Platinum-Triggered Bond-Cleavage of Pentynoyl Amide and N-Propargyl Handles for Drug-Activation

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ABSTRACT: The ability to create ways to control drug activation at specific tissues while sparing healthy tissues remains a major challenge. The administration of exogenous target-specific triggers offers the potential for traceless release of active drugs on tumor sites from antibody–drug conjugates (ADCs) and caged prodrugs. We have developed a metal-mediated bond-cleavage reaction that uses platinum complexes [K₂PtCl₄ or Cisplatin (CisPt)] for drug activation. Key to the success of the reaction is a water-promoted activation process that triggers the reactivity of the platinum complexes. Under these conditions, the decaging of pentynoyl tertiary amides and N-propargyls occurs rapidly in aqueous systems. In cells, the protected analogues of cytotoxic drugs 5-fluorouracil (S-FU) and monomethyl auristatin E (MMAE) are partially activated by nontoxic amounts of platinum salts. Additionally, a noninternalizing ADC built with a pentynoyl traceless linker that features a tertiary amide protected MMAE was also decaged in the presence of platinum salts for extracellular drug release in cancer cells. Finally, CisPt-mediated prodrug activation of a propargyl derivative of S-FU was shown in a colorectal zebrafish xenograft model that led to significant reductions in tumor size. Overall, our results reveal a new metal-based cleavable reaction that expands the application of platinum complexes beyond those in catalysis and cancer therapy.

INTRODUCTION

The targeting of potent drugs with tumor-specific ligands is an essential feature of drug delivery and cancer therapy. Notable in this field are antibody–drug conjugates (ADCs) that use an antibody to transport a drug to cancerous cells and endogenously release it by hydrolysis (low pH, reduction of disulfide bonds) or by proteolysis (e.g., cathepsin B protease). Although the cleavage of ADC linkers with endogenous triggers is the simplest method for drug release, external small-molecule triggers for extracellular drug release may be more advantageous because they avoid any disparity in cleavage rates caused by variable biology across subjects, and drug release is not dependent on the concentrations of cellular triggers. In fact, ADCs built with protease cleavable linkers for drug release have been shown recently to not depend on the cathespin B protease function for efficient and targeted cancer-cell killing. The promise of controlled prodrug activation has fueled research into new triggers that enable bond-cleavage reactions to unlash bioorthogonal protecting groups, which deactivate otherwise potent drugs. Robillard and co-workers pioneered the development of tetrazine-triggered drug delivery from ADCs. By using a noninternalizing ADC consisting of a diabody conjugated to trans-cyclooctene-linked drug monomethyl auristatin E (MMAE), the allylic carbamate-containing linker can rapidly react with a tetrazine through an inverse-electron-demand Diels–Alder reaction. The drug is released within the extracellular tumor environment and showed efficacy in delaying tumor growth in xenograft mice models. Other chemical- or light-mediated decaging reactions have also been developed with an array of applications that range from in situ activation of prodrugs to the gain-of-function study on proteins.

Although the recent results with noninternalizing ADCs for click-triggered drug release show promise, there are several issues that remain to be improved, such as the lack of tumor-selectivity of the chemical triggers or their short in vivo retention times that may result in reduced release of the cytotoxic payloads and thus lower efficacy. Additionally, in such applications the tumor payload concentration is determined by the cell-surface antigen expression, which in some cases may be too low to achieve a useful therapeutic effect. 

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Metal-mediated decaging of prodrugs has been more extensively reported than small-molecule-mediated decaging. Unlike chemical triggers, transition metals can be catalytic, which allows their use in substoichiometric amounts. In these cases, only very small amounts of catalytic metal are required to achieve the desired pharmacologic effect, thereby reducing toxicity and side reactions. This feature was recently demonstrated by Weissleder and co-workers using palladium nanoparticles that accumulate in tumor cells and serve as cellular catalysts for the activation of different model prodrugs and resulted in tumor growth inhibition.

Palladium-mediated decaging is indeed the most studied method for prodrug activation, which relies on the cleavage of terminal propargylic and allylic carbamates moieties introduced into small molecule drugs. Recently, our group developed an internal bifunctional thioether propargyl carbamate linker with a conjugating unit for protein modification and MMAE for palladium-mediated drug release from a nanobody-drug conjugate in cellular systems. Other metals, such as ruthenium and gold, have also been explored for cleavage and drug release. One of the latest additions to this field was the recent report by Peng Chen and co-workers on a copper-releasable reaction for protein gain-of-function and drug activation. Together, these examples highlight the potential of metal-mediated cleavage as a means to achieve controlled and chemically defined drug release.

Whereas the utility of the above-mentioned metals for decaging applications has been extensively demonstrated, other metals have not yet been sufficiently explored. For instance, platinum is widely used in catalysis, but has found few applications in chemical biology, possibly as a result of its intrinsic cytotoxicity. However, in the context of cancer therapy, we hypothesized that the use of platinum complexes [e.g., Cisplatin (CisPt) used in the clinic] as catalysts for cleavage reactions could be propitious for bioorthogonal activation of prodrugs in tumor cells.

"Bioorthogonal" is perhaps unfitting terminology for a compound known to react with water, nucleic acids, amino acids, and proteins. However, CisPt is one of the most commonly used chemotherapy drugs, being used to treat up to 20% of cancer patients. CisPt was deemed a suitable reagent for the development of a drug decaging reaction because it is highly reactive (half-life in humans of ~30 min), accumulates in the tumor, and most importantly, is not present in human biology.

In this way, the activation of prodrugs at the tumor site when the chemical trigger has already accumulated may represent a major achievement. It may be conceivable for metal concentrations to reach 0.25 to 3.7 μg/g of tumor. For a 1 cm³ tumor (approximately 1 g wet weight), the concentration of CisPt is estimated to be 0.83–12.3 μM.

Therefore, we were interested in investigating new bioorthogonal cleavage reactions catalyzed by platinum for applications in prodrug activation. Here, we demonstrate that pentonyl tertiary amide and N-propargyl handles introduced into small-molecule drugs are successfully decaged in aqueous solution and cell media using nontoxic amounts of platinum salts. This strategy was successfully applied to small molecule prodrug activation (MMAE and 5-FU) and further extended to drug release from a noninternalizing ADC in cancer cells. Finally, we show that CisPt-mediated bond cleavage can be used to activate a 5-FU prodrug in a zebrafish xenograft model for treatment of colorectal cancer.
successfully trialed with K$_2$PtCl$_6$ as a representative Pt(IV) species (Figure 1d, Entries 5 and 6; Figure S7) and NaAuCl$_4$ (Figure S8) with good yields. Overall, these results are important because they demonstrate a decaging reaction of stable protected tertiary amides by using substoichiometric amounts of platinum complexes that could function in water, open air, and without need of extreme temperatures or complex ligands. It is important to note that even after all of the starting material has been consumed (as evidenced by loss of terminal alkyne proton), not all of it has decomposed to release the amine (Figure 1e). This lack of conversion is likely due to side reactions, and nucleophilic attack on the alkyne, which seems the most plausible mechanism. To elucidate this a pentynoyl secondary amide (compound S2) was reacted with K$_2$PtCl$_4$ and NaAuCl$_4$ under similar conditions (Figures S9 and S10). We found that the reaction proceeds with much lower extents of decaging, likely due to the amide nitrogen competing as a nucleophile to yield a stable cyclized product, and thus a smaller yield of released amine.

**Mechanistic and Kinetic Studies of the Platinum-Mediated Decaging Reaction.** To further study the platinum decaging reaction, the pentynoyl tertiary amide was conjugated to a naphthalimide-based fluorophore to generate fluorescent quenched probe 7 (Figure 2a, see SI for synthetic details). The reaction was then monitored by the increase in fluorescence upon removal of the protecting group to form fluorescent probe 8. With 50 equiv of K$_2$PtCl$_4$ or CisPt, we found that the fluorescence was restored over a period of 200 min for K$_2$PtCl$_4$ and 300 min for CisPt (Figure 2b), with complete consumption of 7 and formation of corresponding "turned ON" fluorophore, as indicated by LC−MS analysis (Figure S11). For both metals the conversion was accompanied by an initial steady state followed by a marked increase of the fluorescence, which suggested the formation of an activated intermediate. Indeed, it is known that platinum complexes form a series of reactive intermediates by successive replacement of the chloro ligands by water or hydroxyl groups. We hypothesized that formation of such an aqua intermediate early on could be responsible for the activation of the platinum complexes. This hypothesis was verified using LC−MS studies to follow formation of K$_2$PtCl$_4$− and CisPt−aqua complexes over time, which occurred within 6 h (Figures S12 and S13). Consistent with this hypothesis, platinum salts failed to form the aqua complexes when incubated in the presence of N,N-dimethylformamide (DMF). On the basis of...
these observations, we further studied the kinetics of the releasing reaction after formation of the aqua complexes (6 h in water/DMF at 37 °C). As expected, activation of the platinum salts significantly accelerated the turn-on half-time from $t_{1/2} = 171$ min to $t_{1/2} = 30$ min for K$_2$PtCl$_4$ and from $t_{1/2} = 276$ min to $t_{1/2} = 60$ min for “CisPt” (Figure 2c and Table S1). Accordingly, if the reaction was performed in pure DMF, then formation of the decaged probe was not observed (50 equiv of K$_2$PtCl$_4$ or CisPt for 14 h at 37 °C). This result is in agreement with previous LC–MS studies that suggested the requirement of water to generate the active catalyst. The activation of metal chloride in aqueous solvents has few precedents but has been reported for gold complexes. Finally, the compatibility and efficiency of the reactions were tested under physiological conditions. The activated aqua complexes were shown to persist in complete DMEM cell media for at least 16 h at 37 °C, as assessed by LC–MS analysis, although a significant decrease in their concentration was observed over time (Figure S15). Later, the reactions were shown to proceed in cell media with conversions of 69% for K$_2$PtCl$_4$ (50 equiv) and 17% for CisPt (150 equiv) after 14 h at 37 °C (Figures S16 and S17). Similarly, the reaction was also trialed in high salt concentration buffers with high efficiency ($t_{1/2} = 36$ min for 50 equiv of K$_2$PtCl$_4$ and $t_{1/2} = 105$ min for 100 equiv of CisPt, 37 °C in E3 medium, Figures S18 and S19).

Having found an efficient platinum complex for decaging pentynyl tertiary amides, we turned our attention to the determination of the rate constant of the reaction (Figure 2d). By fitting the appearance of 8 in the presence of increasing amounts of metal complexes and by using pseudo-first order conditions, the reactions were found to have second order rate constants of 0.230 ± 0.004 M$^{-1}$ s$^{-1}$ for K$_2$PtCl$_4$ and 0.080 ±

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**Figure 2.** Examination of the platinum-catalyzed bioorthogonal cleavage reaction. a. Naphthalimide-based fluorogenic probes were used to study the cleavage efficiency of the platinum reaction for decaging alkyne-containing molecules. The caged naphthalimide derivatives exhibited high stability in solution and cell media and their quenched fluorescence could be reactivated upon removal of the caging group ($\lambda_{ex} = 445$ nm, $\lambda_{em} = 545$ nm). b. Changes in fluorescence intensity during the time course of the decaging reaction between fluorogenic probe 7 and platinum salts (K$_2$PtCl$_4$/CisPt). c. Determined half-time for the reaction of 7 with activated and nonactivated platinum salts. d. Decaging kinetics for the pentynoyl amide fluorophore. Rate constants were determined under pseudo first order conditions with a 50 μM final concentration of probe 7 and 10–50 equiv of aqua platinum metals. e. Kinetics profiles of the decaging reaction in the presence of the metal poisons CS$_2$ and EDTA. Error bars represent ± s.d. (n = 3). All experiments were repeated 3 independent times. f. Calculated mechanism for the depropargylation reaction catalyzed by Pt with model substrate 4a. Calculations were performed with an implicit solvent model for water. Geometries and frequencies were calculated with the functional revPBE and, to obtain very accurate energetics, single point energy calculations with DLPNO–CCSD(T) and counterpoise corrections were employed to suppress basis set superposition errors.
0.002 M$^{-1}$ s$^{-1}$ for CisPt (Figure 2d). These reaction rates are similar to those reported for other metal-assisted decaging reactions.\textsuperscript{57}

To determine the nature of active species involved in the decaging reaction, we performed kinetic experiments with carbon disulfide (Figure 2e, Table S2). CS$_2$ acts as a catalyst poison for homogeneous and heterogeneous Pt(0) reactions, although Pt(II) species are unaffected. As seen in Figure 2e, the reaction rates are similar to those with or without CS$_2$. This result can be attributed to the noninvolvement of Pt(0) species in the reaction. However, the reaction rates were significantly affected by the addition of ethylenediamine tetracetic acid (EDTA; Figure 2e), possibly due the participation of Pt(II) in the reaction.

We also performed computational studies to help understand the reaction mechanism (Figure 2f). These studies suggest that the most probable operating reaction pathway of substrate 4a is a stepwise process involving the coordination of substrate molecule to Pt(II), followed by an intramolecular attack of the carbonyl oxygen of the Pt-coordinated substrate (CS$_8$) to the pentynyl moiety, which gives five-membered ring intermediate CP. Different pathways to decomposition of CP were explored (Figure S20); the lowest energy one was the hydration of CP leading to formation of intermediate Q$_4$, which readily decomposes to liberate free amine 6a. The metal complex is then recovered in a subsequent step by hydrolysis of 5 (Figure 2f). The complete calculation for the first reaction turnover of the main mechanism is depicted in Figure S21 and SI Movie. This mechanism is further supported by the identification of intermediate species CS$_8$ by LC–MS (Figures S22–S26). The main difference observed for the reaction with substrate 3a relative to 4a was the higher free energy of activation ($\Delta G^\ddagger$ = 2.75 kcal mol$^{-1}$) for the intramolecular attack of the carbonyl oxygen at the pentynyl moiety. However, both substrates share the same energy barrier for hydration of CP and release of 6a (Figure S21).

**Extending the Decaging Reaction to N-Propargyl Group.** Following the discovery of a platinum-cleavable group, we hoped to extend the scope of handles that could be used for decaging. Metal-mediated decaging of N-propargyl handles has been widely explored to modulate the cytotoxic activity of antineoplastic drugs in a controlled manner.\textsuperscript{22,25} On this basis, we investigated the possibility of using N-propargyl groups introduced on drugs of interest for prodrug activation using platinum triggers. First, and similarly to the pentynyl amide reaction, an N-propargyl group was used to protect a secondary amine on a naphthalimide derivative to generate fluorogenic probe 9 (Figure 2a). As described above, we tested the reactivity of K$_2$PtCl$_4$ and CisPt before and after formation of the aqua complexes (6 h incubation at 37 °C in DMF/water). Once again, dissociation of the chloride anions in water was found to be crucial for triggering the reactivity of platinum complexes. Indeed, we found that reactions with aqua complexes are faster according to the calculated half-time for the “fluorescent reactions” (from $t_{1/2}$ = 200 ± 3 min to $t_{1/2}$ = 27 ± 3 min for K$_2$PtCl$_4$ and from $t_{1/2}$ = 628 ± 51 min to $t_{1/2}$ = 303 ± 34 min for CisPt; Table S3). The fluorescence-based assay was also employed to calculate the second-order rate constant for the reaction. Accordingly, the calculated rate constant was 0.120 ± 0.001 M$^{-1}$ s$^{-1}$ for K$_2$PtCl$_4$ and 0.0160 ± 0.0004 M$^{-1}$ s$^{-1}$ for CisPt (Figure S27). These results show that N-propargyls decage slower than pentynyl amides. As a reference, the same study was performed with palladium complex Pd(OAc)$_2$, which behaved slightly better than the platinum salts, to promote formation of 10 with a second-order rate constant of 0.39 ± 0.015 M$^{-1}$ s$^{-1}$ (Figure S28). The reaction was also subjected to CS$_2$ and EDTA poisoning. CS$_2$ had no effect but EDTA completely inhibited the reaction, which indicates the participation of Pt(II) species (Figure S29 and Table S4). On the basis of these results and LC–MS analysis after 2 h of reaction between K$_2$PtCl$_4$ and probe 9 (Figure S30), we propose that the first turnover of the reaction proceeds as recently disclosed for palladium depolymerylation,\textsuperscript{26} i.e., (i) co-ordination of Pt(II) to alkylene moiety, (ii) attack of a H$_2$O molecule at the propargyl terminal carbon to form an enol, (iii) tautomeration to a more stable aldehyde complex, and (iv) C–N bond cleavage by either hydrolysis or β-N elimination followed by hydration of Pt-complex (Figure S31). Finally, we investigated the ability of platinum salts to remove the propargyl protecting group in cells (DMEM) and zebrafish (E3) media. The reaction with the fluorogenic probe was monitored for K$_2$PtCl$_4$ and CisPt for 14 h at 37 °C. Efficiencies in E3 media were generally high with the reaction complete in 60 and 150 min for K$_2$PtCl$_4$ and CisPt, respectively (Figure S32). In DMEM, cleavage was less efficient with conversion yields of 67% for K$_2$PtCl$_4$ (50 equiv) and 30% for CisPt (150 equiv) after 14 h at 37 °C (Figure S33).

**Platinum-Mediated Decaging in Living Cells.** To verify whether platinum-mediated depolymerylation would function in cell culture, a pentynyl amide derivative of antineoplastic drug MMAE was synthesized. MMAE is the drug present in the ADC brentuximab vedotin that is in clinical use to treat patients with relapsed Hodgkin lymphoma and systemic anaplastic large-cell lymphoma,\textsuperscript{60} and remains the drug of choice for antibody-targeted therapies. In addition, a N-propargyl 5-fluorouracil (pFU) derivative was also tested, which was found to be efficiently decaged and activated with gold nanoparticles\textsuperscript{31} and palladium complexes.\textsuperscript{25} When MMAE-am was treated in DMF/water (1:1) for 4 h with 10 equiv of K$_2$PtCl$_4$, complete consumption of MMAE-am was seen by LC–MS with 37% release of MMAE along with the formation of the intermediate Q$_4$, (Figures S20 and S34). In a similar fashion, decaging of pFU proceeds with yields of 46% ± 2 and 72% ± 2 for K$_2$PtCl$_4$ and CisPt, after 14 h reaction with 2 equiv, at room temperature and 37 °C, respectively (Figures S35–S37). These prodrugs (MMAE-am 11 and pFU 12, see the SI for synthetic details) were reacted with platinum salts in cell culture in the hope of observing a “turn-on” of toxicity. Unfortunately, the chemotherapeutic CisPt has a narrow window of nontoxic concentrations for efficient decaging in cells.\textsuperscript{33} Indeed, CisPt was demonstrated to be toxic in HeLa cells at concentrations as low as 2.5 μM (Figure S38). On the contrary, platinum salts K$_2$PtCl$_4$ and K$_2$PtCl$_6$ did not significantly influence the viability of HeLa cells at concentrations below 50 μM (Figure S38). With both prodrugs, an increase of about 2-fold in toxicity could be observed for some of the tested concentrations when reacted with K$_2$PtCl$_4$ over 3 days in cell culture (Figure 3a and 3b; e.g., 1 nM of MMAE-am and 50 μM of pFU). In contrast, no decrease of cytotoxicity was observed in cells treated independently with sFU 13, a non-decaging control derivative, or in combination with K$_2$PtCl$_4$ (Figure 3b). These control studies indicate that sFU was not generated because the alkyl handle does not undergo decaging by K$_2$PtCl$_4$.\textsuperscript{33}
for 9 reaction is more susceptible to the presence of nucleophiles in ADC design.60 Ideally a CisPt-cleavable ADC would be adapted for this purpose because MMAE is a common payload release from an ADC. The caging group of MMAE-am the tertiary amide caging group for chemically controlled drug—active drug in cells to induce cell death. Culture and could achieve release of su

It is important to note that for both prodrugs the addition of K2PtCl4 did not restore their toxicity to the level observed for unmodified MMAE and 5-FU drugs (Figures S39 and S40). Although a 2-fold increase in toxicity for the prodrug activation may look modest, it is important to mention that this is considered relevant given the slow reaction rates possible at the low concentration of K2PtCl4 complex tolerated by cells. Indeed, this low reagent concentration was necessary to ensure the platinum complex remained nontoxic. On top of this, in vitro studies with probes 7 and 9 revealed that the presence of nucleophiles (e.g., glutathione) ends in lower conversions into the corresponding decaged products. It should be noted, however, that even in the presence of high concentrations of glutathione (e.g., 1.5 mM) the reaction still proceeds with moderate rates (t1/2 = 197 min for 7 + K2PtCl4; t1/2 = 246 min for 9 + K2PtCl4; Table S5). Regarding CisPt, we found that the reaction is more susceptible to the presence of nucleophiles (e.g., t1/2 = 921 min for 7 + CisPt in the presence of 0.5 mM of glutathione; Table S6). This deactivation of the metals in the presence of nucleophiles is in line with the modest decaging yields observed in the cell studies. This is an issue that could be further improved, for example, by using platinum-based nanoparticles known to have reduced toxicity and higher payload concentrations or by using platinum complexes stabilized with different organic ligands in a way to optimize the metal reactivity.65 Our data, however, demonstrate that decaging reactions with platinum complexes are possible in cell culture and could achieve release of sufficient amounts of the active drug in cells to induce cell death.

**Platinum Decaging of ADC.** Next, we decided to extend the tertiary amide caging group for chemically controlled drug-release from an ADC. The caging group of MMAE-am 11 was adapted for this purpose because MMAE is a common payload in ADC design.66 Ideally a CisPt-cleavable ADC would be stable to cleavage by endogenous extra- or intracellular conditions. For this reason, we decided to use a carbonyl acrylic bioconjugation handle coupled to MMAE for antibody modification (Figure 4a; SI for synthesis).

To test the susceptibility of the conjugating linker to platinum decaging, compound 14 and K2PtCl4 (10 equiv) were incubated in DMF/water (1:1) at 37 °C for 18 h and analyzed by LC−MS (Figure S41). Release of MMAE was observed with complete consumption of 14 along with two potential

**Figure 3.** Platinum-mediated decaging in cells. HeLa cells were incubated with different concentrations of MMAE-am 11a or pFU 12b for 3 days with or without K2PtCl4 (20 μM, twice a day). Compound 13, a nondecaging alkyl-FU derivative, was used as a negative control. Toxicity was determined by AlamarBlue assay. Error bars represent ± s.d. (n = 3). Each experiment was repeated three times. The statistical significance of the differences between groups was evaluated with the unpaired t test. Statistical results: ns > 0.05, **P ≤ 0.01, ***P ≤ 0.001 and ****P ≤ 0.0001.

**Figure 4.** Platinum-mediated drug decaging from a noninternalizing ADC. a. Cysteine-selective and irreversible modification of the noninternalizing antibody F16 (anti tenasin-C) in IgG format with MMAE conjugating linker 14. IgG(F16) contains a single reactive cysteine at the C-terminal extremity of the light chain ideal for cysteine-specific modification. Briefly, a solution of F16 (7.1 μM) in sodium phosphate buffer (NaPi) pH 7.4 was treated with 14 (40 equiv) in MeCN to a final concentration of 10% v/v. The reaction was heated to 37 °C for 1 h, and reaction progress was monitored by LC−MS. The ADC was purified by dialysis into fresh NaPi buffer pH 7.4 with a 10 kDa MWCO overnight. b. Deconvoluted ESI−MS mass spectrum of the light-chain of F16. c. Deconvoluted ESI−MS mass spectrum of the light-chain of F16-14 that shows an exact drug-to-light-chain ratio of 1. d. Schematic of the platinum-mediated decaging of MMAE from a noninternalizing ADC. e. Cell viability of HeLa cells after treatment with F16-14 and subsequent decaging efficiency upon treatment with 20 μM K2PtCl4 twice daily. Cell viability was measured at day 3 by using AlamarBlue reagent. The statistical significance of the differences between groups was evaluated by using the unpaired t test. A p value ≤ 0.05 (***) was considered statistically significant. Error bars represent ± s.d. (n = 3). Experiments were performed three times.

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intermediates (Figure S41). We then went on and selected the noninternalizing F16 antibody for modification, which is specific to the alternatively spliced A1 domain of tenasin-C, found overexpressed in most solid tumors. A noninternalizing ADC ensures that as little ADC as possible will be metabolized by the cells and that the maximum possible drug release is due to extracellular decaging with platinum complexes. Site-selective conjugation is expected to occur at the engineered cysteine residues in each light-chain of F16 enabling the construction of a chemically defined ADC. Furthermore, the newly formed C−S bond between the linker and the antibody is stable and does not undergo thiol-exchange reactions as in the case of frequently used maleimides. Complete conversion to a homogeneous ADC was achieved after reaction of F16 for 1 h at 37 °C with the carbonyl acryl MMAE drug linker 14 in sodium phosphate buffer pH 7.4 as assessed by LC−MS (Figure 4b,c). Importantly, the heavy chain remained unmodified as expected considering the absence of reactive cysteines in the structure (Figures S42 and S43). Next, we performed the decaging in cells to release MMAE from the ADC (Figure 4d). With a cancer cell line (HeLa cells) as a model, we found F16−14 to be more toxic to cells at submicromolar concentrations in the presence of nontoxic amounts of the platinum complex K2PtCl4 (Figure S44). This tertiary amide decaging reaction should stimulate platinum-mediated MMAE delivery from antibodies in the context of targeted cancer therapeutics. Furthermore, a small model protein (ubiquitin-K63C) engineered with a single cysteine residue was modified with linker 14 for ease of analysis by LC−MS. When attempting decaging in vitro with CisPt, loss of MMAE followed by further degradation of the linker could be observed by LC−MS, which provides further evidence for the efficient release of the secondary amine drug from the protected tertiary amide protected conjugate (Figures S44−S47).

Cisplatin-Mediated Prodrug Decaging in Vivo. To test the in vivo efficacy of pFU and its combinatorial effect with CisPt, we used the zebrafish larvae xenograft model. This model is a fast in vivo platform with resolution to analyze crucial hallmarks of cancer, such as metastatic and angiogenic potentials but it is also highly sensitive to discriminate differential anticancer therapy responses with single-cell resolution. We first attempted to visualize the CisPt reaction by decaging fluorogenic probe 9 in larval zebrafish (Figure 5). This probe shows an increase in fluorescence of 22-fold upon removal of the propargyl group (Figure S48). For in vivo imaging, a set of zebrafish larvae were incubated with probe 9 for 24 h, washed with 1 h in embryonic medium and then further incubated with dimethyl sulfoxide (DMSO) or CisPt for 24 h (Figure 5a). Probe 9 and CisPt were used at the highest nontoxic concentration to the zebrafish embryos (9, 1 μM; CisPt, 34 μM; Figure S49). As shown in Figure 5b, the control group displays nearly no background fluorescence, but the CisPt-treated group showed an increased fluorescence (Figure 5c). This implies that probe 9 and CisPt are tissue-permeable and capable of reacting in vivo.

Before measuring efficacy of CisPt depropargylation, we assessed the maximum tolerated concentration for each compound: pFU 12, cFU 13, CisPt, pFU + CisPt, and cFU + CisPt in nontumor zebrafish larvae (Figure S49). Next, colorectal cancer (CRC) HCT116 zebrafish xenografts were generated as previously described. Briefly, 24 h post injection (hpi), xenografts were randomly distributed into different treatments: DMSO (control), pFU (1.65 mM), cFU (1.65 mM), CisPt (0.034 mM), pFU + CisPt (1.65 mM + 0.034 mM), and cFU + CisPt (1.65 mM + 0.034 mM). Xenografts were analyzed at 4, 6, and 7 days post injection (dpi), i.e., 3, 5, and 6 days post treatment (dpt), respectively (Figure 6). At 3dpt (4dpi) (Figure 6I), in the single treatments with pFU or CisPt, we could not observe any significant reduction of mitotic index (Figure 6I, m), induction of apoptosis (activated caspase 3, Figure 6I, n) or reduction of tumor size (Figure 6I, o). In contrast, the combinatorial treatment—pFU + CisPt—induced a significant antitumor synergistic effect manifested by a ~2 fold increase in apoptosis (Figure 6I, n; DMSO versus pFU + CisPt **P = 0.0033; pFU versus pFU + CisPt ****P = 0.0006) accompanied by 25% reduction of tumor size (Figure 6I, o; DMSO versus pFU + CisPt *P = 0.0279; Figure 6I, a) and 38% reduction of tumor size (Figure 6I, r; DMSO versus pFU + CisPt ****P < 0.0001; Figure 6II, c versus h). Finally, the 6 days of treatment (7 dpi) culminates in a ~45% tumor shrinkage (Figure 6III, s; DMSO versus pFU + CisPt ****P = 0.0010; Figure 6III, i versus l).

Importantly, by comparing the combined treatment of the nondecaging compound cFU with CisPt to the prodrug pFU with CisPt, it is clear that pFU was able to induce a more significant cytostatic (block proliferation) and cytotoxic effect (apoptosis and reduction of tumor size) than the control cFU at both 5 dpf (6 dpi) and 6 dpf (7 dpi; Figures S50 and S51). Also, the combined effect of pFU + CisPt was more pronounced than the combination of S-FU + CisPt, regarding proliferation (DMSO versus pFU + CisPt ****P < 0.0001; DMSO versus FU + CisPt *P = 0.0104; Figure S50i) and tumor size (DMSO versus pFU + CisPt ****P < 0.0001; DMSO versus FU + CisPt *P = 0.0273; Figure S50k). This might be related with the increased permeability of pFU (versus FU), which results in a more efficient intracellular delivery of FU after Pt decaging. In conclusion, our results show the efficient activation of the anticancer pFU in the

![Figure 5](https://dx.doi.org/10.1021/jacs.0c01622)

**Figure 5.** CisPt Decages the Fluorogenic Probe 9 in vivo. Zebrafish larvae were exposed to 9 diluted in embryonic medium for 24 h, followed by a 1 h wash in embryonic medium. Larvae were randomly distributed into two conditions: DMSO or CisPt for 24 h (a). Confocal image of zebrafish larvae exposed to 9 + DMSO (b) and 9 + CisPt (c).
Figure 6. CisPt-mediated prodrug decaging in zebrafish xenografts. HCT116 human CRC cells were fluorescently labeled with lipophilic CM-DiI (shown in red) and injected into the perivitelline space (PVS) 2 days post fertilization (4dpf) Tg(Fli1:eGFP) zebrafish larvae. Zebrafish xenografts were randomly distributed into treatment groups, daily treated with DMSO, CisPt, pFU, and pFU+CisPt and analyzed at 4, 6, or 7dpi for proliferation, apoptosis and tumor size. At 4 dpi, 6 dpi, and 7dpi, zebrafish xenografts were imaged by stereoscope (a−l) and by confocal microscopy (a′−l′ DAPI plus DiI, a′′−l′′ maximum projection of activated caspase 3). Proliferation (mitotic figures: m, p, *P = 0.0104, ***P = 0.0004, ****P < 0.0001; s, **P = 0.0023, ***P = 0.0002), apoptosis (activated caspase 3: n, **P = 0.0033, ***P = 0.0006; q, *P = 0.0126, ****P < 0.0001; t, **P = 0.0068) and tumor size (n° of tumor cells: o, *P = 0.0279; r, ****P < 0.0001; u, *P = 0.0411, **P = 0.0010) were analyzed and
CONCLUSIONS

In summary, we present a new decaging reaction of alkynes with platinum complexes for the release of secondary amines from otherwise stable tertiary amides, both in mammalian cell culture and in living organisms. This reaction was shown to proceed by platinum-mediated intramolecular cyclization mechanism. Our data suggest that water, a necessary solvent in chemical biology applications, is working as a metal-activating agent. Molecular electronic structure calculations further corroborated the mechanism of the reaction which was also supported by LC–MS characterization of the intermediates. The reaction can proceed catalytically under certain conditions and was later extended to N-propargyl groups with comparable efficacies to that of palladium-mediated depropargylation. The caging group was adapted for the synthesis of a noninternalizing ADC, which results in drug release upon treatment with platinum complexes in cancer cells. The reaction was also adapted and demonstrated to function in a colorectal cancer zebrafish xenograft model with nontoxic amounts of CisPt to activate a prodrug of anticancer agent 5-FU, which led to a significant tumor reduction in vivo.

The work disclosed here represents a significant addition to the toolbox of decaging strategies for chemical biology applications. Indeed, the platinum-mediated cleavable reaction can be accomplished in aqueous systems having high concentrations of salts with high yields and reaction rates, similar to those observed for the standard palladium decaging metal. The reaction is, however, susceptible to the presence of nucleophiles resulting in slower rates (~6–15 times slower). We further demonstrate the compatibility of the reaction in cellular environments. Although the reaction is suitable for drug activation on cells inducing cytotoxicity, the presence of a range of biomolecules/nucleophiles significantly reduces the overall yield. These results are suggestive of instability of the Pt complexes, probably by formation of bioinorganic complexes. Although the active aqua Pt species have a limited lifetime in cell media, they persist long enough to be partially effective.

Our work was conceived on the hypothesis that platinum complexes could be used for prodrug activation on tumors during CisPt chemotherapy. The instability of the platinum complexes in physiological/biological conditions preclude the application envisioned. Further studies are needed to obtain Pt complexes compatible for such in vivo applications, but these results set the stage for future developments on platinum-mediated decaging reactions.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c01622.

Detailed methods, characterization data and additional figures (PDF)

Movie with metadynamics calculations (MOV)

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Figure 6. continued

quantified. Graphs represent fold induction (normalized values to controls) of Avg ± SEM. The number of xenografts analyzed is indicated in the representative images and each dot represents one zebrafish xenograft. Statistical analysis was performed using an unpaired test. Statistical results: ns > 0.05, *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001, and ****P ≤ 0.0001. All images are anterior to the left, posterior to right, dorsal up, and ventral down. Scale bar 50 μm.
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