Specific inhibition of the Survivin-CRM1 interaction by peptide-modified molecular tweezers

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Survivin’s dual function as apoptosis inhibitor and regulator of cell proliferation is mediated via its interaction with the export receptor CRM1. This protein–protein interaction represents an attractive target in cancer research and therapy. Here, we report a sophisticated strategy addressing Survivin’s nuclear export signal (NES), the binding site of CRM1, with advanced supramolecular tweezers for lysine and arginine. These were covalently connected to small peptides resembling the natural, self-complementary dimer interface which largely overlaps with the NES. Several biochemical methods demonstrated sequence-selective NES recognition and interference with the critical receptor interaction. These data were strongly supported by molecular dynamics simulations and multiscale computational studies. Rational design of lysine tweezers equipped with a peptidic recognition element thus allowed to address a previously unapproachable protein surface area. As an experimental proof-of-principle for specific transport signal interference, this concept should be transferable to any protein epitope with a flanking well-accessible lysine.

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molecular recognition since such interfaces are usually large (>1000 Å²), well solvated, and display a rugged topology. Thus, until today interfering molecules have in most cases been identified by extensive library screening.

Supramolecular chemistry provides orthogonal artificial elements for protein recognition and, in combination with computational modeling, allows a deeper understanding of the underlying non-covalent interactions. Inter alia, calixarenes, cucurbiturils, molecular tweezers, and GCP motifs recognize well-solvated amino acid residues on protein surfaces and have been successfully used to target protein surfaces and to interfere with PPIs. On the same protein, i.e., ubiquitin, a recent comparative study revealed that these host molecules occupy different areas and seem to exhibit complementary recognition profiles.

Despite these promising features and first applications in cells and animals, the selective recognition of protein elements by supramolecular host systems remains highly challenging. A few recent examples include synthetic ligands for peptide motifs on proteins and for a specific protein context. We here present an advanced approach by combination of a supramolecular host molecule with a well-defined biomolecular interaction. This ditopic hybrid allows us to complex a single critical amino acid motif together with its direct environment on the protein surface, which leads to powerful competition with the natural binding partner. Specifically, we generate a covalent conjugate between a lysine-selective molecular tweezer and a self-complementary peptide and target a critical interface important for the survival of cancer cells, i.e., the nuclear export signal (NES), located on an ordered but somewhat dynamic loop on the Survivin surface.

Survivin is mostly absent in normal resting adult tissues, but highly upregulated in almost all cancer types. Its overexpression is associated with resistance against chemotherapeutic drugs and radiotherapy, frequent recurrences, and a decreased patient survival. Despite its small size (142 aa, 16.5 Da) and its lack of enzymatic activity, Survivin is fulfilling a well-characterized dual role within the cell. As the smallest member of the inhibitor of apoptosis protein family, Survivin on the one hand plays a role in countering programmed cell death. As part of the chromosomal passenger complex (CPC), Survivin is on the other hand crucial for mitotic regulation promoting cell proliferation. For both functions, an interaction with the nuclear export receptor CRM1 mediated by Survivin’s highly conserved, leucine-rich NES is pivotal. Thus, interference with the Survivin–CRM1 interaction can inhibit cancer cell proliferation.

The development of a small molecule, which specifically binds to the NES on Survivin’s surface, would represent a valuable approach to inhibit the Survivin–CRM1 interface. This is a challenge for supramolecular chemistry: we can direct an amino acid-selective host molecule to the functionally relevant epitope on Survivin by combining it with a recognition unit for the NES? This key region is located on an ordered loop and flanked by well-accessible basic amino acids, K90/91 and K103/R106. We therefore designed supramolecular tweezers that only target lysine (Lys) and arginine (Arg) residues and that are equipped with a compact binder to the natural dimer interface overlapping with the NES.

Molecular tweezers possess a torus-shaped arrangement of alternating benzene and norbornadiene rings, which form an electron-rich unipolar cavity—ideally suited to pull the cationic side chains of Lys and Arg inside. Two (hydrogen)phosphate groups lock the included side chain in an ion pair. This unique binding mechanism operates well under physiological conditions and has already been exploited for protease inhibition, prevention of protein aggregation, and modulation of PPIs on shallow grooves. In order to turn these molecular tools into specific Survivin ligands, it was necessary to identify a binding motif for the NES region and to develop a synthesis for efficient tweezers.
Subsequently, peptide and monobutynyl tweezer were coupled in a THF/water solvent mixture with ascorbic acid and CuSO_4 \cdot 5H_2O. Both click reactions proceeded smoothly and gave the coupling products in good yields and excellent purity after HPLC purification (SI2). In addition, FAM-labels were introduced into the peptide-modified tweezers via a C-terminal (Fluorescein-labeled lysine)-glycine dipeptide fragment (SI3). Reaction monitoring was facilitated by the appearance of additional ^1H NMR signals at 5.2 ppm (CH_2-triazole) and 8.0 ppm (CH_arom) indicative of triazole formation. The final peptide tweezers displayed very good water solubility, because they carry multiple negative charges both in the tweezer and in the peptidic part. No self-inclusion of the tweezer moieties was observed in the form of potential upfield shifts in the ^1H NMR spectra, also ruling out the formation of unproductive tweezer dimers (SI2).

**Characterization of binding by isothermal titration calorimetry.** The interaction between the tweezers and Survivin was studied by isothermal titration calorimetry (ITC) and NMR spectroscopy. So far, the only published NMR structure originated from a truncated Survivin (aa 1-120) with improved solubility^38. Indeed, MD simulations of full-length Survivin revealed a highly flexible C-terminal α-helix fragment around residue 120 (SI4), which might explain poor expression of the full-length protein. Since the truncated construct still contains all relevant parts of the protein and gives excellent NMR spectra (SI5), we chose it for all our in vitro experiments and denote it as Survivin120 in the following results.

ITC titration of Survivin120 to the tweezers resulted in exothermic binding isotherms (Fig. 3) with dissociation constants KD of 38 µM for the unmodified tweezer TW, 24 µM for TW-ELTL as well as a KD of 19 µM for TW-ELTLGEFL. These values are in good agreement with the unmodified tweezer TW (formerly CLR01) affinity toward lysine (K_D 17 µM)^39. Fitting the ITC titrations with Survivin120 to a one set of sites model allowed us to derive all relevant thermodynamic data including stoichiometries and changes in Gibbs free energy (G), enthalpy (H), and entropy (S) as summarized in a supplementary table (SI7). Importantly, the unmodified tweezer produced a

**Fig. 1 Design of peptide-modified supramolecular tweezers.** a Representation of Survivin’s dimer interface based on PDB-ID: 1XOX (https://www.rcsb.org/structure/1xox)^38. Both monomers, depicted in blue and gray, mainly interact via the ELTL sequence (contact region of both monomers overlapping with the NES, represented in cyan and green). This sequence was chosen as second binding motif for the peptide-modified tweezer molecules. b Representation of TW-ELTL (shown in d) bound to Survivin. TW-ELTL (yellow) binds the anchor lysine residue K103 (violet) on Survivin’s surface while the peptide motif ELTL (yellow) interacts with the ELTL region of the Survivin monomer (cyan). This is the same region of the dimer interface represented in Fig. 1a, overlapping with the NES (cyan). The chemical structures of the unmodified tweezer molecule TW (c), an asymmetrical tweezer molecule linked to the short peptide ELTL (TW-ELTL) (d), and an asymmetrical tweezer molecule linked to the elongated peptide ELTLGEFL (TW-ELTLGEFL) (e) are depicted.
20:1 stoichiometry (correlating well with 16 accessible lysine and arginine residues in Survivin120), whereas the modified tweezers displayed 2:1 ligand/protein ratios. This stoichiometry might be plausible with regard to the antiparallel orientation of the Survivin monomers in the respective homodimer, probably allowing the conjugated peptides to also align in both directions with the tweezer either binding to one or the other neighboring anchor lysine. In addition, we carried out reverse titrations by adding increasing amounts of unmodified tweezer TW vs. TW-ELTL into a constant concentration of Survivin (SI8). Interestingly, a sharp kink was produced at a 4–5-fold tweezers excess in both cases, indicating that 4–5 tweezer molecules can be accommodated on the protein surface. However, further addition of unmodified tweezers produced a second substantial exothermic titration step, indicating further unspecific binding to accessible lysine and arginine residues, while the peptide tweezer had already almost reached saturation. Due to the biphasic character these binding curves do not allow the determination of binding constants. However, our data indicate that the introduction of a peptide mimicking Survivin’s natural dimer interface enhanced the tweezer affinity toward Survivin only slightly, but greatly increased its regioselectivity. Although the dimer stability of Survivin120 is not known, it likely represents the binding partner for all tweezers, because apart from the NES region V89-L102 some remote residues P4-W10 contribute additional, mostly aromatic, interactions. Even in the dimer, however, the self-complementary peptide loops (V89-L102) are dynamic and can be expected to expose the NES region temporarily (see discussion of the simulations below).

**Mapping tweezer binding sites by NMR.** To map tweezer binding to distinct amino acid residues of Survivin, we performed NMR titrations adding tweezers up to equimolar amounts to 15N-labeled Survivin120 (SI5). Binding of a ligand causes signal shifts and often reduced signal intensities for the residues involved in the interaction (Fig. 4). Titration of the unmodified tweezer resulted in reduced signal intensities and shifts around the basic amino acids K91, K103, and R106 (Fig. 4a, b). K90 also lies in the regions identified; however, it is not assigned in the spectrum. In addition, signal intensities decrease in the same regions (Fig. 4d) confirming these as potential tweezer binding sites. Unfortunately, the applied NMR method does not allow us to differentiate between tweezer binding to residues K90 vs. K91 and K103 vs. R106 since they are in too close proximity. Titration of TW-ELTL into a constant concentration of Survivin (SI8). Interestingly, a sharp kink was produced at a 4–5-fold tweezers excess in both cases, indicating that 4–5 tweezer molecules can be accommodated on the protein surface. However, further addition of unmodified tweezers produced a second substantial exothermic titration step, indicating further unspecific binding to accessible lysine and arginine residues, while the peptide tweezer had already almost reached saturation. Due to the biphasic character these binding curves do not allow the determination of binding constants. However, our data indicate that the introduction of a peptide mimicking Survivin’s natural dimer interface enhanced the tweezer affinity toward Survivin only slightly, but greatly increased its regioselectivity. Although the dimer stability of Survivin120 is not known, it likely represents the binding partner for all tweezers, because apart from the NES region V89-L102 some remote residues P4-W10 contribute additional, mostly aromatic, interactions. Even in the dimer, however, the self-complementary peptide loops (V89-L102) are dynamic and can be expected to expose the NES region temporarily (see discussion of the simulations below).

**Fig. 2 Synthetic strategy leading to monofunctionalized molecular tweezers.** a Introduction of one butynyl phosphate ester arm on the parent diacetoxy tweezer (TCA coupling) followed by click chemistry with N-terminal azidopeptide; b additional introduction of a fluorescence label by adding a C-terminal LysFAM-Gly sequence to the clicked peptide. Blue: phosphate; red: alkyne/triazole; green: azide/triazole; olive: FAM label. DIPEA N,N-diisopropylethylamine, THF tetrahydrofuran.
conjugate on Survivin’s surface. The overall line shape of the HSQC spectra indicates that Survivin120 likely remains dimeric upon tweezer binding. If the dimer would dissociate upon ligand binding, a sharpening of the signals and thus an intensity increase would occur due to slower T2 relaxation. Rather the contrary is observed, especially at ligand:protein ratios >1:1, which indicates beginning aggregation. Nevertheless, anchoring the peptide–tweezers nearby the NES with direct peptide–NES interactions is expected to shield the natural binding site and significantly weaken the Survivin–CRM1 interaction.

For a better understanding of the complexation process, we performed MD and Gaussian accelerated molecular dynamics (GaMD) simulations as well as quantum mechanics/molecular mechanics (QM/MM) calculations (see SI10 for computational details). TW–lysine interactions were calculated in the Survivin120 monomer (protomer A, structure with PDB-ID 1XOX [https://www.rcsb.org/structure/1xox])38. The monomer, unlike the dimer structure, displays an exposed NES region, which is a key feature for the activity of Survivin. Thus, it represents the most suitable model to study the interactions of the tweezers with the lysine residues in this region. Four well-accessible lysine residues (K23, K90, K91, and K103) were selected on the lysine residues in this region. Four well-accessible lysine residues (K23, K90, K91, and K103) were selected on the monomer of Survivin120 and a modified tweezer with the elongated peptide, TW-ELTLGEFL, on K103. Not surprisingly, the GaMD simulations also showed that TW-ELTLGEFL is able to form more noncovalent interactions with the NES region than the shorter TW-ELTL. This difference becomes most apparent in the ELTL region of the dimer interface (involving two hydrogen bonds) has a very low prevalence along the simulation. In the most frequent scenario, no hydrogen bonds are formed, indicating weak and labile interactions between the interface fragments (SI14). Therefore, we also performed GaMD simulations of the monomer of Survivin120 and a modified tweezer with the elongated peptide, TW-ELTLGEFL, on K103. At this point, it is important to notice that the above-discussed structural details are in very good agreement with the shift and intensity changes observed by NMR experiments, with one exception: they require an exposed NES region such as in monomeric Survivin. This apparent discrepancy can be rationalized by the conformational flexibility at the weakly associated dimeric NES region. We therefore also performed GaMD simulations of the Survivin120 dimer in an explicit solvent box. The frequency of hydrogen bond formation during the simulation within the dimer interface was analyzed (Fig. 5a, b). The results show that the strongest interaction at the 95ELTL98-region of the dimer interface (involving two hydrogen bonds) has a very low prevalence along the simulation. In the most frequent scenario, no hydrogen bonds are formed, indicating weak and labile interactions between the interface fragments in both monomers. Not surprisingly, the GaMD simulations also showed that TW-ELTLGEFL is able to form more noncovalent interactions with the NES region than the shorter TW-ELTL. This difference becomes most apparent in the ELTL region of the dimer (95ELTL98) (Fig. 5c). The peptide substituents of the modified tweezers are rather flexible, allowing for frequent interactions with the NES. It seems that the rigid triazole linker acts as an anchor point and facilitates these interactions (SI15)—a synergistic effect between the peptide motif and the (otherwise inactive) linker fragment. As expected, the longer, more flexible
peptide chain of TW-ELTLGEFL explores a larger conformational space and establishes more attractive interactions with the NES, explaining its superior performance in binding experiments. Our calculations show weak and dynamic interactions at the dimer interface. Such features might be leveraged by the TW-anchored peptide to form contacts with one of the Survivin protomers and, hence, shield the corresponding NES region. Without further structural information we cannot exclude that the peptide might also bind in a way that stabilizes the dimer interface and thus prevents CRM1 from binding a Survivin monomer.

**Table 1** The relative energies of the QM region indicate that the tweezer–lysine complexes are most stable in positions 103 and 91.

| TW-ELTL | Relative energy QM region (kcal/mol) |
|--------|-------------------------------------|
| K103   | 0 ± 1                                |
| K91    | 1 ± 14                               |
| K90    | 27 ± 6                               |
| K23    | 68 ± 3                               |

**Tweezer interference with the export complex assembly in vitro.** We next investigated whether tweezers inhibit the Survivin–CRM1 complex formation in vitro. The effects of the unmodified and peptide-linked tweezers were analyzed via pull-down experiments with 293T lysates containing overexpressed HA-tagged Survivin142, recombinant GST-CRM1 as bait protein, and tweezers (Fig. 6). The parent and ELTL-linked tweezer required 10–50 µM to disrupt the Survivin–CRM1 complex. TW-ELTLGEFL was already effective at 1–10 µM and thus turned out to be more potent. In order to provide experimental evidence for the sequence selectivity of the TW-ELTLGEFL ligand, a scrambled analog was synthesized and clicked to the tweezer, resulting in the hybrid molecule TW-LFEEGLLT (SI1, SI2, and SI10). Intriguingly, this ligand was about one order of magnitude less effective than the one bearing the original NES-derived sequence and rather comparable to the unconjugated tweezer TW with regard to the pull-down analyses (Fig. 6a and SI18). Moreover, ITC titrations with TW-LFEEGLLT revealed much lower heat changes, and the corresponding $K_D$ value dropped from 19 ± 3 for TW-ELTLGEFL to 68 ± 22 µM for the scrambled peptide conjugate (SI16 and S17). Thus, it displays significantly (~three times) lower affinity toward the wildtype protein, indicating that the correct self-complementary NES sequence is indeed essential.
for efficient ditopic recognition. Additional NMR studies demonstrated that the tweezer with the scrambled peptide sequence is still able to bind to the two sites K90/K91 and K103/R106 like the unmodified TW (SI17). However, no large perturbations as for TW-ELTL or TW-ELTLGEFL are observed in between residues 91–103, indicating that the peptide moiety does not form specific contacts with Survivin. Instead, the slight chemical shift perturbations might be due to the spatial proximity of the peptide moiety to the anchoring residue. All those results strongly indicate that the additional binding energy from the NES interface was lost due to scrambling and thus point toward specific recognition as the reason for increased selectivity and

Fig. 5 GaMD simulations and QM/MM calculations on the tweezers–Survivin interaction provide deeper insights into the binding event. a Main hydrogen bonds (HB, red) established between Survivin monomers (gray, violet) at the ELTL interface (contact region of both monomers overlapping with the NES, represented in cyan and green). The leucine residues engaged in these interactions are shown in CPK representation. b Occurrence of hydrogen bonds at the dimer interface. c Frequency of noncovalent contacts between the tweezers, bound to K103, and the NES as well as with the ELTL region (95ELTL98) of the dimer interface.

Fig. 6 The assembly of the export complex is disturbed by unmodified and peptide-linked tweezer molecules. a 293T cell lysate with overexpressed Survivin142-HA was preincubated with either unmodified tweezer (TW), TW-ELTL, TW-ELTLGEFL, or a scrambled peptide-modified tweezer, TW-LFEEGLLT, at concentrations ranging from 0.01 to 200 µM. GST-CRM1 was mixed with either non- or preincubated cell lysates in the presence of recombinant RanQ69L and dGTP to allow complex assembly. GST-CRM1 and interacting proteins were pulled by GSH-Sepharose beads. Proteins in input and beads samples were analyzed via immunoblotting with antibodies specific for GST or HA. For each tweezer, samples derive from the same experiment and gels/blots were processed in parallel. Direct comparison for this exact concentration range was performed once for TW, TW-ELTL, and TW-ELTLGEFL and for TW-LFEEGLLT in three technical replicates. b, c Atto488-labeled Survivin120 was preincubated with CRM1_1-1062VLV430AAA mutant in a ratio of 1:5 and titrated with supramolecular tweezers up to approx. 180 µM. Fluorescence anisotropy was measured (n = 1) (b), and IC50 values were determined from the resulting curves (c). TW light blue/ triangles, TW-ELTL blue/squares, TW-ELTLGEFL dark blue/circles. Source data are provided as a Source Data file.
interference with the Survivin–CRM1 interaction. Of note, very high tweezer concentrations (100–200 μM) also weakened GST-CRM1 binding to GSH-Sepharose beads, most likely by low-affinity binding to the GST protein. However, direct ITC titrations gave only small heat changes and revealed that this interaction is endothermic (S19).

We also used fluorescence anisotropy experiments to investigate the ability of the tweezer molecules to disrupt the Survivin–CRM1 complex. Atto488-labeled Survivin120 was preincubated with the CRM1_1–1062VLV430AAA mutant that binds Survivin irrespective of RanGTP and titrated with tweezers. The presence of all tweezers significantly lowered the fluorescence anisotropy, indicating a potent disruption of the Survivin–CRM1 complex (Fig. 6b, c). IC50 values were determined at 53 μM for the unmodified tweezer, 39 μM for TW-ELTL, and 12 μM for TW-ELTLGEFL. Thus, pull-down and fluorescence anisotropy experiments both indicated that the peptide modification increases the inhibitory potential of the tweezers for the Survivin–CRM1 interaction.

Confirmation of the tweezer binding site. If the tweezer-based inhibitors indeed bind to lysines/arginines flanking the NES (K90, K91, K103, R106), their mutation to, e.g., threonine, should inhibit the NES and thus shield it against CRM1. To the best of our knowledge, this is the first so far successful example that an amino acid binder (lysine tweezer) is directed to a specific epitope of a protein—in our case by conjugation to the self-complementary dimer interface comprising the NES sequence. The underlying rational design was supported by ITC titrations and NMR spectroscopy that produced maximum chemical shift perturbations on four lysine/arginine residues flanking the NES. Detailed MD and GaMD simulations complemented with QM/MM calculations revealed K103 as a preferred binding site and supported that even in the dynamic dimer the NES signal is partially exposed to approaching ligands. Pull-down experiments and fluorescence anisotropy titrations both indicated that the peptide modification indeed increases the inhibitory potential and specificity of the tweezers for the Survivin–CRM1 interaction. Importantly, a tweezer molecule equipped with a scrambled peptide motif was significantly less effective with regard to binding and inhibition of the relevant Survivin–CRM1 interaction. Again, this points toward a specific recognition as the origin for increased selectivity of our hybrid ligand. Finally, a double lysine mutant of Survivin (K90/103T) further substantiated the identity of our hybrid ligand, because here the potent tweezer conjugates lost most of their inhibitory potential (ITC, pull-down, FT). Our sophisticated synthetic approach allows the formation of labeled tweezer peptide conjugates for advanced binding experiments, which may also find applications in fluorescence imaging.

Even though the peptide modification increased the ability of the tweezers to shield the NES, it had only moderate impact on the binding affinity. Obviously, additional binding energy must be generated by tailored recognition units incorporated into the tweezer conjugates. A second tweezer unit at the opposed end of the peptide may serve this purpose, or alternatively more powerful supramolecular NES binders of synthetic or natural origin. In this study, we established that a supramolecular amino acid binder can be designed for an exposed surface epitope on a given protein target. Our strategy involves the combination of a lysine–selective tweezer with a peptide recognition element for the desired binding epitope. This was accomplished by trichloroacetonitrile-assisted monoesterification of a single tweezer phosphate with butynol and subsequent click reaction with an azide-modified peptide without the need of any protecting group. Attachment of a single peptide arm renders the hybrid tweezer selective for the peptide loop representing the NES in Survivin which is self-complementary and flanked by well-accessible lysine residues. The design was guided by MD and GaMD simulations as well as QM/MM calculations of the putative tweezer–protein complexes. Structural evidence was provided by 2D NMR proteins and is therefore not specific for Survivin. Here, we present prototypes for an alternative strategy: to address Survivin’s NES with specific supramolecular tweezer conjugates, which dock onto the overlapping natural dimer interface on Survivin’s protein surface with low micromolar affinities. This alternative is now accessible by click reactions from an alkyne-modified parent tweezer. We emphasize that this synthetic strategy greatly expands the design of modified tweezers, because it can be applied to tweezers with one or two phosphate arms and is not restricted to peptides. Virtually any additional functional unit can now be attached to the tweezers by click chemistry: fluorescence labels, chemically reactive groups, peptidic and other recognition units, Au nanoparticles (via C-terminal cysteines), and scaffolds with various alkyne for multivalency have already been introduced37,38. This functionalization synthetic strategy now opens up a pioneering class of advanced tweezer derivatives with two or more functions.

Moreover, as a proof-of-principle, our experiments confirm that binding of peptide-equipped tweezers occurs in Survivin’s NES region overlapping the dimer interface and therefore impairs the interaction with CRM1. To the best of our knowledge, this is the first so far successful example that an amino acid binder (lysine tweezer) is directed to a specific epitope of a protein—in our case by conjugation to the self-complementary dimer interface comprising the NES sequence. The underlying rational design was supported by ITC titrations and NMR spectroscopy that produced maximum chemical shift perturbations on four lysine/arginine residues flanking the NES. Detailed MD and GaMD simulations complemented with QM/MM calculations revealed K103 as a preferred binding site and supported that even in the dynamic dimer the NES signal is partially exposed to approaching ligands. Pull-down experiments and fluorescence anisotropy titrations both indicated that the peptide modification indeed increases the inhibitory potential and specificity of the tweezers for the Survivin–CRM1 interaction. Importantly, a tweezer molecule equipped with a scrambled peptide motif was significantly less effective with regard to binding and inhibition of the relevant Survivin–CRM1 interaction. Again, this points toward a specific recognition as the origin for increased selectivity of our hybrid ligand. Finally, a double lysine mutant of Survivin (K90/103T) further substantiated the identity of our hybrid ligand, because here the potent tweezer conjugates lost most of their inhibitory potential (ITC, pull-down, FT). Our sophisticated synthetic approach allows the formation of labeled tweezer peptide conjugates for advanced binding experiments, which may also find applications in fluorescence imaging.

Discussion Inhibition of the essential Survivin–CRM1 interaction is of great interest because it regulates cell proliferation and mediates a cyto-protective function27,30. However, the development of CRM1 binders bears the disadvantage that it affects a large number of cargo
spectroscopy, affinities were determined by ITC titrations. The hybrid tweezers were able to disrupt the essential complex between Survivin and its export receptor CRM1 in cell lysates as demonstrated with pull-down assays and in vitro as shown by fluorescence anisotropy measurements. Labeled tweezer hybrids revealed strongly diminished affinities to a Survivin double mutant that lacked the NES-flanking lysines and confirmed the selectivity for the respective epitope on Survivin. We thus accomplished the proof-of-principle of epitope targeting by supramolecular binders. Further optimization should improve the performance of our ligands by, e.g., by replacing the peptide unit resembling the natural dimer interface of Survivin with much more powerful interaction partners from the CPC, such as Boréalin fragments. Alternatively, we plan to employ dimeric tweezers with an internal peptide unit—in order to exploit two attachment sites to the NES region. Lysine 90 and 103 have been proven very well suited for this purpose and work into this direction is underway in our laboratory.

In the future, supramolecular inhibition of the CRM1–Survivin interaction should be transferred into the cellular context in order
Methods
principles in basic and applied biomedical research. Due to its excellent biocompatibility and very good tolerance toward peptide side chains, the copper-catalyzed Huisgen cyclodaddition was employed to couple an alkyne peptide with an azide-peptide. This to end the unsymmetrical monophosphate monobutynylphosphate tweezer was synthesized according to our recently published protocol for the synthesis of unsymmetrical diphosphate monoesters via the trichloroacetonitrile method. All peptides were prepared with a final coupling of azidoacetic acid to their N-terminal. Cleavage from the resin and purification by preparative HPLC yielded pure peptides. Subsequent click reactions between free azido peptides and alkyne tweezers were carried out in a mixture of water and THF (1:1). The catalyst was prepared in situ by reaction of copper sulfate with sodium ascorbate in the presence of DIPEA base. The resulting hybrid molecules were precipitated by acidification with HCl, followed by removal of THF in vacuo and filtration. The crude product was pulled CRM1 intensity, the latter is normalized by the GST-CRM1 interaction.

To generate Survivin point mutants, critical lysines were changed by site-directed mutagenesis with the Q5 Site-Directed Mutagenesis Kit from New England Biolabs. The eukaryotic expression vector pc3-Survivin142-HA was analogously constructed by PCR amplification using an appropriate template and primers containing BamHI/Nhel restriction sites (SI20). The PCR product was cloned into the vector pcdNA3.1 as a C-terminus fusion with an HA expression tag and transfected as described.

Plasmids. Bacterial expression vectors encoding Survivin120 variants, CRM1, and RanG69L were constructed by polymerase chain reaction (PCR) amplification using appropriate templates and primers containing Apal/BamHI restriction sites (SI20). PCR products were cloned into the vector pET41-GST-PreSc as an N-terminal fusion with GST and a PreScission protease cleavage site as described (SI20). To generate Survivin-ELTL conjugates, Survivin was changed by site-directed mutagenesis with the Q5 Site-Directed Mutagenesis Kit from New England Biolabs. The heat of dilution was subtracted as constant. For thermodynamic data derived from the graphs see SI7. FAM-labeled unmodified tweezer molecule or ELTL/ ELTLGEFL peptides w/o tweezer. GST-Survivin120- or GST-Survivin120-K90/103T-loaded beads were mixed with CRM1 and RanG69L prey proteins as well as dGTP to allow export complex assembly. Proteins in input and bead samples were analyzed via immunoblotting with antibodies specific for CRM1 or GST. WT, wildtype. One representative example of two independent biological replicates is shown. Samples derive from the same experiment and gels/blots were processed in parallel. Quantification of two independent pull-down experiments. After subtraction of the CRM1 negative control from the pulled CRM1 intensity, the latter is normalized by the GST-Survivin intensity and afterwards normalized by the CRM1 intensity without tweezer incubation.

Export complex assembly is only compromised by peptide tweezers in the wildtype Survivin120, but not in the mutant. No ligand: black; TW: light blue; TW-ELTL: blue; TW-ELTLGEFL: dark blue; ELTL peptide: light gray; ELTLGEFL peptide: dark gray. Source data are provided as a Source Data file.

Methods
Synthesis of tweezer conjugates. Due to its excellent biocompatibility and very good tolerance toward peptide side chains, the copper-catalyzed Huisgen cyclodaddition was employed to couple an alkyne peptide with an azide-peptide. This to end the unsymmetrical monophosphate monobutynylphosphate tweezer was synthesized according to our recently published protocol for the synthesis of unsymmetrical diphosphate monoesters via the trichloroacetonitrile method. All peptides were prepared with a final coupling of azidoacetic acid to their N-terminal. Cleavage from the resin and purification by preparative HPLC yielded pure peptides. Subsequent click reactions between free azido peptides and alkyne tweezers were carried out in a mixture of water and THF (1:1). The catalyst was prepared in situ by reaction of copper sulfate with sodium ascorbate in the presence of DIPEA base. The resulting hybrid molecules were precipitated by acidification with HCl, followed by removal of THF in vacuo and filtration. Unreacted starting materials could be separated from the products by RP-18 column chromatography or preparative HPLC.

Peptide synthesis. All peptides were synthesized using automated, microwave assisted, SPPS. The synthesis was carried out on a CEM peptide synthesizer using a Wang resin (4-hydroxybenzyl alcohol (PHB) on polystyrene) already equipped with the C-terminal amino acid. Coupling was effected with HCTU. In the final step, 2-azidoacetic acid was coupled to the free N-terminus of the peptide; then the entire peptide was cleaved off the resin with TFA, TIS, and water. The peptide in the cleavage solution was poured onto ice-cooled diethyl ether and stored in the freezer for 1 h to precipitate. Each peptide was pre-purified by centrifugation and washed again with diethyl ether. Subsequently, preparative purification was carried out by means of HPLC. Purification was performed on a preparative HPLC system from Jasco with a UV/Vis detector (UV-975, DG-2080-53 solvent degasser, UFLC-980-025 3-channel solvent mixer, peak-detection at 210 nm). The instrument is equipped with a reverse-phase column from Macherey-Nagel (Modell EC 250/4 Nucleosil 100-3 C18). Linear gradients of acetonitrile and water with presence of 0.1 TFA were applied.

Click coupling. Monophosphate monobutynylphosphate tweezer (5.0 mg, 6.4 μmol) was dissolved in 2 mL THF/H2O (1:1) in a 5-mL round-bottom flask together with the respective N-terminal Azac peptide (23 μmol). Fresh distilled DIPEA (11.3 μL) was added to the previously degassed solution. Subsequently, the copper sulfate solution (8.3 mg CuSO4·5H2O, 33 μmol in 1 mL water) was mixed with the sodium ascorbate solution (13 mg C6H8Na2O6, 66 μmol in 1 mL water) and the catalytic brew was immediately added to the reaction solution. The reaction mixture was stirred for 16 h at room temperature and subsequently quenched by addition of 1 M HCl (5 mL), resulting in formation of a colorless (yellow in the case of phenylthiohydroxamate derivatives) precipitate of white material. The suspension was extracted with chloroform (3× 5 mL). The aqueous phase was filtered and the collected solid was washed with water (2× 1 mL). The crude product was rinsed with distilled THF from the fritted funnel and the desired TW-peptide conjugate was obtained as a colorless (or yellow) solid after evaporation to dryness (6 μmol, 94%). LC traces of all final products (peptides, FAM-labeled peptides, tweezer molecules) can be found in the supplementary information (SI19).

Protein expression and purification. GST-tagged proteins were expressed in Escherichia coli solubL21 cultivated in LB media containing 50 μg/mL kanamycin. The expression was induced with 1 mM IPTG at an OD600 of 0.6–0.8. Bacteria were pelleted and lysed with lysozyme, followed by sonication in TRIS/NaCl in 1 M Tris-HCl buffer, pH 7.5, containing 1 mM DTT. Isotopically 15N-labeled GST tag was expressed in E. coli BL21-CodonPlus-RIL from Agilent Technologies using a pTG20 vector obtained from Dr. Sonia Banuelos (Department of Biochemistry and Molecular Biology, Biofisika Institute, University of the Basque Country, Leioa, Spain). Bacteria were cultivated in LB media containing 100 μg/mL carbenicillin for the pTG20 vector and 50 μg/mL chloramphenicol for maintaining the pACYC plasmid in the BL21-Codon Plus strain. The expression was induced with 0.1 mM IPTG at an OD600 of 0.5. Bacteria were pelleted and lysed with lysozyme and subsequent sonication in TRIS/NaCl (pH 7.4) supplemented with 1 mM PMSF. The GST-tagged proteins were then immobilized on GSTrap 4B columns. The GST-Tag was optionally cleaved with PreScission protease on column, depending on the experiments performed afterwards. The protein was then loaded on a Hitrap Q HP column and eluted with a 0.025–1 M NaCl gradient in 50 mM Tris-HCl buffer, pH 7.5, containing 1 mM DTT.

His-tagged CRM1 was expressed in E. coli BL21-ADE3 and transfected as described.

Isotopically 15N-labeled GST-tagged Survivin120 was expressed in E. coli solubL21 by growing a culture in 4 L LB medium at 37°C. At an OD600 of 1.0–1.2, the bacteria were pelleted and resuspended in 1 L M9 minimal medium supplemented with 1 g/L 15N-ammonium chloride. After incubation for 30 min at 37°C, expression was induced with 0.2 mM IPTG and the protein was expressed for 20 h at 30°C. The cells were harvested and lysed by sonication in PBS (pH 7.4) supplemented with 1 mM PMSF.

15N-GST-Survivin120 was purified via a GSTrap 4B affinity column. The GST-tag was cleaved with PreScission protease for 6 h at 4°C. Subsequent preparative size exclusion chromatography was performed with a HiLoad 26/600 Superdex 75 pg column and a
downstream mounted GSH-column from GE Healthcare in 50 mM KPi pH 7.4 with 150 mM KCl and 2 mM DTT. The pure protein was concentrated, and the buffer was exchanged to NMR buffer (50 mM KPi pH 7.4, 90 mM KCl, 2 mM DTT) using Vivaspin Ultrafiltration filters with a molecular weight cutoff of 10 kDa.

Isotopic titration calorimetry (ITC). ITC was performed with a MicroCal iTC200 from Malvern Panalytical in PBS, pH 7.4 at 25 °C with molecular tweezers in the cell and Survivin120 in the titration syringe. The protein was dialyzed overnight against PBS buffer. Ligands were dissolved in water, then 300 µM were titrated with 300 µM Survivin120 WT or 321 µM Survivin120 K90/103T. For reverse titrations, either 33.3 or 34.4 µM Survivin120 was titrated with 5 mM TW or TW-ELTL in the syringe, respectively. All titrations were performed in PBS, pH 7.4 at 25 °C. Then, 1.5 µL injections were used with 120 s spacing time between injections. The injection rate was set to 0.5 µL/s and the reference power was 5 µcal/s. ITC thermograms were fitted to a one set of sites andW who signal overlapped with the signal at the end titration were excluded from the intensity analysis. Spectra were processed with Topspin 3.5 (Bruker) and analyzed in CARA (version 1.9.1.7). Chemical shift perturbation and relative signal intensities were calculated from the raw chemical shift data and peak intensities using Excel 2016 (Microsoft) and plotted with GraphPad Prism 5.

NMR samples of Survivin120 mutants in NMR buffer (500 µM unlabeled Survivin120 mutants in NMR buffer (50 mM KPi pH 6.5, 90 mM KCl, 2 mM DTT) with overexpressed Survivin120-HA was preincubated without ligand or with 500 µM (wildtype) or 500 µM (K90/103T) point mutant was pre-bound to equilibrated GSH-beads in 500 µL pull-down buffer, containing additionally either no ligand or 50 µM unmodified tweezers TW, TW-ELTL, or TW-LFEEGLT for 1 h under rotation. After washing and blocking, GSH-beads were incubated with a protein mixture consisting of 2 mM dGTP, 50 µg CRM1, and 50 µg RanQ69L for 2 h under rotation.

Samples of input and beads taken during pull-down experiments were run on 12.5 SDS gels and transferred onto 0.2 µM PVDF membranes (Amersham Hybond P 0.2) with a molecular weight cutoff of 10 kDa. A 10 ng molecular tweezers (200 nM) were titrated with Survivin120 wildtype or K90/103T mutant until a final concentration of approximately 100 µM tweezers was reached. Data were transformed in Logs and IC50 was fitted using the equation in GraphPad Prism 8:

\[
y = A_{\min} + \frac{(A_{\max} - A_{\min})}{1 + 10^{(\frac{x - x_{\text{IC50}}}{\text{log}_10(10)}})}
\]

where \(A_{\min}\) is the anisotropy in the absence of tweezer, \(A_{\max}\) is the anisotropy at the end titration, and \(x\) is the concentration of tweezer, \(x_{\text{IC50}}\) was constrained for each data set, whereas \(A_{\max}\) and \(IC50\) were fitted.

For binding studies, 10 nM Survivin120 wildtype or 340 𝜇M Survivin120 wildtype or 340 𝜇M Survivin120 K90/103T was reached. Data were normalized to the measured anisotropy \(A_{\text{in}}\) in the absence of protein. Using a single-site binding model, the fluorescence anisotropy data were fitted to the equation:

\[
y = \left(1 + \frac{L}{K_D}\right) - \frac{L}{K_D + L}
\]

where \(A_{\text{in}}\) is the anisotropy at saturation, \(L\) is the concentration of the fluorescent ligand, \(x\) is the concentration of protein titrated, and \(K_D\) is the dissociation constant.

Statistical analysis. ITC thermograms were fitted to a one set of sites model with the software Origin provided with the instrument. Heat of dilution was subtracted as constant from each data point. From three independent experiments (n = 3), one representative example was depicted, and thermodynamic data were derived thereof. For pull-down experiments, only the representative quantification of the depicted western blot is shown. Note of low protein yields of the mutant CRM1, 1062VLV430AAA did not allow replicates of the respective experiment. Fluorescence anisotropy images were generated with GraphPad Prism 5.0. Depicted error bars represent the standard deviation of three independent titrations.

Computational details. For MD simulations of full-length Survivin, an initial structure for Survivin was generated in Modeller v9.17 using the Uniprot protein sequence O15392-1 [https://www.uniprot.org/uniprot/O15392] (human BIRC5 isoform alpha) as target and PDB entries 1E3I [https://www.rcsb.org/structure/1E3I] and 1FLH [https://www.rcsb.org/structure/1FLH] as templates. The best model was selected to minimize the DOPE and molpdf scores and was validated with PROCHECK (v.3.5.1) from the online Swiss-Model Workspace. The root-mean-square deviation (RMSD) between the model and templates is less than 0.7 Å for the crystal structures (1E3I, 1FLH, 3UEG, 3UEF, 3UEI) and 2.0 Å for the NMR solution (1X0X).

MD simulations were run with Gromacs 4.6.7 using the Amber ff9SB force field extended with ZAFF to model the zinc finger53. Topology files were created with the TLEap module of Amber v12.34 and converted to Gromacs topologies by ACy MPEG. Proteins were solvated in a dodecanebox of SPC/E water molecules with a 10 Å minimum separation between the protein and the box boundaries. The system was neutralized by addition of Na+ and Cl− ions to a final ionic strength of 0.15 M. The simulation system was energy minimized by steepest-descent to a total force of 2000 eV, equilibrated for 5 ns in the NVT ensemble with restrained heavy atoms, and for 5 ns in the NPT ensemble without restraints. Production simulations were run in the NPT ensemble for a total of 310 ns (3 × 50 ns, 2 × 80 ns). Temperature was stabilized at 300 K in the NVT and NPT ensembles by the v-rescale thermostat, while the pressure was stabilized at 1 atm in the NPT ensemble by the Berendsen barostat (equilibration) or Parrinello–Rahman barostat (data production). Simulations were carried out on a GPU (GeForce 970 and GeForce 1070, CUDA 6.5) using a time step of 2 fs, the Verlet integration method, and an 8-atom cutoff for nonbonded interactions.
Supplementary Information

Data supporting the findings of this manuscript are available from the corresponding authors upon reasonable request. A reporting summary for this article is available as a Reporting Summary (SI10).

Data availability

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Author contributions

Knauer group: A. M. and S. B. cloned, expressed, and purified proteins, and performed ITC titrations as well as pull-down experiments. S. B. developed the protocols for protein expression and purification as well as for pull-down and initial ITC experiments. A. M. developed, optimized, and performed fluorescence anisotropy experiments, optimized the ITC protocols, and analyzed the ITC data together with J. H. A. M. and S. B. wrote the manuscript together with T. S., E. S.-G. and S. K. K., who designed the study and supervised the experiments together with C. V. Schrader group: C. H. designed the modified tweezer constructs from computational visualization; he also developed the synthetic protocol for the generation of tweezer conjugates by click chemistry and synthesized the first generation tweezer conjugates. I. H. synthesized FAM-labeled tweezers, modified the tweezers with the scrambled peptide, and determined affinities by ITC. T. S. and A. M. wrote the paper together with S. K. K. and E. S.-G. Sánchez-García group: Y. B. R.-B. and J. M.-P. carried out the QM/MM calculations, the classical MD simulations, and GaMD simulations of Survivin dimer and Survivin monomer without and with tweezers, processed the data, and analyzed the results together with E. S.-G., who designed the computational work and wrote the paper together with S. K. K. and T. S. Hoffmann group: J.-N. Grad carried out MD simulations on the full-length Survivin dimer and interpreted the data together with D. H. Bayer group: M. P. expressed and purified 15N-labeled Survivin. C. B. and M. P. performed the NMR titrations. P. B. and C. B. supervised the NMR experiments and interpreted the NMR data.

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Competing interests

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