

Cite this as: Yan HUANG, Jialin YE, Lan XU, Xingjie HU, Nan CHEN, 2026. Dual-functional nanoplatform for simultaneous degradation of circRNA *CDR1as* and real-time monitoring of miR-7 in live cells. *Journal of Zhejiang University-SCIENCE B*, 27(7):761-774. <https://doi.org/10.1631/jzus.B2500787>

Dual-functional nanoplatform for simultaneous degradation of circRNA *CDR1as* and real-time monitoring of miR-7 in live cells

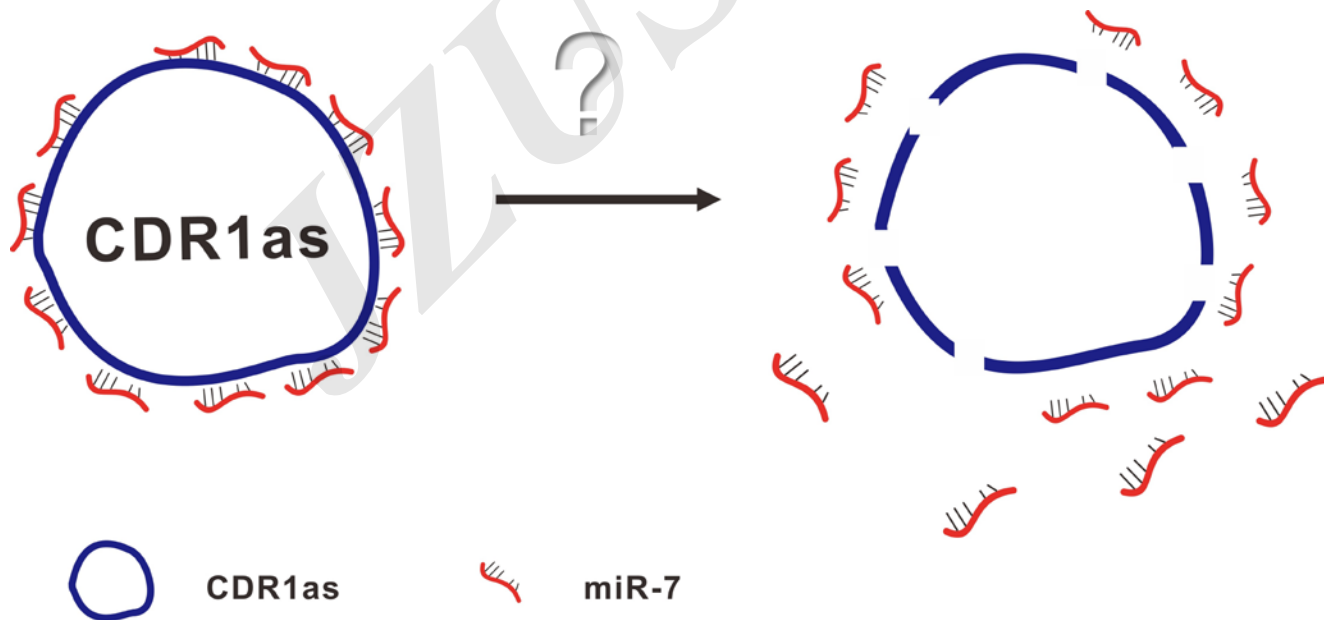
Yan HUANG, Jialin YE, Lan XU, Xingjie HU*, Nan CHEN*

- College of Chemistry and Materials Science, Shanghai Normal University, Shanghai 200234, China
- School of Public Health, Shanghai Jiao Tong University School of Medicine, Shanghai 201318, China

Key words: Circular RNA (circRNA); Zeolitic imidazolate framework-8 (ZIF-8); MicroRNA-7 (miR-7); DNAzyme; Gene regulation

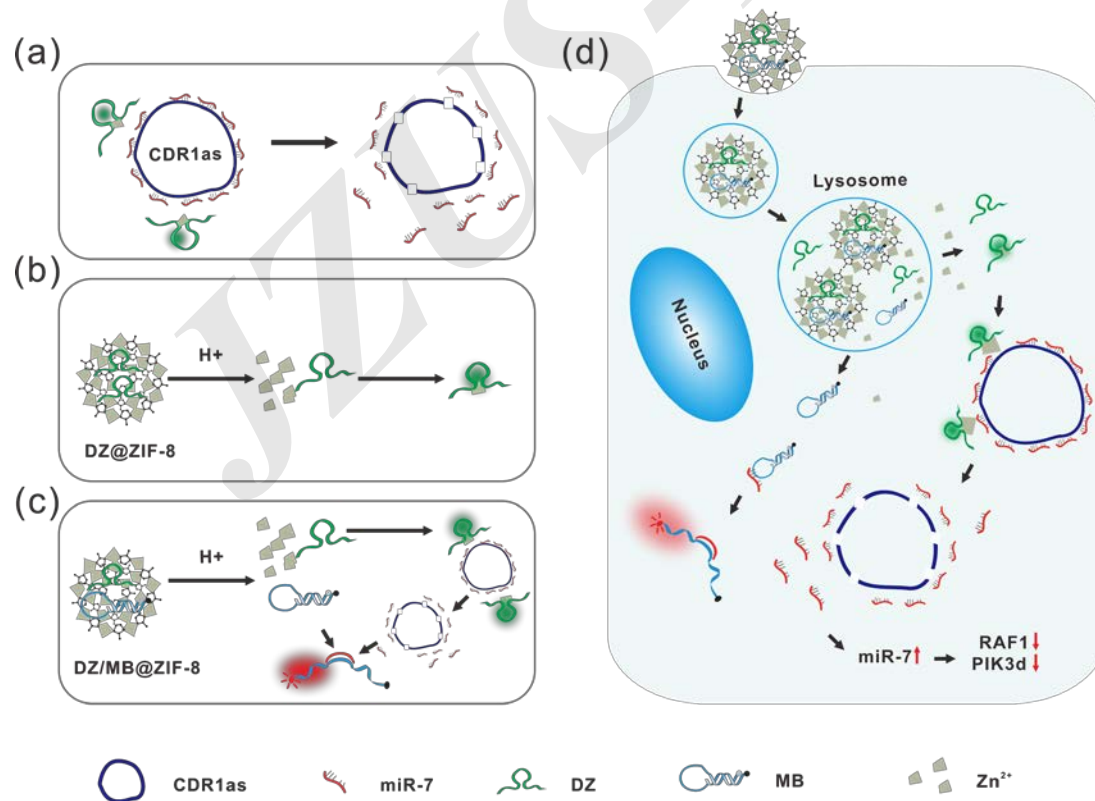
Background & Challenge

- The circRNA CDR1as functions as an oncogenic sponge for microRNA-7 (miR-7), making it a compelling therapeutic target.
- Current circRNA manipulation tools face limitations such as low intracellular delivery efficiency and complex protein components.
- Traditional methods cannot assess therapeutic effects in real time.



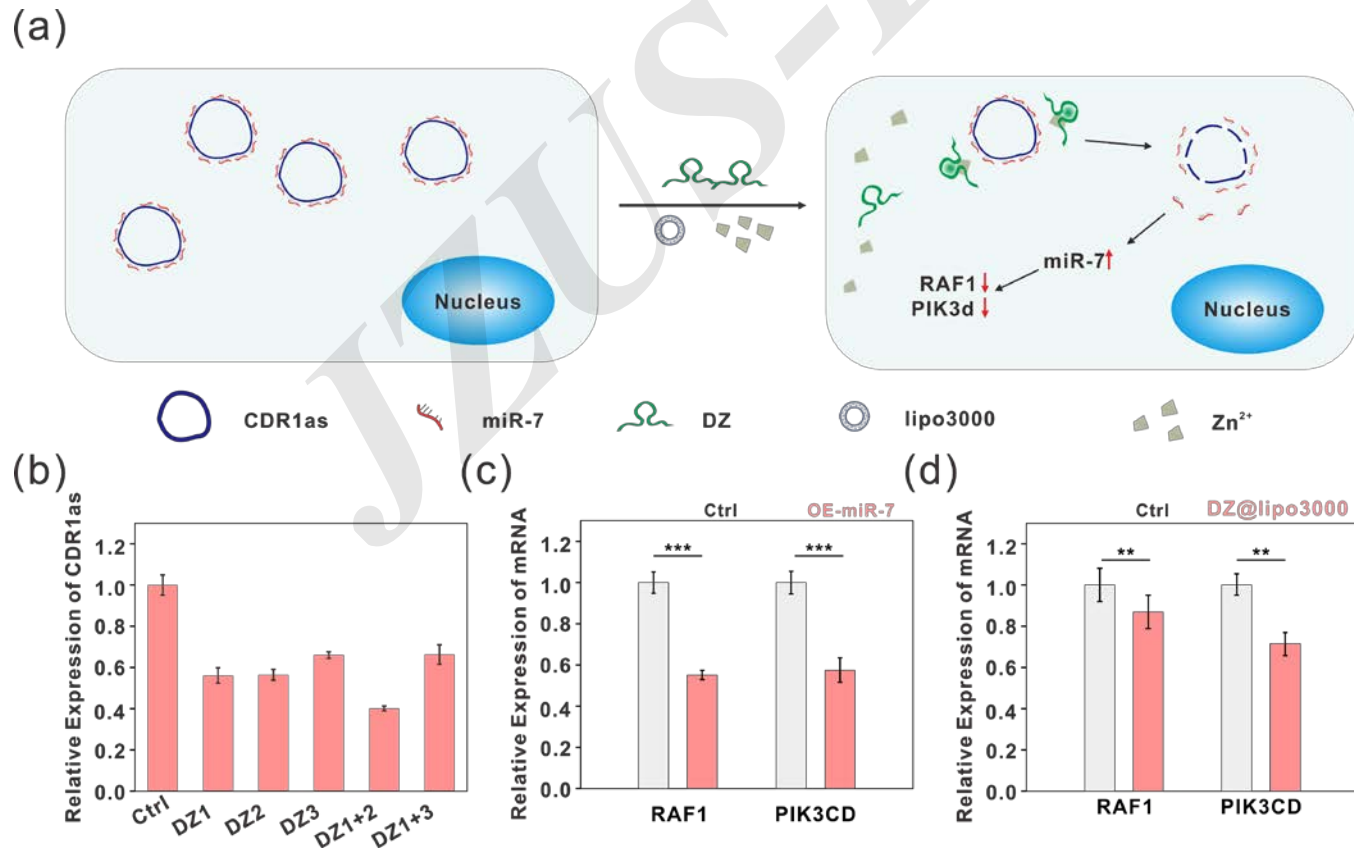
Design & Innovation

- Developed a novel Zeolitic Imidazolate Framework-8 (ZIF-8)-based nanoplatform DZ/MB@ZIF-8.
- Co-encapsulates DNazymes for the catalytic cleavage of CDR1as and a molecular beacon (MB) for reporting miR-7 activity.
- Acidic microenvironment triggers disassembly, releasing therapeutic/sensing components and essential Zn^{2+} cofactors for DNzyme activation.



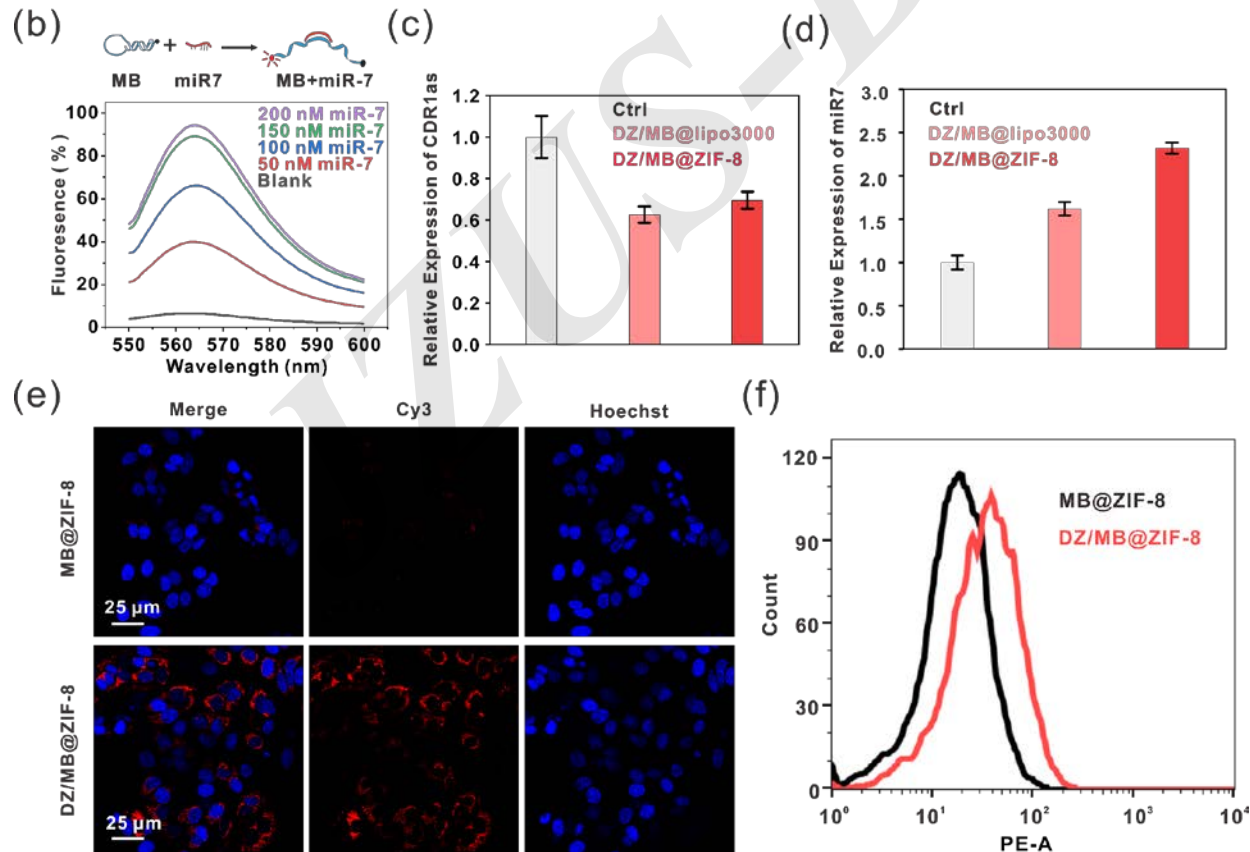
Key Result 1 – Efficient Gene Regulation

- Demonstrates efficient degradation of CDR1as without the need for external Zn^{2+} supplementation.
- Liberates miR-7 and significantly inhibits the expression of its downstream oncogenic targets, RAF1 and PIK3CD.
- Combined DNAzyme sets (DZ1+2) exhibited synergistic cleavage efficiency.



Key Result 2 – Real-Time Live-Cell Monitoring

- The therapeutic effect is directly correlated with a turn-on fluorescent signal from the MB.
- Enables real-time, live-cell readout of circRNA regulation.
- Exhibited excellent universality and was successfully applied across multiple cell lines.



Conclusion & Impact

- Establishes a versatile theranostic strategy that merges targeted gene regulation with intrinsic biosensing.
- Offers a powerful platform for probing circRNA function and advancing RNA-centric therapeutics.