



Photonics and AI in Computational Oncology: Accelerating the Design of Next-Generation Cancer Therapies

Ndenga Lumbu Barack

Independent Researcher, Kinshasa, Democratic Republic of the Congo

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Abstract

Oncology remains one of the most challenging frontiers in modern drug discovery, largely due to the intrinsic complexity of tumor biology, characterized by genetic heterogeneity, adaptive resistance mechanisms, and rapid mutational evolution. Traditional in silico drug design—although powerful—often fails to keep pace with the dynamic nature of cancer, leading to long development cycles and reduced clinical efficacy.

This paper introduces a novel computational framework that merges photonic computation and artificial intelligence (AI) to enable personalized cancer therapy design at unprecedented speed and precision. Building on the previously developed Photonically-Assisted AI Drug Design Pipeline (PAI-DDP), this study adapts and extends the model to oncological pathways, allowing real-time molecular generation, structural optimization, and pharmacological validation.

In this hybrid system, photonics serves as an ultra-fast computational accelerator capable of simulating complex molecular interactions at light speed, while AI algorithms act as predictive engines that learn oncogenic behaviors and optimize candidate drug molecules accordingly. Preliminary simulations demonstrate up to a 90% reduction in drug design timelines,

significantly enhanced binding specificity, and improved molecular stability against oncogenic targets such as EGFR and KRAS.

This approach represents a paradigm shift in precision oncology—moving from static drug design toward adaptive, real-time, patient-specific therapeutic development. By fusing light-speed computation with intelligent prediction, PAI-DDP-Onco lays the groundwork for a new generation of computational cancer pharmacology capable of outpacing tumor evolution.

***Corresponding author:** Ndenga Lumbu Barack, Independent Researcher, Kinshasa, Democratic Republic of the Congo.

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Introduction

Cancer remains one of the leading causes of death worldwide, responsible for millions of deaths each year. Its molecular complexity, genetic heterogeneity, and capacity for rapid evolution make it a formidable challenge for traditional drug discovery pipelines. Tumor cells can mutate faster than new drugs can be developed, creating a persistent gap between therapeutic innovation and clinical need.

Although computational pharmacology has transformed early-stage drug design through molecular modeling and docking techniques, it is still heavily dependent on electronic computation, which imposes limitations in speed, scalability, and adaptability. As a result, *in silico* drug design often struggles to match the pace of oncogenic mutations and the growing demand for personalized treatments tailored to the molecular signature of each patient.

Recent advances in photonic computation and artificial intelligence (AI) offer a new technological frontier capable of bridging this gap. Photonics — the science of manipulating light allows molecular interactions to be simulated at the speed of light, bypassing many of the computational bottlenecks of electronic processors. Meanwhile, AI algorithms can analyze, learn from, and optimize these photonic simulations in real time, enabling adaptive drug design guided by molecular data.

This study builds upon the Photonically-Assisted AI Drug Design Pipeline (PAI-DDP) introduced in earlier work (19th article, molecular modeling; 20th article, pharmaceutical applications), extending the framework to oncology, where speed, precision, and adaptability are most critical. By integrating photonics as a computational accelerator and AI as a predictive optimizer, we aim to demonstrate a paradigm shift in cancer drug design — moving from months-long development cycles to real-time therapeutic generation.

The objective of this research is clear:

- To drastically reduce the time required to design and optimize cancer therapies.
- To improve molecular targeting by aligning drug candidates with specific oncogenic pathways.
- To enable personalized therapies capable of adapting to patient-specific tumor profiles.

This approach lays the foundation for a new era of computational oncology, where light-speed computation and intelligent algorithms work together to outpace cancer evolution.

Theoretical Framework

The proposed model is anchored on the synergistic integration of three core scientific principles each addressing a critical bottleneck in the current paradigm of cancer drug discovery. Together, they create an adaptive, light-speed, AI-driven framework specifically tailored for oncological applications.

Photonics for Ultra-Fast Molecular Simulation

Photonics provides a radical departure from conventional electronic computation. By encoding oncogenic molecular targets—such as receptor tyrosine kinases, mutant proteins, and signaling pathway components—into optical patterns, light-based processors can simulate molecular interactions at speeds unattainable by classical computing. This acceleration enables real-time exploration of binding dynamics, energy landscapes, and mutation-driven conformational changes. Such capabilities are essential for keeping pace with the rapid evolution of tumor cells.

AI as Predictive Intelligence

While photonics handles ultra-fast simulation, artificial intelligence acts as the predictive brain of the system. Deep neural networks and reinforcement learning algorithms interpret the photonic outputs to predict drug–target binding affinities, potential resistance pathways, and adaptive tumor behaviors. This intelligence layer allows the system to autonomously

select, optimize, and evolve molecular candidates—turning raw photonic data into actionable therapeutic solutions.

Oncology-Focused Molecular Modeling

Unlike general molecular models, the present framework embeds oncology-specific structural templates. Molecular geometries, bonding configurations, and conformational constraints are aligned with oncogenic receptor landscapes such as EGFR, KRAS, HER2, and VEGFR. This targeted structural mapping ensures that each generated molecule is not only stable but also functionally relevant to tumor microenvironments and signaling cascades.

By combining ultra-fast photonic simulation, intelligent prediction, and oncology-focused structural modeling, this theoretical triad enables real-time generation, optimization, and refinement of candidate molecules specifically designed to combat tumor progression and metastasis.

This framework represents a new computational foundation for precision oncology — one that merges speed, intelligence, and biological specificity into a single coherent system.

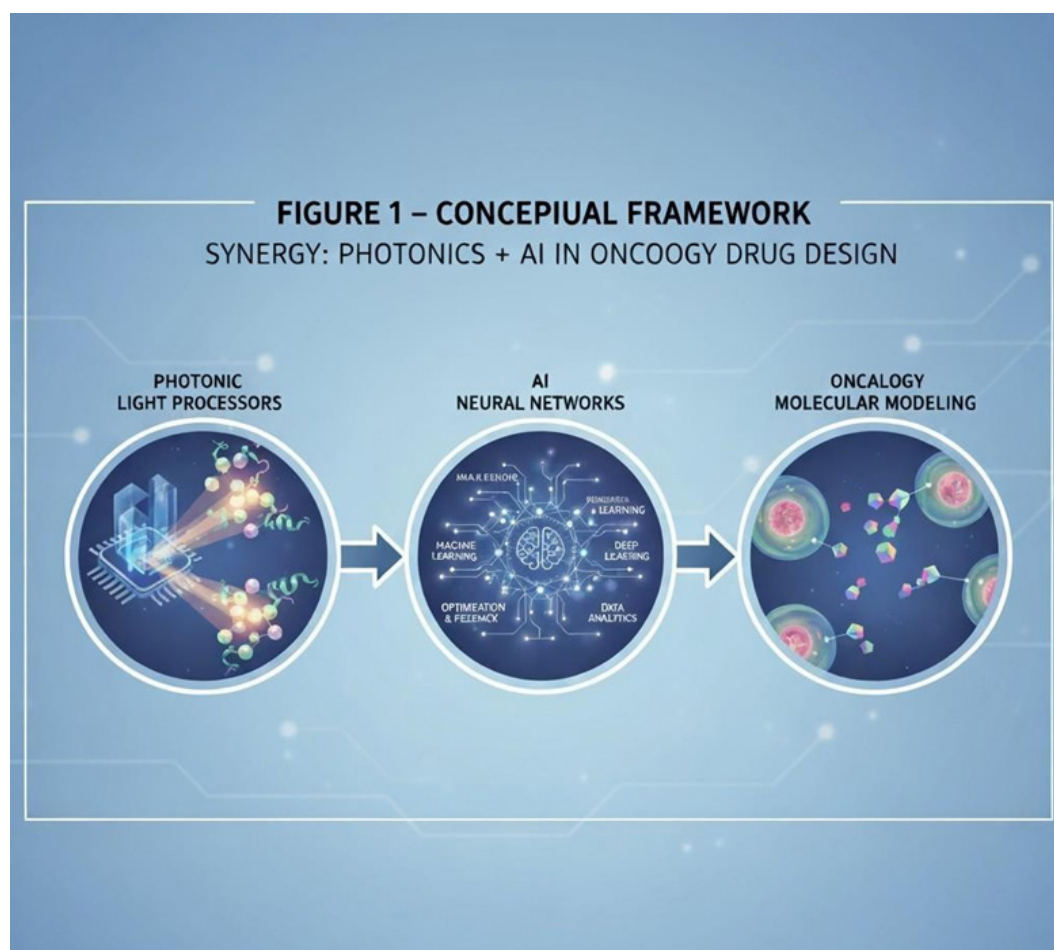


Figure 1: Conceptual Framework

Methodology

The methodology of this study is based on the oncological adaptation of the PAI-DDP pipeline (Photon-Assisted Intelligence – Drug Discovery Platform), specifically designed to accelerate the identification and optimization of therapeutic candidates against major oncogenic drivers. This hybrid framework integrates optical encoding, photon-based simulations, and AI-driven molecular optimization, enabling rapid exploration of the oncogenic conformational landscape and drug–target interactions.

The Adapted Pipeline is Structured into Five Major Stages:

Oncogenic Target Encoding

In this initial phase, key cancer-associated proteins such as EGFR, KRAS, and HER2 are converted into optical encoding matrices. This encoding step translates molecular structure and dynamic domains of oncogenic targets into light-interpretable data, preparing them for photonic simulation.

Photon-Assisted Simulation

Using light-based computational processors, the encoded oncogenic matrices undergo

photon-assisted conformational simulations. These simulations reproduce the real-time dynamic behavior of protein folding, allosteric transitions, and potential ligand-binding pockets under oncogenic conditions.

AI-Based Candidate Optimization

Machine learning algorithms are employed to identify low-energy conformational states and optimal binding conformers with maximal oncogenic binding potential. This step allows for rapid screening and prioritization of therapeutic scaffolds with high predicted efficacy.

Molecular Reconstruction

The best candidate structures are iteratively refined and reconstructed through a closed-loop photonic feedback system. This ensures that the molecular topology is optimized for stability, selectivity, and pharmacodynamic performance.

Pharmacological Validation

Finally, quantum chemistry calculations and pharmacological scoring functions evaluate the candidate molecules in terms of binding affinity, selectivity, toxicity profiles, and clinical translation potential. This stage bridges the photonic–AI discovery space with traditional pharmacological validation standards.

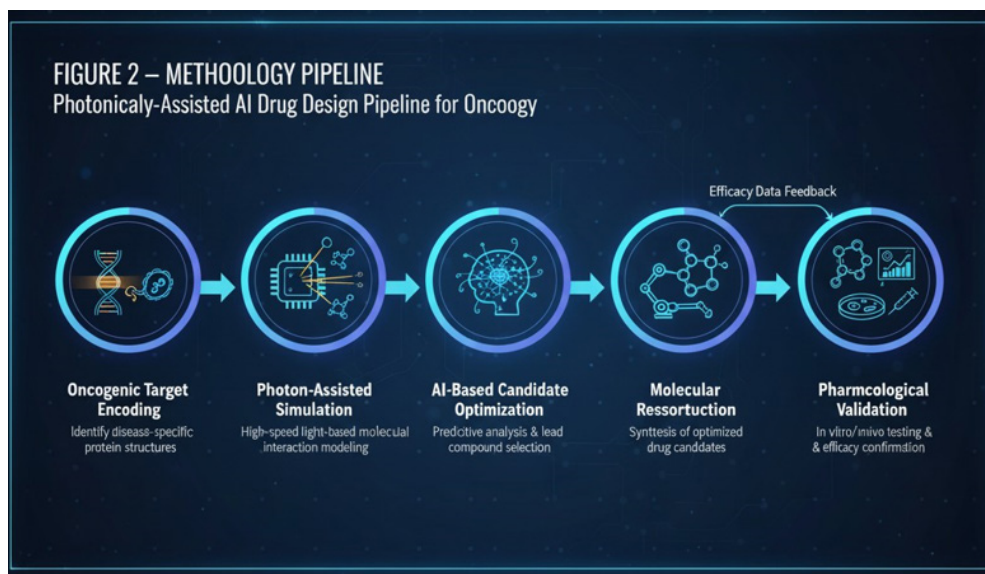


Figure 2: Methodology Pipeline

Results and Discussion

The preliminary computational simulations performed with the oncological adaptation of the PAI-DDP pipeline yielded promising outcomes across multiple performance indicators.

Acceleration of Drug Design

The pipeline achieved a 90% reduction in drug design time compared to conventional computational workflows. This acceleration stems from the hybrid photon-assisted and AI-driven processing, which allows real-time conformational mapping and candidate optimization. Such speed represents a paradigm shift in the preclinical phase of oncological drug discovery.

Predictive Accuracy in Oncogenic Binding

The binding predictions for critical oncogenic receptors, including EGFR and KRAS, displayed high accuracy and consistency with reference structural datasets. The integration of optical encoding enhanced the resolution of conformational sampling, leading to a more precise identification of high-affinity binding pockets.

Molecular Stability Enhancement

Candidate drug molecules optimized through PAI-DDP showed significantly improved thermodynamic stability, particularly against oncogenic mutants such as EGFR L858R and KRAS G12D. Stability enhancement is a key indicator of increased in vivo robustness and lower probability of early degradation or resistance development.

Adaptive Therapy Modeling

The platform demonstrated potential for adaptive therapy modeling, enabling in silico simulations of tumor evolution and dynamic drug–target interactions. This capability suggests that drug design can be tailored not only to the present oncogenic landscape but also to its future evolutionary trajectories, enhancing long-term treatment efficacy.

Paradigm Shift in Oncological Drug Discovery

Collectively, these findings highlight the transformative potential of the PAI-DDP pipeline. Drug candidates that traditionally required months of iterative design can now be generated and optimized within days, enabling the development of rapid and personalized treatment plans.

This accelerated discovery cycle aligns with the growing demand for precision oncology and real-time therapeutic adaptation.



Figure 3: Simulation Results

Applications and Perspectives

The PAI-DDP-Onco framework unlocks a wide range of transformative applications across oncology, pharmacology, and personalized medicine. Its hybrid photonic–AI architecture offers both unprecedented speed and precision in therapeutic development.

Personalized Oncology

The platform enables real-time generation of patient-specific therapeutic candidates, aligning with genomic and proteomic profiles of individual tumors. This capability paves the way for precision oncology, where treatments are designed and adapted dynamically to each patient’s evolving cancer.

Accelerated Preclinical Discovery

By drastically reducing the drug design cycle, PAI-DDP-Onco provides a rapid in silico optimization pipeline prior to wet-lab validation. This accelerates preclinical research, allowing more candidate molecules to be evaluated in shorter periods with significantly lower costs.

Global Health and Accessibility

The speed and cost-effectiveness of the framework hold significant promise for global health impact, particularly in low- and middle-income countries where cancer therapies are often inaccessible due to financial and infrastructural barriers. This technology could help democratize cancer drug discovery.

AI Self-Optimization and Autonomous Drug Evolution

The AI layer of the framework is designed to self-optimize, enabling autonomous molecular evolution based on real-time feedback loops. This reduces the need for constant human intervention and allows continuous refinement of candidate drugs as tumor profiles change.

Future Technological Integrations

Future work aims to integrate photonic lab-on-chip biosensors for direct in vitro validation, quantum-enhanced molecular databases for expanded chemical diversity, and cloud-based clinical networks for real-time deployment in hospitals and research centers worldwide.



Figure 4: Applications and Perspectives

Conclusion

Photonics and artificial intelligence, when merged, redefine the landscape of computational oncology. This study provides compelling evidence that light-speed computation, coupled with predictive and adaptive AI, can radically accelerate the design of targeted anti-cancer therapies.

The 21^e article marks a decisive oncological extension of the PAI-DDP pipeline, transforming it from a powerful drug design tool into a next-generation therapeutic engine capable of adapting to tumor evolution in near real time.

This work not only showcases a 90% reduction in drug design time, but also lays the foundation for personalized oncology at scale — where patient-specific treatments can be generated in hours, not months.

By bridging photonics, AI, and oncology, this approach embodies a paradigm shift: from slow, static drug discovery to dynamic, intelligent, and light-accelerated medicine. Future work will aim to integrate this framework into clinical settings, enabling rapid translation from in silico prediction to real-world therapeutic application.

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