

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

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⁵⁶ **Chapter 1**

⁵⁷ **Introduction**

⁵⁸ **1.1 Overview**

⁵⁹ The rapid advancement of next-generation sequencing (NGS) technologies has
⁶⁰ transformed genomic research by enabling high-throughput and cost-effective
⁶¹ DNA analysis (Metzker, 2010). Among current platforms, Illumina sequencing
⁶² remains the most widely adopted, capable of producing millions of short reads
⁶³ that can be assembled into reference genomes or analyzed for genetic variation
⁶⁴ (Bentley et al., 2008; Glenn, 2011). Despite its high base-calling accuracy,
⁶⁵ Illumina sequencing is prone to artifacts introduced during library preparation,
⁶⁶ particularly polymerase chain reaction (PCR)-induced chimeras, which are ar-
⁶⁷ tificial hybrid sequences that do not exist in the true genome (Judo, Wedel, &
⁶⁸ Wilson, 1998).

⁶⁹ 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 and filtering 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 and filter PCR-induced chimeric reads using both alignment-based and sequence-derived statistical features. The tool aims to provide bioinformatics laboratories, partic-

95 ularly the Philippine Genome Center Visayas (PGC Visayas), with an efficient
96 solution for improving mitochondrial genome reconstruction.

97 1.2 Problem Statement

98 While NGS technologies have revolutionized genomic data acquisition, the ac-
99 curacy of mitochondrial genome assembly remains limited by artifacts produced
100 during PCR amplification. These chimeric reads can distort assembly graphs and
101 cause misassemblies, with particularly severe effects in small, circular mitochon-
102 drial genomes (Boore, 1999; Cameron, 2014). Existing assembly pipelines such
103 as GetOrganelle, MITObim, and NOVOPlasty assume that sequencing reads are
104 free of such artifacts (Dierckxsens et al., 2017; Hahn et al., 2013; Jin et al., 2020).
105 At PGC Visayas, several mitochondrial assemblies have failed or yielded incom-
106 plete contigs despite sufficient coverage, suggesting that undetected chimeric reads
107 compromise assembly reliability. Meanwhile, existing chimera detection tools such
108 as UCHIME and VSEARCH were developed primarily for amplicon-based com-
109 munity analysis and rely heavily on reference or taxonomic comparisons (Edgar,
110 Haas, Clemente, Quince, & Knight, 2011; Rognes, Flouri, Nichols, Quince, &
111 Mahé, 2016). These approaches are unsuitable for single-species organellar data,
112 where complete reference genomes are often unavailable. Therefore, there is a
113 pressing need for a reference-independent, data-driven tool capable of detecting
114 and filtering PCR-induced chimeras in mitochondrial sequencing datasets.

₁₁₅ **1.3 Research Objectives**

₁₁₆ **1.3.1 General Objective**

₁₁₇ This study aims to develop and evaluate a machine learning-based pipeline (Mi-
₁₁₈ toChime) that detects PCR-induced chimeric reads in *Sardinella lemuru* mito-
₁₁₉ chondrial sequencing data in order to improve the quality and reliability of down-
₁₂₀ stream mitochondrial genome assemblies.

₁₂₁ **1.3.2 Specific Objectives**

₁₂₂ Specifically, the study aims to:

- ₁₂₃ 1. construct simulated *Sardinella lemuru* Illumina paired-end datasets contain-
₁₂₄ ing both clean and PCR-induced chimeric reads,
- ₁₂₅ 2. extract alignment-based and sequence-based features such as k-mer compo-
₁₂₆ sition, junction complexity, and split-alignment counts from both clean and
₁₂₇ chimeric reads,
- ₁₂₈ 3. train, validate, and compare supervised machine-learning models for classi-
₁₂₉ fying reads as clean or chimeric,
- ₁₃₀ 4. determine feature importance and identify indicators of PCR-induced
₁₃₁ chimerism,
- ₁₃₂ 5. integrate the optimized classifier into a modular and interpretable pipeline
₁₃₃ deployable on standard computing environments at PGC Visayas.

134 1.4 Scope and Limitations of the Research

135 This study focuses on detecting PCR-induced chimeric reads in Illumina paired-
136 end mitochondrial sequencing data from *Sardinella lemuru*. The decision to re-
137 strict the taxonomic scope to a single species is based on four considerations:
138 (1) to limit interspecific variation in mitochondrial genome size, GC content, and
139 repetitive regions so that differences in read patterns can be attributed more di-
140 rectly to PCR-induced chimerism; (2) to align the analysis with relevant *S. lemuru*
141 sequencing projects at PGC Visayas; (3) to take advantage of the availability of *S.*
142 *lemuru* mitochondrial assemblies and raw datasets in public repositories such as
143 the National Center for Biotechnology Information (NCBI), which facilitates refer-
144 ence selection and benchmarking; and (4) to develop a tool that directly supports
145 local studies on *S. lemuru* population structure and fisheries management.

146 The study emphasizes `wgsim`-based simulations and selected empirical mito-
147 chondrial datasets from *S. lemuru*. It excludes naturally occurring chimeras, nu-
148 clear mitochondrial pseudogenes (NUMTs), and large-scale assembly rearrange-
149 ments in nuclear genomes. Feature extraction is restricted to low-dimensional
150 alignment and sequence statistics, such as k-mer frequency profiles, GC content,
151 read length, soft and hard clipping metrics, split-alignment counts, and map-
152 ping quality, rather than high-dimensional deep learning embeddings. This de-
153 sign keeps model behaviour interpretable and ensures that the pipeline can be
154 run on standard workstations at PGC Visayas. Testing on long-read platforms
155 (e.g., Nanopore, PacBio) and other taxa is outside the scope of this project; the
156 implemented pipeline is evaluated only on short-read *S. lemuru* datasets.

¹⁵⁷ 1.5 Significance of the Research

¹⁵⁸ This research provides both methodological and practical contributions to mi-
¹⁵⁹tochondrial genomics and bioinformatics. First, MitoChime filters PCR-induced
¹⁶⁰ chimeric reads prior to genome assembly, with the goal of improving the con-
¹⁶¹tiguity and correctness of *Sardinella lemuru* mitochondrial assemblies. Second,
¹⁶² it replaces informal manual curation with a documented workflow, improving au-
¹⁶³tomation and reproducibility. Third, the pipeline is designed to run on computing
¹⁶⁴ infrastructures commonly available in regional laboratories, enabling routine use
¹⁶⁵ at facilities such as PGC Visayas. Finally, more reliable mitochondrial assemblies
¹⁶⁶ for *S. lemuru* provide a stronger basis for downstream applications in the field of
¹⁶⁷ fisheries and genomics.

¹⁶⁸ **Chapter 2**

¹⁶⁹ **Review of Related Literature**

¹⁷⁰ This chapter presents an overview of the literature relevant to the study. It
¹⁷¹ discusses the biological and computational foundations underlying mitochondrial
¹⁷² genome analysis and assembly, as well as existing tools, algorithms, and techniques
¹⁷³ related to chimera detection and genome quality assessment. The chapter aims to
¹⁷⁴ highlight the strengths, limitations, and research gaps in current approaches that
¹⁷⁵ motivate the development of the present study.

¹⁷⁶ **2.1 The Mitochondrial Genome**

¹⁷⁷ Mitochondrial genome (mtDNA) is a small, typically circular molecule found in
¹⁷⁸ most eukaryotes. It encodes essential genes involved in oxidative phosphorylation
¹⁷⁹ and energy metabolism. Because of its conserved structure and maternal inher-
¹⁸⁰ itance, mtDNA has become a valuable genetic marker for studies in evolution,
¹⁸¹ population genetics, and phylogenetics (Anderson et al., 1981; Boore, 1999). In

182 animal species, the mitochondrial genome ranges from 15–20 kilobase and contains
183 13 protein-coding genes, 22 tRNAs, and two rRNAs arranged compactly without
184 introns (Gray, 2012). In comparison to nuclear DNA the ratio of the number
185 of copies of mtDNA is higher and has relatively simple organization which make
186 it particularly suitable for genome sequencing and assembly studies (Dierckxsens
187 et al., 2017). Moreover, mitochondrial genomes provide crucial insights into evo-
188 lutionary relationships among species and are increasingly used for testing new
189 genomic assembly and analysis methods.

190 **2.1.1 Mitochondrial Genome Assembly**

191 Mitochondrial genome assembly refers to the reconstruction of the complete mito-
192 chondrial DNA (mtDNA) sequence from raw or fragmented sequencing reads. It is
193 conducted to obtain high-quality, continuous representations of the mitochondrial
194 genome that can be used for a wide range of analyses, including species identi-
195 fication, phylogenetic reconstruction, evolutionary studies, and investigations of
196 mitochondrial diseases. Because mtDNA evolves relatively rapidly and is mater-
197 nally inherited, its assembled sequence provides valuable insights into population
198 structure, lineage divergence, and adaptive evolution across taxa (Boore, 1999).
199 Compared to nuclear genome assembly, assembling the mitochondrial genome is
200 often considered more straightforward but still encounters distinct technical chal-
201 lenges such as sequencing errors, low coverage regions, and chimeric reads that can
202 distort the final assembly, leading to incomplete or misassembled genomes. These
203 errors can propagate into downstream analyses, emphasizing the need for robust
204 chimera detection and sequence validation methods in mitochondrial genome re-

205 search.

206 2.2 PCR Amplification and Chimera Formation

207 Polymerase Chain Reaction (PCR) plays an important role in next-generation
208 sequencing (NGS) library preparation, as it amplifies target DNA fragments for
209 downstream analysis. However, the amplification process can also introduce arti-
210 facts that affect data accuracy, one of them being the formation of chimeric se-
211 quences. Chimeras typically arise when incomplete extension occurs during a PCR
212 cycle. This causes the DNA polymerase to switch from one template to another
213 and generate hybrid recombinant molecules (Judo et al., 1998). Artificial chimeras
214 are produced through such amplification errors, whereas biological chimeras oc-
215 cur naturally through genomic rearrangements or transcriptional events. These
216 biological chimeras can have functional roles and may encode tissue-specific novel
217 proteins that link to cellular processes or diseases (Frenkel-Morgenstern et al.,
218 2012).

219 In the context of amplicon-based sequencing, PCR-induced chimeras can sig-
220 nificantly distort analytical outcomes. Their presence artificially inflates estimates
221 of genetic or microbial diversity and may cause misassemblies during genome re-
222 construction. (Qin et al., 2023) has reported that chimeric sequences may account
223 for more than 10% of raw reads in amplicon datasets. This artifact tends to be
224 most prominent among rare operational taxonomic units (OTUs) or singletons,
225 which are sometimes misinterpreted as novel diversity, which further causes the
226 complication of microbial diversity analyses (Gonzalez, Zimmermann, & Saiz-

227 Jimenez, 2004). Moreover, the likelihood of chimera formation has been found to
228 vary with the GC content of target sequences, with lower GC content generally
229 associated with a reduced rate of chimera generation (Qin et al., 2023).

230 **2.2.1 Effects of Chimeric Reads on Organelle Genome As-**
231 **sembly**

232 In mitochondrial DNA (mtDNA) assembly workflows, PCR-induced chimeras pose
233 additional challenges. Assembly tools such as GetOrganelle and MitoBeam, which
234 operate under the assumption of organelle genome circularity, are vulnerable when
235 chimeric reads disrupt this circular structure. Such disruptions can lead to assem-
236 bly errors or misassemblies (Bi et al., 2024). These artificial sequences interfere
237 with the assembly graph, which makes it more difficult to accurately reconstruct
238 mitochondrial genomes. In addition, these artifacts propagate false variants and
239 erroneous annotations in genomic data. Hence, determining and minimizing PCR-
240 induced chimera formation is vital for improving the quality of mitochondrial
241 genome assemblies, and ensuring the reliability of amplicon sequencing data.

242 2.3 Existing Traditional Approaches for Chimera

243 Detection

244 Several computational tools have been developed to identify chimeric sequences in
245 NGS datasets. These tools generally fall into two categories: reference-based and
246 de novo approaches. Reference-based chimera detection, also known as database-
247 dependent detection, is one of the earliest and most widely used computational
248 strategies for identifying chimeric sequences in amplicon-based microbial commu-
249 nity studies. These methods rely on the comparison of each query sequence against
250 a curated, high-quality database of known, non-chimeric reference sequences to
251 determine whether the query can be more plausibly explained as a composite or
252 a mosaic of two or more reference sequences rather than as a genuine biological
253 variant (Edgar et al., 2011).

254 On the other hand, the de novo chimera detection, also referred to as reference-
255 free detection, represents an alternative computational paradigm that identifies
256 chimeric sequences without reliance on external reference databases. Instead of
257 comparing each query sequence to a curated collection of known, non-chimeric
258 sequences, de novo methods infer chimeras based on internal relationships among
259 the sequences present within the dataset itself. This approach is particularly
260 advantageous in studies of novel, under explored, or taxonomically diverse mi-
261 crobial communities where comprehensive reference databases are unavailable or
262 incomplete (Edgar, 2016; Edgar et al., 2011). The underlying assumption on this
263 method operates on the key biological principle that true biological sequences are
264 generally more abundant than chimeric artifacts. During PCR amplification, au-
265 thentic sequences are amplified early and tend to dominate the read pool, while

266 chimeric sequences form later resulting in the tendency to appear at lower relative
267 abundances compared to their true parental sequences. As such, the abundance
268 hierarchy is formed by treating the most abundant sequences as supposed parents
269 and testing whether less abundant sequences can be reconstructed as mosaics of
270 these dominant templates. In addition to abundance, de novo algorithms assess
271 compositional and structural similarity among sequences, examining whether cer-
272 tain regions of a candidate sequence align more closely with one high-abundance
273 sequence and other regions with a different one.

274 Both reference-based and de novo approaches are complementary rather than
275 mutually exclusive. Reference-based methods provide stability and reproducibility
276 when curated databases are available, whereas de novo methods offer flexibility
277 and independence for novel or highly diverse communities. In practice, many
278 modern bioinformatics pipelines combine both paradigms sequentially: an initial
279 de novo step identifies dataset-specific chimeras, followed by a reference-based pass
280 that removes remaining artifacts relative to established databases (Edgar, 2016).
281 These two methods of detection form the foundation of tools such as UCHIME
282 and later UCHIME2, exemplified by the dual capability of providing both modes
283 within a unified computational framework.

284 2.3.1 UCHIME

285 Developed by Edgar et al. (Edgar et al., 2011), UCHIME is one of the most widely
286 used computational tools for detecting chimeric sequences in amplicon sequencing
287 data. The UCHIME algorithm detects chimeras by evaluating how well a query
288 sequence (Q) can be explained as a mosaic of two parent sequences (A and B)

289 from a reference database. The query sequence is first divided into four non-
290 overlapping segments or chunks. Each chunk is independently searched against a
291 reference database that is assumed to be free of chimeras. The best matches to
292 each segment are collected, and from these results, two candidate parent sequences
293 are identified, typically the two sequences that best explain all chunks of the query.
294 Then a three-way alignment among the query (Q) and the two parent candidates
295 (A and B) is done. From this alignment, UCHIME attempts to find a chimeric
296 model (M) which is a hypothetical recombinant sequence formed by concatenating
297 fragments from A and B that best match the observed Q

298 Chimeric Alignment and Scoring

299 To decide whether a query is chimeric, UCHIME computes several alignment-
300 based metrics between Q, its top hit (T, the most similar known sequence), and
301 the chimeric model (M). The key differences are measured as: dQT or the number
302 of mismatches between the query and the top hit as well as dQM or the number
303 of mismatches between the query and the chimeric model. From these, a chimera
304 score is calculated to quantify how much better the chimeric model fits the query
305 compared to a single parent. If the model's similarity to Q exceeds a defined
306 threshold (typically $\geq 0.8\%$ better identity), the sequence is reported as chimeric.
307 A higher score indicates stronger evidence of chimerism, while lower scores suggest
308 that the sequence is more likely to be authentic.

309 In de novo mode, UCHIME applies an abundance-driven strategy. Only se-
310 quences at least twice as abundant as the query are considered as potential parents.
311 Non-chimeric sequences identified at each step are added iteratively to a growing

312 internal database for subsequent queries.

313 **Limitations of UCHIME**

314 Although UCHIME was a significant advancement in chimera detection, it has
315 notable limitations. According to (Edgar, 2016) and the UCHIME practical notes
316 (Edgar, n.d), many of the accuracy results reported in the original 2011 paper
317 were overly optimistic due to unrealistic benchmark designs that assumed com-
318 plete reference coverage and perfect sequence quality. In practice, UCHIME's
319 accuracy can decline when (1) the reference database is incomplete or contains
320 erroneous entries; (2) low-divergence chimeras are present, as these closely resem-
321 ble genuine biological variants; (3) sequence datasets include residual sequencing
322 errors, leading to spurious alignments or misidentification; and (4) the abundance
323 ratio between parent and chimera is distorted by amplification bias. Additionally,
324 UCHIME tends to misclassify sequences as non-chimeric when parent sequences
325 are missing from the database. These limitations motivated the development of
326 UCHIME2.

327 **2.3.2 UCHIME2**

328 To overcome the limitations of its predecessor, UCHIME2 (Edgar, 2016) intro-
329 duced several methodological and algorithmic refinements that significantly en-
330 hanced the accuracy and reliability of chimera detection. One major improve-
331 ment lies in its approach to uncertainty handling. In earlier versions, sequences
332 with limited reference support were often incorrectly classified as non-chimeric,

333 increasing the likelihood of false negatives. UCHIME2 addresses this issue by
334 designating such ambiguous sequences as “unknown,” thereby providing a more
335 conservative and reliable classification framework.

336 Another notable advancement is the introduction of multiple application-
337 specific modes that allow users to tailor the algorithm’s performance to the
338 characteristics of their datasets. The following parameter presets: denoised,
339 balanced, sensitive, specific, and high-confidence, enable researchers to optimize
340 the balance between sensitivity and specificity according to the goals of their
341 analysis.

342 In comparative evaluations, UCHIME2 demonstrated superior detection per-
343 formance, achieving sensitivity levels between 93% and 99% and lower overall
344 error rates than earlier versions or other contemporary tools such as DECIPHER
345 and ChimeraSlayer. Despite these advances, the study also acknowledged a fun-
346 damental limitation in chimera detection: complete error-free identification is
347 theoretically unattainable. This is due to the presence of “perfect fake models,”
348 wherein genuine non-chimeric sequences can be perfectly reconstructed from other
349 reference fragments. This underscore the uncertainty in differentiating authentic
350 biological sequences from artificial recombinants based solely on sequence similar-
351 ity, emphasizing the need for continued methodological refinement and cautious
352 interpretation of results.

353 **2.3.3 CATch**

354 Early chimera detection programs such as UCHIME (Edgar et al., 2011) relied on
355 alignment-based and abundance-based heuristics to identify hybrid sequences in
356 amplicon data. However, researchers soon observed that different algorithms often
357 produced inconsistent predictions. A sequence might be identified as chimeric by
358 one tool but classified as non-chimeric by another, resulting in unreliable filtering
359 outcomes across studies.

360 To address these inconsistencies, (Mysara, Saeys, Leys, Raes, & Monsieurs,
361 2015) developed the Classifier for Amplicon Tool Chimeras (CATCh), which rep-
362 resents the first ensemble machine learning system designed for chimera detection
363 in 16S rRNA amplicon sequencing. Rather than depending on a single detec-
364 tion strategy, CATCh integrates the outputs of several established tools, includ-
365 ing UCHIME, ChimeraSlayer, DECIPHER, Pintail, and Perseus. The individual
366 scores and binary decisions generated by these tools are used as input features for
367 a supervised learning model. The algorithm employs a Support Vector Machine
368 (SVM) with a Pearson VII Universal Kernel (PUK) to determine optimal weight-
369 ings among the input features and to assign each sequence a probability of being
370 chimeric.

371 Benchmarking in both reference-based and de novo modes demonstrated signif-
372 icant performance improvements. CATCh achieved sensitivities of approximately
373 85 percent in reference-based mode and 92 percent in de novo mode, with corre-
374 sponding specificities of approximately 96 percent and 95 percent. These results
375 indicate that CATCh detected 7 to 12 percent more chimeras than any individual
376 algorithm while maintaining high precision. Integration of CATCh into amplicon-

377 processing pipelines also reduced operational taxonomic unit (OTU) inflation by
378 23 to 35 percent, producing diversity estimates that more closely reflected true
379 community composition.

380 2.3.4 ChimPipe

381 Among the available tools for chimera detection, ChimPipe is a bioinformatics
382 pipeline developed to identify chimeric sequences such as fusion genes and
383 transcription-induced chimeras from paired-end RNA sequencing data. It uses
384 both discordant paired-end reads and split-read alignments to improve the ac-
385 curacy and sensitivity of detecting fusion genes, trans-splicing events, and read-
386 through transcripts (Rodriguez-Martin et al., 2017). By combining these two
387 sources of information, ChimPipe achieves better precision than methods that
388 depend on a single type of signal.

389 The pipeline works with many eukaryotic species that have available genome
390 and annotation data, making it a versatile tool for studying chimera evolution
391 and transcriptome structure (Rodriguez-Martin et al., 2017). It can also predict
392 multiple isoforms for each gene pair and identify breakpoint coordinates that are
393 useful for reconstructing and verifying chimeric transcripts. Tests using both
394 simulated and real datasets have shown that ChimPipe maintains high accuracy
395 and reliable performance.

396 ChimPipe’s modular design lets users adjust parameters to fit different se-
397 quencing protocols or organism characteristics. Experimental results have con-
398 firmed that many chimeric transcripts detected by the tool correspond to func-

399 tional fusion proteins, showing its value for understanding chimera biology and
400 its potential applications in disease research (Rodriguez-Martin et al., 2017).

401 **2.4 Machine Learning Approaches for Chimera 402 and Sequence Quality Detection**

403 Traditional chimera detection tools rely primarily on heuristic or alignment-based
404 rules. Recent advances in machine learning (ML) have demonstrated that mod-
405 els trained on sequence-derived features can effectively capture compositional and
406 structural patterns in biological sequences. Although most existing ML systems
407 such as those used for antibiotic resistance prediction, taxonomic classification,
408 or viral identification are not specifically designed for chimera detection, they
409 highlight how data-driven models can outperform similarity-based heuristics by
410 learning intrinsic sequence signatures. In principle, ML frameworks can inte-
411 grate diverse indicators such as k-mer frequencies, GC-content variation, and
412 split-alignment metrics to identify subtle anomalies that may indicate a chimeric
413 origin (Arango et al., 2018; Liang, Bible, Liu, Zou, & Wei, 2020; Ren et al., 2020).

414 **2.4.1 Feature-Based Representations of Genomic Se- 415 quences**

416 In genomic analysis, feature extraction converts DNA sequences into numerical
417 representations suitable for ML algorithms. A common approach is k-mer fre-
418 quency analysis, where normalized k-mer counts form the feature vector (Vervier,

419 2015). These features effectively capture local compositional patterns that often
420 differ between authentic and chimeric reads. In particular, deviations in k-mer
421 profiles between adjacent read segments can serve as a compositional signature
422 of template-switching events. Additional descriptors such as GC content and
423 sequence entropy can further distinguish sequence types; in metagenomic classifi-
424 cation and virus detection, k-mer-based features have shown strong performance
425 and robustness to noise (Ren et al., 2020; Vervier, 2015). For chimera detection
426 specifically, abrupt shifts in GC or k-mer composition along a read can indicate
427 junctions between parental fragments. Windowed feature extraction enables mod-
428 els to capture these discontinuities that rule-based algorithms may overlook.

429 Machine learning models can also leverage alignment-derived features such as
430 the frequency of split alignments, variation in mapping quality, and local cover-
431 age irregularities. Split reads and discordant read pairs are classical signatures
432 of genomic junctions and have been formalized in probabilistic frameworks for
433 structural-variant discovery that integrate multiple evidence types (Layer, Hall, &
434 Quinlan, 2014). Similarly, long-read tools such as Sniffles employ split-alignment
435 and coverage anomalies to accurately localize breakpoints (Sedlazeck et al., 2018).
436 Modern aligners such as Minimap2 (Li, 2018) output supplementary (SA tags) and
437 secondary alignments as well as chaining and alignment-score statistics that can
438 be summarized into quantitative predictors for machine-learning models. These
439 alignment-signal features are particularly relevant to PCR-induced mitochondrial
440 chimeras, where template-switching events produce reads partially matching dis-
441 tinct regions of the same or related genomes. Integrating such cues within a
442 supervised-learning framework enables artifact detection even in datasets lacking
443 complete or perfectly assembled references.

444 A further biologically grounded descriptor is micro-homology length at puta-
445 tive junctions. Micro-homology refers to short, shared sequences (often in the
446 range of a few to tens of base pairs) that are near breakpoints and mediate
447 non-canonical repair or template-switch mechanisms. Studies of double strand
448 break repair and structural variation have demonstrated that the length of micro-
449 homology correlates with the likelihood of micro-homology-mediated end joining
450 (MMEJ) or fork-stalled template-switching pathways (Sfeir & Symington, 2015).
451 In the context of PCR-induced chimeras, template switching during amplifica-
452 tion often leaves short identical sequences at the junction of two concatenated
453 fragments. Quantifying the longest exact suffix–prefix overlap at each candidate
454 breakpoint thus provides a mechanistic signature of chimerism and complements
455 both compositional (k-mer) and alignment (SA count) features.

456 2.5 Synthesis of Chimera Detection Approaches

457 To provide an integrated overview of the literature discussed in this chapter, Ta-
458 ble 2.1 summarizes the major chimera detection studies, their methodological
459 approaches, and their known limitations. This consolidated comparison brings to-
460 gether reference-based approaches, de novo strategies, alignment-driven tools, en-
461 semble machine-learning systems, and general ML-based sequence-quality frame-
462 works. Presenting these methods side-by-side clarifies their performance bound-
463 aries and highlights the unresolved challenges that persist in mitochondrial genome
464 analysis and chimera detection.

Table 2.1: Summary of Existing Methods and Research Gaps

Method/Study	Scope/Approach	Limitations
Reference-based Chimera Detection	Compares query sequences against curated, non-chimeric reference databases; identifies mosaic sequences by evaluating similarity to known templates.	Depends heavily on completeness and quality of reference databases; often fails when novel taxa or missing parent sequences are present; reduced accuracy for low-divergence chimeras.
De novo Chimera Detection	Identifies chimeras using only internal dataset relationships; relies on abundance patterns and compositional similarity; reconstructs sequences as mosaics of high-abundance parents.	Assumes true sequences are more abundant—fails when amplification bias distorts abundance; struggles with evenly abundant parental sequences; can misclassify highly similar true variants.
UCHIME	Alignment-based chimera detection; segments query sequence, identifies parent candidates, performs 3-way alignment, and computes chimera scores; supports both reference-based and de novo modes.	Accuracy inflated in original benchmarks; suffers under incomplete databases; poor performance on low-divergence chimeras; sensitive to sequencing errors; misclassifies when parents are missing.
UCHIME2	Improved uncertainty handling; classifies ambiguous sequences as unknown; offers multiple sensitivity/specificity modes; more robust with incomplete references; higher sensitivity (93–99%).	Cannot achieve perfect accuracy due to “perfect fake models”; genuine variants may be indistinguishable from artificial recombinants; theoretical detection limit remains.
CATCh	First ML ensemble tool for 16S chimera detection; integrates outputs of UCHIME, ChimeraSlayer, DECIPHER, Pintail, Perseus via SVM classifier; significantly improves sensitivity and specificity.	Depends on performance of underlying tools; ML model limited to features they output; ensemble can still misclassify in datasets with extreme novelty or low coverage.
ChimPipe	Pipeline for detecting fusion genes and transcript-derived chimeras in	Designed for RNA-seq, not amplicons; needs high-quality genome

465 Across existing studies, no single approach reliably detects all forms of chimeric
466 sequences, particularly those generated by PCR-induced template switching in
467 mitochondrial genomes. Reference-based tools perform poorly when parental se-
468 quences are absent; de novo methods rely strongly on abundance assumptions;
469 alignment-based systems show reduced sensitivity to low-divergence chimeras; and
470 ensemble methods inherit the limitations of their component algorithms. RNA-
471 seq-oriented pipelines likewise do not generalize well to organelle data. Although
472 machine learning approaches offer promising feature-based detection, they are
473 rarely applied to mitochondrial genomes and are not trained specifically on PCR-
474 induced organelle chimeras. These limitations indicate a clear research gap: the
475 need for a specialized, feature-driven classifier tailored to mitochondrial PCR-
476 induced chimeras that integrates k-mer composition, split-alignment signals, and
477 micro-homology features to achieve more accurate detection than current heuristic
478 or alignment-based tools.

⁴⁷⁹ **Chapter 3**

⁴⁸⁰ **Research Methodology**

⁴⁸¹ **3.1 Research Activities**

⁴⁸² As illustrated in Figure 3.1, this study will carry out a sequence of computa-
⁴⁸³ tional procedures designed to detect PCR-induced chimeric reads in mitochon-
⁴⁸⁴ drial genomes. The process begins with the collection of mitochondrial reference
⁴⁸⁵ sequences from the NCBI database, which will serve as the foundation for gener-
⁴⁸⁶ ating simulated chimeric reads. These datasets will then undergo bioinformatics
⁴⁸⁷ pipeline development, which includes alignment, k-mer extraction, and homology-
⁴⁸⁸ based filtering to prepare the data for model construction. The machine-learning
⁴⁸⁹ model will subsequently be trained and tested using the processed datasets to
⁴⁹⁰ assess its accuracy and reliability. Depending on the evaluation results, the model
⁴⁹¹ will either be refined and retrained to improve performance or, if the metrics meet
⁴⁹² the desired threshold, deployed for further validation and application.

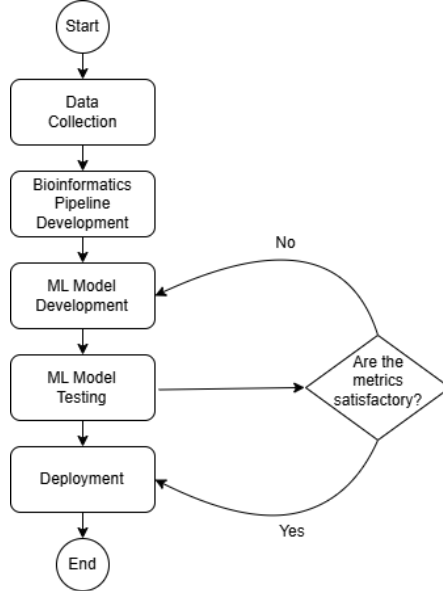


Figure 3.1: Process Diagram of Special Project

⁴⁹³ 3.1.1 Data Collection

⁴⁹⁴ The mitochondrial genome reference sequences of *Sardinella lemuru* will be col-
⁴⁹⁵ lected from the National Center for Biotechnology Information (NCBI) database.
⁴⁹⁶ The downloaded files will be in FASTA format to ensure compatibility with bioin-
⁴⁹⁷ formatics tools and subsequent analysis. The gathered sequences will serve as the
⁴⁹⁸ basis for generating simulated chimeric reads to be used in model development.

⁴⁹⁹ The expected outcome of this process is a comprehensive dataset of *Sardinella*
⁵⁰⁰ *lemuru* mitochondrial reference sequences that will serve as the foundation for
⁵⁰¹ the succeeding stages of the study. This step is scheduled to start in the first
⁵⁰² week of November 2025 and is expected to be completed by the last week of
⁵⁰³ November 2025, with a total duration of approximately one (1) month.

504 **Data Preprocessing**

505 Sequencing data will be simulated using the reference sequences collected from
506 NCBI. Using `wgsim`, a total of 10,000 paired-end reads (R1 and R2) will be gen-
507 erated from the reference genome and designated as clean reads. These reads will
508 be saved in FASTQ (`.fastq`) format. From the same reference, a Bash script will
509 be created to deliberately cut and reconnect portions of the sequence, introducing
510 artificial junctions that mimic chimeric regions. The manipulated reference file,
511 saved in FASTA (`.fasta`) format, will then be processed in `wgsim` to simulate
512 an additional 10,000 paired-end chimeric reads, also stored in FASTQ (`.fastq`)
513 format. The resulting read files will be aligned to the original reference genome
514 using SAMtools, generating SAM (`.sam`) or BAM (`.bam`) alignment files. During
515 this alignment process, clean reads will be labeled as “0,” while chimeric reads will
516 be labeled as “1” in a corresponding CSV (`.tsv`) file. This results in a balanced
517 dataset with an equal number of clean and chimeric reads. This is important to
518 prevent model bias and ensure that the machine learning classifiers can learn to
519 detect chimeras accurately.

520 The expected outcome of this process is a complete set of clean and chimeric
521 paired-end reads prepared for subsequent analysis and model development. This
522 step is scheduled to start in the first week of November 2025 and is expected
523 to be completed by the last week of November 2025, with a total duration of
524 approximately one (1) month.

525 3.1.2 Bioinformatics Tools Pipeline

526 A bioinformatics pipeline will be developed and implemented to extract the nec-
527 essary analytical features. This pipeline will serve as a reproducible and modular
528 workflow that accepts FASTQ and BAM inputs, processes these through a series
529 of analytical stages, and outputs tabular feature matrices (TSV) for downstream
530 machine learning. All scripts will be version-controlled through GitHub, and
531 computational environments will be standardized using Conda to ensure cross-
532 platform reproducibility. To promote transparency and replicability, the exact
533 software versions, parameters, and command-line arguments used in each stage
534 will be documented. To further ensure correctness and adherence to best practices,
535 bioinformatics experts at the Philippine Genome Center Visayas will be consulted
536 to validate the pipeline design, feature extraction logic, and overall data integrity.
537 This stage of the study is scheduled to begin in the last week of November 2025
538 and conclude by the last week of January 2026, with an estimated total duration
539 of approximately two (2) months.

540 The bioinformatics pipeline focuses on three principal features from the sim-
541 ulated and aligned sequencing data: (1) supplementary alignment count (SA
542 count), (2) k-mer composition difference between read segments, and (3) micro-
543 homology length at potential junctions. Each of these features captures a distinct
544 biological or computational signature associated with PCR-induced chimeras.

545 **Alignment and Supplementary Alignment Count**

546 This will be derived through sequence alignment using Minimap2, with subsequent
547 processing performed using SAMtools and `pysam` in Python. Sequencing reads
548 will be aligned to the *Sardinella lemuru* mitochondrial reference genome using
549 Minimap2 with the `-ax sr` preset (optimized for short reads). The output will
550 be converted and sorted using SAMtools, producing an indexed BAM file which
551 will be parsed using `pysam` to count the number of supplementary alignments
552 (SA tags) per read. Each read's mapping quality, number of split segments,
553 and alignment characteristics will be recorded in a corresponding TSV file. The
554 presence of multiple alignment loci within a single read, as reflected by a nonzero
555 SA count, serves as direct computational evidence of chimerism. Reads that
556 contain supplementary alignments or soft-clipped regions are strong candidates
557 for chimeric artifacts arising from PCR template switching or improper assembly
558 during sequencing.

559 **K-mer Composition Difference**

560 Chimeric reads often comprise fragments from distinct genomic regions, resulting
561 in a compositional discontinuity between segments. Comparing k-mer frequency
562 profiles between the left and right halves of a read allows detection of such abrupt
563 compositional shifts, independent of alignment information. This will be obtained
564 using Jellyfish, a fast k-mer counting software. For each read, the sequence will
565 be divided into two segments, either at the midpoint or at empirically determined
566 breakpoints inferred from supplementary alignment data, to generate left and right
567 sequence segments. Jellyfish will then compute k-mer frequency profiles (with $k =$

568 5 or 6) for each segment. The resulting k-mer frequency vectors will be normalized
569 and compared using distance metrics such as cosine similarity or Jensen–Shannon
570 divergence to quantify compositional disparity between the two halves of the same
571 read. The resulting difference scores will be stored in a structured TSV file.

572 **Micro-homology Length**

573 The micro-homology length will be computed using a custom Python script that
574 detects the longest exact suffix–prefix overlap within ± 30 base pairs surround-
575 ing a candidate breakpoint. This analysis identifies the number of consecutive
576 bases shared between the end of one segment and the beginning of another. The
577 presence and length of such micro-homology are classic molecular signatures of
578 PCR-induced template switching, where short identical regions (typically 3–15
579 base pairs) promote premature termination and recombination of DNA synthesis
580 on a different template strand. Quantifying micro-homology allows assessment of
581 whether the suspected breakpoint reflects PCR artifacts or true biological variants.
582 Each read will therefore be annotated with its corresponding micro-homology
583 length, overlap sequence, and GC content.

584 After extracting the three primary features, all resulting TSV files will be
585 joined using the read identifier as a common key to generate a unified feature ma-
586 trix. Additional read-level metadata such as read length, mean base quality, and
587 number of clipped bases will also be included to provide contextual information.
588 This consolidated dataset will serve as the input for subsequent machine-learning
589 model development and evaluation.

590 **3.1.3 Machine-Learning Model Development**

591 This study will explore multiple machine-learning approaches to detect PCR-
592 induced chimeras from mitochondrial Illumina reads: Support Vector Machines
593 (SVM) to separate reads with complex patterns, decision trees to capture hier-
594 archical interactions among SA count, k-mer composition, and micro-homology
595 length, logistic regression as a linear baseline, Random Forest (RF) to improve
596 stability and reduce variance, and gradient boosting (e.g., XGBoost) to model
597 non-linear relationships among the extracted features. Using these approaches
598 enables a balanced assessment of predictive performance and interpretability.

599 The dataset will be divided into training (80%) and testing (20%) subsets.
600 The training data will be used for model fitting and hyperparameter optimization
601 through five-fold cross-validation, in which the data are partitioned into five folds;
602 four folds are used for training and one for validation in each iteration. Perfor-
603 mance metrics will be averaged across folds, and the optimal parameters will be
604 selected based on mean cross-validation accuracy. The final models will then be
605 evaluated on the held-out test set to obtain unbiased performance estimates.

606 Model development and evaluation will be implemented in Python (ver-
607 sion 3.11) using the `scikit-learn` and `xgboost` libraries. Standard metrics
608 including accuracy, precision, recall, F1-score, and area under the ROC curve
609 (AUC) will be computed to quantify predictive performance. Feature-importance
610 analyses will be performed to identify the most discriminative variables contribut-
611 ing to chimera detection.

612 **3.1.4 Validation and Testing**

613 Validation will involve both internal and external evaluations. Internal validation
614 will be achieved through five-fold cross-validation on the training data to verify
615 model generalization and reduce variance due to random sampling. External
616 validation will be achieved through testing on the 20% hold-out dataset derived
617 from the simulated reads, which will serve as an unbiased benchmark to evaluate
618 how well the trained models generalize to unseen data. All feature extraction and
619 preprocessing steps will be performed using the same bioinformatics pipeline to
620 ensure consistency and comparability across validation stages.

621 Comparative evaluation across all candidate algorithms, including SVM, de-
622 cision trees, logistic regression, Random Forest, gradient boosting, and others,
623 will determine which models demonstrate the highest predictive performance and
624 computational efficiency under identical data conditions. Their metrics will be
625 compared to identify the which algorithms are most suitable for further refine-
626 ment.

627 **3.1.5 Documentation**

628 Comprehensive documentation will be maintained throughout the study to en-
629 sure transparency, reproducibility, and scientific integrity. All stages of the re-
630 search—including data acquisition, preprocessing, feature extraction, model train-
631 ing, and validation—will be systematically recorded. For each analytical step, the
632 corresponding parameters, software versions, and command-line scripts will be
633 documented to enable exact replication of results.

634 Version control and collaborative management will be implemented through
635 GitHub, which will serve as the central repository for all project files, including
636 Python scripts, configuration settings, and Jupyter notebooks. The repository
637 structure will follow standard research data management practices, with clear
638 directories for datasets, processed outputs, and analysis scripts. Changes will be
639 tracked through commit histories to ensure traceability and accountability.

640 Computational environments will be standardized using Conda, with environ-
641 ment files specifying dependencies and package versions to maintain consistency
642 across systems. Experimental workflows and exploratory analyses will be con-
643 ducted in Jupyter Notebooks, which facilitate real-time visualization, annotation,
644 and incremental testing of results.

645 For the preparation of the final manuscript and supplementary materials,
646 Overleaf (LaTeX) will be utilized to produce publication-quality formatting, con-
647 sistent referencing, and reproducible document compilation. The documentation
648 process will also include a project timeline outlining major milestones such as
649 data collection, simulation, feature extraction, model evaluation, and reporting to
650 ensure systematic progress and adherence to the research schedule.

651 **3.2 Calendar of Activities**

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

Table 3.1: Timetable of Activities

Activities (2025)	Nov	Dec	Jan	Feb	Mar	Apr	May
Data Collection and Simulation	● ● ● ●						
Bioinformatics Tools Pipeline	● ●	● ● ● ●	● ● ● ●				
Machine Learning Development			● ●	● ● ● ●	● ● ● ●	● ●	
Testing and Validation						● ●	● ● ● ●
Documentation	● ● ● ●	● ● ● ●	● ● ● ●	● ● ● ●	● ● ● ●	● ● ● ●	● ● ● ●

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