Abstract: DNA-encoded combinatorial synthesis provides efficient and dense coverage of chemical space around privileged molecular structures. The indole side chain of tryptophan plays a prominent role in key, or “hot spot”, regions of protein–protein interactions. A DNA-encoded combinatorial peptoid library was designed based on the Ugi four-component reaction by employing tryptophan-mimetic indole side chains to probe the surface of target proteins. Several peptoids were synthesized on a chemically stable hexathymidine adapter oligonucleotide “hexT”, encoded by DNA sequences, and substituted by azide-alkyne cycloaddition to yield a library of 8112 molecules. Selection experiments for the tumor-relevant proteins MDM2 and TEAD4 yielded MDM2 binders and a novel class of TEAD–YAP interaction inhibitors that perturbed the expression of a gene under the control of these Hippo pathway effectors.

The development of small molecules that inhibit protein–protein interactions (PPIs) often suffers from a lack of starting points for compound design, even though many target proteins contain small-molecule binding sites.[1] PPI inhibitors harbor vast potential to understand biological systems and for drug development.[2] For instance, PPIs such as the MDM2–p53 and TEAD–YAP interactions are involved in malignant diseases.[3] Dysregulated PPIs of transcriptional enhancer factor-1 domains (TEAD1-4) with co-transcription factor YAP (Yes-associated protein), late Hippo signaling effectors, are involved in important oncogenic mechanisms.[4–9] Inhibition of the TEAD–YAP PPI has been achieved in vitro with peptides that addressed “interface 3” (Figure 1a).[6,7] In silico small-molecule screening yielded TEAD–YAP inhibitors 1 and 2 (Figure 1b).[8] Intriguingly, TEAD is palmitoylated in a cavity, called a “central pocket”, which contributes to protein stability.[9] Compounds that bound to this pocket such as niflumic acid (4), flufenamic acid (5), and TED-347 (6) displaced a YAP-derived peptide from hTEAD4, while quinolinol 7 augmented YAP–TEAD activity (Figure 1c).[10] DNA-encoded libraries (DELs) have delivered a few PPI inhibitors.[11,12] They enable deep sampling of chemical space around key or “anchor” motifs. Here, we designed a DNA-encoded peptidomimetic library focused on the tryptophan side-chain motif—the indole moiety (Figure 1d). Tryptophan plays a prominent role in key, or “hot spot”, regions of protein–protein interactions. A DNA-encoded combinatorial synthesis provides efficient and dense coverage of chemical space around privileged molecular structures. The indole side chain of tryptophan plays a prominent role in key, or “hot spot”, regions of protein–protein interactions. A DNA-encoded combinatorial peptoid library was designed based on the Ugi four-component reaction by employing tryptophan-mimetic indole side chains to probe the surface of target proteins. Several peptoids were synthesized on a chemically stable hexathymidine adapter oligonucleotide “hexT”, encoded by DNA sequences, and substituted by azide-alkyne cycloaddition to yield a library of 8112 molecules. Selection experiments for the tumor-relevant proteins MDM2 and TEAD4 yielded MDM2 binders and a novel class of TEAD–YAP interaction inhibitors that perturbed the expression of a gene under the control of these Hippo pathway effectors.

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Thus, it has been exploited as an “anchor motif” for the design of PPI inhibitors.\textsuperscript{14,15} In this strategy, the anchor analogue—a substructure that is a chemical mimic of a specific amino acid residue (here tryptophan)—will be used to provide focused libraries with an increased probability of bioactivity. The anchor motif strategy has been repeatedly used to discover potent PPI inhibitors.\textsuperscript{16} Our DNA-encoded peptidomimetic library was selected on the p53-binding domain of MDM2 as the archetypal target for indole-based peptidomimetics.\textsuperscript{14} As a second promising target for this library we selected the YAP-interacting domain of (human) hTEAD4, because it contains a key tryptophan-binding site in “interface 3”, and its central pocket has been demonstrated to accommodate heteroaromatic structures (Figure 1 a).\textsuperscript{10,16}

We elected the Ugi four-component reaction for the initial step in the design of an encoded library because it combines selectable linker moieties to the DNA, diversity elements, and handles for library expansion into a peptoid backbone (Figure 1 d). DNA-encoded library synthesis was initiated by a Ugi reaction (U-4CR, Figure 2 a) on the chemically stable, solid-phase-coupled hexathymidine adapter “hexT”, with a broad range of reaction conditions tolerated (Figures 2 b and S1).\textsuperscript{17} This strategy allows for the synthesis of target molecules from bulk hexT-coupled starting materials in parallel, and it is more efficient than coupling individual multicomponent reaction products to DNA codes; all hexT products were isolated, thus providing fidelity. Carboxylic acid hexT1 and indole-carboxaldehydes hexT2 and hexT3 were reacted either with an alkyne-substituted amine (hexT1, hexT2, and hexT3) or an alkyne-substituted carboxylic acid (hexT2) that served as handles for the library expansion step. Three tryptophan-mimicking indole carbaldehydes and one tyrosine-mimicking p-hydroxybenzaldehyde placed the anchor motif distal from the DNA. A set of 18 isocyanides...
were used as diversity elements R¹ in the Ugi reaction (Figure 2b, Tables S1 and S2). Reaction optimization efforts identified temperatures of 80°C and high concentrations of isocyanides as requisite for peptoid synthesis. We synthesized in total 78 hexT peptoids, which were then ligated in one pot to peptoid backbone- and isocyanide-encoding DNA barcodes (Figures 2c and S4).¹⁷ Pooling, splitting, a second barcode ligation step, and copper(I)-promoted alkyne-azide cycloaddition with 104 azides,¹⁸ —produced in situ from the corresponding halides beforehand— finalized the 8112-membered “tiDEL” (thymidine-initiated DEL; Figure S2 and Table S3). The library was selected against streptavidin for library validation (Figures 3b and S7), against the p53-binding domain of MDM2, and against the YAP-interacting domain of human TEAD4 (hTEAD4). The sequencing data were progressed by the in-house-programmed algorithm ECEC (encoded compound enrichment calculator) based on R (Figures 3a, S5, and S6). Two-dimensional plots visualized enrichment factors of selection experiments versus bead-only control selections to facilitate compound identification. For the MDM2 target, the most highly enriched peptoids contained 6-chloroindole derivatives irrespective of their positioning on the backbone (Figure 3c). Interestingly, a single building block from the 104 diverse azides was enriched, the 2,4-dimethylphenylacetamide A⁴⁸. Compound ⁸ was selected based on enrichment factor calculations. It showed a plausible in silico binding mode, and MST experiments confirmed binding to MDM2, thus validating the library design concept (Figures 4a,b and S9).¹⁵ Selection experiments for hTEAD4 identified peptoid hexT21-A⁵⁶ as the most enriched compound (Figures 3d and S8). In this peptoid, a 6-chloroindole was flanked by a C-terminal hydrocarbon and a triazole-linked imidazopyridine. We synthesized a small series of compounds inspired by hexT21-A⁵⁶/⁹ and investigated their binding to depalmitoylated hTEAD4 by nanodifferential scanning fluorimetry (Figures 4c,d and S10). Stabilization of hTEAD4 was observed for peptoids ⁹—¹¹, which differed in their C-terminal alkyl amides, with a tert-butyl amide leading to the highest ΔTm value. Exchanging the succinate linker by acetamide ¹⁴ reduced the ΔTm value, thereby suggesting that the linker was involved in protein binding.¹⁹ The imidazopyridine substituent could be exchanged by 5-phenyl-oxazole-2-yl (to give ¹²), as suggested by the enrichment plot (Figure S8). We next studied the biological consequences of the compound–hTEAD4 interaction. Compound ⁹ inhibited the palmitic acid–hTEAD4 interaction with an IC₅₀ value of 0.41 μM, while compound ¹⁰ showed a much weaker inhibition, which suggests a different binding mode (Figure 4e). Both compounds were then evaluated for inhibition of the YAP–hTEAD4 interaction (YAP ⁵₀–₁₀₀, Figure 4f). They inhibited the PPI with IC₅₀ values of 6.75 μM (⁹) and 5.65 μM (¹⁰).

Finally, we tested the cellular activity of compound ⁹ by measuring transcript levels of CTGF, a gene under control of the hippo pathway effectors TEAD–YAP (Figure 4g). HEK293 cells were treated with compound ⁹ alone, and with a combination of compound ⁹ and the hippo signaling inhibitor XMU-MP-1. XMU-MP-1 blocks MST1/2 kinases which are upstream components in the Hippo pathway. This inhibition results in inactivation of downstream kinases LATS1/2, and subsequent translocation of YAP into the nucleus, where it forms a transcriptional complex with TEAD, thereby leading to gene expression. The addition of compound ⁹ to HEK293 cells did not alter the CTGF transcript levels, whereas it caused significant reduction in gene expression after inhibition of Hippo signaling by XMU-MP-1. This observation was in line with an on-target mechanism and suggested a potential implication for treating tumors driven by abnormal Hippo pathway signaling.

Initiating encoded library synthesis with an Ugi multicomponent reaction step that turned simple starting materials...
into peptoid side chains provided flexibility in the library design around privileged “anchor” motifs such as tryptophan mimics. This library design uncovered chemical aspects of challenging target proteins from relatively few encoded compounds. Currently, we are elucidating the binding mode of compounds \textit{9} and \textit{10}, and we are synthesizing analogues to better understand the structure–activity relationships of these TEAD–YAP inhibitors.

\section*{Acknowledgements}

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\section*{Conflict of interest}

The authors declare no conflict of interest.

\textbf{Keywords:} combinatorial chemistry · DNA-encoded library · peptidomimetics · protein–protein interaction inhibition · Ugi reaction

\begin{figure}
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\includegraphics[width=\textwidth]{figure4}
\caption{Validation of MDM2 and hTEAD4 binders. a) Docking of \textit{8} into the MDM2–p53 interaction site. b) Effect of the off-DNA-synthesized \textit{8} bound to MDM2. c) Chemical structures of biologically evaluated compounds \textit{9}, \textit{12}–\textit{15}. d) Validation of compound binding to hTEAD4 by nanoDSF. e) Inhibition of palmitic acid binding to the hTEAD4 central pocket measured by fluorescence polarization. f) Inhibition of YAP binding to hTEAD4 measured by fluorescence polarization. g) Evaluation of the cellular activity of \textit{9} by measurement of CTGF transcript expression levels.}
\end{figure}