTU or denoising pipelines due to increased false positives/negatives.
<b>CATCh</b>	First ensemble ML model for 16S chimera detection; integrates outputs of UCHIME, ChimeraSlayer, DECIPHER, Pintail, and Perseus using an SVM to boost overall prediction accuracy.	Performance constrained by underlying tools; ML model cannot capture features not present in component algorithms; may misclassify in highly novel or low-coverage datasets.
<b>ChimPipe</b>	Pipeline for detecting biological chimeras in RNA-seq using discordant paired-end reads and split-read alignments; identifies isoforms and breakpoint coordinates.	Requires high-quality genome and annotation; tailored to RNA-seq rather than amplicons; computationally intensive; limited to organisms with available reference genomes.

457 Across existing studies, no single approach reliably detects all forms of chimeric  
458 sequences, and the reviewed literature consistently shows that chimeras remain a  
459 persistent challenge in genomics and bioinformatics. Although the surveyed tools  
460 are not designed specifically for organelle genome assembly, they provide valua-  
461 ble insights into which methodological strategies are effective and where current  
462 approaches fall short. These limitations collectively define a clear research gap:  
463 the need for a specialized, feature-driven detection framework tailored to PCR-  
464 induced mitochondrial chimeras. Addressing this gap aligns with the research  
465 objective outlined in Section 1.3, which is to develop and evaluate a machine  
466 learning-based pipeline (MitoChime) that improves the quality of downstream  
467 mitochondrial genome assembly. In support of this aim, the subsequent chapters  
468 describe the design, implementation, and evaluation of the proposed tool.

# <sup>469</sup> Chapter 3

## <sup>470</sup> Research Methodology

<sup>471</sup> This chapter outlines the steps involved in completing the study, including data  
<sup>472</sup> gathering, generating simulated mitochondrial Illumina reads, preprocessing and  
<sup>473</sup> indexing the data, developing a feature extraction pipeline to obtain read-level fea-  
<sup>474</sup> tures, applying machine learning algorithms for chimera detection, implementing  
<sup>475</sup> feature selection methods, and validating and comparing model performance.

### <sup>476</sup> 3.1 Research Activities

<sup>477</sup> As illustrated in Figure 3.1, this study carried out a sequence of procedures to  
<sup>478</sup> detect PCR-induced chimeric reads in mitochondrial genomes. The process began  
<sup>479</sup> with collecting a mitochondrial reference sequence of *Sardinella lemuru* from the  
<sup>480</sup> National Center for Biotechnology Information (NCBI) database, which was used  
<sup>481</sup> as a reference for generating simulated clean and chimeric reads. These reads  
<sup>482</sup> were subsequently indexed and mapped. The resulting collections then passed

483 through a feature extraction pipeline that computed k-mer profiles, supplementary  
484 alignment (SA) features, and microhomology information to prepare the data  
485 for model construction. The machine learning models were trained using the  
486 processed input, evaluated using cross-validation and held-out testing, tuned for  
487 improved performance, and then subjected to feature importance and feature  
488 selection analyses before final validation.

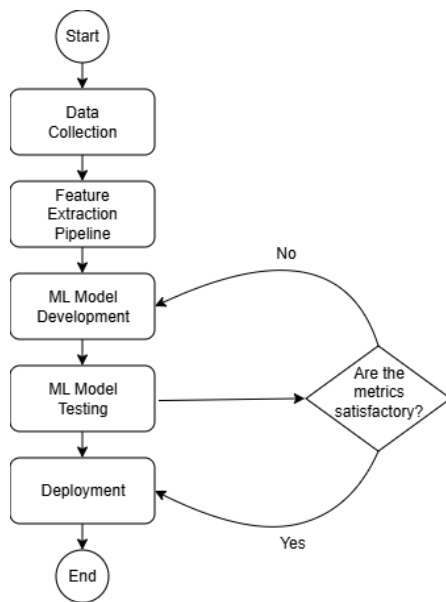


Figure 3.1: Process diagram of the study workflow.

### 489 3.1.1 Data Collection

490 The mitochondrial genome reference sequence of *S. lemuru* was obtained from the  
491 NCBI database (accession number NC\_039553.1) in FASTA format and was used  
492 to generate simulated reads.

493 This step was scheduled to begin in the first week of November 2025 and  
494 expected to be completed by the end of that week, with a total duration of ap-

495 proximately one (1) week.

## 496 Data Preprocessing

497 All steps in the simulation and preprocessing pipeline were executed using a cus-  
498 tom script in Python (Version 3.11). The script runs each stage, including read  
499 simulation, reference indexing, mapping, and alignment processing, in a fixed se-  
500 quence.

501 `wgsim` (Version 1.13) was used to simulate 10,000 paired-end fragments, pro-  
502 ducing 20,000 reads (10,000 forward and 10,000 reverse) from the original refer-  
503 ence (`original_reference.fasta`) and designated as clean reads. The tool was  
504 selected because it provides fast generation of Illumina-like reads with controllable  
505 error rates, using the following command:

```
506 wgsim -1 150 -2 150 -r 0 -R 0 -X 0 -e 0.05 -N 10000 \
507     original_reference.fasta ref1.fastq ref2.fastq
```

508 Chimeric sequences were then generated from the same reference FASTA  
509 file using a separate Python script. Two non-adjacent segments were ran-  
510 domly selected such that their midpoint distances fell within specified minimum  
511 and maximum thresholds. The script attempted to retain microhomology to  
512 mimic PCR-induced template switching. The resulting chimeras were written  
513 to `chimera_reference.fasta` and processed with `wgsim` to simulate 10,000  
514 paired-end fragments, generating 20,000 chimeric reads (10,000 forward reads in  
515 `chimeric1.fastq` and 10,000 reverse reads in `chimeric2.fastq`) using the same  
516 command format as above.

517        Next, a `minimap2` index of the reference genome was created using:

```
518  minimap2 -d ref.mmi original_reference.fasta
```

519        Minimap2 (Version 2.28) was used to map simulated clean and chimeric reads  
520        to the original reference. An index (`ref.mmi`) was first generated to enable efficient  
521        alignment, and mapping produced the alignment features used as input for the  
522        machine learning model. The reads were mapped using the following commands:

```
523  minimap2 -ax sr -t 8 ref.mmi ref1.fastq ref2.fastq > clean.sam
```

```
524  minimap2 -ax sr -t 8 ref.mmi \  
525      chimeric1.fastq chimeric2.fastq > chimeric.sam
```

526        The resulting clean and chimeric SAM files contain the alignment positions of  
527        each read relative to the original reference genome. These files were then converted  
528        to BAM format, sorted, and indexed using `samtools` (Version 1.20):

```
529  samtools view -bS clean.sam -o clean.bam  
530  samtools view -bS chimeric.sam -o chimeric.bam  
531  
532  samtools sort clean.bam -o clean.sorted.bam  
533  samtools index clean.sorted.bam  
534  
535  samtools sort chimeric.bam -o chimeric.sorted.bam  
536  samtools index chimeric.sorted.bam
```

537        The total number of simulated reads was expected to be 40,000. The final col-  
538        lection of reads contained 19,984 clean reads and 20,000 chimeric reads (39,984 en-  
539        tries in total), providing a roughly balanced distribution between the two classes.  
540        After alignment with `minimap2`, only 19,984 clean reads remained because un-  
541        mapped reads were not included in the BAM file. Some sequences failed to align  
542        due to the error rate defined during `wgsim` simulation, which produced mismatches  
543        that caused certain reads to fall below the aligner’s matching threshold.

544        This whole process was scheduled to start in the second week of November 2025  
545        and was expected to be completed by the last week of November 2025, with a total  
546        duration of approximately three (3) weeks.

### 547        3.1.2 Feature Extraction Pipeline

548        This stage directly followed the alignment phase, utilizing the resulting BAM files  
549        (specifically `chimeric.sorted.bam` and `clean.sorted.bam`). A custom Python  
550        script was created to efficiently process each primary-mapped read to extract  
551        the necessary set of features, which were then compiled into a structured feature  
552        matrix in TSV format. The pipeline’s core functionality relied on the `Pysam`  
553        (Version 0.22) library for parsing BAM structures and `NumPy` (Version 1.26) for  
554        array operations and computations. To ensure correctness and adherence to best  
555        practices, bioinformatics experts at PGC Visayas were consulted to validate the  
556        pipeline design, feature extraction logic, and overall data integrity.

557        This stage of the study was scheduled to begin in the last week of Novem-  
558        ber 2025 and conclude by the first week of December 2025, with an estimated

559 total duration of approximately two (2) weeks.

560 The pipeline focused on three feature families that collectively capture bi-  
561 ological signatures associated with PCR-induced chimeras: (1) supplementary  
562 alignment (SA) and alignment-structure metrics, (2) k-mer composition differ-  
563 ence, and (3) microhomology around putative junctions. Additional alignment  
564 quality indicators such as mapping quality were also included.

## 565 **Supplementary Alignment and Alignment-Structure Features**

566 Split-alignment information was derived from the SA tag embedded in each pri-  
567 mary read of the BAM file. This tag is typically associated with reads that map to  
568 multiple genomic locations, suggesting a chimeric structure. To extract this infor-  
569 mation, the script first checked whether the read carried an **SA:Z** tag. If present,  
570 the tag string was parsed using the function **parse\_sa\_tag**, yielding metadata for  
571 each alignment containing the reference name, mapped position, strand, mapping  
572 quality, and number of mismatches.

573 After parsing, the function **sa\_feature\_stats** was applied to establish the fun-  
574 damental split indicators, **has\_sa** and **sa\_count**. Along with these initial counts,  
575 the function aggregated metrics related to the structure and reliability of the  
576 split alignments, including the number of alignment segments, strand consistency,  
577 minimum, maximum, and mean distance between split segments, and summary  
578 statistics of mapping quality and mismatch counts across segments.

579 **K-mer Composition Difference**

580 Comparing k-mer frequency profiles between the left and right halves of a read  
581 allows for the detection of abrupt compositional shifts, independent of alignment  
582 information.

583 The script implemented this by inferring a likely junction breakpoint using the  
584 function `infer_breakpoints`, prioritizing the boundaries defined by soft-clipping  
585 operations. If no clipping was present, the midpoint of the alignment or the read  
586 length was used as a fallback. The read sequence was then divided into left and  
587 right segments at this inferred breakpoint, and k-mer frequency profiles ( $k =$   
588 6) were generated for both halves, ignoring any k-mers containing ambiguous N  
589 bases. The resulting k-mer frequency vectors were normalised and compared using  
590 the functions `cosine_difference` and `js_divergence` to quantify compositional  
591 discontinuity across the inferred breakpoint.

592 **Microhomology**

593 The process of extracting the microhomology feature also started by using  
594 `infer_breakpoints` to identify a candidate junction. Once a breakpoint was  
595 established, the script scanned a  $\pm 40$  base-pair window surrounding the break-  
596 point and applied the function `longest_suffix_prefix_overlap` to identify the  
597 longest exact suffix-prefix overlap between the left and right read segments. This  
598 overlap, representing consecutive bases shared at the junction, was recorded as  
599 `microhomology_length` in the dataset. The 40 base-pair window was chosen  
600 to ensure that short shared sequences at or near the breakpoint were captured

601 without including distant sequences that are unlikely to be mechanistically  
602 related.

603 Additionally, the GC content of the overlapping sequence was calculated using  
604 the function `gc_content`, which counts guanine (G) and cytosine (C) bases within  
605 the detected microhomology and divides by the total length, yielding a proportion  
606 between 0 and 1 that was stored under the `microhomology_gc` attribute. Micro-  
607 homology was quantified using a 3–20 bp window, consistent with values reported  
608 in prior research on PCR-induced chimeras. A k-mer length of 6 was used to cap-  
609 ture patterns within the 40 bp window surrounding each breakpoint, providing  
610 sufficient resolution to detect informative sequence shifts.

### 611 3.1.3 Machine Learning Model Development

612 After feature extraction, the per-read feature matrices for clean and chimeric  
613 reads were merged into a single dataset. Each row corresponded to one paired-  
614 end read, and columns encoded alignment-structure features (e.g., supplementary  
615 alignment count and spacing between segments), CIGAR-derived soft-clipping  
616 statistics (e.g., left and right soft-clipped length, total clipped bases), k-mer com-  
617 position discontinuity between read segments, microhomology descriptors near  
618 candidate junctions, and alignment quality (e.g., mapping quality). The result-  
619 ing feature set comprised 23 numeric features and was restricted to quantities  
620 that can be computed from standard BAM/FASTQ files in typical mitochondrial  
621 sequencing workflows.

622 The labelled dataset was randomly partitioned into training (80%) and test

623 (20%) subsets using stratified sampling to preserve the 1:1 ratio of clean to  
624 chimeric reads. Model development and evaluation were implemented in Python  
625 (Version 3.11) using the `scikit-learn`, `xgboost`, `lightgbm`, and `catboost` li-  
626 braries. A broad panel of classification algorithms was then benchmarked on the  
627 training data to obtain a fair comparison of different model families under identical  
628 feature conditions. The panel included a trivial dummy classifier,  $L_2$ -regularized  
629 logistic regression, a calibrated linear support vector machine (SVM),  $k$ -nearest  
630 neighbours, Gaussian Naïve Bayes, decision-tree ensembles (Random Forest, Ex-  
631 tremely Randomized Trees, and Bagging with decision trees), gradient boosting  
632 methods (Gradient Boosting, XGBoost, LightGBM, and CatBoost), and a shallow  
633 multilayer perceptron (MLP).

634 For each model, five-fold stratified cross-validation was performed on the train-  
635 ing set. In every fold, four-fifths of the data were used for fitting and the remaining  
636 one-fifth for validation. Mean cross-validation accuracy, precision, recall, F1-score  
637 for the chimeric class, and area under the receiver operating characteristic curve  
638 (ROC–AUC) were computed to summarize performance and rank candidate meth-  
639 ods. This baseline screen allowed comparison of linear, probabilistic, neural, and  
640 ensemble-based approaches and identified tree-based ensemble and boosting mod-  
641 els as consistently strong performers relative to simpler baselines.

#### 642 **3.1.4 Model Benchmarking, Hyperparameter Optimiza- 643 tion, and Evaluation**

644 Model selection and refinement proceeded in two stages. First, the cross-validation  
645 results from the broad panel were used to identify a subset of competitive mod-

els for more detailed optimization. Specifically, ten model families were carried forward:  $L_2$ -regularized logistic regression, calibrated linear SVM, Random Forest, ExtraTrees, Gradient Boosting, XGBoost, LightGBM, CatBoost, Bagging with decision trees, and a shallow MLP. This subset spans both linear and non-linear decision boundaries, but emphasizes ensemble and boosting methods, which showed superior F1 and ROC–AUC in the initial benchmark.

Second, hyperparameter optimization was conducted for each of the ten selected models using randomized search with five-fold stratified cross-validation (`RandomizedSearchCV`). For tree-based ensembles, the search space included the number of trees, maximum depth, minimum samples per split and per leaf, and the fraction of features considered at each split. For boosting methods, key hyperparameters such as the number of boosting iterations, learning rate, tree depth, subsampling rate, and column subsampling rate were tuned. For the MLP, the number and size of hidden layers, learning rate, and  $L_2$ -regularization strength were varied. In all cases, the primary optimisation criterion was the F1-score of the chimeric class, averaged across folds.

For each model family, the hyperparameter configuration with the highest mean cross-validation F1-score was selected as the best-tuned estimator. These tuned models were then refitted on the full training set and evaluated once on the held-out test set to obtain unbiased estimates of performance. Test-set metrics included accuracy, precision, recall, F1-score for the chimeric class, and ROC–AUC. Confusion matrices and ROC curves were generated for the top-performing models to characterise common error modes, such as false negatives (missed chimeric reads) and false positives (clean reads incorrectly labelled as chimeric). The final model or small set of models for downstream interpretation was chosen based on

671 a combination of test-set F1-score and ROC–AUC.

672 **3.1.5 Feature Importance, Feature Selection, and Inter-**  
673 **pretation**

674 To relate model decisions to biologically meaningful signals, feature-importance  
675 analyses were performed on the best-performing tree-based models. Two comple-  
676 mentary approaches were used. First, built-in importance measures from ensemble  
677 methods (e.g., split-based importances in Random Forest and Gradient Boosting)  
678 were examined to obtain an initial ranking of features based on their contributon  
679 to reducing impurity. Second, model-agnostic permutation importance was com-  
680 puted on the test set by repeatedly permuting each feature column while keeping  
681 all others fixed and measuring the resulting decrease in F1-score. Features whose  
682 permutation led to a larger performance drop were interpreted as more influential  
683 for chimera detection.

684 For interpretability, individual features were grouped into conceptual families:  
685 (i) supplementary alignment and alignment-structure features (e.g., SA count,  
686 spacing between alignment segments, strand consistency), (ii) soft-clipping fea-  
687 tures (e.g., left and right soft-clipped length, total clipped bases, inferred break-  
688 point position), (iii) k-mer composition discontinuity features (e.g., cosine dis-  
689 tance and Jensen–Shannon divergence between k-mer profiles of read segments),  
690 (iv) microhomology descriptors (e.g., microhomology length and local GC content  
691 around putative breakpoints), and (v) other alignment quality features (e.g., map-  
692 ping quality). This analysis provided a basis for interpreting the trained models  
693 in terms of known mechanisms of PCR-induced template switching and for iden-

694 tifying which alignment-based and sequence-derived cues are most informative for  
695 distinguishing chimeric from clean mitochondrial reads.

696 Building on these importance results, an explicit feature selection step was  
697 implemented using CatBoost as the reference model, since it was among the top-  
698 performing classifiers. Permutation importance scores were re-estimated for Cat-  
699 Boost on the held-out test set using the F1-score of the chimeric class as the  
700 scoring function. Negative importance scores, which indicate that permuting a  
701 feature did not reliably harm performance, were set to zero and interpreted as  
702 noise. The remaining non-negative importances were sorted in descending order  
703 and converted into a cumulative importance curve by expressing each feature's  
704 importance as a fraction of the total positive importance.

705 A compact feature subset was then defined by selecting the smallest number of  
706 features whose cumulative importance reached at least 95% of the total positive  
707 importance. This procedure yielded a reduced set of four strongly predictive  
708 variables dominated by soft-clipping and k-mer divergence metrics (for example,  
709 total clipped bases and k-mer divergence between read halves).

710 To quantify the impact of this reduction, CatBoost was retrained using only  
711 the selected feature subset, with the same tuned hyperparameters as the full 23-  
712 feature model, and evaluated on the held-out test set. Performance of the reduced  
713 model was then compared to that of the full model in terms of F1-score and ROC-  
714 AUC to assess whether dimensionality could be reduced without appreciable loss  
715 in predictive accuracy.

716 In addition, an ablation experiment was performed to specifically evaluate  
717 the contribution of explicit microhomology features. The microhomology vari-

ables (`microhomology_length` and `microhomology_gc`) were removed from the full feature set to obtain a 21-feature configuration. CatBoost was refitted on this microhomology-ablated feature set, using the same tuned hyperparameters, and evaluated on the held-out test set. Comparing the full, reduced-subset, and microhomology-ablated variants allowed the study to quantify both the degree of redundancy among features and the practical contribution of microhomology to classification accuracy.

Taken together, the feature importance and feature selection analyses provided a more parsimonious model variant and a clearer interpretation of which alignment-based and sequence-derived signals are most informative for detecting PCR-induced chimeras.

### 3.1.6 Validation and Testing

Validation involved both internal and external evaluations. Internal validation was achieved through five-fold stratified cross-validation on the training data to verify model generalization and reduce variance due to random sampling. External testing was performed on the 20% hold-out dataset from the simulated reads, providing an unbiased assessment of model generalization. Feature extraction and preprocessing were applied consistently across all splits.

Comparative evaluation was performed across all candidate algorithms and CatBoost feature-set variants to determine which models demonstrated the highest predictive performance and computational efficiency under identical data conditions. Their metrics were compared to identify which algorithms and feature

740 configurations were most suitable for further refinement and potential integration  
741 into downstream mitochondrial assembly workflows.

### 742 3.1.7 Documentation

743 Comprehensive documentation was maintained throughout the study to ensure  
744 transparency and reproducibility. All stages of the research, including data gath-  
745 ering, preprocessing, feature extraction, model training, feature selection, and  
746 validation, were systematically recorded in a `README` file in the GitHub reposi-  
747 tory. For each analytical step, the corresponding parameters, software versions,  
748 and command line scripts were documented to enable exact replication of results.

749 The repository structure followed standard research data management prac-  
750 tices, with clear directories for datasets and scripts. Computational environments  
751 were standardised using Conda, with an environment file (`environment.yml`)  
752 specifying dependencies and package versions to maintain consistency across sys-  
753 tems.

754 For manuscript preparation and supplementary materials, Overleaf (LATEX)  
755 was used to produce publication-quality formatting and consistent referencing.

## 756 3.2 Calendar of Activities

757 Table 3.1 presents the project timeline in the form of a Gantt chart, where each  
758 bullet point corresponds to approximately one week of planned activity.

Table 3.1: Timetable of activities.

<b>Activities (2025)</b>	<b>Nov</b>	<b>Dec</b>	<b>Jan</b>	<b>Feb</b>	<b>Mar</b>	<b>Apr</b>	<b>May</b>
Data Collection and Simulation	• • • •						
Feature Extraction Pipeline	•	•					
Machine Learning Development		•	• •	• • • •	• • • •	• •	
Testing and Validation						• •	• • • •
Documentation	• • • •	• • • •	• • • •	• • • •	• • • •	• • • •	• • • •

759 **Chapter 4**

760 **Results and Discussion**

761 This chapter presents the performance of the proposed feature set and machine  
762 learning models for detecting PCR-induced chimeric reads in simulated mito-  
763 chondrial Illumina data. The behaviour of the extracted features is first examined  
764 through descriptive and correlation analyses, followed by a comparison of baseline  
765 and tuned classifiers. The chapter then examines model performance in detail and  
766 investigates the contribution of individual features and feature families, including  
767 the impact of feature selection on classification performance.

768 The final dataset contained 31,986 reads for training and 7,997 reads for test-  
769 ing, with classes balanced (approximately 4,000 clean and 4,000 chimeric reads in  
770 the test split).

## 771 4.1 Descriptive Analysis of Features

### 772 4.1.1 Summary Statistics Per Class

773 Summary statistics were computed separately for clean reads (class 0) and  
774 chimeric reads (class 1) to characterize the distributional behavior of the features.  
775 For each feature, the mean, standard deviation, median, first and third quartiles  
776 (Q1, Q3), interquartile range (IQR), minimum, maximum, and sample size ( $n$ )  
777 were calculated.

778 Only a subset of the features is summarized in the main text to highlight key  
779 trends, and not all summary statistics columns are shown for brevity. The com-  
780 plete set of per-class summary statistics for all features is provided in Appendix A  
781 (Table A.1).

### 782 Alignment and Supplementary Alignment Features

783 Features related to supplementary alignments show strong separation between  
784 classes. Chimeric reads exhibit supplementary alignments, as reflected by higher  
785 values of `has_sa`, `sa_count`, and `num_segments`, whereas clean reads consistently  
786 show a single alignment segment with no supplementary mappings. This behavior  
787 is consistent with the expected structure of chimeric reads and indicates that  
788 alignment-based features are highly informative.

789 **Clipping-Based Features**

790 Clipping-related features, including `softclip_left`, `softclip_right`, and  
791 `total_clipped_bases`, display higher means and broader distributions in chimeric  
792 reads. Clean reads are dominated by zero or near-zero clipping, while chimeric  
793 reads exhibit increased clipping and greater variability, which reflects the presence  
794 of split alignments.

795 **K-mer Distribution Features**

796 K-mer-based features, such as `kmer_js_divergence` and `kmer_cosine_diff`, show  
797 only modest differences between clean and chimeric reads. Chimeric reads show  
798 slightly higher average divergence, but substantial overlap with clean reads means  
799 this feature alone cannot reliably distinguish the classes.

800 **Microhomology Features**

801 Microhomology-related features (`microhomology_length` and `microhomology_gc`)  
802 exhibit nearly identical summary statistics across both classes. The majority of  
803 reads in both classes contain short or zero-length microhomologies, resulting in  
804 minimal separation. This means that microhomology serves as a weak standalone  
805 indicator and is more appropriately treated as supporting evidence.

806 Overall, the summary statistics indicate that alignment-based and clipping-  
807 based features provide the strongest class separation, k-mer features contribute  
808 limited but complementary signal, and microhomology features exhibit minimal

809 discriminative power on their own. These observations motivate the combined  
810 multi-feature approach used in subsequent modeling and evaluation.

Table 4.1: Summary statistics of selected key features by class.

Feature	Class	Mean	Std	Median	IQR
has_sa	chimeric	0.406	0.491	0.0	1.0
has_sa	clean	0.000	0.000	0.0	0.0
num_segments	chimeric	1.406	0.491	1.0	1.0
num_segments	clean	1.000	0.000	1.0	0.0
softclip_left	chimeric	12.55	21.90	0.0	19.0
softclip_left	clean	0.23	1.54	0.0	0.0
softclip_right	chimeric	12.90	22.12	0.0	19.0
softclip_right	clean	0.21	1.51	0.0	0.0
total_clipped_bases	chimeric	25.44	25.48	19.0	48.0
total_clipped_bases	clean	0.44	2.16	0.0	0.0
kmer_js_divergence	chimeric	0.974	0.025	0.986	0.043
kmer_js_divergence	clean	0.976	0.025	0.986	0.040
kmer_cosine_diff	chimeric	0.974	0.026	0.986	0.042
kmer_cosine_diff	clean	0.976	0.025	0.986	0.041
microhomology_length	chimeric	0.458	0.755	0.0	1.0
microhomology_length	clean	0.462	0.758	0.0	1.0
microhomology_gc	chimeric	0.172	0.361	0.0	0.0
microhomology_gc	clean	0.172	0.361	0.0	0.0

### 811 4.1.2 Boxplots By Class

812 Boxplots were generated for each feature, with the x-axis representing the class  
813 clean reads and chimeric reads and the y-axis representing the feature value. Fig-  
814 ure 4.1 presents a panel of selected key features, while boxplots for all numeric  
815 features are provided in Appendix B.

816 For clipping-related features, chimeric reads exhibit higher medians and longer  
817 upper whiskers than clean reads, indicating increased variability and the presence

818 of split alignments.

819 Supplementary alignment features show that clean reads are largely zero,  
820 whereas chimeric reads display a wider distribution, reflecting frequent supple-  
821 mentary alignments.

822 K-mer metrics show a slight upward shift for chimeric reads, but substantial  
823 overlap with clean reads indicates modest discriminative power.

824 Microhomology features have nearly overlapping distributions for both classes,  
825 consistent with their low standalone predictive importance.

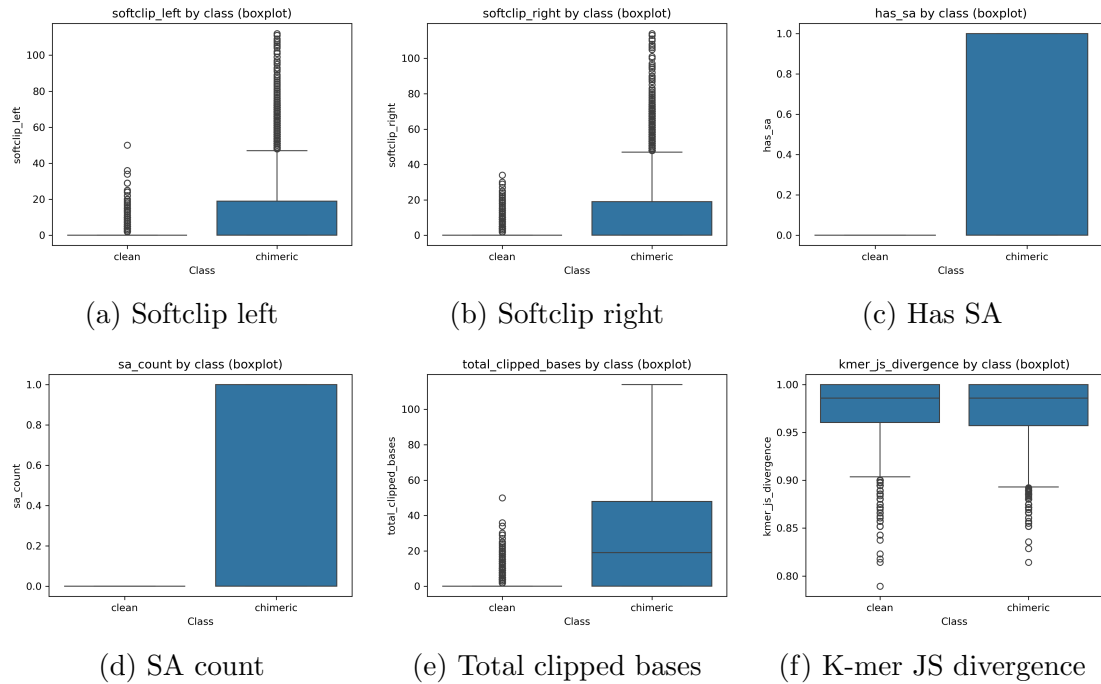


Figure 4.1: Boxplots of key features by class

826 **4.1.3 Correlation Analysis of Extracted Features**

827 A feature correlation heatmap (Figure 4.2) was generated to examine relation-  
828 ships among the extracted variables and to identify patterns of redundancy and  
829 independence within the feature set. The analysis shows that alignment- and  
830 clipping-related features form a strongly correlated cluster, including indicators  
831 of supplementary alignments, alignment segment counts, positional differences,  
832 and soft-clipping measures. These features capture related aspects of alignment  
833 fragmentation, which is a known characteristic of chimeric reads, and several show  
834 moderate correlations with the class label, supporting their relevance for distin-  
835 guishing chimeric from clean reads. In contrast, general read- and alignment-  
836 quality metrics, such as read length, base quality, and mapping quality, exhibit  
837 weak correlations with most split-alignment features, indicating that they pro-  
838 vide distinct information rather than overlapping with alignment-derived signals.  
839 Sequence-based features display a similar pattern of independence, as k-mer di-  
840 vergence metrics show weak correlations with other feature groups, while micro-  
841 homology features exhibit generally low correlations with both alignment- and  
842 k-mer-based features. Overall, the correlation structure highlights intentional re-  
843 dundancy within alignment-derived features and clear separation between feature  
844 families, supporting the use of features that capture different aspects of chimeric  
845 read characteristics to improve chimera classification.

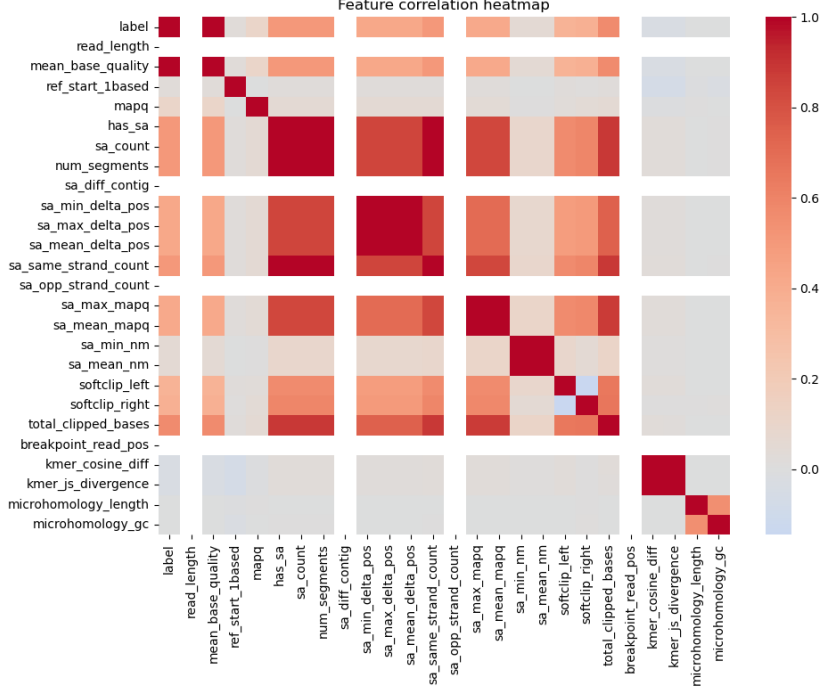


Figure 4.2: Feature correlation heatmap showing relationships among alignment-derived and sequence-derived features.

## 846 4.2 Baseline Classification Performance

847 Table 4.2 summarises the performance of eleven classifiers trained on the engi-  
 848 neered feature set using five-fold cross-validation and evaluated on the held-out  
 849 test set. All models were optimised using default hyperparameters, without ded-  
 850 icated tuning.

851 The dummy baseline, which always predicts the same class regardless of the  
 852 input features, achieved an accuracy of 0.50 and test F1-score of 0.67. This re-  
 853 flects the balanced class distribution and provides a lower bound for meaningful  
 854 performance.

855 Across other models, test F1-scores clustered in a narrow band between ap-  
 856 proximately 0.74 and 0.77 and ROC–AUC values between 0.82 and 0.84. Gradi-  
 857 ent boosting, CatBoost, LightGBM, XGBoost, bagging trees, random forest, and  
 858 multilayer perceptron (MLP) all produced very similar scores, with CatBoost and  
 859 gradient boosting slightly ahead (test F1  $\approx$  0.77, ROC–AUC  $\approx$  0.84). Linear  
 860 models (logistic regression and calibrated linear SVM) performed only marginally  
 861 worse (test F1  $\approx$  0.74), while Gaussian Naive Bayes lagged behind with substan-  
 862 tially lower F1 ( $\approx$  0.65) despite very high precision for the chimeric class.

Table 4.2: Performance of baseline classifiers on the held-out test set.

model	test_accuracy	test_precision	test_recall	test_f1	test_roc_auc
dummy_baseline	0.500000	0.500000	1.000000	0.667000	0.500000
logreg_l2	0.789000	0.945000	0.614000	0.744000	0.821000
linear_svm_calibrated	0.789000	0.945000	0.614000	0.744000	0.820000
random_forest	0.788000	0.894000	0.654000	0.755000	0.834000
extra_trees	0.788000	0.901000	0.647000	0.753000	0.824000
gradient_boosting	0.802000	0.936000	0.648000	0.766000	0.840000
xgboost	0.800000	0.929000	0.650000	0.765000	0.839000
lightgbm	0.799000	0.926000	0.650000	0.764000	0.838000
catboost	0.803000	0.936000	0.650000	0.767000	0.839000
knn	0.782000	0.892000	0.642000	0.747000	0.815000
gaussian_nb	0.741000	0.996000	0.483000	0.651000	0.819000
bagging_trees	0.792000	0.900000	0.657000	0.760000	0.837000
mlp	0.789000	0.931000	0.625000	0.748000	0.819000

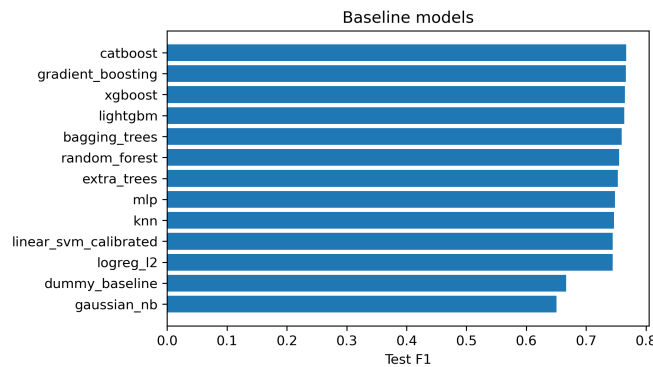


Figure 4.3: Test F1 of all baseline classifiers, showing that no single model clearly dominates and several achieve comparable performance.

### 863 4.3 Effect of Hyperparameter Tuning

864 To assess whether performance could be improved further, ten model families un-  
865 derwent randomised hyperparameter search. The tuned metrics are summarised  
866 in Table 4.3. Overall, tuning yielded modest but consistent gains for tree-based en-  
867sembles and boosting methods, while leaving linear models essentially unchanged  
868 or slightly worse.

869 CatBoost, gradient boosting, LightGBM, XGBoost, random forest, bagging  
870 trees, and MLP all experienced small increases in test F1 (typically  $\Delta F1 \approx 0.002$ –  
871  $0.009$ ) and ROC–AUC (up to  $\Delta AUC \approx 0.008$ ). After tuning, CatBoost remained  
872 the best performer with test accuracy 0.80, precision 0.92, recall 0.66, F1-score  
873 0.77, and ROC–AUC 0.84. Gradient boosting achieved almost identical perfor-  
874 mance (F1 0.77, AUC 0.84). Random forest and bagging trees also improved to  
875 F1 scores around 0.76 with AUC  $\approx 0.84$ .

Table 4.3: Performance of tuned classifiers on the held-out test set.

model	test_accuracy	test_precision	test_recall	test_f1	test_roc_auc
logreg_l2_tuned	0.788000	0.946000	0.612000	0.743000	0.818000
linear_svm_calibrated_tuned	0.788000	0.944000	0.612000	0.743000	0.818000
random_forest_tuned	0.797000	0.915000	0.655000	0.763000	0.842000
extra_trees_tuned	0.794000	0.910000	0.652000	0.760000	0.837000
gradient_boosting_tuned	0.802000	0.928000	0.654000	0.767000	0.843000
xgboost_tuned	0.799000	0.922000	0.653000	0.765000	0.839000
lightgbm_tuned	0.801000	0.930000	0.651000	0.766000	0.842000
catboost_tuned	0.802000	0.924000	0.658000	0.769000	0.844000
bagging_trees_tuned	0.798000	0.922000	0.650000	0.763000	0.842000
mlp_tuned	0.790000	0.934000	0.625000	0.749000	0.821000

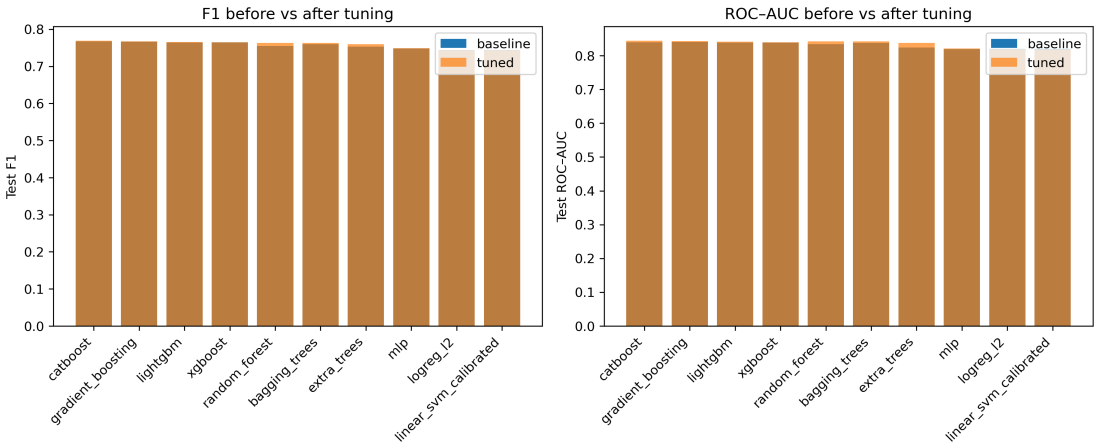


Figure 4.4: Comparison of test F1 (left) and ROC–AUC (right) for baseline and tuned models.

Because improvements are small and within cross-validation variability, tuning was interpreted as stabilising and slightly refining the models rather than completely altering their behaviour or their relative ranking.

## 4.4 Detailed Evaluation of Representative Models

For interpretability and diversity, four tuned models were selected for deeper analysis: CatBoost (best-performing boosted tree), scikit-learn gradient boosting (canonical gradient-boosting implementation), random forest (non-boosted ensemble baseline), and  $L_2$ -regularised logistic regression (linear baseline). All models were trained on the engineered feature set and evaluated on the same held-out test data.

#### **4.4.1 Confusion Matrices and Error Patterns**

Classification reports and confusion matrices for the four models reveal consistent patterns. CatBoost and gradient boosting both reached overall accuracy of approximately 0.80 with similar macro-averaged F1 scores ( $\sim 0.80$ ). For CatBoost, precision and recall for clean reads were 0.73 and 0.95, respectively, while for chimeric reads they were 0.92 and 0.66 ( $F1 = 0.77$ ). Gradient boosting showed nearly identical trade-offs.

Random forest attained slightly lower accuracy (0.80) and chimeric F1 (0.76), whereas logistic regression achieved the lowest accuracy among the four (0.79) and chimeric F1 (0.74), although it provided the highest chimeric precision (0.95) at the cost of lower recall (0.61).

Across all models, errors were asymmetric. False negatives (chimeric reads predicted as clean) were more frequent than false positives. For example, CatBoost misclassified 1,369 chimeric reads as clean but only 215 clean reads as chimeric. This pattern indicates that the models are conservative and prioritise avoiding false chimera calls at the expense of missing some true chimeras. Consultation with PGC Visayas indicated that this conservative behavior is generally acceptable, though further evaluation and testing will be required to assess its impact on downstream analyses.

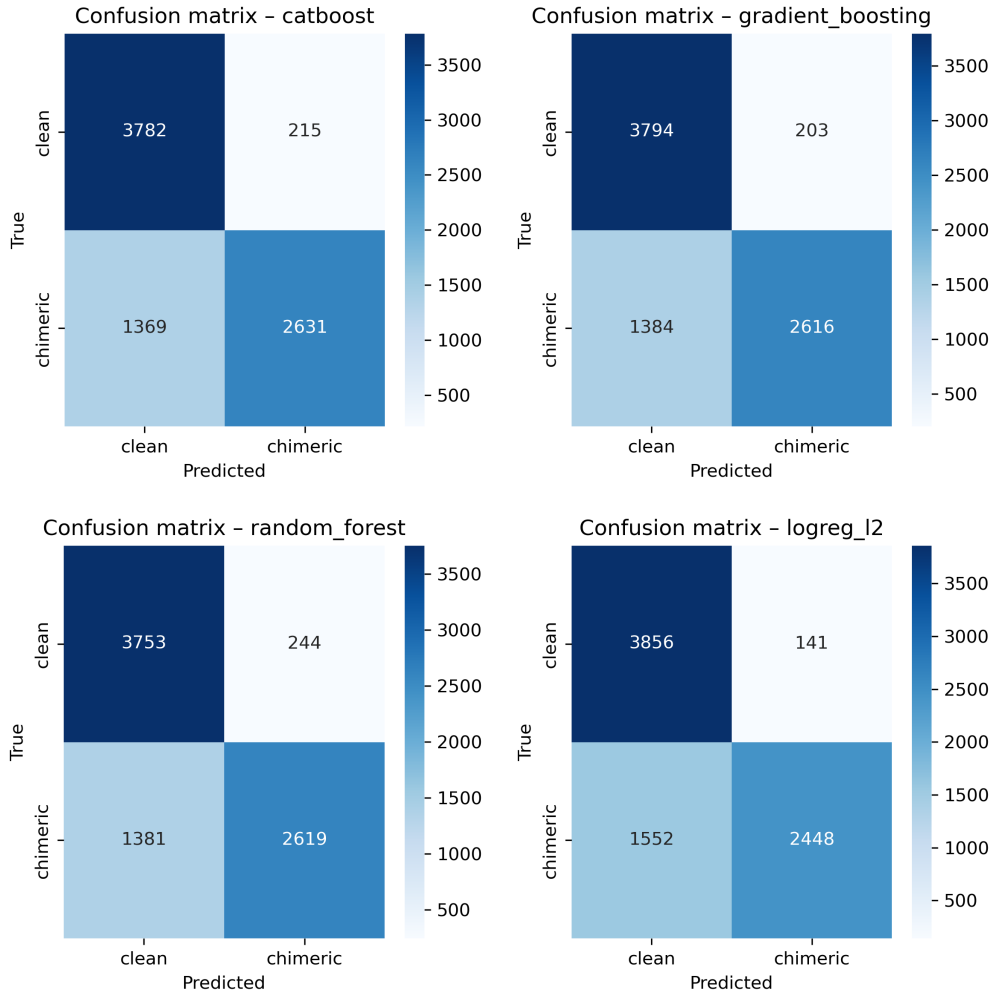


Figure 4.5: Confusion matrices for the four representative models on the held-out test set.

#### 906 4.4.2 ROC and Precision–Recall Curves

907 Receiver operating characteristic (ROC) and precision–recall (PR) curves as  
 908 shown in Figure 4.6 further support the similarity among the top models. The  
 909 three tree-based ensembles (CatBoost, gradient boosting, random forest) achieved  
 910 ROC–AUC values of approximately 0.84 and average precision (AP) around 0.88.

911 Logistic regression performed slightly worse ( $AUC \approx 0.82$ ,  $AP \approx 0.87$ ) but still  
912 substantially better than the dummy baseline.

913 The PR curves show that precision remains above 0.9 across a broad range  
914 of recall values (up to roughly 0.5–0.6), after which precision gradually declines.  
915 This behaviour indicates that the models can assign very high confidence to a  
916 subset of chimeric reads, while more ambiguous reads can only be recovered by  
917 accepting lower precision.

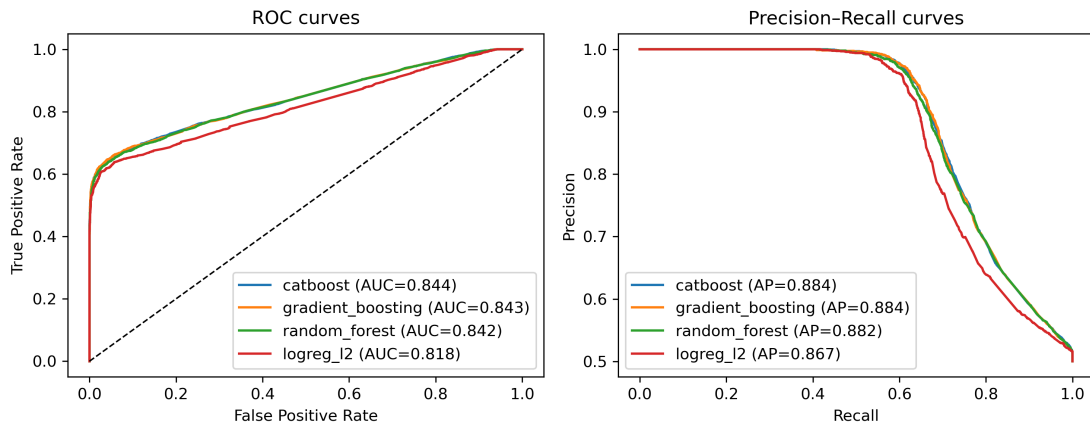


Figure 4.6: ROC (left) and precision–recall (right) curves for the four representative models on the held-out test set.

## 918 4.5 Feature Importance

### 919 4.5.1 Permutation Importance of Individual Features

920 To understand how each classifier made predictions, feature importance was quan-  
921 tified using permutation importance. This analysis was applied to four represen-  
922 tative models: CatBoost, Gradient Boosting, Random Forest, and L<sub>2</sub>-regularized

923 Logistic Regression.

924 As shown in Figure 4.7, the total number of clipped bases consistently pro-  
925 vides a strong predictive signal, particularly in Random Forest, Gradient Boosting,  
926 and L<sub>2</sub>-regularized Logistic Regression. CatBoost differs by assigning the highest  
927 importance to k-mer divergence metrics such as `kmer_js_divergence`, which cap-  
928 ture subtle sequence changes resulting from structural variants or PCR-induced  
929 chimeras. Soft-clipping features (`softclip_left` and `softclip_right`) provide  
930 more information around breakpoints, complementing these primary signals in all  
931 models except Gradient Boosting. L<sub>2</sub>-regularized Logistic Regression relies more  
932 on alignment-based split-read metrics.

933 Overall, these results indicate that accurate detection of chimeric reads relies  
934 on both alignment-based signals and k-mer compositional information. Explicit  
935 microhomology features contribute minimally in this analysis, and combining both  
936 alignment-based and sequence-level features enhances model sensitivity and speci-  
937 ficity.

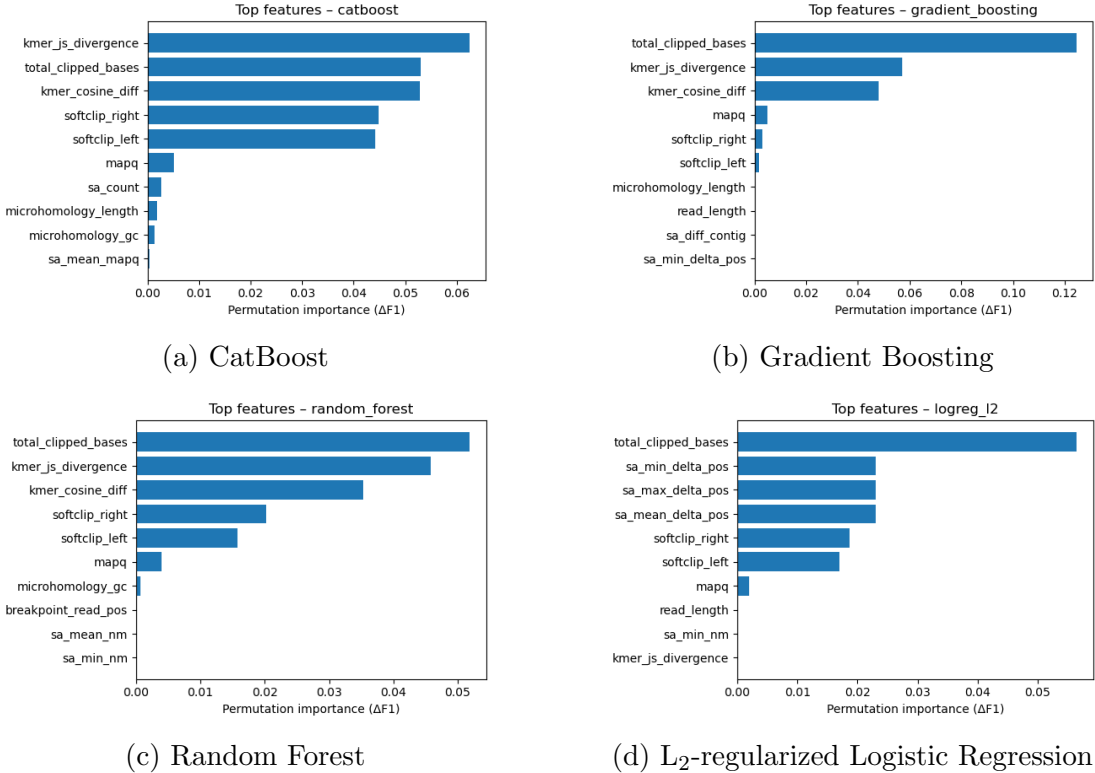


Figure 4.7: Permutation-based feature importance for four representative classifiers.

### 938 4.5.2 Feature Family Importance

939 To evaluate the contribution of broader signals, features were grouped into  
 940 five families: SA\_structure (supplementary alignment and segment met-  
 941 rics, e.g., has\_sa, sa\_count, sa\_min\_delta\_pos, sa\_mean\_nm, etc.), Clipping  
 942 (softclip\_left, softclip\_right, total\_clipped\_bases, breakpoint\_read\_pos),  
 943 Kmer\_jump (kmer\_cosine\_diff, kmer\_js\_divergence), Micro\_homology (microhomology\_length, microhomology\_gc), and Other (e.g., mapq).

945 Aggregated analyses reveal consistent patterns across models. In CatBoost,  
 946 the Clipping family has the largest cumulative contribution (0.14), followed

947 by Kmer\_jump (0.12), with Other features contributing minimally (0.005) and  
948 SA\_structure (0.003) and Micro\_homology (0.003) providing minimal predictive  
949 power. Gradient Boosting shows a similar trend, with Clipping (0.13) domi-  
950 nating, Kmer\_jump (0.11) secondary, and the remaining families contributing  
951 negligibly. Random Forest integrates both Clipping (0.088) and Kmer\_jump  
952 (0.08) effectively, while SA\_structure, Micro\_homology, and Other remain minor  
953 contributors. L<sub>2</sub>-regularized Logistic Regression emphasizes Clipping (0.09)  
954 and SA\_structure (0.07), with Kmer\_jump and Micro\_homology having minimal  
955 impact.

956 Both feature-level and aggregated analyses indicate that detection of chimeric  
957 reads in this dataset relies primarily on alignment irregularities (Clipping) and  
958 k-mer compositional shifts (Kmer\_jump), which often arise from PCR-induced  
959 template switching events, while explicit microhomology features contribute min-  
960 imally.

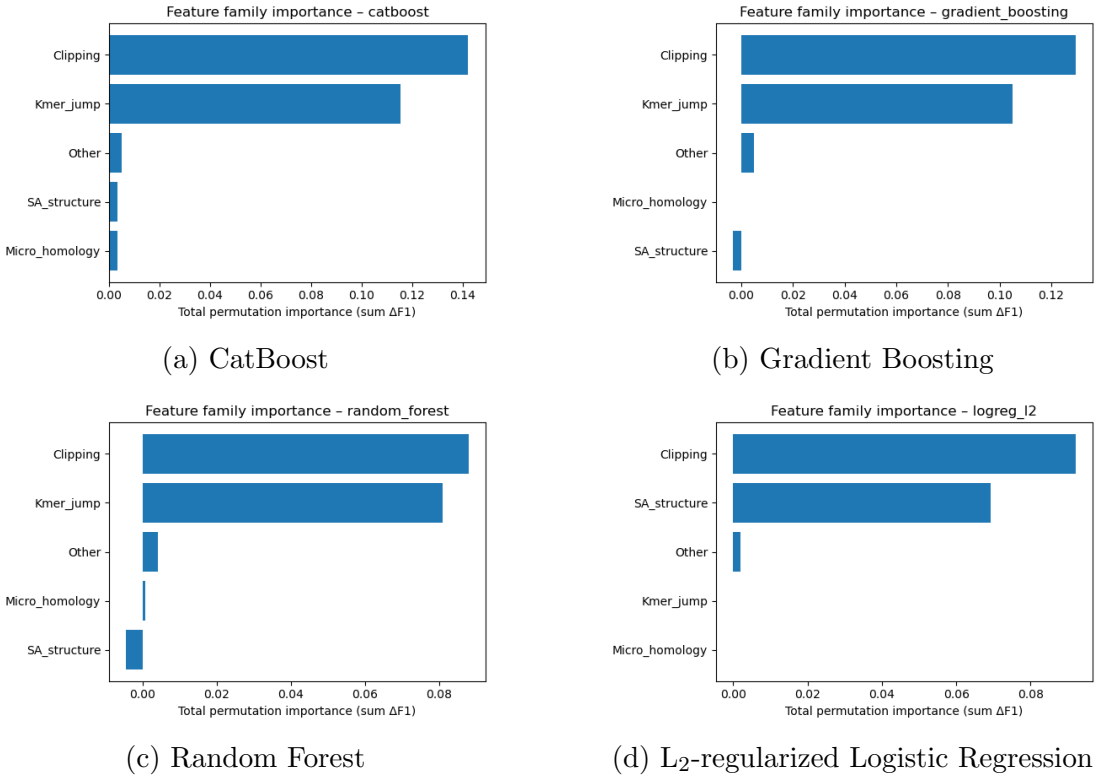


Figure 4.8: Aggregated feature family importance across four models.

## 961 4.6 Feature Selection

962 Feature selection was performed to identify the smallest subset reaching 95% cu-  
 963 mulative importance. Three models were evaluated as references: the full model  
 964 with all 23 features, a reduced model with the top- $k$  features, and an ablation  
 965 model excluding microhomology features, using a tuned CatBoost classifier to  
 966 assess feature contributions and overall classification performance.

### 967 4.6.1 Cumulative Importance Curve

968 The cumulative importance curve was computed using the tuned CatBoost clas-  
969 sifier. Figure 4.9 illustrates the contribution of features sorted by importance.  
970 The curve rises steeply for the first few features and then gradually plateaus,  
971 indicating that a small number of features capture most of the model’s pre-  
972 dictive power. A cumulative importance of 95% is reached at  $k = 4$  features,  
973 which are `total_clipped_bases`, `kmer_js_divergence`, `kmer_cosine_diff`, and  
974 `softclip_left`.

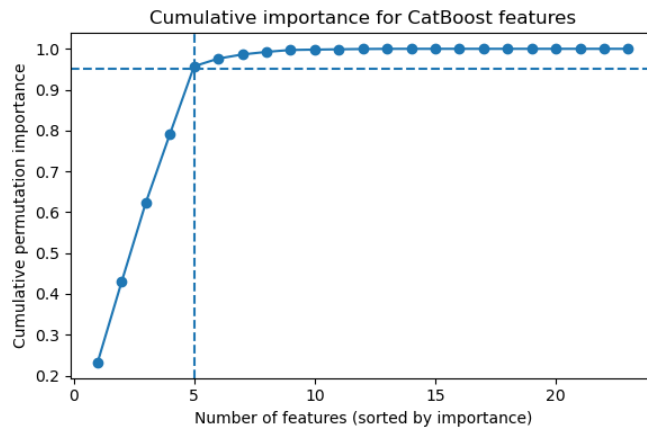


Figure 4.9: Cumulative importance curve of features sorted by importance.

### 975 4.6.2 Performance Comparison Across Feature Sets

976 Classification performance was compared across three feature sets using a tuned  
977 CatBoost classifier. The full model, incorporating all 23 engineered features,  
978 achieved an F1 score of approximately 0.7686 and a ROC–AUC of 0.8436.  
979 A reduced model using only the top four features (`total_clipped_bases`,  
980 `kmer_js_divergence`, `kmer_cosine_diff`, and `softclip_left`) achieved nearly

981 equivalent performance with an F1 of 0.7670 and a ROC–AUC of 0.8353. An  
 982 ablation model excluding microhomology features (`microhomology_length` and  
 983 `microhomology_gc`) also performed comparably, with an F1 of 0.7679 and ROC–  
 984 AUC of 0.8447. These results indicate that clipping and k-mer features capture  
 985 almost all predictive signal, while microhomology features are largely redundant  
 986 in this dataset.

Table 4.4: Test set performance of three feature set variants using tuned CatBoost.

Variant	No. of Features	Test F1	ROC–AUC
Full CatBoost	23	0.7686	0.8436
Selected (top-4)	4	0.7670	0.8353
No microhomology	21	0.7679	0.8447

987 Figure 4.10 presents a bar chart comparing F1 and ROC–AUC across the  
 988 three variants, with the x-axis showing the model variants and two bars per group  
 989 representing the F1 and ROC–AUC values.

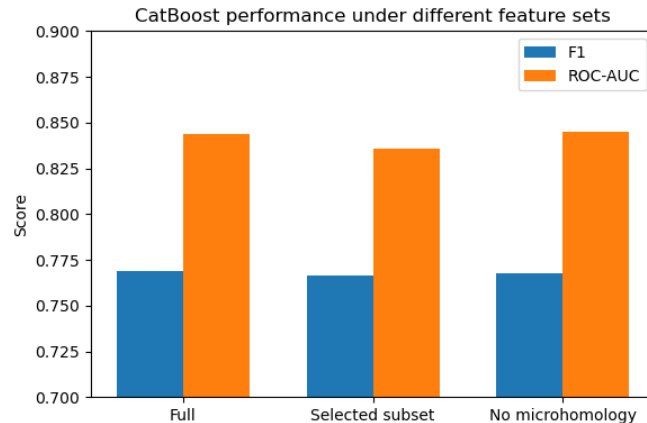


Figure 4.10: Comparison of F1 and ROC–AUC for the full, top-4 selected, and no-microhomology feature set variants.

990 **4.6.3 Interpretation and Final Feature Set Choice**

991 The full 23-feature model is retained as the primary configuration for the re-  
992 mainder of the study, while the four-feature subset serves as a lightweight al-  
993 ternative. Clipping features reflect alignment junctions and mapping disruptions  
994 typical of chimeric reads, and k-mer divergence captures changes in sequence com-  
995 position across breakpoints. Microhomology features appear largely redundant,  
996 as their signal is either indirectly represented by clipping and k-mer features or  
997 not strongly expressed in the simulation dataset.

998 **4.7 Summary of Findings**

999 All evaluated machine learning models substantially outperformed the dummy  
1000 baseline, demonstrating that the engineered feature set contains meaningful  
1001 signals for detecting PCR-induced chimeric reads. Across classifiers, the best-  
1002 performing models achieved test F1-scores of approximately 0.77 and ROC–AUC  
1003 values around 0.84 on held-out simulated mitochondrial reads, indicating reli-  
1004 able discrimination between clean and chimeric sequences. Among the tested  
1005 approaches, tree-based ensemble and boosting methods consistently showed the  
1006 strongest and most stable performance. In particular, CatBoost and Gradient  
1007 Boosting ranked among the top models across multiple evaluation metrics,  
1008 both before and after hyperparameter tuning. These results suggest that non-  
1009 linear ensemble methods are well suited to capturing the interaction between  
1010 alignment-derived and sequence-derived features in this setting.

1011 Analysis of feature behaviour revealed clear differences in how effectively fea-

ture groups distinguished clean and chimeric reads. Alignment- and clipping-based features, such as soft-clipping measures and total clipped bases, showed strong separation between clean and chimeric reads and emerged as the most informative signals. K-mer divergence features provided additional but weaker separation, contributing complementary information beyond alignment irregularities. In contrast, microhomology features and several supplementary alignment (SA) structure metrics exhibited minimal class separation and contributed little to overall predictive performance.

Feature selection results further supported these observations. A reduced subset of four features, dominated by clipping-based and k-mer divergence metrics, achieved nearly identical performance to the full 23-feature model. Moreover, removing explicit microhomology features did not degrade performance and in some cases resulted in slightly improved metrics, suggesting that these features are largely redundant under the simulated conditions tested.

Overall, these findings suggest that alignment-based and k-mer-based features provide sufficient signal to detect PCR-induced chimeric reads in simulated mitochondrial data, supporting the use of a compact and interpretable machine learning approach as a pre-assembly chimera detection step.

<sup>1030</sup> **Appendix A**

<sup>1031</sup> **Complete Per-Class Summary**

<sup>1032</sup> **Statistics**

Table A.1: Complete per-class summary statistics for all extracted features.

Feature	Class	Mean	Std	Median	Q1	Q3	IQR	Min	Max	n
breakpoint_read_pos	chimeric	75.000	0.000	75.000	75.000	75.000	0.000	75.000	75.000	20000
breakpoint_read_pos	clean	75.000	0.000	75.000	75.000	75.000	0.000	75.000	75.000	19983
has_sa	chimeric	0.406	0.491	0.000	0.000	1.000	1.000	0.000	1.000	20000
has_sa	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
kmer_cosine_diff	chimeric	0.974	0.026	0.986	0.958	1.000	0.042	0.817	1.000	20000
kmer_cosine_diff	clean	0.976	0.025	0.986	0.959	1.000	0.041	0.814	1.000	19983
kmer_js_divergence	chimeric	0.974	0.025	0.986	0.957	1.000	0.043	0.811	1.000	20000
kmer_js_divergence	clean	0.976	0.025	0.986	0.959	1.000	0.040	0.817	1.000	19983
mapq	chimeric	59.987	0.355	60.000	60.000	60.000	0.000	43.000	60.000	20000
mapq	clean	59.663	2.036	60.000	60.000	60.000	0.000	0.000	60.000	19983
mean_base_quality	chimeric	40.000	0.000	40.000	40.000	40.000	0.000	40.000	40.000	20000
mean_base_quality	clean	13.000	0.000	13.000	13.000	13.000	0.000	13.000	13.000	19983
microhomology_gc	chimeric	0.172	0.361	0.000	0.000	0.000	0.000	0.000	1.000	20000
microhomology_gc	clean	0.172	0.361	0.000	0.000	0.000	0.000	0.000	1.000	19983
microhomology_length	chimeric	0.458	0.755	0.000	0.000	1.000	1.000	0.000	5.000	20000
microhomology_length	clean	0.462	0.758	0.000	0.000	1.000	1.000	0.000	5.000	19983

Continued on next page

Feature	Class	Mean	Std	Median	Q1	Q3	IQR	Min	Max	n
num_segments	chimeric	1.406	0.491	1.000	1.000	2.000	1.000	1.000	2.000	20000
num_segments	clean	1.000	0.000	1.000	1.000	1.000	0.000	1.000	1.000	19983
read_length	chimeric	150.000	0.000	150.000	150.000	150.000	0.000	150.000	150.000	20000
read_length	clean	150.000	0.000	150.000	150.000	150.000	0.000	150.000	150.000	19983
ref_start_1based	chimeric	8428.635	4248.348	8433.000	5013.000	11786.250	6773.250	1.000	16521.000	20000
ref_start_1based	clean	8200.121	4626.918	8240.000	3639.000	11565.000	7926.000	1.000	16521.000	19983
sa_count	chimeric	0.406	0.491	0.000	0.000	1.000	1.000	0.000	1.000	20000
sa_count	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_diff_contig	chimeric	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	20000
sa_diff_contig	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_max_delta_pos	chimeric	1573.531	2364.996	0.000	0.000	2826.250	2826.250	0.000	16519.000	20000
sa_max_delta_pos	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_max_mapq	chimeric	14.104	21.424	0.000	0.000	27.000	27.000	0.000	60.000	20000
sa_max_mapq	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_mean_delta_pos	chimeric	1573.531	2364.996	0.000	0.000	2826.250	2826.250	0.000	16519.000	20000
sa_mean_delta_pos	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_mean_mapq	chimeric	14.104	21.424	0.000	0.000	27.000	27.000	0.000	60.000	20000
sa_mean_mapq	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983

Continued on next page

Feature	Class	Mean	Std	Median	Q1	Q3	IQR	Min	Max	n
sa_mean_nm	chimeric	0.022	0.319	0.000	0.000	0.000	0.000	0.000	6.000	20000
sa_mean_nm	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_min_delta_pos	chimeric	1573.531	2364.996	0.000	0.000	2826.250	2826.250	0.000	16519.000	20000
sa_min_delta_pos	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_min_nm	chimeric	0.022	0.319	0.000	0.000	0.000	0.000	0.000	6.000	20000
sa_min_nm	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_opp_strand_count	chimeric	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	20000
sa_opp_strand_count	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
sa_same_strand_count	chimeric	0.406	0.491	0.000	0.000	1.000	1.000	0.000	1.000	20000
sa_same_strand_count	clean	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	19983
softclip_left	chimeric	12.546	21.898	0.000	0.000	19.000	19.000	0.000	150.000	20000
softclip_left	clean	0.225	1.543	0.000	0.000	0.000	0.000	0.000	56.000	19983
softclip_right	chimeric	12.896	22.123	0.000	0.000	19.000	19.000	0.000	150.000	20000
softclip_right	clean	0.212	1.513	0.000	0.000	0.000	0.000	0.000	55.000	19983
total_clipped_bases	chimeric	25.442	25.481	19.000	0.000	48.000	48.000	0.000	150.000	20000
total_clipped_bases	clean	0.437	2.157	0.000	0.000	0.000	0.000	0.000	110.000	19983

1034 **Appendix B**

1035 **Boxplots for All Numeric**

1036 **Features by Feature Family**

1037 **B.0.1 SA Structure (Supplementary Alignment and Seg-**  
1038 **ment Metrics)**

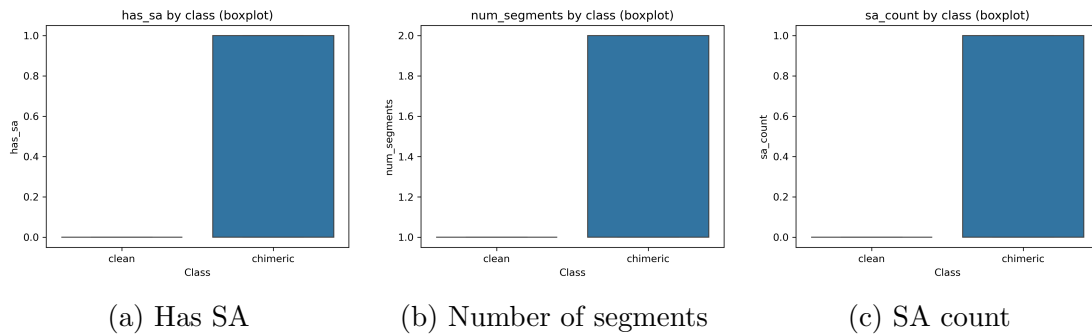


Figure B.1: Boxplots of SA Structure features by class (1/2).

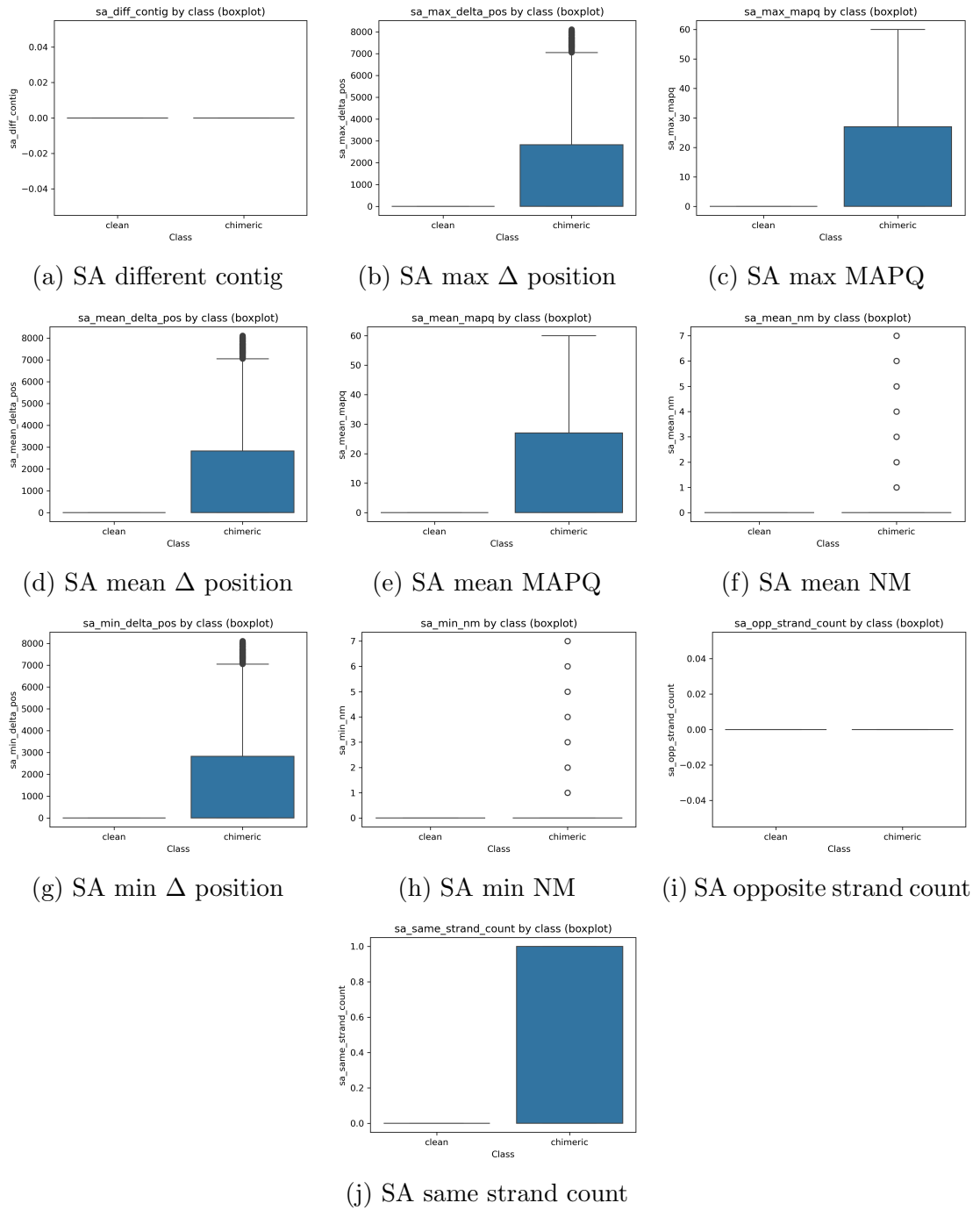


Figure B.2: Boxplots of SA Structure features by class (2/2).

1039 **B.0.2 Clipping-Based Features**

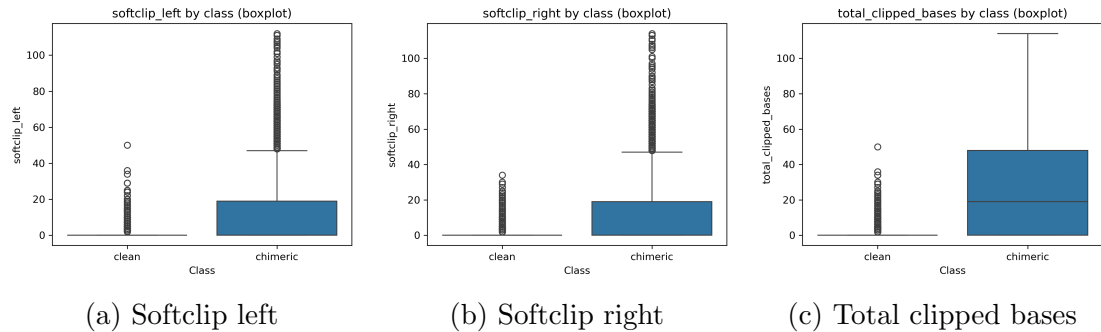


Figure B.3: Boxplots of clipping-based features by class.

1040 **B.0.3 K-mer Features**



Figure B.4: Boxplots of k-mer features by class.

1041 **B.0.4 Microhomology Features**



Figure B.5: Boxplots of microhomology features by class.

1042 **B.0.5 Others**

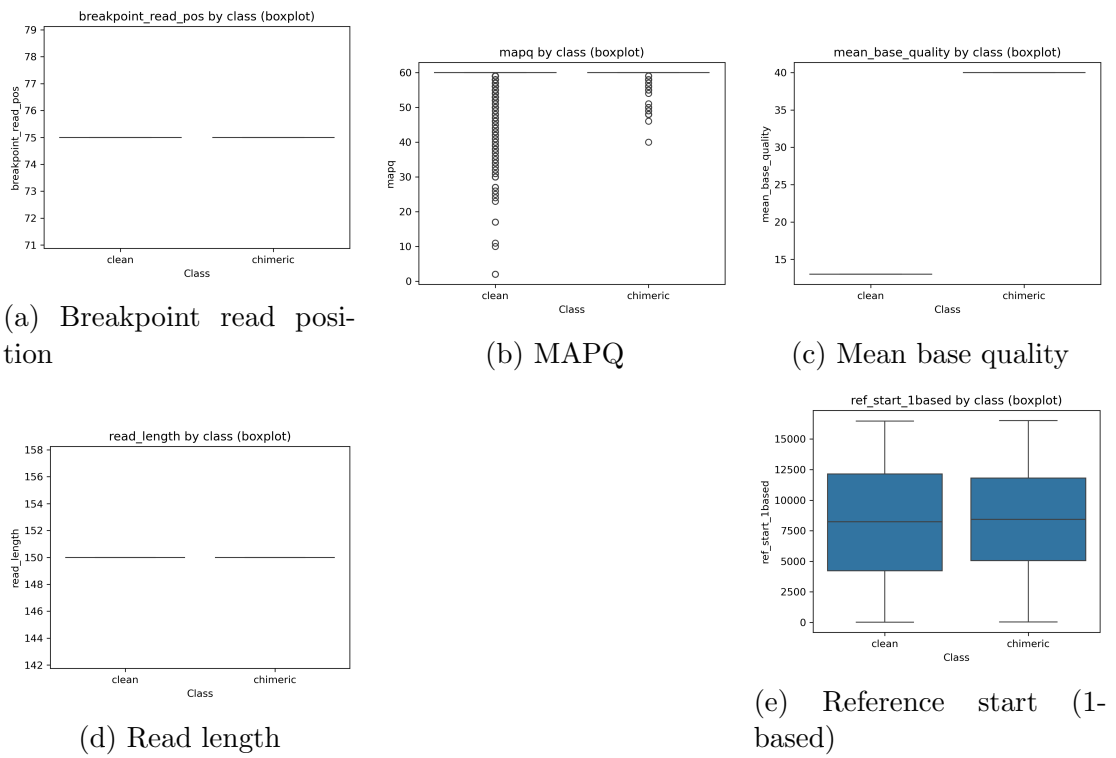


Figure B.6: Boxplots of other numeric features by class.

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