

# Digital Discovery

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# 1 **AI-Driven Antiviral Natural Products Drug Development: A**

## 2 **Technical Overview**

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View Article Online  
DOI: 10.1039/D5DD00504C

29

## 30 **Abstract**

31 The emergence of viral pandemics and rapid pathogen evolution presents formidable  
32 challenges for conventional antiviral development, including prolonged timelines, high  
33 costs, and susceptibility to resistance mechanisms. Natural products (NPs) offer  
34 promising antiviral potential through structural diversity and multi-target synergism,  
35 while their development faces critical bottlenecks in structural characterization, target  
36 identification, and synthetic optimization. Given the current situation, artificial  
37 intelligence (AI), particularly machine learning (ML) and deep learning (DL), is  
38 revolutionizing drug development by transforming data analysis and predictive  
39 modeling. This review explores AI applications across the antiviral NPs drug  
40 development continuum, providing insights for AI-driven pharmaceutical research.

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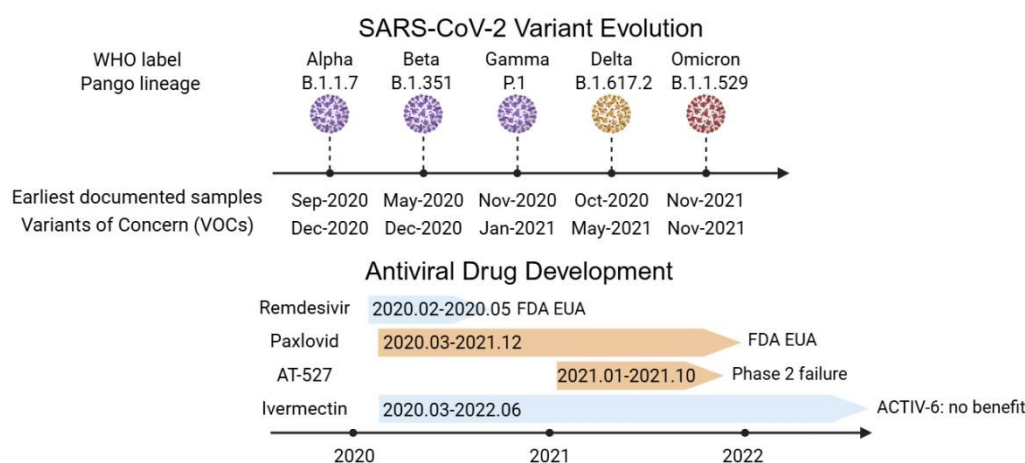
## 42 **1. Introduction**

43

44 In recent years, many new and re-emerging viral pathogens—such as severe acute  
45 respiratory syndrome coronavirus (SARS-CoV), Middle East respiratory syndrome  
46 coronavirus (MERS-CoV), and SARS-CoV-2—have continued to pose a risk to global  
47 public health(1). Antiviral medicine development is associated with high costs and  
48 extended timescales(2). Moreover, the current pace of pharmaceutical research and  
49 development critically lags behind the exponentially growing need for rapid therapeutic  
50 interventions during emergent pandemic scenarios (**Fig.1**). The remdesivir  
51 development timeline exemplifies both the opportunities and persistent limitations in  
52 this race against time. Initially explored in 2009 for hepatitis C and later Ebola, it gained  
53 emergency use authorization in May 2020 for COVID-19—only half a year after the



54 SARS-CoV-2 outbreak, showcasing a smooth translation from scientific discovery to  
 55 emergency response(3). Yet, as depicted in Fig. 1, even this rapid repurposing occurred  
 56 amid ongoing viral evolution, with variants like Alpha and Beta emerging before full  
 57 clinical implementation, highlighting that such successes depend on pre-existing  
 58 scaffolds and may not suffice for de novo threats. For novel viral threats where no such  
 59 prior knowledge exists, the de novo drug development process remains substantially  
 60 slower than the pace of outbreaks (4). Due to the virus's unique genetic system (lack of  
 61 complex genetic information synthesis proofreading system), the virus can rapidly  
 62 mutate and evade drug treatment. Consider the H1N1 influenza variants: Ingenious  
 63 mutations in the Sb region of the HA protein serve like a molecular camouflage kit,  
 64 allowing them to evade vaccine-educated antibodies and leaving vaccination  
 65 campaigns in the lurch(5). Such complex pressures require more sophisticated  
 66 therapeutic strategies, which demand transformed drug development paradigms.



67  
 68 **Fig. 1 Asynchrony Between SARS-CoV-2 Variant Evolution and Antiviral Drug**  
 69 **Development.** This figure illustrates the evolutionary dynamics of major SARS-CoV-  
 70 2 variants during the COVID-19 pandemic, aligned with the development and clinical  
 71 implementation timelines of four representative small-molecule antivirals. The upper  
 72 axis delineates SARS-CoV-2 evolutionary dynamics using a generational color scheme:  
 73 purple signifies the initial waves of Variants of Concern (Alpha, Beta, and Gamma)  
 74 marked by early increases in transmissibility; yellow identifies the Delta variant as a



75 pivotal transition toward significantly higher viral loads and pathogenicity, and red  
76 represents the Omicron lineage, which constitutes a fundamental paradigm shift in  
77 immune evasion and mutation density. Each variant is annotated with its earliest  
78 documented detection date and official WHO Variant of Concern (VOC) designation,  
79 with earliest documented detections referring to retrospectively identified sequences  
80 rather than real-time discovery, to highlight the inherent delay in global surveillance  
81 and response. Antivirals in the lower axis were selected according to two defining  
82 dimensions: the development pathway (repurposing of established agents versus de  
83 novo structural design) and the clinical resolution (regulatory authorization versus  
84 termination due to futility). Blue bands (Remdesivir and Ivermectin) represent the drug  
85 repurposing strategy, aimed at immediate deployment based on known safety profiles.  
86 In contrast, orange bands (Paxlovid and AT-527) signify de novo discovery programs.  
87 Despite such technological acceleration, vertical alignment across the axes reveals that  
88 by the time these high-potency agents reached their respective clinical endpoints, the  
89 viral landscape had already transitioned through multiple generational cycles,  
90 illustrating the persistent structural lag between therapeutic intervention and emergent  
91 pandemic needs. This figure highlights the asynchrony between viral evolution and  
92 therapeutic development, underscoring the challenges in maintaining efficacy against  
93 phylogenetically divergent lineages. Figure created with BioRender.com.

94 Natural products—complex metabolites produced by plants, fungi, animals, and  
95 microorganisms—exhibit the highest chemical diversity in nature(6), and are an  
96 important source of antiviral medicines. Many licensed treatments and prospects stem  
97 directly or indirectly from plants, microbes, and marine animals. For instance,  
98 artemisinin's antimalarial potency signified a milestone for natural-product-based anti-  
99 infective discovery(7); diammonium glycyrrhizinate from licorice root is authorized in  
100 China and Japan as an adjuvant for chronic hepatitis B(8); and numerous antivirals (e.g.,  
101 acyclovir, ganciclovir, vidarabine, zidovudine) emerged from natural leads via  
102 structural optimization(9). Despite this promise, development confronts obstacles:  
103 limited access to bioactive substances (only ~1% of microbial species are



104 culturable(10)); complex metabolite isolation and characterization (bioassay-guided  
105 fractionation and structural elucidation are often essential); uncertain pharmacological  
106 mechanisms (e.g., the multi-component synergy of Lianhua Qingwen capsules remains  
107 incompletely defined(11)); and complex synthesis (e.g., up to 30 enzymatic cascade  
108 steps to generate vinblastine(12)).

109 AI provides transformative solutions for these challenges. Advanced algorithms—  
110 including Transformer architectures and graph neural networks (GNNs)—have made  
111 great strides in accuracy for predicting drug-target interaction(13). In addition, the  
112 family of biomedical data is exploding (e.g. the ChEMBL compound library  
113 contains >20.3 million bioactivity measurements and >2.4 million unique  
114 compounds(14). Furthermore, the Traditional Chinese Medicine Systems  
115 Pharmacology Database (TCMSP) includes 29,384 components, 3,311 with targets,  
116 and 837 linking to diseases(15), thus simplifying model training. On the other hand,  
117 GPU clusters speedup computational tasks, improving by several orders of magnitude  
118 the speed of training large-scale AI models, such as DL architectures for drug-target  
119 interaction prediction, when compared with CPUs(16, 17).

120 The recent emergence of AI-driven platforms has already yielded significant  
121 breakthroughs in antiviral discovery. For instance, a sophisticated AI pipeline recently  
122 identified established antiretrovirals, such as bictegravir and etravirine, as potent broad-  
123 spectrum inhibitors against monkeypox virus and related poxviruses(18). While such  
124 successes underscore the transformative potential of AI in accelerating drug  
125 repurposing and novel application discovery, there remains a need for a more granular  
126 technical overview focused specifically on the end-to-end integration of AI across the  
127 entire antiviral natural product development continuum.

128 In this Review we summarize some of the critical applications of AI in upstream  
129 (resource mining and target identification), midstream (drug candidate screening and  
130 optimization), and downstream (preclinical and clinical stages). We subsequently  
131 explore the current technological limitations and emerging possibilities. Finally, we  
132 discuss the outline future directions for the field. We hope to highlight a new era of



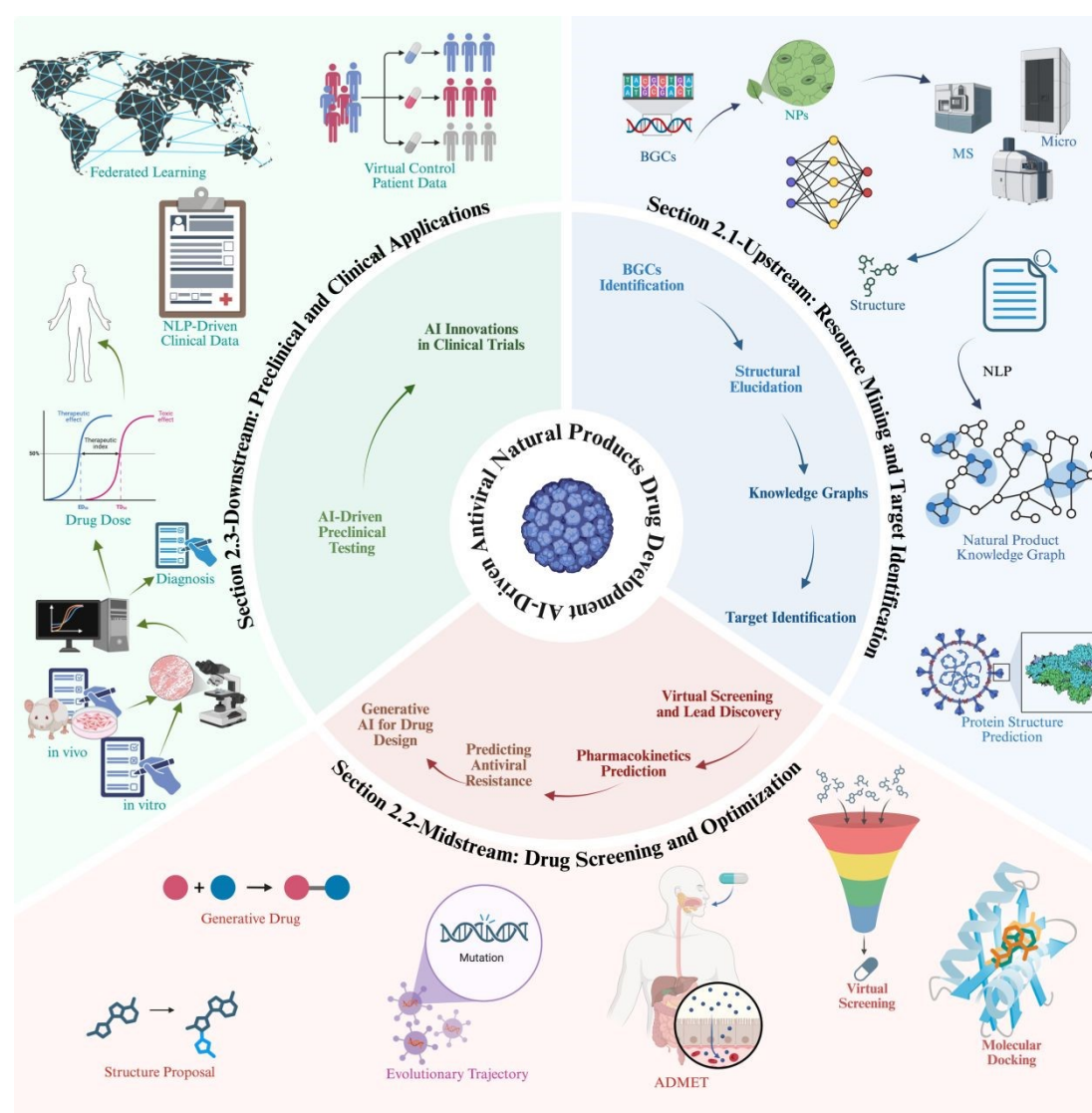
133 technology, efficiency and precision in drug development that is expected to speed  
 134 delivery of new and improved medicines to patients.

135

## 136 2. AI applications in antiviral natural products drug development

137

138 AI technologies are increasingly integrated into various stages of natural product-based  
 139 antiviral drug development, forming a systematic and intelligent pipeline from resource  
 140 mining and target identification to preclinical and clinical applications (Fig. 2).



**Fig. 2 AI-Driven Strategies for Antiviral Natural Products Drug Development Across the Full Pipeline.** This figure illustrates the comprehensive application of



AI throughout the full pipeline of antiviral natural products drug development. It is divided into three main stages: Upstream (Resource Mining and Target Identification), Midstream (Drug Screening and Optimization), and Downstream (Preclinical and Clinical Applications). In the upstream stage, AI facilitates biosynthetic gene clusters (BGCs) identification, structural elucidation of NPs, construction of knowledge graphs, protein structure prediction, and target identification. In the midstream stage, AI supports virtual screening, molecular docking, ADMET prediction, evolutionary trajectory analysis, antiviral resistance prediction, and generative drug design. In the downstream stage, AI enhances preclinical testing, dose optimization, diagnostic assistance, NLP-driven clinical data mining, federated learning, virtual control cohort construction, and innovations in clinical trials. Different color-coded sections visualize key processes across stages, highlighting the integrative role of AI in advancing natural product-based antiviral drug discovery. Figure created with BioRender.com.

141

## 142 **2.1 Upstream: Resource Mining and Target Identification**

### 143 **2.1.1 AI for Genome Mining and Biosynthetic Gene Clusters (BGCs)**

144 Multiple antiviral NPs are the products of secondary metabolites prescribed in  
145 microbial(19, 20) and plant genomes(21-23). AI is enhancing the search for these  
146 biosynthetic gene clusters. Standard genome-mining tools, such as anti-SMASH,  
147 employ rule-based pattern matching to identify known classes of BGCs, and this results  
148 in under-representative identification of clusters that are “atypical” clusters and in  
149 significant false-negative rates(24). Recognizing small sequence motifs beyond human-  
150 defined elements, DL approaches can be used to discover novel BGCs. For example,  
151 the RNN-based DeepBGC found novel BGCs in *Streptomyces* genomes(25). Such AI  
152 models leverage gene context, conserved domains, and amino acid properties  
153 generation to illuminate potentially hidden BGCs that may yield new antiviral drugs.



154 Bridging genetic and metabolomic data, these growing algorithms can correlate  
155 putative gene clusters with actual molecules, completing the link from genes to the  
156 NPs they produce. Genome-to-metabolite prediction is essential for discovering new  
157 antiviral chemicals that are stored in nature's genomic libraries.

### 158 2.1.2 AI-Assisted Structural Elucidation

159 The unambiguous structure determination of NPs is typically a labor-intensive, time-  
160 consuming process(26). AI enhances understanding of complex spectrometric data,  
161 including mass spectrometry (MS) and liquid chromatography–mass spectrometry (LC-  
162 MS)(27, 28). Methods for AI-assisted structural elucidation can be broadly classified  
163 into three categories: (1) ML/DL-based MS/MS spectrum annotation and substructure  
164 prediction; (2) predictive modeling for LC-MS retention time, peak feature extraction,  
165 and spectrum grouping; and (3) integrated pipelines for advanced techniques like  
166 microcrystal electron diffraction (MicroED). This classification reflects the shift from  
167 rule-based manual analysis to data-driven automation(29).

168 Recent advancements further strengthen this capability: MZmine 3, a scalable  
169 open-source platform, supports integrative processing of multimodal MS data  
170 (including LC-MS and ion mobility), enabling efficient feature detection, visualization,  
171 and annotation tailored to natural product workflows(30). Similarly, studies employing  
172 LC-MS/MS and molecular networking have identified marine-derived secondary  
173 metabolites with anti-SARS-CoV-2 activity, such as homofascaplysin A and aureol,  
174 though structural confirmation remains labor-intensive due to stereochemical  
175 complexity(31). MicroED with streamlined AI pipelines has enabled 3D structure  
176 determination of macrocyclic NPs with antiviral potential, overcoming the need for  
177 large single crystals by utilizing microcrystals—yet sample preparation is challenging,  
178 as NPs often form amorphous or poorly diffracting microcrystals, limiting resolution  
179 and throughput(32). ML-based retention-time prediction further enhances confidence  
180 by avoiding re-isolation of known compounds, but struggles with NPs' high chemical



181 diversity and batch variability(33-35). In summary, AI-facilitated spectrometric data  
182 analysis has begun to streamline structural elucidation in natural product research,  
183 thereby facilitating the identification of potential antiviral candidates. Nevertheless,  
184 overcoming NPs-inherent hurdles—such as mixture complexity, data scarcity for rare  
185 scaffolds, and the substantial validation gap—will be essential to translate these  
186 computational advances into robust, clinically relevant antiviral natural products.

### 187 **2.1.3 Knowledge Graphs and natural language processing (NLP) for Natural** 188 **Product Data**

189 New tools, including AI, are also increasingly utilized in organizing and mining this  
190 vast knowledge repository of NPs and traditional medicine. Knowledge graphs are  
191 structured networks that integrate heterogeneous data sources, such as chemical  
192 structures, biological targets, biosynthetic pathways, and literature references, enabling  
193 cross-domain analysis(36). NLP methods, on the other hand, extract structured  
194 information from unstructured texts, particularly historical herb manuals and medical  
195 literature. knowledge graphs offer high connectivity and query efficiency but face  
196 challenges in entity resolution and data integration due to the heterogeneity of NP  
197 sources (e.g., varying nomenclature and incomplete annotations). NLP excels at text  
198 mining but struggles with ancient language ambiguity, OCR errors, and domain-  
199 specific terminology in traditional medicine texts.

200 In applications, knowledge graphs have demonstrated value. For example, a  
201 knowledge graph on natural products connects segments of tandem MS to predicted  
202 metabolites and those predicted metabolites to potential generating genes, as  
203 implemented in frameworks like the Experimental Natural Products Knowledge Graph  
204 (ENPKG), which integrates multimodal data for plant-derived compounds(37). Recent  
205 efforts further demonstrate this by leveraging AI to associate MS fragmentation patterns  
206 with biosynthetic gene clusters through substructure discovery and BGC-metabolite  
207 mapping(38). This graph can emulate an expert chemist's intuition, mining



208 correlations that yield novel antivirals. In the textual side, NLP methods are being  
209 applied to the vast literature on medicinal plants and traditional therapies. One of them  
210 constructed TCMBank, one of the largest integrative databases associating TCM with  
211 multi-omics data, based on text-mining historical herb manuals and medical texts. The  
212 system, employing advanced NLP techniques initially based on bidirectional LSTM  
213 networks and conditional random fields with subsequent enhancements incorporating  
214 Transformer-based models, amassed structured data on herbs, chemicals and known  
215 effects from dozens of ancient manuscripts(39). This “knowledge reconstruction” re-  
216 works past empirical material into a machine-readable database enabling the AI to  
217 rapidly sift through potential antiviral medicines in conventional literature and then  
218 match them against prevailing biomedical information.

219 By transforming fragmented knowledge into connected, machine-readable data  
220 through knowledge graphs and curated databases, AI enhances upstream discovery of  
221 natural antivirals. Researchers can now query these systems to identify promising  
222 compounds and targets far more efficiently than manual curation. Nevertheless, the  
223 field must address NPs-specific hurdles—such as textual ambiguity in ancient sources,  
224 data heterogeneity, and the persistent validation gap—to ensure reliable, translational  
225 impacts in antiviral drug development.

#### 226 **2.1.4 AI-Driven Target Identification**

227 Another essential upstream step is targeting molecular targets (viral or host) for natural  
228 antivirals. AI accelerates this process through a combination of data-driven and  
229 structure-based approaches, enabling more efficient prediction and validation(40). To  
230 achieve comprehensive coverage, we organize AI-driven target identification into a  
231 trinity framework: Chemical-centric, Systems-centric, and Physics-centric. This  
232 structure addresses ligand-based, network-based, and structure-based paradigms in  
233 pharmacology, while incorporating emerging techniques to fill critical gaps such as  
234 phenotypic profiling, metabolomic interference, and dynamic simulations.



235 (i) Chemical-centric strategies focus on ligand-chemical space using deep transfer  
236 learning and matrix factorization to associate "orphan ligands" with known targets,  
237 including phenotypic/image-based AI (e.g., Cell Painting assays) that enables forward  
238 pharmacology by inferring pathways from cellular morphological fingerprints without  
239 prior ligand-target data(41). For instance, the STarFish platform, a stacked ensemble  
240 model, identifies potential targets for NPs by leveraging known ligand-target data,  
241 achieving high accuracy in multi-target predictions on benchmark datasets(42).  
242 Similarly, DeepPurpose employs DL for drug-target interaction prediction, facilitating  
243 virtual screening of NPs with improved hit rates(43).

244 (ii) Systems-centric strategies integrate knowledge graphs and multimodal GNNs  
245 to reveal hidden nodes in virus-host interaction networks, integrated with AI-driven  
246 metabolomic analysis (e.g., flux balance analysis in integrated metabolic models) to  
247 predict how NPs alter host metabolic environments, uncovering indirect antiviral  
248 targets(44). For example, graph neural networks have been applied to construct and  
249 analyze SARS-CoV-2 knowledge graphs based on virus-host interactions, pathways,  
250 and drug associations, identifying potential host genes and biological processes for  
251 antiviral drug repurposing(45). Tools like TCMBank leverage NLP and knowledge  
252 graphs for traditional medicine data mining, enabling synergistic target discovery(39).  
253 This dimension also extends to genomic surveillance for pathogen evolution, enhancing  
254 target relevance in dynamic viral contexts(46).

255 (iii) Physics-centric strategies employ generative AI for biophysical predictions  
256 and dynamic simulations. AlphaFold 3 exemplifies this by providing ~50% improved  
257 accuracy in predicting protein-ligand and nucleic acid interactions compared to  
258 physics-based docking(47) and RoseTTAFold All-Atom for all-atom modeling of  
259 protein-ligand dynamics(48).

260 Overall, these AI technologies—from knowledge graphs to predictive models—  
261 hold significant potential to streamline upstream discovery by connecting chemical



262 leads with biological targets, paving the way for next-generation antiviral drugs.  
263 Notably, polypharmacology represents a core advantage of NPs: their multi-target  
264 effects enable synergistic therapeutic outcomes, and AI shows emerging potential to  
265 actively design and quantify these synergies or antagonisms for optimized efficacy(49).  
266 However, challenges persist, including data bias in training sets (e.g.,  
267 underrepresentation of rare interactions leading to high false-positive rates in a data-  
268 starved setting) and the experimental-computational gap, necessitating rigorous  
269 validation and diverse datasets to mitigate biases and improve generalization.

## 270 **2.2 Midstream: Drug Screening and Optimization**

### 271 **2.2.1 Virtual Screening and Lead Discovery**

272 In the screening and lead identification phase, AI has significantly enhanced processes  
273 by enabling ultra-efficient virtual screening of enormous chemical libraries. While  
274 traditional high-throughput wet lab screens are often resource-intensive, AI in silico  
275 methods can evaluate billions of compounds more rapidly. However, the effectiveness  
276 of these in silico screenings heavily relies on the quality and maturity of the underlying  
277 machine learning models, such as the accuracy of training data and model  
278 generalization to avoid common pitfalls like overfitting or data biases(50, 51).  
279 Quantitative structure–activity relationship (QSAR) models play a central role in this,  
280 with performance that is nuanced and contingent upon the available data regime. In  
281 high-data regimes, modern approaches utilizing GNNs and transformers excel at  
282 learning complex molecular features from structural information, eliminating the need  
283 for manual descriptor selection and improving prediction accuracy for large  
284 datasets(52). For example, DL-QSAR models integrating molecular fingerprints and  
285 GNN-derived features can accelerate antiviral activity prediction and prioritize natural  
286 product analogs(53). In contrast, in low-data regimes—common in natural product  
287 antiviral discovery—classic techniques such as tree-based models (e.g., random forests)  
288 combined with circular fingerprints often perform comparably or better, offering



289 robustness, simplicity, and superior generalization with fewer samples(51, 54, 55).  
290 Hybrid strategies blending classic and advanced methods may yield optimal results  
291 across diverse scenarios, balancing computational efficiency and reliability.

292 In structure-based virtual screening, traditional physics-based molecular docking  
293 faces prohibitive computational costs when traversing billion-scale chemical spaces,  
294 necessitating AI-driven strategies that optimize both accuracy and throughput while  
295 addressing limitations in generalization, especially for structurally complex NPs. Key  
296 facets of this paradigm shift encompass enhancing evaluation precision through DL-  
297 driven affinity estimation—exemplified by curvature-based GNNs such as CurvAGN,  
298 which capture intricate 3D molecular geometries and multi-scale interactions to  
299 mitigate systematic biases in classical empirical scoring functions(56). To surmount  
300 scalability constraints of exhaustive docking, Deep Docking-type paradigms deploy DL  
301 surrogate models trained on representative subsets to forecast docking scores for the  
302 remainder of the library, thereby excluding over 99% of non-binders without explicit  
303 physics-based computations—a capability indispensable for trillion-scale ultra-large  
304 chemical library screening(57). This acceleration is further augmented by Active  
305 Learning (AL) strategies, which reconfigure virtual screening into a dynamic  
306 'sampling-docking-training-prediction' iterative loop(58). Through iterative selection  
307 of the most informative compounds, AL frameworks identify top-tier leads while  
308 requiring docking of less than 1% of the library, drastically mitigating resource  
309 demands. Integrating these AI accelerators with robust error-control mechanisms, such  
310 as the Conformal Prediction (CP) methodology, assures high sensitivity and reliability  
311 in ligand discovery for challenging targets like G protein-coupled receptors  
312 (GPCRs)(59). Although these innovations markedly expedite antiviral natural product  
313 screening, persistent challenges include limited model generalization to novel scaffolds  
314 and substantial computational infrastructure requirements.

315 AI also permits multi-target screening, interrogating interactions between  
316 compounds and multiple viral proteins (i.e., polymerase and protease inhibitors), and



317 the imposition of broad-spectrum antivirals with tuned polypharmacology(60). The  
318 synergistic use of generative models, CP-guided docking, and multi objective  
319 optimization allows AI-driven virtual screening to accelerate hit discovery while  
320 increasing chemical diversity when compared to brute-force approaches in terms of  
321 both speed and lead quality.

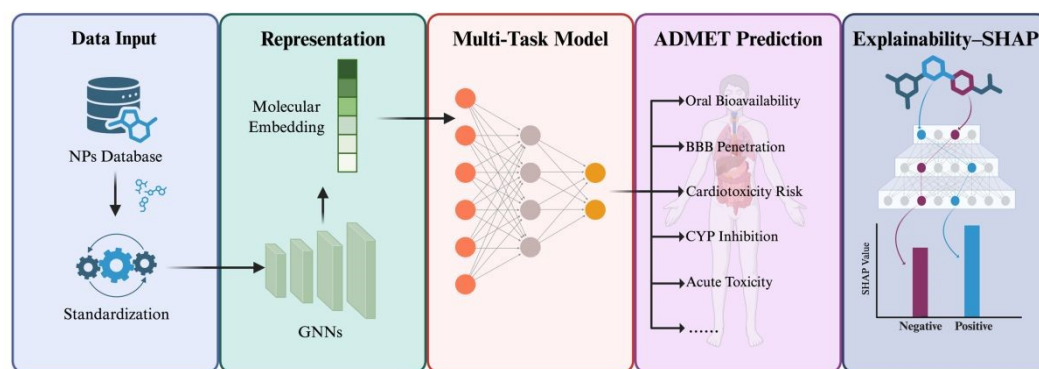
### 322 2.2.2 AI-Enhanced Pharmacokinetics and Toxicity Prediction

323 The optimization of pharmacokinetic (PK) and toxicity profiles has traditionally  
324 represented a high-attrition, late-stage bottleneck in drug development; however, AI is  
325 increasingly transforming this process into an early-stage, parallel Multi-Parameter  
326 Optimization (MPO) paradigm. Rather than treating ADMET (Absorption, Distribution,  
327 Metabolism, Excretion, and Toxicity) as a sequential experimental filter, multi-task  
328 learning algorithms now enable the simultaneous prediction of diverse drug-likeness  
329 endpoints from a single molecular representation (**Fig. 3**)(61, 62). This holistic  
330 evaluation facilitates the navigation of complex trade-offs—such as balancing oral  
331 bioavailability against cardiotoxicity risks—during the hit-to-lead transition(63, 64).  
332 Furthermore, the integration of transfer learning addresses data sparsity challenges in  
333 natural product research by adapting animal-derived datasets to human-specific  
334 predictions, while interpretability tools like SHAP (SHapley Additive exPlanations)  
335 offer mechanistic insights by identifying toxicophoric substructures for targeted  
336 medicinal chemistry modifications(65, 66).

337 The industrial applicability of these AI-driven workflows is supported by  
338 documented improvements in throughput and predictive accuracy. For instance,  
339 platforms such as ADMETLab 3.0, which employ directed message-passing neural  
340 networks, have shown the ability to evaluate over 119 endpoints with AUROC values  
341 up to 0.94, contributing to enhanced efficiency in computational screening compared  
342 to traditional empirical models(67, 68). Regulatory horizon-scanning reports, such as  
343 those from the European Medicines Agency (EMA), underscore the deployment of



344 tools like Toxometris.ai and the Deep-PK framework in toxicity and pharmacokinetic  
 345 predictions and preclinical study designs within pharmaceutical pipelines(59, 69).  
 346 These implementations suggest that AI can enhance predictive capabilities and  
 347 potentially de-risk the development of natural antivirals by supporting a more  
 348 streamlined transition from hit identification to clinical candidate nomination.  
 349 Nonetheless, ongoing challenges in model generalization to novel scaffolds highlight  
 350 the need for rigorous validation and hybrid approaches to ensure translational  
 351 reliability(59, 61).



**Fig. 3 Schematic workflow of AI-driven multi-task learning for ADMET prediction in natural product drug development.** The workflow starts with data input from NPs databases. Data standardization follows, normalizing formats (e.g., SMILES) and curating biases. GNNs then generate molecular embeddings by encoding compounds as graphs, capturing NP-specific features like macrocycles. A multi-task model predicts interrelated endpoints simultaneously, such as oral bioavailability, blood-brain barrier (BBB) penetration (e.g., via permeability coefficients), cardiotoxicity risk (e.g., hERG channel inhibition), cytochrome P450 (CYP) enzyme inhibition (e.g., CYP3A4 isoform specificity), and acute toxicity (e.g., LD50 estimates), using shared representations for efficiency. SHAP, a game-theoretic interpretability framework based on Shapley values from cooperative game theory, quantifies feature contributions (positive or negative) to each prediction,



visualizing substructure impacts (e.g., highlighting aromatic rings contributing to hepatotoxicity) to guide targeted structural modifications in lead optimization. Figure created with BioRender.com.

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DOI: 10.1039/D5DD000504C

### 352 **2.2.3 Predicting and Mitigating Antiviral Resistance**

353 Viruses evolve rapidly, posing a persistent challenge to antiviral drug efficacy as  
354 resistance mechanisms emerge, often rendering treatments obsolete within months or  
355 years. Traditional approaches rely on reactive surveillance and empirical testing, but AI  
356 introduces a proactive paradigm by forecasting evolutionary trajectories, identifying  
357 resistance signatures, and guiding resilient inhibitor design.

358 AI models leverage sequence data and structural predictions to anticipate viral  
359 mutations at binding sites, enabling the development of inhibitors that maintain potency  
360 against future variants. For example, EVEscape computationally produced multi-  
361 mutant SARS-CoV-2 spikes to replicate immune escape, with experimental  
362 validation(70). Predicting mutations at binding sites can guide design of inhibitors  
363 resilient to future variants. AI may also search sequence databases for resistance  
364 signatures and offer chemical modifications to bypass common mechanisms(71). These  
365 methods also facilitate chemical modifications, using generative AI to suggest scaffold  
366 alterations that bypass common resistance pathways, shifting from post-resistance  
367 response to preemptive antiviral engineering.

368 Despite these advances, significant dilemmas arise from the interplay of viral  
369 biology and technological limitations. Viruses' high mutation rates create out-of-  
370 distribution challenges for AI models, where training on historical variants may fail to  
371 generalize to novel, phylogenetically divergent strains, leading to inaccurate  
372 predictions and false confidence in drug resilience(46). Computationally, scaling  
373 simulations for multi-mutant landscapes demands immense resources, often exceeding  
374 available infrastructure and introducing biases from incomplete datasets. NPs' structural  
375 diversity offers a vast pool for discovering new scaffolds less prone to resistance, but



376 integrating this with AI remains underdeveloped due to data scarcity in NP-specific  
377 resistance profiles.

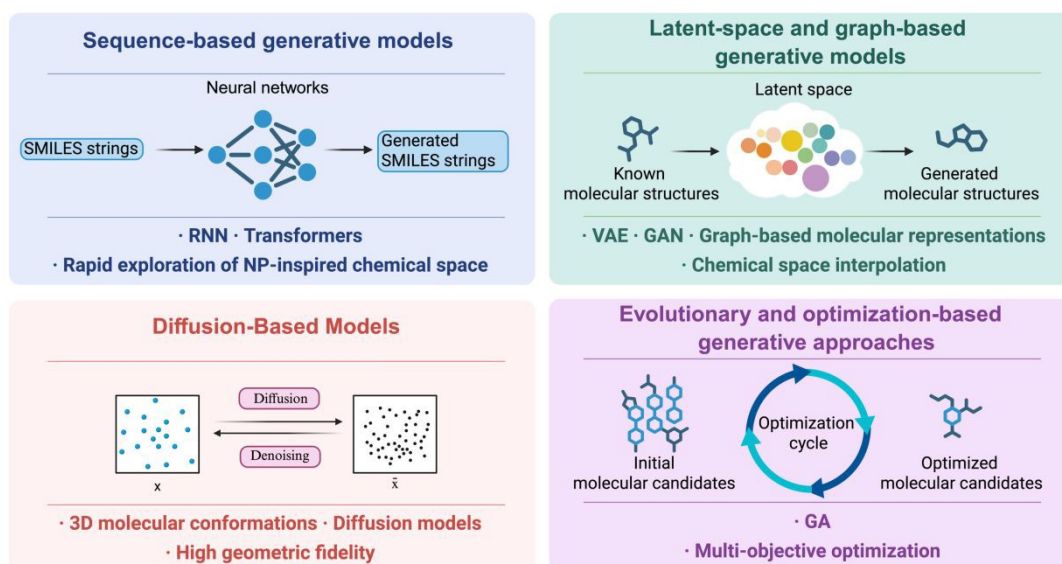
#### 378 **2.2.4 Generative AI for Novel Drug Design and Synthesis Planning**

379 Generative AI serves as an impressive midstream approach in antiviral NPs drug  
380 development, supporting the de novo generation of molecular structures inspired by  
381 NPs' structural diversity and multi-target synergies(72). These models process chemical  
382 patterns from large datasets, such as general chemical databases like ChEMBL (which  
383 includes NPs) or NP-focused libraries like TCMSP and COCONUT, to suggest  
384 candidates that could address viral evolution and resistance issues(73). Generative  
385 frameworks commonly include variational autoencoders (VAEs), generative  
386 adversarial networks (GANs), and diffusion models, often implemented with sequence-  
387 based backbones (e.g., recurrent neural networks (RNNs) or Transformers) or graph-  
388 based structures, allowing exploration of chemical spaces beyond conventional  
389 screening and potentially aiding in timeline efficiency(74). These major generative AI  
390 paradigms can be summarized as follows (**Fig. 4**). They efficiently generate molecular  
391 structures inspired by natural products through mechanisms such as sequence modeling,  
392 latent space exploration, diffusion processes, or evolutionary optimization. Notably,  
393 generative AI generally produces de novo molecules that are NP analogs or mimics,  
394 often struggling to replicate the intricate structural features of authentic NPs, such as  
395 high chirality centers or complex ring systems, yet it can integrate NP-like motifs (e.g.,  
396 macrocycles or polyketide scaffolds) to approximate bioactivity benefits like  
397 polypharmacology, as discussed in recent AI-NP literature(75). This method aims to  
398 align natural-inspired designs with synthetic feasibility, facilitating multi-objective  
399 considerations for aspects like binding affinity, ADMET properties, and accessibility  
400 in antiviral settings.

401 Examining individual architectures, VAEs project molecular structures into latent  
402 spaces for sampling NP-inspired analogs, supporting the development of structurally



403 complex antiviral candidates, though accurately mirroring full NP polypharmacology  
 404 can be difficult(76). GANs utilize adversarial training to yield plausible NP-like  
 405 molecules, such as analogs of artemisinin for possible broad-spectrum antiviral  
 406 exploration, via generator-discriminator refinement(75). Diffusion models construct  
 407 full 3D molecular conformations by progressively denoising atomic coordinates and  
 408 atom types in an equivariant manner, offering advantages in generating geometrically  
 409 accurate NP-mimetic structures that might counter viral mutations(77). RNNs and  
 410 Transformers, as sequence-based backbones, manage formats like SMILES strings to  
 411 produce varied analogs, enabling investigation of NP-adjacent chemical spaces for  
 412 antiviral purposes(78). GAs, meanwhile, simulate evolution via selection and mutation  
 413 of known structures, embedding synthetic accessibility measures to optimize NP  
 414 analogs(74). By linking these architectures with retrosynthesis planning tools like  
 415 AiZynthFinder, generative AI can propose candidate structures alongside practical  
 416 synthesis routes, encouraging an iterative design-make-test process that broadens  
 417 chemical diversity in line with NP bioactivity concepts, although validation against  
 418 diverse viral variants remains a key limitation(79).



**Fig. 4 Overview of major generative AI paradigms applied to natural product–inspired molecular design.** Sequence-based generative models treat molecular



generation as a language modeling problem, in which molecular structures are encoded as SMILES strings and processed by sequence-based neural networks to rapidly explore NP-inspired chemical space. Latent-variable and graph-based generative models embed known molecular structures into a continuous latent space using latent representations and graph-based molecular encodings, enabling interpolation and sampling to generate novel molecular structures. Diffusion-based generative models generate molecular structures by progressively denoising stochastic representations, allowing the reconstruction of high-fidelity three-dimensional molecular conformations. Evolutionary and optimization-based generative approaches iteratively refine molecular candidates through mutation and selection under multi-objective optimization criteria, explicitly incorporating considerations such as chemical properties and synthetic feasibility. Figure created with BioRender.com.

## 419 **2.3 Downstream: Preclinical and Clinical Applications**

### 420 **2.3.1 AI-Driven Preclinical Testing**

421 Before advancing to clinical trials, antiviral natural products must undergo rigorous  
422 preclinical evaluation, including in vitro assays, animal studies, and toxicity  
423 assessments. AI has been integrated into these stages to enhance efficiency, but its  
424 application in NPs—characterized by structural complexity and multi-target  
425 interactions—presents unique challenges. This subsection provides a systematic  
426 overview of AI methods in preclinical testing for antiviral NPs, focusing on  
427 experimental optimization, automated data analysis, and predictive modeling, while  
428 critically discussing limitations and real-world impacts.

429 First, AI facilitates experimental design through optimization algorithms, such as  
430 Bayesian optimization (BO) and active learning. BO iteratively selects experimental  
431 parameters (e.g., compound dosages or combinations) based on prior data to maximize



432 information gain with minimal trials. In the context of antiviral NPs, BO has been  
433 applied to refine in vitro assays for plant-derived compounds like artemisinin analogs  
434 against broad-spectrum antivirals including coronaviruses, achieving comparable  
435 enrichment factors while reducing the experimental footprint by approximately 40-50%  
436 compared to traditional methods(80). Similarly, active learning frameworks combine  
437 machine learning with high-throughput screening to prioritize NPs from microbial  
438 sources, such as polyketides discovered through genome mining with anti-viral  
439 potential(81). These approaches accelerate preclinical workflows by intelligently  
440 navigating the vast chemical space of NPs.

441 Second, AI enhances automated analysis of preclinical data, particularly in imaging  
442 and histopathology. DL models, such as CNNs, automate the scoring of pathology  
443 slides from animal models. For instance, the CSGO (Cell Segmentation with Globally  
444 Optimized boundaries) pipeline has recently been validated for high-throughput,  
445 whole-cell segmentation in Hematoxylin-and-Eosin (H&E)-stained tissues, enabling  
446 precise quantification of inflammatory cell infiltration in lung injury models(82). Such  
447 methodologies provide objective, high-resolution readouts for evaluating the  
448 therapeutic efficacy of NP candidates, such as quercetin, in alleviating virus-induced  
449 pulmonary inflammation. This automation significantly reduces inter-observer  
450 variability and ensures the reproducibility of toxicity and efficacy assessments for  
451 natural antivirals in preclinical animal trials.

452 Third, AI-driven predictive modeling, including physiologically based PBPK  
453 models augmented with ML, bridges in vitro data to in vivo outcomes. These models  
454 predict pharmacokinetics (e.g., clearance, half-life) for NPs by integrating  
455 physicochemical properties and multi-omics data. A notable application involves  
456 forecasting human systemic exposure for plant-derived antivirals like berberine. For  
457 instance, mechanistic PBPK models have been refined to capture complex interactions  
458 between berberine and multiple transporters (e.g., P-gp and OCTs), providing a  
459 quantitative framework to evaluate its therapeutic potential against viral infections,



460 particularly in the context of drug-drug interactions (DDI)(83). Furthermore, ML-  
461 enhanced PBPK has demonstrated significant superiority, reducing prediction errors  
462 (e.g., RMSE) by 20–30% compared to empirical scaling methods, as shown in small-  
463 molecule validations(84). Although primarily validated in targeted therapeutics like  
464 oligonucleotides(85), the transfer learning strategies utilized for cross-species  
465 translation are increasingly being adapted to NPs, facilitating more accurate animal-to-  
466 human extrapolations for complex natural compounds and offering methodological  
467 insights for antiviral drug development.

468 However, despite these advancements, AI in preclinical testing for antiviral NPs  
469 faces significant failure modes. Recent high-profile blind challenges, such as the  
470 CACHE series and the ASAP-Polaris initiatives, have exposed a stark reality: the  
471 majority of AI-prioritized compounds fail to exhibit reproducible activity in wet-lab  
472 assays, with failure rates exceeding 90% in many cases(86, 87). During the COVID-19  
473 pandemic, the “rush to screen” led to an influx of low-quality in silico studies where  
474 herbal compounds like quercetin were frequently identified as hits. Retrospective  
475 analyses now confirm these were largely false positives—often due to the molecules  
476 being Pan-Assay Interference Compounds (PAINS) or the AI overestimating binding  
477 affinities by neglecting complex solvation effects and structural plasticity(87). These  
478 failures underscore a systemic translational gap, necessitating a shift toward more  
479 rigorous, physics-informed AI models for natural products.

### 480 2.3.2 AI Innovations in Clinical Trials

481 Antiviral natural products drug, like other therapeutics, undergo expensive and lengthy  
482 clinical trial phases. Artificial intelligence is beginning to offer targeted improvements  
483 in efficiency and informativeness, though many applications remain in early stages or  
484 face substantial limitations in real-world translation. Several AI-based methodologies  
485 are under exploration in Phase I–III studies for antiviral agents:



486 (i) Federated learning (FL) for multi-center data integration — FL enables  
487 collaborative model training across institutions without centralizing sensitive patient  
488 data, thereby preserving privacy while leveraging diverse cohorts. A landmark example  
489 is its application to predict clinical outcomes in COVID-19 patients from multiple  
490 hospitals, demonstrating improved generalizability across heterogeneous  
491 populations(88). Although this approach has not yet been widely reported for antiviral  
492 NP trials, it holds methodological promise for modeling disease trajectories or  
493 treatment responses in multi-ethnic or multi-risk-group settings, provided data  
494 harmonization and model robustness challenges are addressed.

495 (ii) NLP for unstructured clinical data — NLP algorithms can extract relevant  
496 outcomes, adverse events, or symptom patterns from electronic health records,  
497 physician notes, and free-text entries in near real-time. Systematic reviews indicate that  
498 NLP enhances signal detection in clinical decision support and could support more  
499 responsive monitoring in trials(89). In the context of antiviral NPs, such tools may  
500 facilitate earlier identification of efficacy or safety signals in adaptive trial designs,  
501 although current implementations are largely limited to general medical contexts rather  
502 than NP-specific endpoints.

503 (iii) Synthetic control arms and generative models — Generative AI methodologies,  
504 particularly GANs, is being explored to create synthetic patient data (SPD) to augment  
505 or partially replace traditional control arms. Early benchmarks using tabular clinical  
506 datasets demonstrate that GAN-based frameworks, such as GANerAid, can synthesize  
507 patient-level records that preserve the complex statistical correlations and longitudinal  
508 trajectories of actual trial participants(90). This approach is particularly promising for  
509 establishing synthetic control arms in rare disease or oncology trials, where simulating  
510 survival data (e.g., progression-free survival) can reduce the reliance on large placebo  
511 cohorts while maintaining statistical power(91). However, applications to antiviral NPs  
512 remain nascent, with significant challenges in ensuring biological plausibility—the risk  
513 that GANs may generate pharmacologically impossible trajectories for multi-



514 component natural extracts. Furthermore, regulatory acceptance requires rigorous  
515 validation against "hallucinated" correlations and the avoidance of bias amplification  
516 in complex patient profiles.

### 517 **3. Bottlenecks In Ai-Enabled Antiviral NPs Development**

#### 518 **3.1 Data constraints: NPs complexity and viral dynamics**

519 Data scarcity and heterogeneity are the key obstacles. NPs data are multi-modal,  
520 complex, and unevenly standardized across sources and formats(92). High-quality,  
521 empirically validated annotations for structures, biosynthetic processes, antiviral  
522 activity (including negatives), and toxicity are limited. Multi-target/network  
523 pharmacology mechanisms are tough to capture adequately, hindering mechanistic  
524 modeling. Knowledge integrated in traditional medicine literature is valuable; however,  
525 it is unstructured, archaic, and philosophically complex, challenging NLP and  
526 knowledge-graph development(93).

527 Viral evolution data are limited in sample size and dynamic. For new strains or  
528 unexpected resistance mutations, establishing robust models is challenging(94). Rapid  
529 genomic change needs frequent model updating(46). Many viruses replicate within  
530 liquid–liquid phase-separated (LLPS) condensates, which lack well-defined structural  
531 targets, complicating design; changing sequences may affect condensate  
532 physicochemical features and dynamics(95). Host-target inhibition may generate  
533 compensating responses that current AI rarely can foresee(49). Addressing such  
534 “moving targets” requires models that spot dynamic or allosteric sites or even intervene  
535 in phase behavior, potentially via system-level “digital twins”—well beyond existing  
536 capabilities.

#### 537 **3.2 Model limitations: from pattern recognition to biological understanding**



538 AI's advancements primarily rely on pattern recognition, but deep biological and  
539 chemical understanding for intricate NPs structures and polypharmacology remains  
540 inadequate. NPs feature intricate ring structures, multiple chiral centers, and flexible  
541 conformations; capturing fine-grained conformational dynamics and flexible protein–  
542 ligand interactions is difficult(76). Even employing AlphaFold 3, difficulties exist in  
543 modeling dynamics, allostery, and binding of unusual ligands, which can affect docking  
544 accuracy(96). NPs frequently act via several viral and host targets, forming complex  
545 networks; current AI struggles to interpret synergy or antagonism or predict system-  
546 level ramifications. Multi-component traditional formulations confront extra “black-  
547 box” issues that limit optimization and sensible combination design(97). For swiftly  
548 growing viruses, generalization to out-of-distribution variations is limited; even single-  
549 point mutations might have structural ramifications that models fail to capture(98).

### 550 **3.3 Synthetic accessibility**

551 Bridging computational hits to physical molecules remains a formidable hurdle for NPs.  
552 While NPs benefit from innate bioactivity and initial accessibility through extraction or  
553 fermentation, scaling their production to meet clinical or industrial demands reveals  
554 significant bottlenecks. Direct large-scale extraction from natural sources is often  
555 unsustainable due to resource scarcity, seasonal variability, environmental degradation  
556 (e.g., overharvesting), and extremely low yields (frequently below 0.01% dry  
557 weight)(6). Similarly, native microbial fermentation is hampered by the low  
558 productivity of wild-type strains and difficult-to-control fermentation conditions,  
559 though modern strain improvement and optimization techniques (e.g., fed-batch or  
560 continuous fermentation) can enhance productivity(99).

561 Heterologous biosynthesis has emerged as a transformative strategy, but metabolic  
562 engineering still faces a "scale-up gap," where titer optimization often stalls at the mg/L  
563 level, failing to reach the g/L threshold required for commercial viability(100). For  
564 molecules with extreme structural complexity, semi-synthesis offers a middle ground,



565 yet it remains constrained by limited reagent-accessible space and high operation  
566 costs(101).

## 567 **4. Outlook And Future Directions**

### 568 **4.1 High-quality, multi-modal benchmark datasets:**

569 To overcome data scarcity and fragmented evaluation, the field needs FAIR (Findable,  
570 Accessible, Interoperable, Reusable), high-quality, multi-modal benchmark datasets.  
571 These should integrate NPs structures, biosynthetic pathways, multi-dimensional  
572 bioactivity profiles (including negative results), and ADMET properties, along with  
573 standardized metrics for fair model comparison and industrial translation.

574

### 575 **4.2 Model and algorithmic innovation**

576 Beyond correlational prediction, future AI should prioritize interpretability (e.g.,  
577 explainable AI, XAI) and robust out-of-distribution generalization to novel biological  
578 systems (e.g., new variants). Incorporating causal inference and neuro-symbolic  
579 techniques may facilitate a move from pattern recognition to scientific discovery,  
580 yielding better mechanistic understanding.

581

### 582 **4.3 Stronger experimental closed loops and high-throughput validation**

583 To match AI's high-throughput hypothesis development, experimental validation must  
584 accelerate. Automated platforms (self-driving laboratories) and active learning can  
585 connect in silico forecasts tightly with wet-lab feedback during DBTL cycles,  
586 eliminating the compute-experiment gap.

587

### 588 **4.4 Cross-disciplinary collaboration and ecosystem building**

589 Antiviral NPs discovery needs fundamental integration across chemistry, biology,  
590 medicine, and data science. Training cross-domain competence and promoting  
591 academia-industry-international partnerships will enable secure data exchange (e.g.,



592 privacy-preserving federated learning). Building an automated and intelligent end-to-  
593 end pipeline—from data collection and modeling to candidate generation and  
594 validation—will require continual investment but is essential to realize AI's full  
595 promise in antiviral NPs discovery.

## 596 **5. Conclusion**

597 Research on antiviral drugs today faces significant difficulties: expensive, time-  
598 consuming, and limited against rapidly evolving viruses. NPs are still hindered in their  
599 analysis, target detection, and synthesis because of their structural diversity and multi-  
600 target synergy. AI is a powerful new paradigm that is changing how the drug discovery  
601 process is conducted and through the integration of data and algorithmic learning.

602 In the upstream phase, AI helps to refine genome mining and the modeling of  
603 biosynthetic gene clusters, while knowledge graphs and natural language processing  
604 aid in extracting insight from conventional medical databases. AI-driven virtual  
605 screening and generative design during the midstream phase significantly expand  
606 chemical space exploration, yielding multi-target antiviral candidates with optimal  
607 pharmacokinetics and resilience against resistance. On the downstream, AI optimizes  
608 preclinical models and clinical trial efficiency using techniques such as federated  
609 learning and synthetic control arms, which accelerates translational outcomes.

610 There are many critical issues, including data scarcity, limited model  
611 interpretability, and synthetic accessibility challenges, necessitating integrated  
612 solutions to advance the field. The complexity of NPs and viral dynamics demands  
613 FAIR, multi-modal benchmark datasets that unify structural, biosynthetic, and  
614 bioactivity data, while interpretable AI and causal inference are critical to move beyond  
615 pattern recognition toward mechanistic understanding of polypharmacology and  
616 dynamic targets like liquid-liquid phase-separated condensates. Automated platforms,  
617 such as self-driving laboratories, must bridge the compute-experiment gap through



618 high-throughput Design-Build-Test-Learn cycles, embedding synthetic feasibility early  
619 in the design process. By fostering cross-disciplinary collaboration and privacy-  
620 preserving data-sharing ecosystems, the field can build an intelligent, end-to-end  
621 pipeline, transforming reactive antiviral NP discovery into a proactive, resilient strategy  
622 against evolving viral threats.

623 Although not a cure all, AI represents the best catalyst for converting NP-based  
624 antiviral discovery from empirical serendipity to predictive engineering. If adopted  
625 more widespread—as long as there's solid validation and international data-sharing  
626 agreements—the virus's arsenal of antiviral responses may be unleashed. We are on the  
627 cusp of the fourth drug discovery revolution enabled by symbiotic human-AI  
628 collaboration, where the speed of machines meets the smarts of chemical evolution to  
629 move beyond the capacity for viral adaptation.

### 630 **Acknowledgments**

631 We would like to express our profound gratitude to all participants for their invaluable  
632 contributions to this research. Additionally, we appreciate the assistance of  
633 BioRender.com in creating the figure for this study.

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View Article Online  
DOI: 10.1039/D5DD00504C

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## Data Availability Statement

View Article Online  
DOI: 10.1039/D5DD00504C

This article is a review, and as such, does not report any new primary data. All data and information discussed are available in the original publications and sources cited within the manuscript's references section

