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34

35 Abstract

36 Histone modifications and chromatin-binding proteins are important regulators 37 of gene expression in eukaryotes and have pivotal roles in fungal pathogenicity and 38 development. However, profiling these modifications or proteins across the genome in 39 fungi is still challenging, due to the technical limitations of the traditional widely-used 40 ChIP-Seq method. Here, we present an optimized CUT&Tag-Seq protocol 41 (fCUT&Tag-Seq) specifically designed for filamentous fungi and dimorphic fungi. 42 Our approach involves the preparation of protoplasts and nuclear extraction to 43 enhance antibody accessibility, along with formaldehyde crosslinking to improve 44 protein-DNA binding efficiency. We then successfully applied fCUT&Tag-Seq to 45 accurately profile multiple histone modifications like H3K9me3, H3K27me3, 46 H3K4me3, and H3K18ac, across different plant pathogenic or model fungal species 47 including Verticillium dahliae, Neurospora crassa, Fusarium graminearum, and 48 Sporisorium scitamineum. Compared to the traditional ChIP-Seq, our method showed 49 superior signal-to-noise ratios, higher reproducibility, and enhanced detection 50 sensitivity. Furthermore, we extended this method to profile chromatin-binding 51 proteins, such as the histone acetyltransferase Gcn5. This study establishes 52 fCUT&Tag-Seq as a robust and useful tool for fungal epigenetic research, enabling 53 detailed exploration of chromatin dynamics and advancing our understanding of 54 fungal gene regulation, development, and pathogenicity.

55

56 Introduction

57 Chromatin structure and modifications are essential in regulating gene 58 expression in eukaryotic organisms. The basic unit of chromatin, the nucleosome, 59 consists of DNA wrapped around histone octamers composed of core histones H2A, 60 H2B, H3, and H4[1]. The N-terminal tails of these histories can undergo various 61 post-translational modifications (PTMs), such as methylation, acetylation. 62 phosphorylation, ubiquitination, sumoylation, and ADP-ribosylation[2][3]. These 63 modifications can alter chromatin structure and impact vital biological processes, such 64 as gene expression, DNA repair, and chromosome organizationError! Reference 65 source not found.[4].

66 Among these modifications, histone methylation is critical in regulating gene 67 expression. For instance, trimethylation of histone H3 at lysine 9 (H3K9me3) and 68 lysine 27 (H3K27me3) is highly associated with heterochromatin formation and gene 69 silencing. In contrast, trimethylation of histone H3 at lysine 4 (H3K4me3), which 70 typically occurs at transcription initiation sites, is commonly enriched near gene 71 promoters and is associated with active transcription[6]-[8]. 72 Furthermore, chromatin-binding proteins, such as the histone acetyltransferases and 73 transcription factors, play crucial roles in modifying histones or binding to promoters, 74 thus regulating gene expression, development, stress responses, and pathogenicity in 75 eukaryotes, including fungi [9][10].

Traditional methods for studying histone modifications and chromatin-binding
proteins have relied heavily on Chromatin Immunoprecipitation (ChIP) techniques,
developed in the mid-1980s. The subsequent advent of ChIP followed by

79 high-throughput sequencing (ChIP-Seq) enables comprehensive identification of 80 histone modification or protein-binding sites across the whole genome. While 81 ChIP-Seq has become a foundational approach for genome-wide chromatin analysis, 82 it still has several limitations. The method requires large amounts of input material, 83 may suffer from epitope masking, and often produces high background noise with low 84 signal-to-noise ratios[11]-[14]. These challenges are particularly pronounced when 85 studying filamentous fungi, where rigid cell walls and secondary metabolites reduce 86 antibody accessibility and affect immunoprecipitation efficiency[15].

87 The recent development of Cleavage Under Targets and Tagmentation 88 (CUT&Tag) provids a more efficient method for profiling chromatin features[15]. 89 This technique employs a tethered Tn5 transposase directed by specific antibodies to 90 simultaneously cleave DNA at target sites and integrate sequencing adapters. 91 CUT&Tag-Seq offers several advantages over traditional ChIP-Seq, including higher 92 resolution mapping with lower cell input requirements (as few as 60 cells) and 93 reduced background noise. While CUT&Tag has been successfully applied to 94 mammalian, plant cells, and yeast, its application in filamentous fungi has been 95 limited due to technical challenges posed by the fungal cell wall[17].

96 In this study, we present an optimized CUT&Tag-Seq method (fCUT&Tag-seq) 97 adapted for plant pathogenic or model fungi, including various species such 98 as Verticillium dahliae, Neurospora crassa, Fusarium graminearum, and Sporisorium 99 scitamineum. Our study addresses the challenges posed by fungal cell walls through 100 protoplast preparation and nuclear extraction, enhancing antibody accessibility and 101 DNA availability. Additionally, for chromatin-binding proteins of low abundance, we 102 incorporated a formaldehyde crosslinking step to strengthen protein-DNA interactions. 103 We apply this method to diverse fungal species, focused on mapping the distribution 104 of histone modifications (H3K9me3, H3K4me3, H3K27me3, and H3K18ac) and 105 chromatin-binding proteins such as acetyltransferase SsGcn5. Our results demonstrate 106 that the fCUT&Tag-Seq method can be successfully applied to filamentous fungi 107 using low cell numbers, producing high-quality sequencing libraries with improved 108 signal-to-noise ratios and reproducibility compared to traditional ChIP-Seq methods.

Implementing CUT&Tag-Seq in fungal studies offers a powerful tool for studying fungal epigenetics and gene regulation. This method reduces the required cell input and experimental duration, and enhances data quality, facilitating more detailed and accurate analyses of chromatin dynamics in fungi.

113

114 **Results**

115 Optimization and Implementation of CUT&Tag-Seq Method for Fungal Systems

116 We developed an optimized fCUT&Tag-Seq specifically adapted for fungal 117 systems, building upon the traditional approach using the Vazyme Hyperactive 118 Universal CUT&Tag Assay Kit for Illumina Pro (Cat. TD904). Our optimization 119 addresses the unique challenges posed by fungal cell architecture while maintaining 120 the high sensitivity and specificity characteristic of CUT&Tag methodology. The 121 modified protocol incorporates several key steps for fungal applications (Fig. 1). To 122 overcome the barrier presented by the fungal cell wall, we added a controlled 123 enzymatic digestion step to generate protoplasts, followed by gentle nuclear 124 extraction to preserve chromatin integrity. The isolated nuclei are then specifically 125 captured by Concanavalin A-coated magnetic beads, which provide a stable platform 126 for subsequent reactions while minimizing sample loss.

127 The workflow proceeds through several precisely controlled steps (Fig. 1): (1) 128 Nuclear membrane permeabilization is enhanced through carefully optimized 129 digitonin treatment, facilitating antibody access while maintaining nuclear structural 130 integrity. (2) Primary antibodies specific to the target histone modifications are 131 introduced, followed by secondary antibodies that serve to amplify the detection 132 signal. (3) The pA/G-Tn5 transferase complex is recruited to the antibody-bound locations through protein A/G interactions. (4) Upon Mg^{2+} activation, the tethered Tn5 133 134 simultaneously cleaves adjacent DNA and integrates sequencing adapters, enabling 135 efficient library preparation through subsequent PCR amplification. After library 136 construction, next-generation sequencing and data analysis can be performed, to map 137 the distribution pattern of histone modifications.





139 Figure 1. Schematic overview of the fCUT&Tag-Seq workflow for filamentous fungi.

140 Illustration of the key steps in the optimized fCUT&Tag-Seq protocol in filamentous fungi: (1) Preparation of

- 141 fungal protoplasts through cell wall digestion; (2) Nuclear extraction from protoplasts; (3) Capture of nuclei using
- 142 Concanavalin A-coated magnetic beads; (4) Addition of primary antibody specific to the histone modification or

143 protein of interest; (5) Binding of secondary antibody; (6) Addition of pA/G-Tn5 transferase complex; (7)

144 Mg²⁺-activated DNA cleavage and adapter integration; (8) PCR amplification for library construction. The

145 protocol incorporates digitonin treatment to enhance nuclear permeability and can be applied to both histone

146 modification and chromatin-binding protein analysis.

147

fCUT&Tag-Seq Demonstrates Superior Quality Control Metrics and Target Enrichment Compared to ChIP-Seq in V. dahliae

To evaluate if we could use fCUT&Tag-Seq for profiling histone modifications in filamentous fungi, we first selected the plant pathogenic fungus *V. dahliae* as a model system. This pathogenic fungus, known for its broad host range, causes severe wilt, yellowing, and vascular browning in infected plants, often resulting in significant crop losses[18]. Given the known roles of H3K9me3 and H3K27me3 in gene silencing and pathogenicity [2][19], we focused our analysis on these two modifications. To enhance our understanding of *V. dahliae*'s genetic background, we sequenced the whole genome of *V. dahliae* V592 strain using Pacbio HiFi platform (**Fig. S1**). Using the genome of this specific strain as the reference for mapping can avoid errors introduced by inter-species genomic differences, and also capture species-specific epigenetic modifications of genes, thereby providing more accurate information for following analyses.

162 We then compared traditional ChIP-Seq data from the SRA database (Home -163 SRA - NCBI) and fCUT&Tag-Seq datasets based on multiple quality control metrics. 164 Initial quality control analysis of fCUT&Tag-Seq libraries revealed characteristic 165 nucleosomal ladder patterns through agarose gel electrophoresis, indicating successful 166 fragmentation and library preparation (Fig. 2A). Fragment length distribution analysis 167 further confirmed the expected size distribution pattern typical of nucleosome-sized 168 fragments (Fig. 2B), validating the technical success of our protocol. Comparative 169 analysis between fCUT&Tag-Seq and traditional ChIP-Seq demonstrated several key 170 advantages of our optimized method. Most notably, fCUT&Tag-Seq exhibited 171 significantly lower duplication rates (8.73%) compared to ChIP-Seq (88.45%), 172 indicating substantially improved library complexity and data utilization efficiency 173 (Fig. 2C). Following alignment to the genome and subsequent filtering, we conducted 174 comprehensive quality metrics analysis using ChIPQC. The fCUT&Tag-Seq data 175 showed superior Signal Space Distance (SSD) scores, reflecting enhanced enrichment 176 efficiency. Furthermore, the Fraction of Reads in Peaks (FRiP) consistently exceeded 177 7.8% in fCUT&Tag-Seq samples, with H3K9me3 modification showing particularly 178 improved signal-to-noise ratios compared to ChIP-Seq (Fig. 2D). Peak calling analysis 179 revealed comparable patterns between fCUT&Tag-Seq and ChIP-Seq data (Fig. 2E 180 and 2F), validating the biological relevance of our results. However, CUT&Tag-seq 181 identified a greater number of peaks and showed improved signal intensity (Fig. 2E), 182 indicating enhanced sensitivity and specificity in detecting histone modifications. The 183 improved signal-to-noise ratio was particularly evident in genome-wide coverage 184 plots, where fCUT&Tag-Seq showed more distinct enrichment patterns and clearer peak boundaries (Fig. 2F). These comprehensive quality metrics demonstrate that our fCUT&Tag-Seq protocol not only meets standard quality benchmarks but also offers superior performance compared to traditional ChIP-Seq methods in terms of data quality, enrichment efficiency, and technical reproducibility. The improved metrics suggest that this method provides a more reliable and efficient approach for studying histone modifications in *V. dahliae*.



191

192 Figure 2. Comprehensive quality control analysis of CUT&Tag-seq compared to ChIP-seq in Verticillum

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193 dahliae.
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194 A. Nucleosomal ladder pattern analysis of CUT&Tag-seq libraries by agarose gel electrophoresis. Libraries were 195 generated from H3K9me3 and H3K27me3 histone modification profiling experiments in Verticillum dahliae. B. 196 Fragment length distribution analysis of sequencing reads from fCUT&Tag-Seq and ChIP-Seq experiments. Green 197 histograms represent H3K9me3 libraries (replicate 1 and replicate 2), while blue histograms show H3K27me3 198 libraries (replicate 1 and replicate 2). C. Bar plot representing the percentage of reads mapped to peak regions in 199 the fCUT&Tag-Seq and ChIP-Seq datasets (rep1 and rep2). CUT&Tag had higher enrichment efficiency than 200 ChIP-seq. D. Comparative analysis of key quality metrics between CUT&Tag-seq and ChIP-seq methods. Metrics 201 include mapping rates (proportion of reads aligned to the reference genome), duplication rates (percentage of PCR

duplicates), and FRiP scores (Fraction of Reads in Peaks). Data demonstrate superior quality metrics for
CUT&Tag-seq across all parameters. E. Signal intensity profiles across peaks in fCUT&Tag-Seq versus ChIP-Seq
datasets, indicating higher signal-to-noise ratios in fCUT&Tag-Seq. The upper panel displays the H3K9me3
modification, while the lower panel shows the H3K27me3 modification. F. Genomic distribution analysis of
identified peaks comparing CUT&Tag-Seq and ChIP-seq methods, showing increased sensitivity, specificity, lower
background, and better repeatability in CUT&Tag-seq experiments. The displayed region is randomly selected.
The upper panel represents the H3K9me3 modification, and the lower panel shows the H3K27me3 modification.

209

210 fCUT&Tag-Seq Method Efficiently Profiles Histone Modifications in V. dahliae

Following validation of our optimized protocol, we applied fCUT&Tag-Seq to investigate the genome-wide distribution of two key repressive histone modifications, H3K9me3 and H3K27me3, in *V. dahliae*. To evaluate the specificity of our method, we analyzed both wild-type strains and their corresponding methyltransferase mutants (Vd $\Delta kmt1$ and Vd $\Delta ezh2$). We incorporated spike-in controls for data normalization and used IgG antibodies as negative controls to account for background signal.

Genome-wide profiling revealed robust enrichment of both H3K9me3 and H3K27me3 modifications in wild-type strains (**Fig. 3A and 3B**). In contrast, control experiments using non-specific IgG antibodies showed minimal signal across the genome, confirming the specificity of our approach (**Fig. S2A**). Importantly, we observed significantly reduced levels of H3K9me3 and H3K27me3 in their respective methyltransferase mutants, Vd $\Delta kmt1$ and Vd $\Delta ezh2$, consistent with the loss of these enzymatic activities (**Fig. 3A and 3B**).

224 Analysis of the genomic distribution of these modifications revealed distinct 225 patterns. H3K9me3 peaks were predominantly located in promoter regions (68.81%) 226 and intergenic spaces (25.28%), while H3K27me3 showed similar enrichment 227 patterns with 87.56% of peaks in promoter regions and 8.58% in intergenic regions 228 (Fig. 3C). These distribution patterns align with the known roles of these 229 modifications in transcriptional regulation. Comparison of signal intensities between 230 fCUT&Tag-Seq and ChIP-Seq data demonstrated superior enrichment in 231 CUT&Tag-seq experiments, particularly across gene bodies and their 3-kb flanking 232 regions (**Fig. S2B**).

To assess the reproducibility of our modified protocol, we performed correlation analyses between biological replicates. fCUT&Tag-Seq demonstrated excellent reproducibility, with Pearson correlation coefficients of 0.99 and 1.0 for H3K9me3 and H3K27me3 modifications, respectively. These values exceeded those obtained from traditional ChIP-Seq experiments (0.91 and 0.89), highlighting the enhanced reliability of our optimized method (**Fig. 3D**).

These results collectively demonstrate that our fCUT&Tag-Seq protocol enables highly specific and reproducible mapping of histone modifications in *V. dahliae*, providing a robust tool for investigating chromatin-based regulation in this important plant pathogen.





244 Figure 3. Genome-wide profiling of histone modifications in V. dahlae using fCUT&Tag-seq

245A. Genome browser view of H3K9me3 distribution in wild-type and Vd $\Delta kmt1$ strains. The tracks show specific246H3K9me3 enrichment in WT and a significant reduction in the Vd $\Delta kmt1$ mutant. B. Genome browser view of

247 H3K27me3 distribution in wild-type and Vd/ezh2 strains. The tracks demonstrate specific H3K9me3 enrichment 248 in WT and a significant reduction in the $Vd \Delta ezh2$ mutant. C. Comparative analysis of genomic feature distribution 249 for H3K9me3 and H3K27me3 modifications between fCUT&Tag-seq and ChIP-seq approaches. The left panel 250 shows the distribution of H3K9me3 peaks across different genomic elements, and the right panel shows 251 H3K27me3 peak distribution, demonstrating the comprehensive coverage achieved by fCUT&Tag-seq. D. Pearson 252 correlation analysis of signals between biological replicates for H3K9me3 and H3K27me3 modifications in 253 CUT&Tag-seq (replicate 1 and replicate 2) and ChIP-seq experiments (replicate 1 and replicate 2). Pearson 254 correlations were calculated using deepTools, with the multiBamSummary tool followed by plotCorrelation. Read 255 counts were divided into 1,000-bp bins across the genome for the analysis. Higher correlation in CUT&Tag-seq 256 showed better reproducibility.

257

fCUT&Tag-Seq Method Shows Broad Applicability Across Model Filamentous Fungi

To assess whether our fCUT&Tag-seq protocol could be extended beyond *V. dahliae*, we tested it on two additional filamentous fungi: *N. crassa* and *F. graminearum*. Both species serve as valuable model organisms for studying fungal biology and pathogenicity, making them ideal candidates for validating broad applicability. Following the same experimental workflow, we focused on histone modifications H3K9me3 in *N. crassa*, and H3K4me3 in *F. graminearum*.

266 *N. crassa*, a member of the Ascomycota phylum, is widely recognized as a model 267 organism due to its ease of cultivation and haploid genetics, which facilitate 268 straightforward genetic analysis. Over the past few decades, it has been instrumental 269 in studying various epigenetic modifications, including DNA methylation and gene 270 silencing mechanisms[20][22]. We examined the levels of H3K9me3 in both the 271 wild-type strain and the H3K9me3 methyltransferase mutant Nc $\Delta dim5$. Quality 272 control assessments, including agarose gel electrophoresis and sequencing data 273 analysis, confirmed that the fCUT&Tag-Seq libraries met the required standards (Fig. 274 **4A and 4B**). The quality metrics indicated excellent performance for the *N. crassa* 275 wild-type samples (Fig. 4C). Notably, the peak distribution patterns observed were 276 consistent with those identified through ChIP-Seq, yet CUT&Tag-Seq demonstrated 277 higher signal intensity. The correlation between CUT&Tag-Seq and ChIP-Seq data 278 reached 0.73 (Fig. 4D and 4E), further validating the reliability of our method. 279 Additionally, biological replicates exhibited strong correlation, indicating high 280 reproducibility (Fig. 4F). Comparative analysis revealed that H3K9me3 was 281 significantly enriched in the wild-type strain, while the Nc/dim5 mutant displayed 282 markedly lower modification levels (Fig. 4G). As expected, H3K9me3 modifications 283 were predominantly located in distal intergenic regions and promoter regions (Fig. 284 **4H**).







A. Nucleosomal ladder pattern analysis of fCUT&Tag-Seq libraries by agarose gel electrophoresis. Libraries were
 generated from two replicates of H3K9me3 profiling experiments in *N. crassa*. B. Fragment length distribution
 analysis of sequencing reads from fCUT&Tag-seq experiments. Histograms display the expected nucleosomal
 fragmentation patterns for 2 replicates. C. Comparative analysis between fCUT&Tag-Seq and ChIP-Seq data for

291 H3K9me3 modification in N. crassa. D. Genome browser view of peak distribution analysis comparing 292 CUT&Tag-seq and ChIP-seq results. The upper panel shows H3K9me3 enrichment in ChIP-seq, and the lower 293 panel displays the modification distribution from CUT&Tag-Seq. E. Correlation matrix analysis between 294 CUT&Tag-Seq and ChIP-seq data, indicating strong agreement. Pearson correlations were calculated using 295 deepTools (multiBamSummary followed by plotCorrelation), based on read counts divided into 1000-bp bins 296 across the genome. F. Pearson correlation analysis of H3K9me3 signals between biological replicates in ChIP-Seq 297 and fCUT&Tag-Seq, demonstrating higher reproducibility of fCUT&Tag-Seq. G. Genome browser view of the 298 H3K9me3 peaks accross representative genomic regions in wild-type and Nc/dim5 strains in N. crassa. H. Feature 299 distribution comparison of H3K9me3 modification between ChIP-seq and CUT&Tag-seq, showing similar 300 distribution patterns between the two methods.

301

302 Similar results were obtained for F. graminearum, the causative agent of 303 fusarium head blight in wheat [23]. In addition to analyzing H3K9me3 and 304 H3K27me3 modifications, we also assessed H3K4me3 levels, which are associated 305 with gene activation. The sequencing quality metrics of fCUT&Tag-Seq for F. 306 graminearum were superior to those of ChIP-Seq, with a greater percentage of reads 307 mapping to peak regions (Fig. 5A and 5B). The Signal Space Distance (SSD) score 308 and the percentage of reads in peaks (Rip%) reached 21 and 46, respectively (Fig. 5B). 309 These findings confirm the robustness of our method, as the wild-type strain exhibited 310 significant enrichment (Fig. 5C). Notably, peak enrichment was predominantly 311 observed in the promoter regions (96.51%) (Fig. 5D). And the Pearson correlation 312 coefficient was 0.99, indicating a strong correlation between the sample groups (Fig. 313 **5E**).

Overall, these results demonstrate that our fCUT&Tag-Seq method is not only effective for detecting histone modifications in *V. dahliae* but also applicable to other model fungi, including *N. crassa* and *F. graminearum*. This versatility underscores the potential of CUT&Tag-Seq as a powerful tool for studying epigenetic modifications across a broad range of fungal species.



319

320 Figure 5. Genome-wide profiling of H3K4me3 modifications in *F. graminearum* using fCUT&Tag-seq

321 analysis

322 A. Fragment length distribution of sequencing libraries for fCUT&Tag-Seq. The left panel shows the fragment 323 length distribution for the H3K4me3 library in replicate 1, and the right panel shows the distribution for replicate 2. 324 B. Comparative analysis of key quality metrics for ChIP-Seq and fCUT&Tag-Seq datasets, showing the higher 325 quality of CUT&Tag-Seq. C. Genome browser view of H3K4me3 modification peak distribution analysis 326 comparing fCUT&Tag-Seq and ChIP-Seq results. The genome regions were randomly selected. Green peaks 327 represent ChIP-Seq results, and blue peaks represent CUT&Tag-Seq results. D. Genome browser view of the 328 H3K4me3 peak distribution analysis comparing fCUT&Tag-Seq and ChIP-Seq results in two replicates of 329 wild-type strain, showing significantly enrichment of H3K4me3 modification. E. The pie chart and UpSet plot 330 illustrate the distribution of peaks across different genomic features. The results showing that H3K4me3 is 331 predominantly enriched in promoter regions. Distinct colors represent different genomic features. F. Pearson 332 correlation coefficients between sample groups for ChIP-Seq and fCUT&Tag-Seq, confirming high reproducibility 333 for CUT&Tag-seq. The scatter plot displays normalized read counts in 1-kb bins across the genome.

334

335 fCUT&Tag-Seq Method Also Enables Profiling of Chromatin-Binding Proteins

336 To extend the utility of our optimized CUT&Tag-Seq protocol beyond histone 337 modifications, we adapted the method to profile chromatin-binding proteins, which 338 play crucial roles in transcriptional regulation. Given the typically lower abundance of 339 chromatin-binding proteins compared to histones, we incorporated an additional 340 formaldehyde crosslinking step before protoplast preparation. This modification 341 strengthens protein-chromatin interactions, thereby improving detection sensitivity. 342 Following fragmentation, we performed sequential decrosslinking, protease digestion, 343 and RNase treatment to ensure efficient DNA purification and library preparation (Fig. 344 **6**).

345 To validate this adapted fCUT&Tag-seq method, we investigated the histone 346 acetyltransferase Gcn5 in S. scitamineum, a dimorphic pathogenic fungus responsible 347 for sugarcane smut. Sugarcane smut is one of the most significant diseases affecting 348 global sugarcane production worldwide. The disease poses substantial economic 349 threats, highlighting the urgent need for effective prevention and control strategies 350 against S. scitamineum. It has been reported that histone acetylation may play a 351 critical role in fungal pathogenesis [24][27]. To explore the relationship between 352 histone acetylation and fungal pathogenicity, we screened possible histone 353 acetyltransferase candidate genes and introduced a Flag epitope tag into the histone 354 acetyltransferase Gcn5 in two S. scitamineum strains with different mating types 355 (JG35 and JG36), employing our previously reported method [28].

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356

357 Figure 6. Schematic overview of the optimized fCUT&Tag-seq method for detecting chromatin-binding

358 proteins

Workflow diagram illustrating the key steps of the fCUT&Tag-seq protocol for analyzing chromatin-binding
 proteins in fungi, including protoplast preparation, formaldehyde crosslinking, nuclear extraction, and subsequent
 CUT&Tag-Seq procedures.

362

363 Quality control analysis of the resulting CUT&Tag-Seq libraries revealed proper 364 DNA fragmentation patterns by agarose gel electrophoresis and expected nucleosomal 365 size distributions in sequencing reads (Fig. 7A and 7B), confirming successful library 366 preparation. Genome-wide profiling demonstrated robust enrichment of 367 SsGcn5-3Flag binding sites across the S. scitamineum genome (Fig. 7C and 7D). 368 Notably, comparative analysis revealed a strong correlation between SsGcn5 binding 369 sites and regions of H3K18 acetylation (Fig. 7E), consistent with the known catalytic

activity of Gcn5. Feature distribution analysis showed that SsGcn5 binding sites were predominantly located in promoter regions (85.93% of total peaks) (**Fig. 7F**), suggesting a primary role in transcriptional regulation. The high reproducibility of our method was demonstrated by the near-perfect Pearson correlation coefficient (approximately 1.0) between biological replicates (**Fig. 7G**).

Our results confirmed that the fCUT&Tag-Seq method not only excels in detecting histone modifications, but also effectively profiles chromatin-binding proteins in filamentous fungi. By enhancing the detection of low-abundance proteins, this method also provides a powerful tool for exploring chromatin-binding proteins in regulating gene expression and chromatin dynamics across a broad range of fungal species.



381

382 Figure 7. Genome-wide profiling of SsGcn5 chromatin-binding protein and its correlation with H3K18ac

383 modification in S. scitamineum using fCUT&Tag-seq

384 A. Quality assessment of fCUT&Tag-seq libraries by agarose gel electrophoresis. DNA fragment distributions are

385 shown for two biological replicates of JG35 (lanes 1-2) and JG36 (lanes 3-4) mating types. B. Fragment length 386 distribution analysis of sequencing reads demonstrating expected nucleosomal patterns. Blue and pink histograms 387 represent Gcn5-Flag distributions in JG35 and JG36 strains, respectively. C. Comprehensive quality metrics for 388 fCUT&Tag-seq datasets in DNA-binding protein enrichment in JG35 and JG36 strains. D. Genome browser view 389 of SsGcn5-3Flag binding sites identified by CUT&Tag-Seq. Genome regions were selected randomly. E. 390 Correlation analysis between SsGcn5-3Flag binding sites and H3K18ac modification regions, showing the 391 functional relationship between histone acetyltransferase localization and its catalytic activity. F. Genomic feature 392 distribution analysis of SsGcn5-Flag peaks in JG35 and JG36 strains, showing the relative enrichment across 393 different genomic elements (promoters, gene bodies, intergenic regions). G. Correlation analysis between 394 biological replicates for SsGcn5-3Flag CUT&Tag-seq experiments, demonstrating high reproducibility of the 395 method.

396

397 Conclusion & Discussion

398 Over prolonged evolutionary periods, plant pathogenic fungi have developed 399 intricate strategies to penetrate and colonize their hosts. Precise control of pathogenic 400 gene expression is pivotal to these infection processes, influencing fungal 401 development, environmental stress responses, and virulence[29]. Emerging evidence 402 underscores the importance of epigenetic regulation, including histone methylation, 403 histone acetylation, and chromatin remodeling, in orchestrating such transcriptional 404 reprogramming in plant pathogens[30]-[32]. However, until recently, the available 405 tools for epigenetic research in diverse fungal systems were still very limited, making 406 advanced investigations into fungal chromatin biology challenging.

407 CUT&Tag-Seq method, introduced by Kaya-Okur et al.[15] in 2019, offers a 408 powerful alternative to ChIP-Seq through higher resolution and lower background 409 noise. Yet its application had largely been confined to plant, yeast, and animal 410 cells[15][33][34]. In this study, we optimized and extended CUT&Tag-Seq to several 411 important fungal species, including *V. dahliae*, *F. graminearum*, *N. crassa*, and *S.* 412 *scitamineum*. Our fCUT&Tag-Seq method effectively addresses the unique obstacles 413 posed by fungal cell walls by integrating protoplast preparation and gentle nuclear 414 extraction. Additionally, a formaldehyde crosslinking step was introduced to
415 strengthen DNA-protein interactions, particularly beneficial for profiling
416 low-abundance chromatin-binding proteins.

417 Our data revealed that the fCUT&Tag-Seq method accurately and sensitively 418 detects histone modifications (H3K9me3, H3K27me3, H3K4me3, and H3K18ac) as 419 well as chromatin-binding proteins in these fungi. Furthermore, our comparisons with 420 conventional ChIP-Seq protocols highlighted notable improvements in library 421 complexity, signal-to-noise ratios, and correlation across biological replicates. These 422 advantages facilitate comprehensive insights into the epigenetic landscapes governing 423 fungal pathogenicity, development, and physiology.

While our fCUT&Tag-Seq method has demonstrated significant improvements in data quality and sensitivity, it is important to acknowledge the limitations. Challenges remain in enriching low-abundance proteins and chromatin-binding proteins located at the heterochromatin regions, where the antibody accessibilities are largely reduced. Future studies should focus on further optimizing methods for relaxing heterochromatin to enhance the detection of these proteins.

Taken together, we established fCUT&Tag-Seq as a robust and versatile method for fungal epigenetic research. By reducing cell input requirements and enhancing data quality, it enables a deeper exploration of chromatin-based mechanisms that shape fungal behavior and plant-fungus interactions. We anticipate that future studies will leverage this method to uncover new regulatory pathways, thereby advancing the characterization of fungal pathogenicity and guiding innovative approaches for crop disease management.

437

438 Materials and Methods

439 Fungal Strains and Culture Conditions

440 The following fungal strains were used in this study: *V. dahliae* strain V592 (isolated

from cotton in Xinjiang, China) [35][36], N. crassa 87-3 and Nc∆dim5 (provided by

442 Xiao Liu)[37], S. scitamineum strains JG35 and JG36 (provided by Shan Lu)[38], and

443 F. graminearum wild-type strain (provided by Guangfei Tang and Haoxue Xia)[39].

All strains were maintained and cultured according to standard protocols specific toeach species.

446

447 Generation of Mutant and Tagged Fungal Strains

448 For V. dahliae, VdKMT1 (VDAG_07826) and VdEZH2 (VDAG_00983) deletion strains (Vd $\Delta kmt1$ and Vd $\Delta ezh2$) were constructed as previously reported[40]. 1kb 449 450 genomic sequences upstream and downstream of target genes were amplified from 451 V592 DNA with the following primer pairs: VdKMT1-A/ VdKMT1-a, VdKMT1-d/ 452 VdKMT1-B, VdEZH2-A/ VdEZH2-a, VdEZH2-d/ VdEZH2-B (listed in Table S1) 453 using the Phanta Max Super-Fidelity DNA Polymerase (P505, Vazyme) and cloned 454 into the pGKO-HPT / pGKO-NAT vector via homologous recombination. 455 Sequence-verified vectors were then used to transform V. dahliae by Agrobacterium 456 tumefaciens-mediated transformation (ATMT) system. V. dahliae transformants were 457 selected (PDA) on potato dextrose agar plates supplemented with 458 5-fluoro-2'deoxyuridine (5FU) and hygromycin B / Nourseothricin. Putative 459 transformants were screened utilizing PCR to confirm the successful deletion of target 460 genes by primer pairs: V1/Hpt-R and V2/Hpt-F. The primers used in construction are 461 listed in Table S1.

462 For S. scitamineum, the SsGcn5-3Flag strain was constructed using homologous 463 recombination. The engineered vector pEX1-GAP-Flag-HPT, which carries a Flag tag, 464 was initially constructed by modifying the pEX1-GAP-HPT vector. Subsequently, 465 approximately 1.5 kb of upstream (5') and downstream (3') flanking sequences near 466 the target protein's stop codon, excluding the stop codon itself, were amplified using 467 PCR from the wildtype strains JG35 and JG36 with the following primer pairs: 468 SsGcn5-tg-A/SsGcn5-a, SsGcn5-d/SsGcn5-B (listed in Table S1). Furthermore, two 469 truncated fragments of the hygromycin resistance gene HPT, designated HPT-up and 470 HPT-down, were amplified from the pEX1-GAP-Flag-HPT plasmid via PCR with the 471 following primer pairs: HPT-LB-F/HPT-LB-R, HPT-RB-F/HPT-RB-R. Overlap PCR 472 was employed to fuse the upstream DNA sequence with HPT-up with the primer pair

473 SsGcn5-tg-A/ HPT-LB-R and the downstream DNA sequence with HPT-down with
474 the primer pair HPT-RB-F/ SsGcn5-B, and the products were introduced into *S*.
475 *scitamineum* strains JG35 and JG36 using PEG-mediated protoplast transformation.
476 Transformants were selected on YEPSA medium containing 150 μg/mL hygromycin
477 B and verified by PCR using primer pairs SsGcn5-C(V1)/Hpt-LB-R and
478 SsGcn5-D(V2)/Hpt-RB-F.

479 fCUT&Tag-Seq for Histone Modification

480 To detect histone modifications using CUT&Tag-Seq, protoplasts were first 481 prepared. This was achieved by digesting mycelium (V. dahliae, N. crassa and F. 482 graminearum) or spores (S. scitamineum) with 10 mL of enzyme solution (0.25g 483 VinoTaste®Pro enzyme is dissolved in 12.5 mL 0.7 M NaCl for V. dahliae, N. crassa 484 and S. scitamineum's protoplasts preparation; 0.1 g Diselease, 0.2 g Celulase and 0.2 485 g Lysozyme were diluted with 0.7 M NaCl to 20 mL for F. graminearum's protoplasts 486 preparation). The requisite digestion time was contingent upon the specific fungus, 487 with the following durations observed: 3 h for V. dahliae mycelium, F. graminearum 488 and N. crassa mycelium, and 30 min for S. scitamineum spores. Filtering protoplasts 489 with Miracloth (475855-1R; Merck), followed by centrifugation for 10 min at 3800 490 rpm for V. dahliae and N. crassa, 2200 rpm for S. scitamineum and 5000 rpm for F. 491 graminearum, 25 . Protoplasts were resuspended using 10 mL of 0.7 M NaCl, 492 collected by centrifugation. Finally, the number of cells was counted using a 493 hemocytometer.

For extracting nuclei, take 1 million cells for 1.5 mL EP tubes based on the number of cells counted, followed by centrifugation at 3000 rpm at 25 \square for 5 min. The precipitate was resuspended with 100 µL of nuclear extract buffer (TD904; Vazyme) and left on ice for 10 min, followed by centrifugation at 2500 rpm at 25 \square for 6 min. Carefully remove the supernatant and resuspend the nuclei by adding 100 µL of wash buffer (TD904; Vazyme).

For the adsorption of nuclei, activated Concanavalin A-coated magnetic beads
(TD904; Vazyme) and 250,000 resuspended cell nuclei were added to 200 μl 8-row
tubes, along with 75 μl of wash buffer (TD904; Vazyme). The contents were then

503 mixed by inverting the tubes. Leave at room temperature for 10 min, turning up and504 down 2-3 times.

505 In setting up for the antibody reaction, the 8-row tubes from the previous step 506 were placed on a magnetic rack, the 8-row tubes were removed after discarding the 507 supernatant, and 50 µL of antibody buffer (TD904; Vazyme) was added to each 508 sample to resuspend the magnetic bead cell nuclear complex. Add 1 μ L of antibody 509 against trimethylation of histone 3 at lysine 9 (H3K9me3) (Cat39161; Active Motif), 510 trimethylation of histone 3 at lysine 27 (H3K27me3) (Cat39155; Active Motif), and 511 trimethylation of histone 3 at lysine 4(H3K4me3) (Cat39060; Active Motif) according 512 to the ratio used. After mixing upside down and upside down, centrifuge instantly and 513 place in the refrigerator at 4°C overnight.

514 On the second day, the overnight magnetic bead antibody complexes were 515 mixed upside down and centrifuged instantaneously before being placed on a 516 magnetic rack. After clarifying and discarding the supernatant, 0.5 μ L of rabbit 517 antibody (Ab207; Vazyme) was mixed with 49.5 μ L of Dig wash buffer (792 μ L Wash 518 Buffer, 8 µL 5% Digitonin) and added to each sample, which was slowly spun for 1 h 519 at room temperature (9-11 rpm/min). After transient centrifugation on a magnetic rack, 520 the supernatant was clarified and discarded and the beads was washed three times 521 with 200 µL of Dig wash buffer.

For transposase incubation, 2 μ L of Hyperactive pA/G-Transposon (TD904; Vazyme) in 98 μ L Dig-300 buffer (100 μ L 10 × Dig-300 Buffer, 2 μ L 5% Digitonin, 20 μ L 50×protease inhibitor, 878 μ L ddH₂O) was added to each reaction and incubated with gentle rotation (9-11 rpm/min) at room temperature for 1 h. Subsequently, instantaneous centrifugation on a magnetic rack, clarification and discarding of supernatant, and wash three times using 200 μ L Dig-300 buffer to remove unbound pA/G-Transposon.

529 For fragmentation, 10 μL trueprep tagment buffer L (TD904; Vazyme) in 40 μL
530 Dig-300 buffer was added to each reaction and incubated at 37°C for 1 h.

531 For DNA extraction, 2 μ L 10% SDS (TD904; Vazyme) was added to stop the

532 fragmentation reaction, and 0.5 µL DNA spike in (TD904; Vazyme) was added as a 533 reference standard between samples during sequencing, after mixing, 55°C for 10 min. 534 Instantaneous centrifugation was placed on a magnetic rack and the supernatant was 535 transferred to a new 200 μ L 8-row tube. Add 50 μ L the activated DNA extract beads 536 (TD904; Vazyme) to the 8-row tube containing the supernatant, mix upside down, and 537 incubate for 20 min at room temperature, mixing 2-3 times during the incubation. 538 Instantaneous centrifugation, 8-row tubes were placed on a magnetic rack, clarified by 539 discarding the supernatant, washed twice with 200 µL of 1x B&W buffer (TD904; 540 Vazyme), dried on a magnetic rack, and resuspended the magnetic bead DNA 541 complex by adding 30 μ L of RNAase free ddH₂O.

542 Libraries for deep sequencing were constructed by PCR, with the PCR reaction 543 (50 μ L) set up as follows:15 μ L beads with affinity purified DNA, 25 μ L 544 2xCUT&Tag Amplification Mix (TD904; Vazyme), 5 µL N5XX Primer (TD202; 545 Vazyme), 5 µL N7XX Primer (TD202; Vazyme). The adaptor pooling guide strategy 546 from Illumina was followed. Each PCR solution was mixed gently blowing with a 547 pipette. Denaturation: 72°C for 3 min, 95°C for 3 min; Cycling reaction: 98°C for 10 s, 548 60°C for 5 s; Extension: 72°C for 1 min, 4°C hold. The number of cycles of the PCR 549 reaction ranged from 9 to 20.

550 For PCR product purification, 100 µL of VAHTS DNA clean beads (N411; 551 Vazyme) were added to the 8-row tube where the PCR reaction was completed. Mix 552 by vortex oscillation and let stand at room temperature for 5 min. Centrifuge the 553 8-row tube on a magnetic rack, clarify and discard the supernatant, wash twice with 554 200 μ L of 80% ethanol, dry on the magnetic rack, add 22 μ L of RNAase-free ddH₂O 555 to resuspend the magnetic beads PCR product complex, and incubate for 5 min at 556 room temperature. Place on a magnetic rack for clarification, and transfer supernatant 557 to a new 1.5 mL EP tube. Followed by quality control and deep sequencing.

558

559 fCUT&Tag-Seq for chromatin-binding protein proteins

560 To analyze lower-abundance chromatin-binding proteins, such as transcription 561 factors or histone acetyltransferases, a formaldehyde crosslinking step was integrated. Approximately 1 g of fungal spores was fixed with 9 mL 0.1% formaldehyde for 8 min at room temperature. The cross-linking was subsequently stopped by 1 mL of 2 M glycine for 5 min at room temperature. The cross-linked cells were washed three times using 20 mL SCS buffer.

566 Following crosslinking, protoplast formation and nuclei capture with 567 ConA-beads were performed as described above. After incubation with the 568 appropriate chromatin-binding protein antibodies, the pA/G-Tn5 transposome was 569 assembled onto the antibody-bound locus. DNA extraction, and purification were 570 carried out similarly to the procedure for histone modifications. To reverse crosslinks, 571 samples were incubated overnight at 65°C with 3 µL 5 M NaCl. RNA was removed 572 by RNase A treatment (37°C for 30 min), and proteins were digested by Proteinase K 573 (incubation at 53° C for 1 h). Proteases were removal by protease inhibitors (room 574 temperature for 5 min). Purified DNA libraries were further processed by PCR 575 amplification.

576

577 CUT&Tag-seq Data Analysis

578 Raw ChIP-seq data were downloaded from the SRA database using SRA Toolkit 579 (3.1.1). Raw data of ChIP-seq and CUT&Tag were first processed using fastp (0.23.4) 580 [41] to remove adapters and low-quality sequences. The cleaned reads were then 581 aligned to the spike-in sequences using BWA (0.7.18) [42]. After alignment, samtools 582 (1.21) [43] was used to count the number of successfully mapped reads. The sample 583 with the fewest mapped reads was selected as the baseline for spike-in normalization. 584 The normalized data were then re-aligned to the reference genome using BWA 585 (0.7.18), with reads having a quality score below 20 being filtered out using samtools 586 (1.21). The alignment results were then processed using Picard (Picard Tools - By 587 Broad Institute) for sorting and deduplication. Peak calling was performed using 588 MACS2 (2.2.9.1) [44], with IgG controls used to correct for background. The "-B" 589 flag was employed to generate histone modification and control lambda .bdg files, 590 which were subsequently compared using the bdgcmp function in MACS2. ChIPQC 591 (1.42.0, <u>Bioconductor - ChIPQC</u>) was applied for quality control of the CUT&Tag analysis. Graphical representations of the data were generated using Spark.py (2.6.2)
[45]. Peak annotation was performed using ChIPseeker (1.42.0) [46] to assign
genomic features to the identified peaks. Correlation analysis was carried out using
the multiBigwigSummary and plotCorrelation functions of deepTools (3.3.5) [47],
while heatmaps were generated using the computeMatrix and plotHeatmap functions
of deepTools (3.3.5).

598

599 Data Availability

600 All ChIP-seq data were obtained from the SRA database (Home - SRA - NCBI). 601 The ChIP-seq data for H3K9me3 modification in V. dahliae were downloaded from 602 SRR10571946 and SRR10571947, respectively, and data for H3K27me3 modification 603 were downloaded from SRR10571948 and SRR10571949. For H3K9me3 604 modification in N. crassa, the ChIP-seq data were downloaded from SRR1566112 and 605 SRR12229310. The ChIP-seq data for H3K4me3 modification in F. graminearum 606 were downloaded from SRR21677668 and SRR21677669. The reference genomes 607 used in the mapping process are as follows: PRJNA1212841 for V. dahliae V592, 608 GCF_000182925.2 for N. crassa, FgraminearumPH-1 for F. graminearum (available 609 at FungiDB), and GCA_001010845.1 for S. scitamineum. The CUT&Tag data used in 610 this study are available in the NCBI BioProject database under accession number 611 PRJNA1213569 and PRJNA1166993.

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- 758

759 Supplementary Materials:

- Figure S1: *V. dahliae* V592 genome assembly presentation;
- Figure S2: Genome-wide profiling of nonspecific IgG signals and histone modification patterns in *V. dahliae* using fCUT&Tag-seq;
- Figure S3: Original images of Figure 2A, Figure 4A, Figure 7A;
- Table S1: Primers used in this study.

765

766 Author Contributions:

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 and editing;
- 773 **J.Y.**: formal analysis, writing—original draft, and writing-review and editing;
- 774 M.L.: formal analysis, writing—original draft, and writing-review and editing;
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- G.T.: methodology, investigation, data curation, finish the experiments involving
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810 **Conflicts of Interest:**

- 811 The authors declare that there is no conflict of interest.
- 812
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817 Figure S1. V. dahliae V592 genome assembly presentation

A. The output quality of sequencing data. Sample: sample name; SeqNum: number of sequences; SumBase: total
number of data bases; N50Len: N50 length of sequencing data; N90Len: length of sequencing data N90; MeanLen:
mean length of sequencing sequence; MaxLen: maximum length of sequencing sequence. B. Assembly result
statistic. Length: length of the sequence after concatenation; GC (%): GC content of the concatenated sequence. C.
The CIRCOS plot of the genome.





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Figure S2. Genome-wide profiling of nonspecific IgG signals and histone modification patterns in *V. dahliae*

826 using fCUT&Tag-seq

A. Genome browser view of the H3K9me3, H3K27me3, and nonspecific IgG antibody signals across
representative genomic regions in Wild-type strain. The minimal background signal demonstrates the specificity of
the fCUT&Tag-seq method. Data were presented at the same scale as specific antibody signals. B. Heatmap
analysis of H3K9me3 and H3K27me3 signals near protein-coding genes in CUT&Tag-seq and ChIP-seq. Scale
regions were 3,000 bp upstream of the translation starting site (TSS), 3,000 bp downstream of the translation end
site (TES), and a 1,000-bp region on the gene body. The lengths were plotted using the computeMatrix and
plotHeatmap tools in deepTools.

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836 Figure S3: Original images of Figure 2A, Figure 4A, Figure 7A

837 A. Original images of Figure 2A. B. Original images of Figure 4A. C. Original images of Figure 7A