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Bistable Photoswitch Allows in Vivo Control of Hematopoiesis

Lea Albert, Jatin Nagpal, Wieland Steinchen, Lei Zhang, Laura Werel, Nemanja Djokovic, Dusan Ruzic, Malte Hoffarth, Jing Xu, Johanna Kaspareit, Frank Abendroth, Antoine Royant, Gert Bange, Katarina Nikolic, Soojin Ryu, Yali Dou, Lars-Oliver Essen, and Olalla Vázquez*

ABSTRACT: Optical control has enabled functional modulation in cell culture with unparalleled spatiotemporal resolution. However, current tools for in vivo manipulation are scarce. Here, we design and implement a genuine on–off optochemical probe capable of achieving hematopoietic control in zebrafish. Our photopharmacological approach first developed conformationally strained visible light photoswitches (CS-VIPs) as inhibitors of the histone methyltransferase MLL1 (KMT2A). In blood homeostasis, MLL1 plays a crucial yet controversial role. CS-VIP 8 optimally fulfills the requirements of a true bistable functional system in vivo under visible-light irradiation, and with unprecedented stability. These properties are exemplified via hematopoiesis photoinhibition with a single isomer in zebrafish. The present interdisciplinary study uncovers the mechanism of action of CS-VIPs. Upon WDR5 binding, CS-VIP 8 causes MLL1 release with concomitant allosteric rearrangements in the WDR5/RbBP5 interface. Since our tool provides on-demand reversible control without genetic intervention or continuous irradiation, it will foster hematopathology and epigenetic investigations. Furthermore, our workflow will enable exquisite photocontrol over other targets inhibited by macrocycles.

INTRODUCTION

The precise photocontrol of molecular activity to systematically trigger phenotypic traits opens unprecedented venues for in vivo elucidation of complex biological processes. Since the first genetically encoded photoreceptors in neuroscience were developed, the field has evolved. Now photopharmacology controls functional outputs within unmodified targets bypassing gene delivery and has achieved in vivo modulation of processes related to neurobiological phenomena like vision and membrane transport, or, also, cytoskeleton dynamics. However, these examples involve UV light and continuous irradiation and do not achieve the desirable on–off effect of photocages.

Epigenetics coordinates gene expression responsible for balancing hematopoiesis. Its improper orchestration causes aberrant hematopoietic stem cell (HSC) differentiations and malfunction leading to hematological malignancies. The histone methyltransferase MLL1 (KMT2A) entails the so-called MLL1 complex (MLL1, WDR5, RbBP5, Ash2L, Dpy30), where subunit interactions control MLL1’s activity. As chromatin-modifier, the MLL1 complex is vital for sustaining both hematopoiesis and HSC self-renewal. Furthermore, MLL1 is a potent oncogenic driver in hematopoietic cancers by influencing HOX gene expression. Interestingly, MLL1 may maintain its hematopoietic roles using mechanisms that do not depend on its histone methyltransferase (HMT) activity but on protein–protein interactions (PPIs). The functional consequences of PPI modulations within the MLL1 complex remains elusive. To decode the complex biology underlying MLL1-dependent hematopoiesis, we require integrative approaches involving in vivo models, which are interrogated in highly controlled fashion. To our knowledge, there are no precedents of in vivo photocontrol of hematopoiesis. Such tools would elucidate the molecular mechanisms behind blood cell formation and may lead to novel therapeutics for hemopathies.

Herein, we report conformationally strained visible-light photoswitches (CS-VIPs), which allow quantitative functional conversion by visible-light irradiation in cell culture and zebrafish. CS-VIPs undergo exceptionally slow relaxation (five months) in aqueous solution. Employing crystallography, hydrogen–deuterium exchange mass spectrometry (HDX-MS), molecular dynamics (MD), and biological assays, we define their action mechanism on the MLL1 multiprotein complex. Importantly, our most potent photopharmacological agent, CS-VIP 8, is a true on–off switch that enables precise in vivo photocontrol over hematopoietic function without harsh UV light or wash-out.

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RESULTS AND DISCUSSION

Design and Synthesis of the CS-VIPs. Recently, we reported in vitro MLL1 photocontrol, which ultimately affected leukemia cell proliferation. However, the modest activity difference between isomers precluded in vivo experiments. We envisioned that the inclusion of a molecular transducer into a structurally defined scaffold should impart high conformational restraints and, thereby, achieve different biological output between photoisomers. The potent MLL1 inhibitor MM-401 (Figure 1A) is ideal to test if strained photoswitchable cyclopeptides can exert effective photo-modulation of the molecular recognition process of MLL1 affecting in vivo hematopoiesis. As a photoswitch, we chose Hecht’s OF4Azo due to its excellent photochemical properties. Initially, we explored 13 CS-VIPs (Figure 1A), which differed in OF4Azo position, D-phenylglycine (D-Phg) presence, and ring size. D-Phg can epimerize; therefore, we considered its elimination and exchange by D-phenylalanine (D-Phe). Brieﬂy, solid phase peptide synthesis (SPPS; Figure 1B) enabled our linear precursors where OF4Azo was always on-resin incorporated at last, and arginine was Boc-protected to circumvent lability under Fmoc-deprotection and Pbf side reactions, respectively. Afterward, they were either head-to-tail or side-chain-to-tail cyclized at high dilution (1 mM) under irradiation at 520 nm for 3 min to steer the intramolecular product via cis isomerization.

Photochemical Characterization. As an illustrative example, the spectroscopic properties of CS-VIP 8 were analyzed in detail (Figure 2). Its UV–vis spectra displayed the expected absorbance bands of OF4Azo-containing compounds (Figure 2B). Photoisomerization was fully reversible without photodegradation (Figure 2C). Interestingly, the thermodynamically unstable cis isomer is kinetically highly stable in organic solvents (OF4Azo half-life: 700 days at rt in DMSO, 92 h at 60 °C in acetonitrile). In contrast, thermal cis → trans conversion proceeds faster in water (OF4Azo linear peptide cis content 86% at 520 nm photostationary state (PSS); 76% after 7 days). Importantly, when OF4Azo is incorporated into the conformationally restricted CS-VIP 8, its extraordinary stability was not only retained in water, but also surpassed nonrestricted OF4Azo molecules and unsubstituted azobenzene-containing cyclopeptides. Indeed, our photoswitchable peptide maintained its equilibrium state by at least five months (Figure 2D,E). Notably, the 520 nm PSS achieved 95% cis-isomer (cis:trans 19:1), whereas the 405 nm
PSS formed a cis:trans mixture in a ratio of ~1:10. Consequently, a switch between these PSS causes an ~9.5/18-fold change of the cis/trans-isomer concentration, respectively. Finally, the stability against glutathione (GSH) was also improved (Figure S18). Preliminary photochemical studies of the other CS-VIPs showed similar behaviors.

Optimization of CS-VIP Binding to WDR5. Fluorescence polarization (FP) based assays revealed the binding of our CS-VIPs to WDR5 (Table 1, Figure S19). Importantly, εF/Azo incorporation generally retained the parental nanostructure for most CS-VIPs (4--9 and 11). Affinity variations depended on substitution pattern, ring size, and the effective concentration of the binding-active isomer as set by the chosen PSS. CS-VIPs 1–3, without MM-401 isobutyrylated N-terminus displayed either no detectable (CS-VIP 1 and 3) or low micromolar (CS-VIP 2) binding. In silico experiments of CS-VIPs 1 and 3 revealed that the key arginine is not inserted in WDR5 cavity (Figure S36). Our experiments reassert the positive effect of the isobutyrylated N-terminus on WDR5 binding affinity.

Considering CS-VIPs with the same ring size, the phenyl group is crucial for high affinities, i.e., CS-VIP 10–12 devoid of this moiety displayed micromolar affinities and lacked light-dependent switching. In our virtual docking (VD), these binding poses were also ranked lower (Table S6, Figure S37) than the ones bearing D-Phg (CS-VIP 4–6, Figure S37) or D-Phe (CS-VIP 7–9, Figure S38). The D-Phe-containing CS-VIPs displayed the highest affinity differences between isomers. Also, for the inactive CS-VIP 1, the substitution of D-Phg for D-Phe (CS-VIP 2) increased affinity being consistent with VD (Table S6, Figure S36). In contrast to CS-VIP 1 and CS-VIP 3, packing interactions between the arginine and the WDR5 residues F133, F263, and C261 could be predicted for CS-VIP 2 (Figure S36). Consequently, our D-Phe modification not only improved CS-VIP synthetic accessibility but also functional properties.

Optimal ring size is decisive for maximizing both interaction and conformational changes between photoisomers. For us, calculated CHEMPLP values (Table S6) suggested that the 2-carbon linker (ε-Dab) had the ideal length to supply the best CS-VIPs: 5 and 8, which we verified experimentally (Table 1, Figure S19). The latter excels with an almost 10-fold switch of binding affinity. This agrees with the theoretical limit for a compound being only active in the cis-state. To confirm this, we adjusted the FP curve fit parameters to the effective cis concentration of CS-VIP 8 (Figure 2F). We obtained an excellent concordance between the Ki values of the 520 nm PSS and the 405 nm PSS, when considering the residual cis isomer concentration (10%). Besides, when the samples were

Table 1. CS-VIP Inhibition Constants to WDR5 by FP-Based Competitive Assays*

| CS-VIP | PSS at 405 nm Kc [µM] | PSS at 520 nm Kc [µM] | ratio |
|--------|-----------------------|-----------------------|-------|
| 1      | n.c.                  | n.c.                  |       |
| 2      | ~3.35b                | ~1.31b                | 2.56  |
| 3      | n.c.                  | n.c.                  |       |
| 4      | 0.043 ± 0.006         | 0.079 ± 0.016         | 1.84  |
| 5      | 0.051 ± 0.010         | 0.008 ± 0.002         | 6.38  |
| 6      | 0.394 ± 0.023         | 0.239 ± 0.071         | 1.65  |
| 7      | ~0.448b               | 0.104 ± 0.010         | 4.31  |
| 8      | 0.079 ± 0.004         | 0.008 ± 0.0008        | 9.88  |
| 9      | 0.831 ± 0.054         | 0.137 ± 0.012         | 6.07  |
| 10     | ~1.10b                | ~1.52b                | 1.38  |
| 11     | 0.688 ± 0.004         | 0.459 ± 0.060         | 1.50  |
| 12     | ~3.69b                | ~2.90b                | 1.27  |
| 13     | ~19.3b                | ~6.42b                | 3.00  |

*Mean values from at least two independent measurements (in triplicates); [fluorescent tracer] = 20 nM; fluorescent tracer Kc = 0.00104 ± 0.0005 µM. b Estimated values due to a lack of the bottom plateau. c Not calculable.
illuminated after WDR5 complexation (Figure S20), only the trans PSS mixture of CS-VIP 8 isomerized to the cis isomer yielding comparable $K_i$ values (Figure S20). In contrast, cis $\rightarrow$ trans photoisomerization is apparently slightly less effective when CS-VIP 8 is bound to WDR5 ($K_i$ at 405 nm PSS = 64.7 nM; $K_i$ for cis $\rightarrow$ trans = 47.1 nM; Figure S20).

Taken all together, our CS-VIP 8 variant is the most promising candidate for in vivo experiments as it maintains the nanomolar parental WDR5 affinity with an $\sim$10-fold difference between its photostationary isomers. These features significantly improved our previous results with photoresponsive linear peptides. As for the MM-401 enantiomer, the CS-VIP 8 enantiomer, CS-VIP 13, exerted a 244/803-fold diminished binding to WDR5 relative to its $K_i$ values recorded at 405 and 520 nm PSSs, respectively.

Next, we determined the cocrystal structures of the WDR5•CS-VIP 8 complex using different cis/trans ratios during crystallization (Figure 3). One by cocrySTALLization with WDR5 set to a 1:10 cis/trans mixture formed by 405 nm PSS (cis1, PDB: 7AXS), the other with the predominant cis isomer formed at 520 nm PSS (cis2, PDB: 7AXP). Both states share an interaction mode that closely mimics MM-401 (PDB: 4GM9, Figure S27) along with the isobutyl-d-Dab-Arg-Abu-d-Phe motif. Interestingly, the superposition of these cis-CS-VIP 8 complexes (Figure 3A) revealed different orientations of the FzAz and d-Phe. In the cis1 crystal the d-Phe ring enables $\pi$-stacking contacts with Y260 like d-Phg in MM-401 while the cis2 crystal has a higher solvent content and features less H-bonds and no $\pi$-stacking of cis-CS-VIP 8 to Y260 (Figures 3B, S25). UV–vis microspectroscopy displayed only the cis isomer of CS-VIP 8 within cis1 crystals (Figure S28). To reveal structural consequences of cis$\rightarrow$trans isomerization within complexes, we irradiated the cis1 crystallization setups of WDR5•cis-CS-VIP 8 after crystal formation at 405 nm to enforce cis$\rightarrow$trans switching (PDB: 7AXU). UV–vis measurements verified cis $\rightarrow$ trans photoisomerization within the crystals (Figure S28). However, this procedure caused complete diffusion of CS-VIP 8 out of WDR5-binding site as verified by a lack of electron density defining the ligand. (Figure 3D). Finally, we tested whether in crystallo irradiation at 405 nm of isolated cocrystals enables the intermediary states for the off-diffusion of the trans-CS-VIP 8 isomer. In crystallo switching to trans-CS-VIP 8 was observed by microspectroscopy at both 180 (Figure 3E) and 293 K (Figure S28). However, both structures (PDB: 7AXQ, 7AXX) apparently indicated X-ray mediated trans $\rightarrow$ cis back isomerization by displaying again the cis-CS-VIP 8 within the binding site. This phenomenon was confirmed by microspectroscopic analysis after X-ray exposure (Figures 3F, S26, and S28). Similar X-ray induced structural changes within photoisomerizable chromophores have been observed before in the past, e.g. in the phytochromes Cph2 and Pixf. We only observed increased thermal B factors for the $\epsilon FzAz$ group, especially in the cocrystal cryo-trapped at 180 K. These observations corroborated our previous FP experiments, which were additionally substantiated by computational calculations (see S1 in the SI). Overall, our biophysical analyses showed that only the cis-
CS-VIP 8 is active toward WDR5 and that the WDR5 presence increases the \( \text{trans} \rightarrow \text{cis} \) isomerization due to its preference for the bound \text{cis} isomer (Figure S28D).

**MLL1 Inhibition and Leukemogenesis Suppression.** Compounds capable of effectively binding to WDR5 have displayed MLL1 inhibition by blocking MLL1/WDR5 interaction, which decrease leukemia cell proliferation. In vitro functional radiometric HMT-assays with MLL1 complex (MLL1, WDR5, RbBP5, Ash2L) demonstrated that cis-CS-VIP 8 is a potent inhibitor of MLL1 methylation activity with a 12-fold difference depending on illumination (IC\(_{50}\) 405 nm PSS = 9.20 ± 3.04 μM; IC\(_{50}\) 520 nm PSS = 0.792 ± 0.259 μM; Figure S21). Along these lines, we tested CS-VIP 8’s effect on leukemia-related cells (MOLM-13) after 2-day incubation at different PSSs. The poor cellular uptake of CS-VIP 8 demanded pep-1 carrier as transfection method and relatively high concentrations. Isomers are distinct compounds with different pharmacokinetic properties. To exclude that isomer differences could emerge due to artifacts from transfection, we included a washing step, which will confirm that observed cytotoxicity is only attributed to intracellular processes. Compared to the 405 nm PSS, our photoswitchable CS-VIP 8 clearly gained inhibitory potency upon 520 nm irradiation in a dose-dependent manner (Figure 4A), which enabled the determination of its half-maximal inhibitory concentration (IC\(_{50}\)); i.e., 58.6 ± 1.6 μM (Figure 4B). However, under the same conditions, it was not possible to accurately obtain IC\(_{50}\) values for the trans-CS-VIP 8 (Figure 4B), but this must be higher than 150 μM at least. These data clearly demonstrate that CS-VIP 8 exhibits different isomer-dependent activities in MOLM-13 cells. In addition, we synthesized the enantiomer of CS-VIP 8 where the key i-Arg was replaced by i-Lys (CS-VIP 14). This lacked any detectable WDR5 binding (Figure S19N); therefore, CS-VIP 14 is a suitable control to evaluate the effect of the \( \delta F\alpha Azx \) itself without expecting any MLL1-dependent inhibition mechanism. Gratifyingly, the effect of cis-CS-VIP 8 in MOLM-13 cells was the most potent one, and as occurred for trans-CS-VIP 8, the dose–response experiments of both CS-VIP 14 isomers (Figure S22) could not be translated into accurate IC\(_{50}\) values under the same conditions (Figure 4B). These data suggest that the mechanism of the cis-CS-VIP 8 is distinguishable from that of the WDR5-independent CS-VIP 14 analogs as well as trans-CS-VIP 8.

**cis-CS-VIP 8 Affects MLL1-Complex Assembly.** Despite the crucial contribution of MLL1 activity in regulating leukemia transcription programs, its HMT function could be dispensable for normal hematopoiesis. However, depleting MLL1 components such as DPY30 and Ash2L displayed severe defects of lineage specification and differentiation.

Having established the structural basis of cis-CS-VIP 8 binding to WDR5, we employed HDX-MS to corroborate our findings in solution by analyzing WDR5 in either the absence or presence of cis-CS-VIP 8. Hereby, regions of reduced H/D-exchange of WDR5 clustered around the binding site as evidenced by our crystal structures and virtual docking (Figures 5A and S29). We next studied cis-CS-VIP 8’s impact on the assembly of the whole MLL1 complex. Initially, we corroborated the integrity of the recombinant MLL1 complex. Thus, regions of reduced H/D-exchange correlated well with reported interaction interfaces (Figures S30–S34). We then evaluated the H/D-exchange rates of the MLL1 complex with cis-CS-VIP 8 and compared it with untreated MLL1. We obtained altered profiles for all components except DPY30 and mapped those differences onto available crystal structures (Figure 5B). The MLL1 polypeptide, Win, displayed increased H/D-exchange rates as well as the core of the MLL1 SET domain, suggesting interaction loss with WDR5 and RbBP5/Ash2L (Figures 5B and S30). Correspondingly, amino acids 336–354 of RbBP5 bridging between MLL1 and Ash2L (Figures 5B and S30). Conformational changes observed for Ash2L are in the proximity of its RbBP5/MLL1 interface (Figure S33), which further strengthens the notion of MLL1 release. Pull-down assays clearly indicated MLL1 vacating the complex in the presence of cis-CS-VIP 8 (Figure S35). Although cis-CS-VIP 8-dependent H/D-exchange could not be detected at the Win binding site of WDR5, we postulate that this is due to opposing H/D-exchange differences induced by cis-CS-VIP 8 binding (Figure 5A) and Win peptide release, roughly canceling each other. Notably, cis-CS-VIP 8 induced elevated HDX in WDR5 residues 220–230, which are remote from the ligand binding site (Figures 5B and S31) and coincide with RbBP5-binding interface (PDB: 3PF4).

**Figure 4.** Dose–response toxicity of CS-VIPs on leukemia MOLM-13 cells. (A) Cell viability with both CS-VIP 8 isomeric states for 2-day incubation; \( \chi \): corresponding concentration. (B) IC\(_{50}\) determination of CS-VIP 8 and CS-VIP 14 in different isomeric states. All mean data points and standard deviations are derived from two independent experiments, and each concentration is in triplicate; n.c. not calculable.
in SASA were calculated for hydrogens from peptide bonds of WDR5 alone vs WDR5•cis-CS-VIP 8 (Figures S42 and SN2).

Most importantly, conformational analysis of MD trajectories of the MLL1 complex either alone or complexed to cis-CS-VIP 8 substantiated the partial unbinding of RbBP5 from WDR5 residues 220–230 during the course of MD simulation. We found that allosteric effects of cis-CS-VIP 8 caused different dynamical behavior, for the MLL1 polypeptide, which resulted in partial unbinding of RbBP5 (Figures S41 and 43 and SN4). However, we conclude that complete dissociation of WDR5/RbBP5 is unlikely because no difference in deuterium uptake was observed at the second binding site of WDR5 to RbBP5 (residues 240–250, 289) upon cis-CS-VIP 8 addition. As WDR5, RbBP5, and Ash2L still interacted in pull-down assays (Figure S35), this analysis suggests an altered structure of the remaining MLL1-devoid complex with cis-CS-VIP 8.

**Allosteric Communication through WDR5.** Further WDR5 analysis using perturbation-response scanning (PRS) revealed that “sensor regions”, i.e., protein sites that are conformationally changed upon perturbation of effector residues, are positioned along the WDR5/RbBP5 interface, and effectors around its MLL1 binding site as in the central pore of WDR5 (Figure S43 and SN3). Thereby, we imply a direct allosteric communication between MLL1 and RbBP5 binding sites of residue interaction network identified through network analysis. (E) Model of CS-VIP 8 mode of action. Gray boxes highlight altered interaction interfaces.
by MD simulations, communication pathways between MLL1 and RbBP5 were studied. Betweenness centrality (BC) analysis of obtained networks (Figures 5C,D and S47 and SN4) revealed that WDR5 residues D92 and K52 at the MLL1 binding site, as well as RbBP5 binding residue K221, are contributors to the overall information flow along all derived networks (Figures 5C,D). This allosteric communication with the RbBP5 binding site through the central pore of WDR5 was recognized as the main cause of partial RbBP5 unbinding. Additionally, residues around Y191 were also identified to contribute for RbBP5 stabilization (Figure 5D, SN4, and Table S7), which coincides with increase of D-uptake of WDR5 residues 195–205 upon cis-CS-VIP 8 binding, and the reported mutational analysis where Y191F was identified as a cause of low MLL1 activity.34 All together, the activity loss with cis-CS-VIP 8 apparently destabilizes the closed SET1 conformation, which uncovers the action mode of WDR5-binding inhibitors within the MLL1 complex (Figure S5E).

**In Vivo Optochemical Control of Hematopoiesis.** Encouraged by the on-target functional window for CS-VIP 8 isomers and the severe disruption of MLL1 complex, we evaluated its potential for in vivo photomodulation of hematopoiesis. Genetic programs of hematopoiesis are highly conserved across vertebrates; therefore, we considered zebrafish as the best model because it is not only optically transparent but also encodes most of the functional domains of human MLL1 gene products. Furthermore, MLL1 depletion in zebrafish causes severe defects related to hematopoiesis, such as lack of blood flow.35 These phenotypes were detectable from day 2 onward by using morpholino (MO) mediated gene knockdown.35 Indeed, it was between the third and fourth days postfertilization (dpf) when Wan et al.35 observed the maximum reduced blood flow (78%) without lethality. This fact dictated our choice of larvae stage. Thus, once CS-VIP 8 isomers reached their corresponding PSS in pure DMSO, each was diluted with egg water to a final concentration of 500 μM (1.5% DMSO). These solutions were directly added to 3-dpf larvae, which were previously treated with 1-phenyl-2-thiourea (PTU) to improve optical transparency and incubated for 18 h in darkness at 28.5 °C. Strikingly, only the cis-CS-VIP 8 as obtained from the 520 nm PSS caused a complete lack of responsivity upon tail touch in the larvae, compared with untreated fish. Furthermore, under these experimental conditions, abnormal developmental phenotypes like curved body axis and heart edema were observed (Figure 6A,B). Remarkably, larvae incubated with either 1.5% DMSO or CS-VIP 8 from 405 nm PSS (mainly the trans state) behaved as untreated ones, i.e., they were all alive with fast response to touch and without any apparent abnormalities. To quantify and determine whether these observed physiological changes were related to hematopoiesis, we performed three independent experiments, in which every condition was in duplicate containing five larvae per well (sample size, N = 30) and stained larvae with o-dianisidine for hemoglobin detection.36 The staining pattern remained unaffected for the vehicle, i.e., 1.5% DMSO as well as for 97% of larvae incubated with CS-VIP 8 at the 405 nm PSS. In contrast, staining with cis-CS-VIP 8 was highly diminished (60%) or even absent (40%, Figure 6A,B). These reduced-blood flow and abnormal phenotypes coincide with antisense mll1 knockdowns.35 In addition, to provide greater confidence that the observed phenotypes are, indeed, due to Mll1 disruption, we conducted gene expression analysis using reverse transcription quantitative real-time PCR (RT-qPCR). It has been previously reported that MLL1 primarily regulates the expression of the genes HOX7–11 (S′ in the HOX cluster), while expression of genes HOX1–6 (3′ in the HOX cluster) is relatively unaffected by MLL1 activity.7,38 Consequently, we evaluated the expression of both hoxa9 and hoxb5 genes (S′ and 3′ genes in the hox cluster, respectively) after the incubation of zebrafish embryos with CS-VIP 8. In line with our hypothesis, only the treatment with cis-CS-VIP 8 led to a significant downregulation of expression of hoxa9.
paralogs (hoxa9a and hoxa9b) while the expression of hoxb5 remained unaltered (Figure S24). Notably, the embryos incubated with the trans isomer or the vehicle did not display these changes in the gene expression. Together with our structural data, which display specific MLL1 complex disruption, these functional data strongly support that only the cis-CS-VIP 8 can affect hematopoiesis in vivo.

Finally, to release the full potential of our photoresponsive probes, we explored the possibility to externally control the in vivo inhibition of hematopoiesis via on-demand photo-activation. Thus, larvae incubated with the innocuous trans-CS-VIP 8 for 1 h were irradiated in situ at 520 nm for 30 s. After 18 h incubation, we detected effects comparable to the positive control, where larvae were exposed to cis-CS-VIP 8. Precisely, only 3% of larvae displayed an intact o-dianisidine staining pattern while in the remainder labeling was either partially disrupted (30%) or completely affected (67%). Negative controls of untreated fish or vehicle (1.5% DMSO) verified the harmlessness of visible-light irradiation (Figure S23). Taken together, we first developed a light-controlled enzyme modulator capable of achieving in vivo optochemical control of hematopoiesis, to our knowledge.

Concluding Remarks. Our understanding of gene regulatory mechanisms at the molecular level is often limited due to the lack of in vivo structure-function studies that can cope with the redundancy and ubiquitous expression of chromatin-modifying complexes. In this context, our work provides a successful proof-of-concept of in vivo photopharmacology to gain insight into the hematopoietic function of the MLL1 complex. We introduced the family of conformationally strained visible-light photoswitches (CS-VIPs) that can not only tightly interact with WDR5 but also light-control in vivo hematopoiesis in zebrafish without additional genetic modification. Our molecular studies provide a detailed view on the action of MLL1–WDR5 inhibitors in the context of the intact MLL1 complex. Upon CS-VIP 8 binding to WDR5, MLL1 dissociates from the core-complex, while the remaining complex subunits undergo conformational changes, e.g., by rewiring residues interaction networks in the WDR5–RbBP5 binding interface. Given the advent of time-resolved crystallography techniques like serial synchrotron crystallography (SSC)59 our WDR5•CS-VIP 8 cocrystals represent a highly promising model system for studying the cis → trans conversion of cyclic azopeptides in crystallo with their concomitant off/on diffusion from the protein target.

Our in vivo 3-dpf larvae zebrafish experiments displayed abnormal developmental traits such as hematopoiesis staining deficits, complete lack of responsibility, curved body axis, heart edema, and death preferentially upon cis-CS-VIP 8 addition. These phenotypic changes together with our RT-qPCR, which demonstrated robust reduction of hoxa9a/b gene expression while hoxb5a gene one remained unaffected at the same development stage, link the in vivo effects of cis-CS-VIP 8 with the specific inhibition of MLL1 activity. These observations suggest that the toxicity in the leukemia-related cells may be MLL1-dependent as well. However, to fully understand these complex biological mechanisms further in-depth investigations will be conducted where reporter genes assays as well as earlier stages of development will be studied. In our opinion this work marks a first step into the uncharted territory of hematopoiesis photopharmacology as well as in vivo light-controlled epigenetics. Furthermore, although the proof-of-concept application of our CS-VIPs was to unravel MLL1-related hematopoiesis, we expect our compounds to find further applications in diverse areas where in situ temporal control of MLL1 architecture is required or in which, β-propeller proteins, such as WDR5, act as interactions hubs.47 Our methodology goes beyond the traditional reductionist mindset for dissecting biological systems, providing new tools for analyzing the assembly of complex biological machineries. Finally, both the implication of epigenetics in the development of hematopoietic malignancies and the appearance of epigenetic modifiers in clinical trials provides a perfect timeline for exploiting the modulation of hematopoiesis via epigenetic networks.
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Notes

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