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A fluorogenic probe for screening aldehyde dehydrogenase 1A1 (ALDH1A1) inhibitors

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We report the first competitive aldehyde dehydrogenase 1A1 (ALDH1A1) inhibitor featuring a fluorogenic binding response and nanomolar affinity (K_D), enabling real-time screening of non-fluorescent inhibitors with high cellular compatibility and no observed toxicity.

A fluorogenic non-covalent competitive inhibitor is often superior for direct, functional, and imaging-compatible studies of enzyme activity and inhibition, especially in live cells without genetic manipulation (*e.g.* unlike the nanoBRET assay). It offers key advantages in facilitating sensitive real-time measurement of enzymatic activity and is ideal for high-throughput screening to evaluate kinetic parameters of new competitive inhibitors.^{1–3} The fluorescence turn-on mechanism with good signal-to-noise ratio enhances the accuracy and reliability of quantification allowing screening of new candidates through kinetic assays.³

A specific fluorogenic inhibitor for ALDH1A1 (also known as retinal dehydrogenase 1) remains unavailable, although ALDH1A1 is one of 19 isoforms in the aldehyde dehydrogenase (ALDH) family, responsible for the oxidation of aldehydes in diverse metabolic pathways.⁴ Given the importance of ALDH1A1 in various physiological and pathological processes, including but not limited to retinoic acid biosynthesis, neuroprotection against toxic dopamine metabolites, and the maintenance of cancer stem cells (CSCs).^{5–7} ALDH1A1 has been identified as a CSC biomarker in multiple malignancies, including breast, lung, pancreatic, and colorectal cancers, where elevated enzymatic activity correlates with chemotherapy resistance, tumor progression, and poor clinical outcomes.⁴ Inhibitors targeting ALDH1A1 with nanomolar potency have drawn increasing interest due to their potential, in reducing CSC-associated phenotypes, suppressing hepatic glucose production, and attenuating triacylglycerol synthesis in the liver.

These attributes position ALDH1A1 as a promising therapeutic target in oncology and in metabolic disorders (diabetes, dyslipidemia, obesity, inflammation and Sjögren–Larsson syndrome).^{8–12} Detailed understanding of ALDH1A1 enzyme kinetics is important for rational design of selective inhibitors for targeted therapies.^{10,13–17} Thus, a fluorogenic ALDH1A1 inhibitor is an important addition to this family.

Most commercially available fluorescent probes for studying ALDH activity are broadly reactive substrates that measure total isoform activity, with the exception of ‘AlDeSense,’ a fluorogenic substrate specific to ALDH1A1.¹⁸ However, such substrates are not suitable for assessing enzyme kinetics or inhibitor interactions, as they exit the catalytic site after conversion. In contrast, fluorogenic inhibitors with a good binding constant (equilibrium dissociation constant (K_D) in nM range) provide a key advantage by enabling displacement-based assays to screen novel, including non-fluorescent, inhibitors and evaluate binding parameters like K_D , K_i and IC_{50} .

Cell-permeable fluorogenic inhibitors uniquely support both novel inhibitor screening, and live-cell imaging apart from mechanistic dissemination for pathway analysis upon inhibition of the target protein. Hurley *et al.* identified isatin derivatives as potent nanomolar inhibitors of ALDH enzymes and enhanced selectivity for ALDH1A1 over ALDH3A1 by incorporating an *N*-alkyl spacer to accommodate additional functional groups.¹⁴

We report the development of a fluorogenic ALDH1A1 inhibitor synthesized by conjugating a green-emitting naphthalimide fluorophore with an isatin scaffold. The resulting isatin–naphthalimide hybrid shows strong and competitive binding to the ALDH1A1 substrate-binding site. Owing to its competitive and reversible binding, the compound can be displaced by other inhibitors, enabling its use in displacement-based assays for screening binding constants of novel competitive ALDH1A1 inhibitors (Scheme 1).

This study initiated with *in silico* design and investigation of a potential fluorogenic ALDH1A1 inhibitor by linking a green-emitting naphthalimide derivative to an isatin *via* an ethyl linker. Given the possibility that the naphthalimide conjugation might

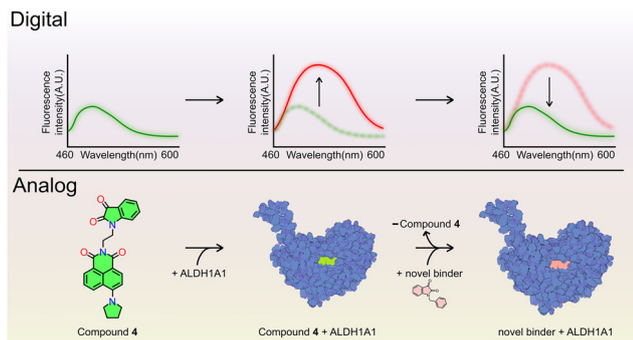
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Scheme 1 Illustration of the fluorogenic mechanism where compound **4** binds competitively to ALDH1A1, enabling displacement-based inhibitor screening.

impair binding affinity, we investigated docking of **4** against the substrate-binding site of ALDH1A1 (PDB ID: 7JWU).

The docking studies show that **4** favors the substrate-binding pocket over the NAD⁺ site, indicating higher affinity and competitive binding (Fig. 1(A)).

Furthermore, docking against other ALDH isoforms (ALDH1A3, ALDH2, and ALDH3A1) suggested potential pan-ALDH inhibition, with strong affinity for ALDH1A1, comparable to the selective inhibitor *N*-benzylisatin (NBI)¹⁴ (Fig. S1). Notably, **4** yielded a higher docking score of 93 ± 6 for ALDH1A1 in

comparison to 67 ± 3 for NBI. Interestingly, in the most favorable docking conformation, the naphthalimide moiety of **4** occupied the ALDH1A1 active site rather than the isatin moiety. This is contrary to the well-documented affinity of isatin for the substrate-binding pocket.¹⁴

To validate the docking conformation, we performed 100 ns molecular dynamics (MD) simulations. The trajectory supported the docking pose, with the ligand gradually migrating deeper into the binding cavity (Fig. 1(B)–(E)). RMSD analysis indicated system equilibration after ~ 20 ns. Binding free energy calculations using the molecular mechanics Poisson–Boltzmann surface area (MMPBSA) method (100 snapshots, final 10 ns) predicted -29.68 ± 2.48 kcal mol⁻¹ (Table S1). Stabilizing interactions included π -stacking of the isatin moiety with Y297 (-2.21 ± 0.05 kcal mol⁻¹) and van der Waals interactions between the naphthalimide core and V460 (-2.13 ± 0.09 kcal mol⁻¹), along with additional π -stacking involving F290, F171, and W178. Destabilizing contributions arose from D122, E289, and G458. Pose validation compared docking scores of isatin and naphthalimide with their alkyl chain derivatives (**5** and **6**), where naphthalimide–alkyl derivatives scored higher than isatin but lower than **4** (Fig. S3 and S4). Further docking of **4** into ALDH1A1 (PDB: 7JWU) with its isatin motif placed in the active site (Fig. S5) yielded a lower binding free energy (-26.87 ± 2.14 kcal mol⁻¹, Table S2) than when the naphthalimide moiety occupied the catalytic site.

Encouraged by these results, we synthesized **4** in a few steps (Scheme S1 and Fig. S6–S19) and isolated it in pure form ($> 95\%$, Fig. S20). The compound exhibited green fluorescence ($(\lambda_{\text{max}}^{\text{abs}}/\lambda_{\text{max}}^{\text{em}} = 450/535$ nm in methanol, quantum yield (Φ) = 0.0056); Fig. S21), high aqueous stability over 72 h (HPLC, single peak at 19 min; Fig. S22), and a lipophilicity of $\log P_{\text{o/w}} = 3.7$ (Fig. S23 and Table S2), consistent with Lipinski's rule of five and suggestive of favorable drug-like properties.

Upon binding to ALDH1A1 (purified from *E. coli* BL21 (pLysS) cells, Fig. S24 and S25) compound **4** exhibited a fluorogenic response characterized by a 3-fold increase in fluorescence intensity, $\Phi = 0.0264$ (ca. 4.7-fold higher than the free state), and a redshift of ~ 20 nm, consistent with its interaction within the hydrophobic active site of the enzyme (Fig. 1(F)). This pronounced fluorescence shift enabled the characterization of pre-steady-state kinetics for **4** (1 μM) binding to ALDH1A1 (100 nM) *via* stopped-flow spectroscopy with a cutoff filter of 515 nm. The absolute binding affinity of **4** yielded $K_{\text{D}}^{\text{abs}} = 102 \pm 4$ nM, derived from the ratio $k_{\text{off}}/k_{\text{on}}$ where $k_{\text{off}} = (2.885 \pm 0.201) \times 10^{-2}$ s⁻¹ and $k_{\text{on}} = (2.84 \pm 0.32) \times 10^{-4}$ nM⁻¹ s⁻¹ (Fig. 1(G) and (H)). The inhibitory potency of pure, stable **4** against ALDH1A1 was evaluated by monitoring NAD⁺ reduction to NADH. Assays were conducted in a buffer containing 20 mM HEPES (pH 7.4), 150 mM NaCl, 1 mM NAD⁺, and 1 mM DTT, with 1 mM benzaldehyde as a substrate and 3.5 μM ALDH1A1 enzyme. NADH formation was measured *via* absorbance at 340 nm over 90 minutes at 25 °C. Compound **4** exhibited potent inhibition with an IC₅₀ of 370 ± 8 nM ($K_{\text{i}} = 63 \pm 14$ nM) (Fig. 2(A)). Considering isatin's previously reported mild inhibitory activity against ALDH1A1,¹⁴ we investigated the inhibitory effect of the naphthalimide moiety alone (compound **2**).

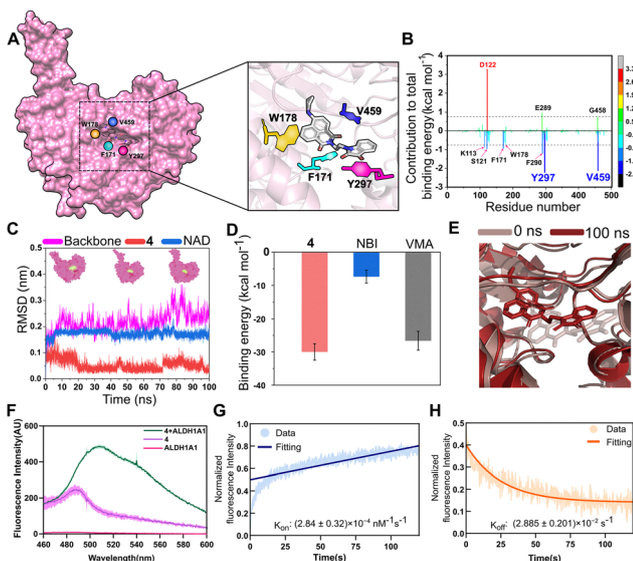


Fig. 1 (A) Structural depiction of the interaction between **4** and the catalytic site of ALDH1A1 (PDB: 7JWU). (B) Evaluation of the contributions of individual residues to the stabilization or destabilization of the ALDH1A1–**4**–NAD complex. (C) RMSD analysis of the system for 100 ns. (D) Comparison of binding energies (mean \pm SD) of **4**, NBI, and VMA (the native ligand of 7JWU). (E) **4** shifts deeper into the catalytic pocket over 100 ns, enhancing binding energy. (F) Change in the fluorescence intensity of **4** when it binds to the active site of ALDH1A1 ($n = 3$). (G) and (H) are the plots of time-dependent association and dissociation binding kinetics of ALDH1A1 and **4**, respectively. The inset represents the single exponential fitting in one-phase association and one-phase decay. The error bar represents the SD from three independent experiments ($n = 3$).



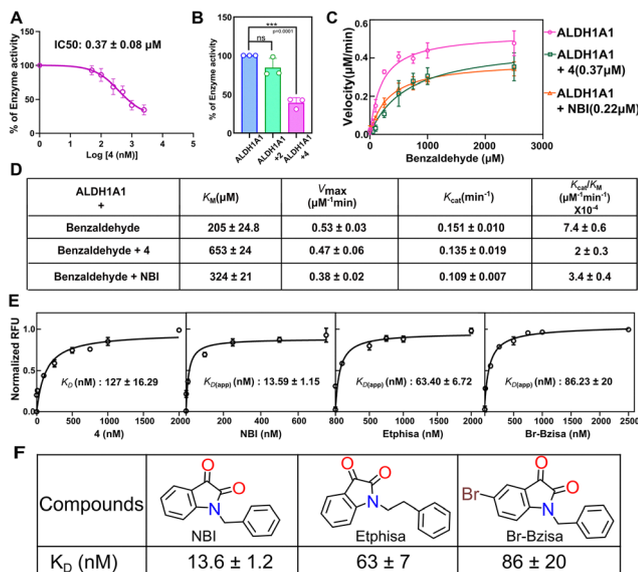


Fig. 2 (A) Enzyme inhibition profile of **4** targeting ALDH1A1 ($n = 3$). (B) Percentage of enzyme activity in the presence of $1 \mu\text{M}$ **2** and **4** showing that **2** is not able to inhibit ALDH1A1 ($n = 3$, ns = non-significant). (C) Comparative Michaelis–Menten kinetics between ALDH1A1 and ALDH1A1 with **4** and NBI using benzaldehyde as a substrate ($n = 3$). (D) The table represents the kinetic parameters of ALDH1A1 in the presence of **4** and NBI at their IC_{50} concentrations, implying that **4** serves as a competitive inhibitor. (E) Fluorescence titration to determine the equilibrium binding constant (K_D^{eq}) of **4** and apparent binding constants K_D^{app} of NBI, Etphisa, and Br-Bzisa by fluorescence displacement assay ($n = 3$). (F) K_D^{app} values of NBI, Etphisa, and Br-Bzisa. Statistical significance was determined as follows: $p < 0.05$ (*), $p < 0.01$ (**), $p < 0.001$ (***), $p < 0.0001$ (****); ns denotes not significant.

We further evaluated the inhibitory effects of compounds **5** and **6**, in which the naphthalimide moiety is linked *via* a propyl or ethyl spacer containing a nitrogen atom (Fig. S26–S30). Compounds **2**, **6** and **7** showed no inhibition, confirming that both the isatin and naphthalimide components contribute synergistically to the inhibitory effect (Fig. 2(B)). Under identical conditions, the ALDH1A1-selective inhibitor NBI displayed an IC_{50} of 220 ± 20 nM ($K_i = 38 \pm 5$ nM) (Fig. S31). Using propionaldehyde as the enzymatic substrate, we assessed ALDH1A1 inhibition by **4** and NBI at $1 \mu\text{M}$.

Both agents potentially suppressed catalytic activity, evidenced by a marked reduction ($> 80\%$) in specific activity relative to controls (Fig. S32). Michaelis–Menten kinetics were assessed by measuring NADH production at varying benzaldehyde concentrations in the presence of **4** and NBI (Fig. S33). The kinetics observed with compound **4** (IC_{50} concentration) indicate a threefold K_m increase, an unchanged V_{max} , and a fourfold decrease in k_{cat}/K_m (Fig. 2(C) and (D)), confirming competitive inhibition of ALDH1A1. In contrast, NBI is identified as a noncompetitive inhibitor using a Line–Weaver Burk plot (Fig. S34). **4** increases K_m without affecting V_{max} , showing classical competitive inhibition at the active site. In contrast, NBI reduces V_{max} with little change in K_m , consistent with non-competitive inhibition. This distinction explains that displacement assays are suitable for compound **4**. Leveraging this mechanism, the equilibrium dissociation constant (K_D^{eq}) of **4** was determined by fluorescence titration to be 127 ± 16.3 nM

(Fig. 2(E)). Thus, the approximate equivalence of K_D^{eq} to the K_D^{abs} confirms the validity of our assay setup. Furthermore, as a fluorescent competitive inhibitor, **4** enables measurement of apparent dissociation constants (K_D^{app}) *via* fluorescence quenching, providing a robust platform for screening new inhibitors and guiding inhibitor design. The compound's fluorescence intensity, sensitivity, and emission wavelength are compatible with standard fluorimeters, facilitating assay development. To demonstrate the use of **4** in evaluating new inhibitors using its fluorogenic properties, we tested three ALDH1A1 inhibitors with known IC_{50} values and evaluated their K_D , including NBI, 1-phenethylindoline-2,3-dione (Etphisa), and 1-benzyl-5-bromoindoline-2,3-dione (Br-Bzisa). ALDH1A1 (500 nM) was pre-incubated with **4** ($1 \mu\text{M}$) at 4°C to achieve active-site saturation. Competitive binding of candidate inhibitors was then assessed *via* titration against the pre-formed ALDH1A1:**4** complex (Fig. S35). The fluorescence intensity of **4** is negligible at 510 nm unless it binds to ALDH1A1, so titration with an inhibitor can be tracked by monitoring the decrease in fluorescence at this wavelength until it plateaus. This allowed determination of the K_D^{app} for each inhibitor (Fig. 2(F) and Table S4). These K_D^{app} values correlated well with their reported IC_{50} values (Fig. S36),¹⁴ validating **4** as an effective probe for screening and ranking the binding affinities of ALDH1A1 inhibitors.

Given the presence of a naphthalimide fluorophore, the aggregation behavior of **4** in aqueous media was further evaluated using dynamic light scattering (DLS) in $1 \times$ PBS (pH 7.4) across concentrations. Aggregation was observed at concentrations $\geq 3 \mu\text{M}$. At $1 \mu\text{M}$, the average particle size remained ~ 50 nm (Fig. S37 and S38). Field emission scanning electron microscopy (FESEM) corroborated the DLS data, revealing small, dispersed aggregates. Thus, the aggregation occurs at concentrations *ca.* 7-fold higher than the K_D value (127 ± 16 nM), suggesting that the measured K_D is genuine and not an artifact of aggregation-induced enzyme inactivity.

ALDH1A1 is primarily cytoplasmic, and co-localization studies confirmed that **4** similarly distributes within cells (Fig. 3(A) and (B)). Fixed-cell immunofluorescence and spinning disk confocal microscopy showed widespread cytoplasmic localization of both **4** and ALDH1A1 (Fig. S39). Live-cell imaging in MDA-MB-231 cells revealed that $1 \mu\text{M}$ **4** exhibited minimal nuclear presence and was equally partitioned between mitochondria and lysosomes (PCC ~ 0.5 ; Fig. 3(A) and (B)), supporting its cytoplasmic accumulation essential for probing ALDH1A1 inhibition and downstream mechanisms.

To study the effect of **4** at the cellular level, we selected three cancer cell lines (MIA PaCa-2, PANC-1, and MDA-MB-231) expressing ALDH1A1 (Fig. S40). **4** exhibited minimal cytotoxicity ($\text{IC}_{50} > 100 \mu\text{M}$) in all of them (Fig. S41 and Table S5), in contrast to the ALDH1A1 inhibitor NBI, which showed a cellular IC_{50} of $18.7 \pm 1.8 \mu\text{M}$ in MIA PaCa-2 cells. MIA PaCa-2 has the highest expression of ALDH1A1 among the three cell lines (Fig. S40). The minimal cytotoxicity of **4** ($> 100 \mu\text{M}$ in MTT assays) suggests reduced off-target effects compared to NBI, apart from the stronger binding affinity toward ALDH1A1. ALDH1A1 inhibition in cells was confirmed using the AldeRed assay, where red fluorescence increases upon ALDH-mediated aldehyde oxidation and



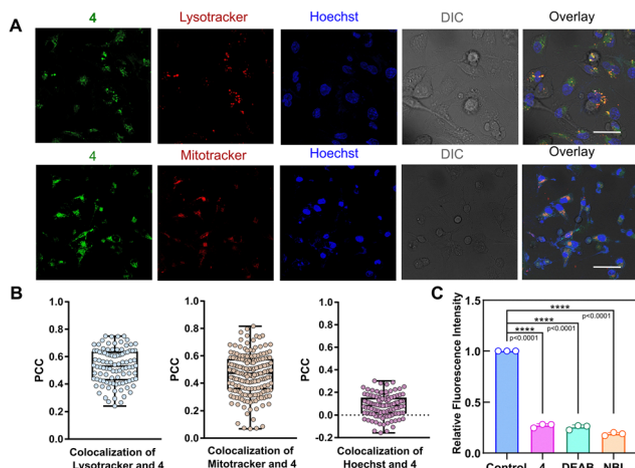


Fig. 3 (A) Co-localization analysis of lysosomal, mitochondrial, and nuclear markers in MDA-MB-231 cells treated with 1 μM of **4** (scale bar = 10 μm). (B) Pearson's correlation coefficient (PCC) values quantify the degree of co-localization with **4** across cellular compartments ($n > 100$). (C) Aldefluor assay representing ALDH activity reduction ($n = 3$) in MIA PaCa-2 in the presence of 1 μM **4**, 15 μM DEAB and 1 μM NBI. Statistical significance was determined as follows: $p < 0.05$ (*), $p < 0.01$ (**), $p < 0.001$ (***), $p < 0.0001$ (****); ns denotes not significant.

retention in ALDH1A1^{high} MIA PaCa-2 cells. Treatment with 1 μM of compound **4** markedly reduced fluorescence (Fig. 3(C) and Fig. S42), indicating effective intracellular ALDH inhibition.

ALDH1A1 regulates cancer stem cell (CSC) traits,^{19–21} and we examined the effect of treatment of MIA PaCa-2 cells, known for high CSC activity,¹⁹ with **4** or NBI on a few key stemness-related genes (Table S5). ALDH1A1 and c-MYC mRNA levels reduced upon treatment with **4** or NBI (Fig. S43 and S44), consistent with ALDH1A1's role in c-MYC regulation via the Wnt/ β -catenin pathway.²² Although increase in compensatory stemness-related genes NOTCH1, SOX2, SOX9, PI3K, and OCT4, highlighted the multifactorial regulation of stemness beyond ALDH1A1 alone.

In summary, **4** is a potent fluorogenic ALDH1A1 inhibitor with nanomolar K_D and low cytotoxicity. MD simulations reveal cooperative binding of its isatin and naphthalimide groups, consistent with competitive inhibition and fluorogenic response. The displacement-based assay, the first of its kind, enables precise evaluation of non-fluorescent ALDH1A1 inhibitors, with K_D values aligning with the reported IC_{50} . Beyond inhibition, **4** suggests that ALDH1A1 inhibition may trigger compensatory stemness mechanisms, highlighting the complexity of ALDH1A1 biology. These results support context-specific therapeutic strategies and provide a foundation for next-generation fluorogenic inhibitors with optimized emission and reduced aggregation. AM thanks SERB (CRG/2021/001118) and IISER Kolkata for support. S. G., B. R., A. M., and S. M. from AM Lab acknowledge CSIR and KVPY; S. S. and S. G. (R.D. Lab) thank UGC and IISER Kolkata for fellowships.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article have been included as part of the SI which contains synthetic schemes, NMR and mass spectra, docking and MD simulation data, kinetic plots, additional figures, and tables. See DOI: <https://doi.org/10.1039/d5cc03947a>.

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