Click to enter: activation of oligo-arginine cell-penetrating peptides by bioorthogonal tetrazine ligations†

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Cell-penetrating peptides are able to transport a wide variety of cargo across cell membranes. Although promising, they are not often considered for therapeutic purposes as they lack controllable activity and cell selectivity. We have developed an activation strategy based on a split octa-arginine cell-penetrating peptide (CPP) that can be activated by means of bioorthogonal ligation. To this end we prepared two non-penetrating tetra-arginine halves, functionalized either with a tetrazine or with a complementary bicyclo[6.1.0]nonyne (BCN) group. We demonstrate that an active octa-arginine can be reconstituted in situ upon mixing the complementary split peptides. The resulting activated peptide is taken up as efficiently as the well-established cell-penetrating peptide octa-arginine. The activation of the oligo-arginines can also be achieved using trans-cyclooctene (TCO) as a ligation partner, while norbornene appears too kinetically slow for use in situ. We further show that this strategy can be applied successfully to transport a large protein into living cells. Our results validate a promising first step in achieving control over cell penetration and to use CPPs for therapeutic approaches.

Previously, we demonstrated an activation strategy for oligo-arginines. For these peptides it is known that a minimum amount of six arginine residues is needed to induce cellular uptake. Making use of this characteristic, a split-CPP approach was designed that involved truncated oligo-arginines consisting of three to five arginine residues and a C-terminal cysteine. These peptides were assembled and activated through disulfide bridge formation with complementary oligo-arginines containing an N-terminal cysteine residue. Whereas the truncated split peptides did not induce uptake in HeLa cells, the assembled octa- and nona-arginines were efficient CPPs. Although we demonstrated that activation of short oligo-arginine peptides is possible by extension of their sequence, its dependency on the formation of disulfide bridges hampers their use for in situ activation due to the dominant oxidative environment and the presence of potentially interfering thiols in many cellular environments. To overcome this limitation, and to circumvent possible non-specific interactions of the split CPPs with endogenous biomolecules, we investigated alternative activation strategies based on the bioorthogonal tetrazine ligation (Fig. 1).

Results and discussion

Design and synthesis

The tetrazine ligation with alkenes or alkynes is based on an inverse electron demand Diels–Alder (iEDDA) reaction and has been used previously for several in vivo applications. For our
in situ activation strategy of CPPs we synthesized several truncated tetra-arginines equipped with a tetrazine moiety or with a complementary alkene and alkyne reactive group. We first prepared a tetra-arginine, of which the C-terminus was connected to a 3-aminobenzyl-6-methyl substituted tetrazine (R4-Tz, Fig. 2). We selected this tetrazine, because of its stability and solubility in aqueous environments. Although the reactivity of this tetrazine is moderate compared to other more electron-deficient less stable tetrazines, the reaction rates are still suitable for biological applications. Further, we labelled the N-terminus with fluorescein isothiocyanate (FITC), linked via a 6-aminohexanoic acid spacer, to enable the visualization of cellular uptake with confocal laser scanning microscopy (CLSM) and quantification using flow cytometry. As an activation partner for the R4-Tz we first explored a tetra-arginine containing the bioorthogonal bicyclo[6.1.0]nonyne (BCN-R4). In addition, we synthesized fluorescently labelled octa-arginine (R8) as a reference CPP and pre-conjugated R4-b-R4 – prepared from R4-Tz and BCN-R4 – that allowed us to determine the activity of the fully reconstituted CPP and find suitable conditions for uptake (Fig. 2).

Uptake of pre-ligated conjugates

The mechanism of uptake of arginine-rich CPPs is dependent on their concentration and their structures, therefore we started by comparing the uptake of CPP R8 with that of the pre-ligated peptide R4-b-R4 at 5 μM, 10 μM, 15 μM and 20 μM using confocal fluorescence microscopy (Fig. 3). As a control, we measured the uptake of the truncated peptide R4-Tz. At low concentrations (5 μM and 10 μM), we did not observe any uptake of the truncated peptide R4-Tz, which was in accordance with earlier findings that demonstrated that at least six arginines within one sequence are required for efficient cellular uptake. At higher concentrations (15 μM and 20 μM) we observed some

![Fig. 1 Overview of the click to enter strategy.](image1)

![Fig. 2 Structures of oligo-arginine peptides used in this study.](image2)

![Fig. 3 CLSM micrographs corresponding to cellular uptake studies in HeLa cells. The concentration of the CPP increases from top to bottom (5, 10, 15 and 20 μM). The experiments were performed with reference peptide R8, pre-ligated R4-b-R4 and truncated R4-Tz at 37 °C, with an incubation time of 30 minutes in serum-containing medium.](image3)
in situ

R4-Tz

While 5 eq. of co-incubating HeLa cells with 5 eq. of bioorthogonal tetrazine ligation. We commenced by us to study the in situ activation of oligo-arginines using the bioorthogonal tetrazine ligation. We commenced by co-incubating HeLa cells with 5 \( \mu M \) R4-Tz in serum-containing culture medium and added 5 eq. of BCN-R4 to allow ligation and formation of R4-b-R4. We included R8 as positive control, while R4-Tz served as negative control. As seen by CLSM, the uptake of in situ activated R4-Tz + BCN-R4 at 30 minutes was similar to the uptake of R8 as well as R4-b-R4, indicating that in situ activation of R4-Tz to R4-b-R4 is possible at these conditions (Fig. 4 and S1†).

Next, we evaluated whether uptake could be improved by increasing the amount of activating agent BCN-R4 or by prolonging the incubation time. We found that addition of 10 eq. of BCN-R4 instead of 5 eq. did not improve uptake (Fig S2†), while reducing the amount of BCN-R4 from 5 eq. to 2.5 eq. resulted in a significant decrease in cellular uptake (Fig. S3†). Subsequently, we assessed the effect of incubation time on in situ activation of R4-Tz. Using similar conditions as above we observed CPP uptake already after 30 minutes and that uptake increased slightly over time. At 60 minutes, similar uptake levels were reached as observed after 90 minutes (Fig. S3†). From these results we concluded that 30–60 minutes of incubation using 5 eq. of BCN-R4 activating agent are needed for sufficient CPP uptake.

To study the effect of the tetrazine ligation rate, we additionally prepared tetra-arginine fragments that were N-terminally equipped with either a norbornene (Norb-R4), or a trans-cyclooctene (TCO-R4) moiety (Fig. 2). Norbornene and TCO are known to display a respectively lower and higher reactivity towards tetrazines compared to BCN. We compared the uptake of the in situ activation of R4-Tz by the tetra-arginines with the different bioorthogonal reagents using CLSM (Fig. 4A). To acquire the uptake efficiency more quantitatively, we additionally measured the cellular fluorescence of the various tetrazine reactions with flow cytometry (Fig. 4B). HeLa cells were incubated with R8 at 5 \( \mu M \) for 30 minutes at 37 °C in serum-containing medium and compared to cells that were incubated for 30 minutes with 5 \( \mu M \) R4-Tz to which 5 equivalents of BCN-R4, Norb-R4, or TCO-R4 were added respectively. As shown in Fig. 4, truncated split R4-Tz again hardly gave any uptake. Interestingly, the R4-Tz peptide incubated with Norb-R4 showed only poor uptake while both activation with BCN-R4 and TCO-R4 gave uptake that was comparable to reference peptide R8. Interestingly, no significant difference was found between BCN and TCO.

Protein cargo

As our activation strategy is feasible using an iEDDA conjugation reaction for a small molecule fluorophore we next explored whether our strategy could also be used for the induced uptake of a much larger protein cargo. To this end we selected fluorescently labelled human serum albumin (HSA) as a model protein. This 66 kDa protein contains a single free cysteine that allowed us to site selectively modify the protein with a tetrazine. This was achieved by first reacting the cysteine of HSA with a bi-functional maleimide-DBCO linker and reaction of free lysines of HSA with

![Figure 4](image-url)
FTTC providing fluorescent HSA-DBCO. A tetra-arginine with an N-terminal azide and a C-terminal tetrazine, N3-R4-Tz, was conjugated in a sequential step yielding HSA-R4-Tz (Fig. 5A). In addition, we prepared an octa-arginine protein conjugate (HSA-R8) using N3-R8, for comparison.

Using HSA as cargo conjugated to a tetrazine oligo-arginine, activation with the BCN-R4 peptide under the conditions employed above, we did not observe significant cellular uptake as evidenced by FACS as well as confocal microscopy. Fortunately, incubation at a slightly higher concentration (10 μM) and increased time (90 min) resulted in clear uptake of the HSA-R4-Tz protein which was in situ activated by conjugation to BCN-R4 (Fig. 5B). Similar uptake levels were observed for HSA-R8 while unreacted HSA-R4-Tz was only poorly taken up. These qualitative results were quantified using flow cytometry using the same conditions (Fig. 5C) and substantiate that our in situ CPP activation strategy is suitable for the delivery of a large protein cargo.

Conclusions

In conclusion, we have successfully developed an in situ activation strategy for oligo-arginine CPPs based on a bio-orthogonal tetrazine ligation between two split tetra-arginine parts. The norbornene-based reaction did not show kinetics suitable for in situ application, but both BCN and TCO were efficient in reconstituting active CPPs. This provides a valuable method for probe design and is the basis for further efforts in this line of research. Moreover, using BCN efficient cellular uptake was observed for a low molecular weight fluorescent probe as well as a large protein cargo. Future efforts in this line of research are to incorporate homing peptides or targeting ligands in our tetra-arginine activation strategy, providing the first steps towards a targeted and site-selective cargo delivery strategy involving activatable CPPs.

Conflicts of interest

There are no conflicts to declare.

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Notes and references