Mathematical framework for activity-based cancer biomarkers

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Detailed Terms
Mathematical framework for activity-based cancer biomarkers

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Advances in nanomedicine are providing sophisticated functions to precisely control the behavior of nanoscale drugs and diagnostics. Strategies that coopt protease activity as molecular triggers are increasingly important in nanoparticle design, yet the pharmacokinetics of these systems are challenging to understand without a quantitative framework to reveal nonintuitive associations. We describe a multicompartment mathematical model to predict strategies for ultrasensitive detection of cancer using synthetic biomarkers, a class of activity-based probes that amplify cancer-derived signals into urine as a noninvasive diagnostic. Using a model formulation made of a PEG core conjugated with protease-cleavable peptides, we explore a vast design space and identify guidelines for increasing sensitivity that depend on critical parameters such as enzyme kinetics, dosage, and probe stability. According to this model, synthetic biomarkers that circulate in stealth but then activate at sites of disease have the theoretical capacity to discriminate tumors as small as 5 mm in diameter—a threshold sensitivity that is otherwise challenging for medical imaging and blood biomarkers to achieve. This model may be adapted to describe the behavior of additional activity-based approaches to allow cross-platform comparisons, and to predict allometric scaling across species.

Results

Multicompartment Model of Synthetic Biomarkers. Synthetic biomarkers are made of NPs conjugated with protease-cleavable peptides. Following administration, a chain of events unfolds to detect disease: (i) NPs accumulate in diseased tissue, (ii) peptides are cleaved by local proteases, (iii) the cleavage fragments are

Significance

The discovery of cancer at an early stage improves treatment outcomes, yet cancer detection thresholds based on measuring the abundance of biomarkers produced by small tumors are biologically limited. Here we develop a mathematical framework to explore the use of activity-based biomarkers for early cancer detection. In contrast to abundance-based biomarkers, activity-based biomarkers rely on the catalytic activity of enzymes to amplify cancer-derived signals and allow detection of small tumors. Using a class of activity-based biomarkers called synthetic biomarkers, we comprehensively explore how detection sensitivities depend on probe design, enzymatic activity, and organ physiology, and how they may be precisely tuned to reveal the presence of small tumors in humans.


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and secreted tumor-associated proteases (e.g., MMP9) according to Michaelis–Menten kinetics (\(k_{\text{cat}}^\text{MMP}, K_{M}^\text{MMP}\)) and are cleaved by tumor-associated proteases (e.g., MMP9) according to Michaelis–Menten kinetics (\(k_{\text{cat}}^\text{MMP}, K_{M}^\text{MMP}\)). The cleaved reporters then diffuse back into blood, where they combine with reporters produced from nonspecific proteases (\(k_{\text{cat}}^\text{blood}, K_{M}^\text{blood}\), and \(k_{\text{cat}}^\text{blood}, E_{\text{blood}}^\text{blood}\)) and secreted tumor-associated proteases (\(k_{\text{cat}}^\text{tumor}, E_{\text{tumor}}^\text{tumor}\)). A fraction of the reporters are reabsorbed by the kidneys (\(k_{\text{absorb}}\)) before clearance into urine (\(k_{\text{urine}}\)). See SI Text for derivation of model equations.

Cleared into urine, and (iv) the urine is analyzed by multiple analytical platforms. In previous studies (3, 21–23), we engineered synthetic biomarkers for use in different diseases and methods of urinalysis. Here we chose to use PEG cores to formally explore the role of scaffold size, fluorogenic substrates to quantify kinetics by fluorescent imaging, and a xenograft model of CRC.

Our multicompartment model (Fig. 1 and SI Text) is made of a blood compartment (compartment i) where synthetic biomarkers circulate with half-life \(t_{1/2}\), and passively diffuse into the tumor compartment (compartment ii) by the rate constant \(k_{\text{in}}\). In the tumor, peptides are cleaved by tumor proteases [e.g., matrix metalloproteinase 9 (MMP9)] following Michaelis–Menten enzyme kinetics (i.e., \(k_{\text{cat}}^\text{MMP}, K_{M}^\text{MMP}\)), and passively diffuse \(k_{\text{cat}}^\text{blood}, E_{\text{blood}}^\text{blood}\)) to the blood compartment (compartment iii). These reporters combine with reporters cleaved in the blood by circulating nonspecific proteases (\(k_{\text{cat}}^\text{blood}, K_{M}^\text{blood}\), and \(k_{\text{cat}}^\text{blood}, E_{\text{blood}}^\text{blood}\)) and tumor-secreted proteases (\(k_{\text{cat}}^\text{tumor}, E_{\text{tumor}}^\text{tumor}\)) before filtering into urine (\(k_{\text{urine}}\)). We assumed negligible nonspecific protease activities within the tumor compartment, and introduced a rate constant, \(k_{\text{absorb}}\), to account for nonspecific binding (e.g., on blood vessels) and kidney reabsorption of our reporter. In each compartment, our model assumes a homogeneous, well-mixed distribution of NPs or reporters. A deterministic, well-mixed assumption is valid for Michaelis–Menten kinetics when a large number of molecules are involved, which is true for this study involving solution concentrations ranging from micromolar to nanomolar. Model parameters are summarized in Table S1.

**Sensing Protease Activity with Peptide—PEG Synthetic Biomarkers.** A key feature of synthetic biomarkers is the use of a nanoscale core to increase the hydrodynamic diameter of surface-conjugated peptides beyond the pore size of the glomerulus, which for NPs is \(\approx 5\) nm in diameter (25); as a result, peptides conjugated to NPs appear in the urine after a cleavage event. To explore this size dependency in a model system, we analyzed amine-terminated eight-arm branched PEG (10 kDa, 20 kDa, and 40 kDa) by dynamic light scattering (DLS) and observed increases in hydrodynamic diameters from \(\approx 3\) nm to \(10\) nm corresponding to their molecular weights (Fig. 2 A and B) that were similar to measurements made in 50% FBS (Fig. S1). Benchmarked against the size of our previous nanoworm (NW) formulation (\(\approx 30\) nm), PEG NPs were smaller by severalfold, which increases passive diffusion rates (18). To determine renal clearance efficiencies, we administered fluorescent PEG and NWs and detected urine levels up to \(\approx 80\%\) of injected dose (ID) for 10 kDa PEG and \(\approx 40\%\) ID for 20 kDa PEG within 60 min (Fig. S2). By contrast, 40 kDa PEG and NWs filtered at negligible levels into the urine (\(\approx 0.09\%\) ID and \(\approx 0.19\%\) ID, respectively), consistent with the size exclusion limit of the glomerulus (Fig. 2C). Based on these results, we selected 40kD PEG to develop our model.

Next, we investigated the use of peptide–PEG NPs to sense protease activity. We selected an MMP substrate PLGVRG (26) and synthesized a fluorogenic derivative, Q1, containing an N-terminal fluorophore (FAM), an internal quencher (CPO-2), and a C-terminal cysteine residue for thiol-mediated coupling onto 40 kDa PEG (Q1 = 5FAM-GGPLGVRGKK(CPQ-2)-PEG2-C). To test subtype cleavage, we incubated Q1–PEG with MMP9, a gelatinase overexpressed by solid tumors, and detected a rapid increase in sample fluorescence (Fig. 2D). In the presence of the MMP inhibitor Marimastat (Fig. 2D, arrow), we observed an immediate arrest in fluorogenesis, confirming protease-dependent activation of Q1–PEG. To validate the use of PEG-based probes to cancer mice, we synthesized F1 (sequence, eGvndneeGfarskGGPLGVRG), a tandem peptide made of our MMP substrate PLGVRG and the protease-resistant d-stereoisomer of glutamate–fibrinopeptide B (Glu–Fib; sequence = eGvndneeGfarsr, lower case = d-isomer) (Fig. 2I). Glu–Fib is a biologically inert peptide we used in past studies to
promote renal clearance of cleaved peptide fragments (3). In mice bearing LS174T CRC tumors and given F1-PEG, we observed significant increases in bladder fluorescence compared with their healthy counterparts (Fig. 2E)—similar to our previous results using MMP-sensitive NWs (3). Together, our data indicated that synthetic biomarkers with PEG cores can sense protease activity and detect CRC from urine.

**Extracting Model Parameters in Vivo.** We next sought to identify the values of the parameters for our system of ordinary differential equations (ODEs). To determine circulation times, we assumed that systemic clearance is due to renal filtration and clearance by the mononuclear phagocyte system (MPS). Thus, to identify clearance rate constants \( (k_{MPS}, k_{ MPS \to tumor}, k_{MPS \to blood}, k_{ tumor \to MPS}) \), we administered fluorescent PEG and Glu–fib and fitted their levels in blood and urine to a simplified three-compartment model (\( n = 3 \); error bars, s.d.; Figs. S3 and S4). Consistent with their relative size, Glu–fib cleared into the urine at a rate approximately fivefold greater than MPS clearance \( (k_{ tumor \to MPS} vs. k_{ MPS \to tumor}) \), but, by contrast, PEG was characterized by an MPS to renal clearance \( (k_{MPS} vs. k_{ MPS \to tumor}) \) ratio of \( \sim 35 \) (Table S1). From these values, we calculated circulation half-lives of \( \sim 18 \) min and \( \sim 18.7 \) h for Glu–fib and PEG, respectively. To determine NP transport rate into tumor \( (k_{ tumor}) \), we administered fluorescent PEG and NWs into mice and analyzed tumor homogenates along with major organs at 0 h, 0.5 h, 1 h, 3 h, 6 h, and 16 h to determine the levels of localized NPs (\( n = 3 \); error bars, s.d.; Figs. S3 and S5). Using a simplified two-compartment model (Fig. S4), we determined a tumor transport rate constant for PEG that was \( \sim 3.5 \)-fold greater than NWs (Table S1) and consistent with its smaller hydrodynamic diameter.

To incorporate the role of tumor proteases, we sought to determine a representative set of Michaelis–Menten constants to describe the rate at which F1-PEG may be cleaved by tumor proteases. For simplification, we focused on MMP9, a gelatinase that is involved in the growth and invasion of solid tumors and secreted by LS174T cells (\( n = 3 \); error bars, s.d.; Fig. S6). To determine the cleavage efficiency, we incubated different concentrations of F1-PEG with MMP9 and extracted initial cleavage velocities to calculate \( k_{cat}^{MMP9} \) and \( k_{MMP9}^{M} \) (\( n = 3 \); error bars, s.d.; Fig. 3C), resulting in a catalytic efficiency \( k_{cat}/k_{M} \sim 2.0 \times 10^{5} \) M\(^{-1}\) s\(^{-1}\) that was similar to published values (27, 28). Similarly, we simplified nonspecific protease activity in blood into a single set of representative Michaelis–Menten constants \( (k_{blood}, k_{ blood}, E_{blood}) \) whose values were determined by fitting our system of ODEs against urine kinetics obtained in healthy mice (Table S1). These values allowed us to fit for the concentration of tumor-specific proteases in the tumor and blood compartments \( (E_{tumor} \text{ and } E_{blood}) \), respectively using tumor-bearing animals (\( n = 3 \); error bars, s.d.; Fig. S3D). Taken together, our experimentally fitted multicompartment model predicted urine pharmacokinetics that closely aligned with our experimental observations.

**Exploring Synthetic Biomarker Properties in Silico.** We next set out to explore the parameters important for detection sensitivity. In healthy blood, circulating proteases such as the coagulation and complement family may cleave peptide substrates. Consequently, to explore the effect of nonspecific protease activity, we varied \( k_{cat}^{blood} \) and \( E_{blood} \) from their starting values and observed that background urine signals were reduced to \( \sim 10\% \) of their original values if either \( k_{cat}^{blood} \) or \( E_{blood} \) was reduced by a factor of 16 (\( n = 3 \); error bars, s.d.; Fig. 3A). Under these conditions, the difference in urine signal (i.e., tumor – control) increased and broadened, shifting the maximum peak intensity from \( \sim 50 – 200 \) min (Fig. 3B). The time point at which our formulation was predicted to reach maximum levels \( \sim 50 \) min, red line, Fig. 3B) was consistent with our previous studies in which we found maximum disease contrast to occur at 1 h (3, 22).

Synthetic biomarkers are formulated from nanomaterials whose properties can be precisely tuned (29); therefore, we explored how these properties may affect detection sensitivity. We increased the NP half-life \( (\tau_{NP}^{half}, Fig. 3C) \) from 18 h to 300 h by factors of 2, which had negligible effects, indicating that our formulation was sufficiently long-circulating to sense protease activity. By contrast, signal curves broadened and increased in magnitude when the administered dose of NPs (\( N \), Fig. 3D) was increased from 2.5 \( \mu \)M (\( \sim 0.5 \) mg/kg) to 20 \( \mu \)M (\( \sim 4 \) mg/kg)—a range of concentrations for iron oxide nanomaterials that are well-tolerated in humans (30, 31). Intuitively, larger substrate reservoirs are created at higher NP doses, consequently prolonging and extending peak signal intensities. Similarly, increasing NP transport rates into tumors \( (k_{NP}^{tumor}) \), Fig. 3E) increased signal intensities, reaching a limit at values three to four decades above the starting rate constant \( (1.4 \text{ min}^{-1}) \). At this point, detection signals are no longer dependent on NP transport but limited by the rate at which tumor proteases cleave the probe. Lastly, we increased the concentration of proteases in the tumor and blood compartments \( (E_{tumor}, E_{blood}) \), Fig. 3F), which shifted signal curves upward and maximum disease contrast earlier. Varying \( k_{cat}^{MMP9} \) resulted in identical results (Fig. 3G) because
protease activity is directly proportional to \( k_{cat} \) and \( E \) in the Michaelis–Menten model. Collectively, our mathematical model allows us to systematically explore the landscape of synthetic biomarker properties and identify critical system parameters that influence detection signals.

**Validation of a Model Prediction.** To validate the predictions of our model, we normalized the urine kinetics to examine the signal-to-noise ratio (SNR). At low concentrations of the administered dose, \( N \), the SNR started at a peak and decreased exponentially (light blue, Fig. 4A), and by high \( N \), the SNR kinetic curves appeared constant (black, Fig. 4A). To explain this nonintuitive behavior, we hypothesized that at low concentrations of \( N \), the NPs are quickly degraded in the blood by tumor-secreted and background proteases, leaving a small fraction of the ID to diffuse into the tumor bed. Therefore, the SNR traces at low \( N \) are largely reflective of blood activity. By contrast, at high \( N \), an NP reservoir is created in the circulation that saturates probe activation by circulating proteases, and maintains a sufficiently large concentration of unactivated NPs to move down their concentration gradient into the tumors. Because this process depends on passive diffusion, the signal generated from tumor proteases is expected to occur later than blood cleavage; therefore, the combined effect of early activation by blood and delayed activation by tumors results in relatively constant SNR kinetic curves. To test this in silico, we examined the bounds of our model at small \( N \) and observed that the signal (tumor + blood) was generated largely by blood but not tumor activation, and rapidly decayed (Fig. S7A). We then tested large \( N \) and observed that the detection signal resulted from a constant contribution from blood (i.e., protease activity saturated) and a delayed activation signal from the tumor (Fig. S7B), consistent with our hypothesis.

One consequence of this model prediction is that within the range of reasonable NP doses (< 10 \( \mu \)M or ~10 mg/kg) that would be expected to be well-tolerated in humans, the combined effect of probe activation by the blood and tumor compartments results in an SNR that would be relatively constant within an hour following administration (Fig. 4A and blue dots in Fig. 4B). To validate this prediction, we administered our probe in two cohorts of tumor-bearing mice at doses separated by an order of magnitude (1 \( \mu \)M and 10 \( \mu \)M, \( n = 5 \) mice, Fig. 4C and D). Consistent with our model, the signal of the urine samples collected from tumor animals was significantly higher compared with the control group, and the difference in SNR was not statistically different (Fig. 4E). One implication of this result is that it may be possible to detect disease by administering our probes at “microdose” levels (< 100 \( \mu \)g in humans), a regulatory pathway created by the US Food and Drug Administration to compress the time frame and translational path to the clinic (32). This approach would require an ultrasensitive analytical method for urine analysis such as digital ELISA (23). Taken together, these experiments validate the potential of our quantitative framework to predict nonintuitive behavior in living systems.

**Strategies for Early Cancer Detection.** Synthetic biomarker-based signals are produced by the catalytic activity of tumor proteases and concentrated into urine; therefore, we hypothesized that this platform may be used to detect early clinical-stage tumors that are otherwise challenging to detect using blood biomarkers. To establish a basis for comparison, we adapted a steady-state ODE model (5) for the blood biomarker carcinoembryonic antigen (CEA), a biomarker for CRC. At steady state, the concentration of CEA depends on its tumor production rate and circulation half-life (SI Text). To estimate the bounds, we assumed an average tumor cell density of 10^6 cells per cubic millimeter, a CEA retention rate of 90% (5), a half-life of 72 h (33) (\( t_{1/2} \)), and minimum (\( P_{MC} \)) and maximum (\( P_{MC}^{MAX} \)) CEA production rates to be 15 pg and 260 ng per 10^6 cells per 10 d, respectively, based on our previous analysis of 24 human CRC cell lines (median value = 1.5 ng per 10^6 cells per 10 d) (5). At steady state, our model predicted that tumors smaller than 10 mm in diameter (5, gray box, Fig. 5A) would result in CEA levels beneath the baseline threshold expected in healthy individuals (34) (5 ng/mL, dashed line, Fig. 5A) and therefore, indistinguishable. By contrast, tumors that were 10 mm, 20 mm, and 50 mm in diameter were predicted to be detectable above normal levels of CEA only if the tumors produced the biomarker at a rate within the top 11%, 32%, or 60% of the range bounded by the maximum (square, Fig. 5A) and minimum (triangle, Fig. 5A) production rates, respectively. These results were consistent with previous mathematical studies, which established a detection limit of 1–2 cm for blood biomarkers (5). We next combined both models to analyze 10-mm-sized LS174T tumors, which produce CEA at a rate 100-fold above the median, and our model predicted that our activity-based probe would result in urine signals that would eclipse the steady-state level of CEA in blood (Fig. 5B). This prediction in humans is consistent with our prior study in mice, where we showed that synthetic biomarkers detected LS174T tumors earlier and with significantly higher predictive power compared with CEA analysis (3).

We next applied our model to calculate its potential to detect a theoretical human solid tumor of 5 mm in diameter, by considering the concentration of proteases expected in a tumor of this size. We chose 5-mm tumors as a target sensitivity threshold to represent a detection limit that is below the current thresholds associated with computed tomography (CT) (~1 cm) (35) or blood biomarkers (~1–2 cm) (4, 5). To estimate protease production rates, we analyzed conditioned media collected from four human CRC lines and detected MMP9 secretion rates ranging from ~1.9 pg/10^6 cells per day to 12.5 pg/10^6 cells per day by ELISA (Fig. 5C). To calculate a range of possible protease concentrations, we assumed an average tumor cell density of ~10^9/mm^3 and a protease retention rate of 90% (i.e., 10% of proteases secreted by tumor cells are shed into the blood) (5) and...
arrived at protease levels ranging from ~0.4 nM to 2.7 nM (gray box, Fig. 5C). We further adjusted the administrated NP dose to reflect the higher total blood volume (~5 L) in humans compared with mice (~2 mL) to maintain an identical starting concentration.

Using these parameters, we explored how detection signals depend on the concentration of proteases produced by tumors. Our selected PEG formulation (red dot, Fig. 5C) produced detection signals in urine at concentrations (~7 μM) that were readily detectable by ELISA (21, 22), which has a lower limit of detection of ~10 pM (dashed line, Fig. 5C). Using 10 pM as the lower detection limit, we decreased protease levels from the empirically fitted value (red) by several decades to reach the range expected for 5-mm tumors (gray box), and observed that our model predicted urine signals to fall below the limit of detection (blue dot) (Fig. 5C). We then tested a 10-fold improved synthetic biomarker formulation by increasing the rates for tumor proteases and penetration by a factor of 10 (~k_{tumor} and k_{NP} respectively) while simultaneously decreasing nonspecific cleavage in blood (~k_{blood}) (Fig. 5D). With these values, the sensitivity markedly increased, elevating urine signals from 5-mm tumors to levels (Fig. 5D, blue dot) above the limit of detection by ELISA. Lastly, to assess the boundaries of our platform, we tested a 100-fold improved formulation (Fig. 5E) and found that urine signals were bounded by the maximum plateau (Fig. 5E, blue dot). Collectively, these data revealed important design variables for our diagnostic, and identified strategies to formulate ultra-sensitive synthetic biomarkers for detecting 5-mm tumors.

Discussion

The development of computational techniques in concert with preclinical testing may provide the abilities to rapidly assess system parameters, identify tunable nanomaterial properties, and predict allometric scaling to humans (11, 16–19). Here, we established a computational framework to describe the pharmacokinetics of activity-based synthetic biomarkers and to quantitatively explore the parameters that are important for sensitive detection of disease.

In this study, we used a xenograft model of cancer to fit our mathematical model. Tumor vessels in both primary tumors and xenografts are poorly formed and fenestrated, and there is abundant evidence demonstrating the effectiveness of passive NP delivery to primary human tumors (36, 37)—results that are consistent with observations that were first borne out in xenograft tumor models. Based on these studies, xenograft tumors may be a reasonable first step for building a quantitative framework. Looking forward, our system of ODEs may be readily adapted for orthotopic or patient-derived xenografts, syngeneic transplant models, or genetically engineered models—because it is built from fundamental principles of mass transport—by identifying the values of the parameters that are specific for these tumors. In addition, our framework may be extended to describe the behavior of activity-based biomarkers such as fluorogenic peptide substrates designed to emit fluorescent signals after cleavage, or chemical substrates that covalently modify target proteases (14, 15). Here the model parameters related to probe pharmacokinetics will be different for each probe, and, within the tumor compartment, additional parameters will need to be incorporated to account for the mechanism by which the probe produces a detection signal. For example, for fluorogenic probes, these include the fluorescent intensity of a quenched and cleaved probe, background fluorescence of the target tissue, and clearance rate of the fluorescent reporter.

The main focus of this model is to address the limit of detection issue; thus we selected MMP9, a protease that is broadly dysregulated in solid cancers. MMPs have been implicated in cancer cell invasion and metastasis for over 35 years (38), and many of the hallmarks of cancer (39) are dependent on MMP activity. An ultrasensitive diagnostic for MMP9 activity may be used in clinical settings when the tumor type is known, such as after primary resection, when the surveillance protocol relies on CT or magnetic resonance imaging (MRI) despite their difficulties detecting metastasis 1–2 cm in diameter (35). MMP9 is expressed to activate VEGF ligands (40) during the angiogenic “switch” that occurs when nascent tumors are 1–2 mm in diameter; therefore, measuring MMP9 activity may allow detection of tumors smaller than the detection threshold of medical imaging. To address specificity in addition to sensitivity, the development of a cancer-type-specific diagnostic would require a multiplexed library of activity-based probes to account for the over 550 proteases expressed by the human genome that are dysregulated with disease or are organ-specific. Previous studies showed that the cleavage signatures of a 10-plex set of activity-based probes detected disease with significantly improved sensitivity and specificity compared with single probes, and identified activity-based signatures for CRC, liver fibrogenesis, and its resolution (3). Expanding our model to include multiple proteases as well as their endogenous inhibitors [e.g., tissue inhibitor of matrix metalloproteinases (TIMPs)] may allow prediction of...
Amine terminated eight-arm PEG (10 kDa, 20 kDa, and 40 kDa; Jenkem Technologies) was dissolved in PBS and passed through a 0.2-μm filter. DLS measurements were conducted on a Zetasizer (Malvern) in PBS and 50% PBS. A 20-fold molar excess of succinimidyl iodoacetate (SIA; Pierce) was reactivated to introduce thiol reactive handles. Excess SIA was removed by FPLC on a Superdex 200 column (AKTA purifier; GE Healthcare). Cysteine-terminated peptides (Q1 = 5FAM-GGPLGVRGKK(CPQ-2)-PEG2-C, CPC Scientific; F1 = eGydneneGyffasedGGGPLGVRGC, lowercase = o-spherostemer, Tufts University Peptide Core Facility) were then reacted with PEG at a 20:1 molar ratio before excess peptides were removed by FPLC. F1-PEG conjugated was further reacted with NHS-Vivotag750 (Perkin-Elmer) for imaging applications. NWs were synthesized according to previously published protocols and functionalized with peptides as above (41). All peptide–NP formulations were stored in PBS at 4 °C.

In Vitro Protease Assays. Q1–PEG (3 μM by peptide) was mixed with human MMP-9 (5 μg/mL working concentration; Enzo Life Science) in a 384-well plate at 37 °C in activity buffer (50 mM Tris, 150 mM NaCl, 5 mM CaCl2, 1 μM ZnCl2) containing 1% BSA and monitored with a microplate reader (SpectrMax Gemini EM). Marimastat (Tocris Bioscience) was dissolved in DMSO and used at a final concentration of 5 μM. Michaelis–Menten constants were determined by acquiring cleavage velocities at different substrate concentrations followed by mathematical fitting using GraphPad 5.0 (Prism).

In Vivo Imaging. All animal work was approved by the Committee on Animal Care at Massachusetts Institute of Technology (protocol 0414-02-17). To produce tumor xenografts, L5174T cancer cells (ATCC) were maintained in RPMI 1640 media supplemented with 10% FBS, and 1% Penicillin/Streptomycin.


For organ distribution, VT-750-labeled PEG or NWs were administered i.v. (5 μM, 200 μL PBS) before necropsy. Major organs were placed in five volumes (v/v) of homogenization buffer (20 mM Tris, 1% SDS, pH 8.0) and homogenized by gentleMACS Octo dissociator (Miltenyi Biotec). VT-750 fluorescence in the supernatant was quantified against free VT-750 by Licor scanner. For half-life measurements, fluorescent NWs or PEG was administered i.v. (5 μM, 200 μL PBS) and ~10 μL of blood was collected by retroorbital draws using microhematocrit tubes (VWR). Samples were transferred into 100 μL of PBS containing 5 mM EDTA and spun at 1,000 g to pellet blood cells. Fluorescent VT-750 levels were quantified against a standard curve generated by a three-point standard curve generated by whole animal imaging (IVIS, Xenogen). For absolute quantification of urine fluorescence, urine samples were collected and quantified against VT-750 Fluorb standards on a Licor Odyssey scanner.

Mathematical Modeling. Our system of ODEs was solved with MatLAB using the differential equation solver ODE15s. The model was fitted to our experimental data by minimizing the square of the difference with a default tolerance of 10-3. See SI Text for model equations.

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