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De novo designed peptides for cellular delivery and subcellular localisation

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Increasingly, it is possible to design peptide and protein assemblies *de novo* from first principles or computationally. This approach provides new routes to functional synthetic polypeptides, including designs to target and bind proteins of interest. Much of this work has been developed *in vitro*. Therefore, a challenge is to deliver *de novo* polypeptides efficiently to sites of action within cells. Here we describe the design, characterisation, intracellular delivery, and subcellular localisation of a *de novo* synthetic peptide system. This system comprises a dual-function basic peptide, programmed both for cell penetration and target binding, and a complementary acidic peptide that can be fused to proteins of interest and introduced into cells using synthetic DNA. The designs are characterised *in vitro* using biophysical methods and X-ray crystallography. The utility of the system for delivery into mammalian cells and subcellular targeting is demonstrated by marking organelles and actively engaging functional protein complexes.

De novo peptide and protein design has advanced to allow the construction of a rapidly increasing variety of stably folded structures and assemblies completely from scratch^{1–3}. This ability ushers in new challenges for *de novo* design. These include endowing synthetic polypeptides with functions such as binding and catalysis and, as most designs to date have been made and tested *in vitro*, porting these molecules into living cells, tissues and organisms. Whilst there have been successes with *in cellulo* and *in vivo* applications of *de novo* design^{4–13}, considerable challenges remain to make this mainstream activity. Here we address the challenge of delivering exogenous cargoes to subcellular targets in mammalian cells. We do this by tagging the cargo with a *de novo* designed cell-penetrating peptide (CPP) that forms a highly specific peptide–peptide interaction with a partner peptide appended to the subcellular target.

Peptide–peptide, peptide–protein and protein–protein interactions (PPIs) are central to most, if not all, biological processes¹⁴. Therefore, disrupting existing PPIs or generating new ones to monitor or intervene in such processes is a major endeavour in peptide and protein design and engineering^{14–16}. Nonetheless, even with bioactive polypeptides that target PPIs in hand, the challenge of delivering these into mammalian cells where they would be most useful remains. Exogenous reagents—usually proteins and larger assemblies—presented to cells can enter actively through endocytosis^{17,18}. In this case, however, the reagents end up in the endolysosomal system, which can inhibit release into the cytosol¹⁹. Alternatively, certain reagents—such as CPPs^{20,21}—can enter cells via direct or passive translocation across biological membranes. Typically, CPPs are basic or cationic peptides of ≤ 30 amino acids in length, and the addition or conjugation of such sequences can be used to effect the import of various cargoes into cells^{22–25}.

The small size of CPPs has advantages and disadvantages: it facilitates cell uptake, but it leaves little room for internal functionalisation. One strategy to address this is to enhance the cell-permeability

of a functional peptide directly by overlaying charged residues at non-target-binding positions²⁶. α -Helical coiled coils (CCs) are a class of peptide assembly with potential to be programmed for both cell penetration and protein recognition^{27,28}. Effectively, CC peptides are bifaceted: the peptide–peptide interfaces are well defined and can be designed rationally or computationally to deliver various oligomers, topologies and partners^{2,3}. This leaves exposed surfaces that can be altered for water or membrane solubility. Moreover, CC peptides are short and synthetically accessible and could fit the CPP paradigm^{29–32}.

Here we describe the design and characterisation of new antiparallel homo- and hetero-dimeric CCs that expand the set of structurally resolved *de novo* CCs^{33–39}. We show that the heterodimer is effective for both delivery into cells and subcellular targeting. In this system, an arginine-rich basic CC peptide is used as the exogenous reagent, as it incorporates both cell-penetrating and target-binding functionalities. The sequence of its complementary acidic partner can be fused to various subcellular proteins and introduced into cells via the transfection of synthetic DNA. We show that this system can tag intracellular proteins, complexes and organelles. Moreover, the two components diversify the labels that can be incorporated for imaging and intervening in biological processes.

Results

Rational *de novo* design of antiparallel coiled-coil dimers. We decided to design a cell-penetrating system using antiparallel rather than parallel CCs. This was because structurally resolved *de novo* designed antiparallel CC dimers are currently poorly represented⁴⁰, which offers an opportunity to develop new design principles and sequences to address our target. The rational design of CC dimers to tetramers centres on the interfacial *g-a-d-e* sites of their *abcdefg*, heptad repeats²⁸ (Fig. 1a). This can be guided by the large number of available natural CC sequences and structures. Therefore, we

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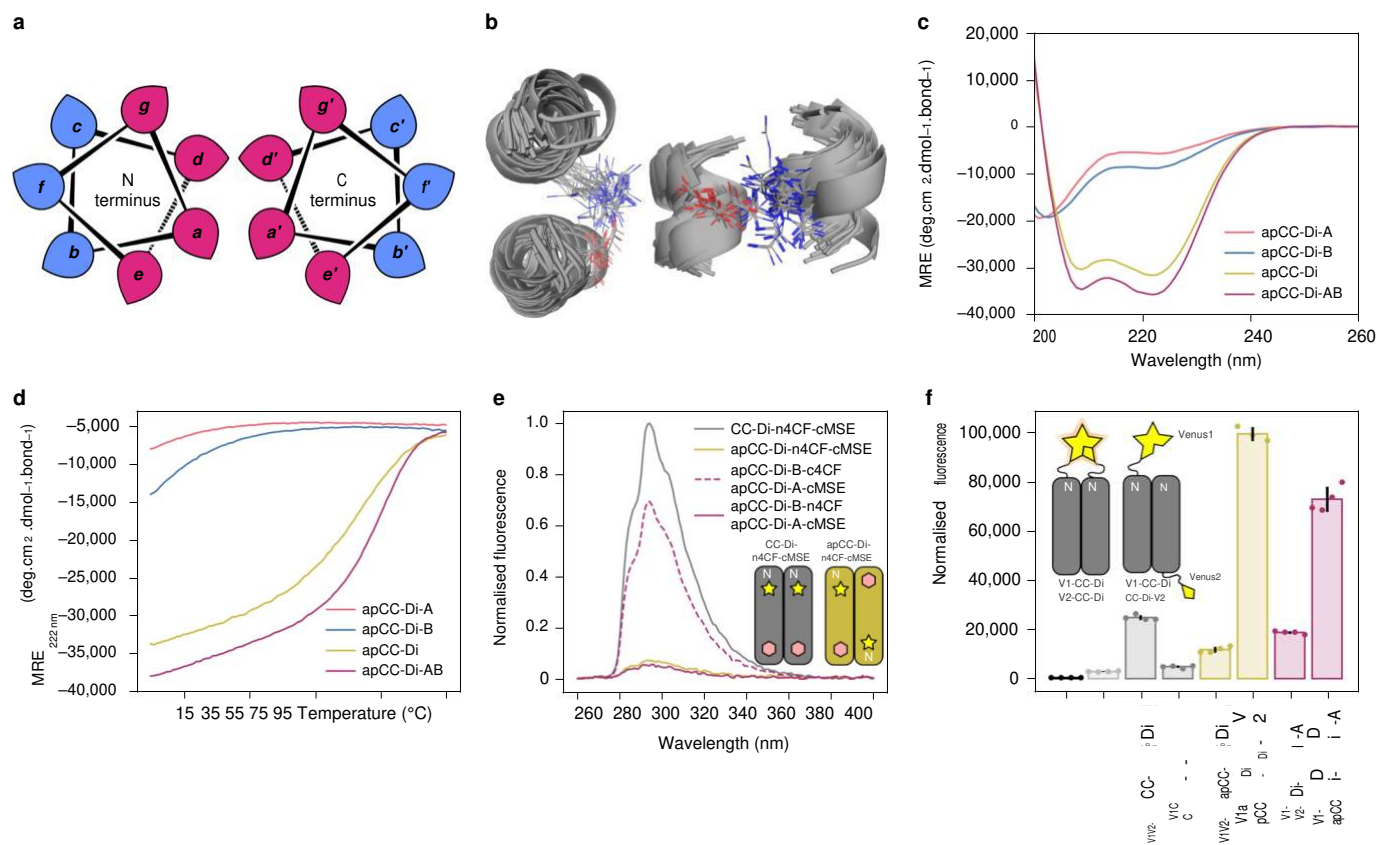


Fig. 1 | Design and biophysical characterisation of antiparallel CC dimers. **a**, Helical wheels for the seven-residue repeat of CC sequences labelled **a** to **g**. Interface positions (**a**, **d**, **e** & **g**) are coloured pink and solvent-exposed positions (**b**, **c** & **f**) are coloured blue. **b**, Examples of arginine to aspartic acid interactions observed in antiparallel CC dimers. **c,d**, Circular-dichroism spectra at 20 °C (**c**) and thermal-denaturation profiles (**d**), for apCC-Di and apCC-Di-AB. Conditions: 100 μ M total peptide, PBS, pH 7.4. **e**, Fluorescence-quenching assay for labelled apCC-Di, apCC-Di-AB, and the control parallel peptide CC-Di. n and c indicate mutations near the N and C termini, respectively; 4CF, 4-cyano-L-phenylalanine fluorophore (yellow star); MSE, L-selenomethionine fluorescence quencher (pink hexagon); CC-Di, parallel CC dimer control. Conditions: 100 μ M concentration of each peptide, PBS, pH 7.4. **f**, BiFC assay for CC-Di, apCC-Di and apCC-Di-AB. V1 and V2 are N- and C-terminal fragments of the Venus fluorescent protein, respectively. Peptide names that precede the Venus fragment name denote fusion to the C terminus of the CC, for example, apCC-Di-B-V2, and vice versa. Fluorescence values are normalised to cell density (OD_{600}) and are presented as mean \pm s.d. from $n \geq 3$ technical replicate measurements (dots).

selected a subset of antiparallel two-helix CCs from the CC + structural database⁴¹ and examined the amino-acid profiles of their repeats (Supplementary Note and Supplementary Figs. 1 and 2). This revealed preferences for isoleucine (Ile, I) and leucine (Leu, L) at the core-forming **a** and **d** positions, respectively. Unfortunately, this combination also specifies parallel CC dimers²⁸. Polar residues at these sites can aid oligomer specification and helix orientation^{28,34,42,43}. Comparison of the profiles for antiparallel and parallel dimers indicated that arginine (Arg, R) was enriched at **a** sites of the former by almost twofold. Inspection of antiparallel structures showed that R@**a** pairs most often (30% of cases) with aspartate (Asp, D) at proximal **e'** sites of the partnering helices resulting in inter-side-chain hydrogen bonds or salt bridges (Fig. 1b). Therefore, in addition to I@**a** and L@**d**, we included a single R@**a** to interact with a complementary D@**e** in four-heptad, 30-residue designs, Table 1. The use of oppositely charged residues at core-flanking **e** and **g** sites is established in CC design²⁸. Therefore, to target an antiparallel homo-dimer, apCC-Di, we made the N-terminal **e** and **g** sites glutamate (Glu, E), the most favoured residues at these sites in CC dimers, and the complementary C-terminal **e** and **g** sites Arg and lysine (Lys, K), respectively. The heterodimer, apCC-Di-AB, required two peptides: apCC-Di-A had all Glu at **e** and **g**; and apCC-Di-B had a combination of Arg and Lys at these sites to: (i) complement the acidic 'A' peptide; and (ii) promote cell penetration. Several sequences were made by solid-phase peptide synthesis and

characterised biophysically (Supplementary Table 1, Supplementary Figs. 3–9,20–28), but we focus here on the most successful designs apCC-Di and apCC-Di-AB below (see Supplementary Note for design iterations).

Structural characterisation of apCC-Di and apCC-Di-AB.

The circular dichroism (CD) spectrum of apCC-Di indicated a fully α -helical structure (Fig. 1c). Furthermore, temperature-dependent CD measurements showed a sigmoidal, reversible, and concentration-dependent unfolding transition as expected for a unique, cooperatively folded species (Fig. 1d). Consistent with this and the design, sedimentation-equilibrium analytical-ultracentrifugation experiments returned dimer molecular weights (Supplementary Figs. 22–28). The apCC-Di-AB design required iteration (Supplementary Note). In isolation, the final peptides, apCC-Di-A and apCC-Di-B, were unfolded (Fig. 1c,d). When mixed 1:1, they gave a fully α -helical species (Fig. 1c), with a reversible thermal unfolding transition (Fig. 1d), and a dimeric molecular weight (Supplementary Fig. 28). We estimated nanomolar affinities for apCC-Di and apCC-Di-AB from the concentration-dependent thermal-denaturation experiments⁴⁴ (Supplementary Figs. 29 and 30).

We crystallised and solved X-ray structures for apCC-Di, apCC-Di-AB, and several variants of these (Table 1 and Supplementary Table 3). The atomic-resolution structures of

Table 1 | Peptide sequences and summary of biophysical data

Name	Sequence				Helicity (%)	T_M (°C)	AUC	XRD
	<i>cdefgab</i>	<i>cdefgab</i>	<i>cdefgab</i>	<i>cdefgab</i>				
apCC-Di	GQLEQELA	ALDQQIA	ALKQRRR	ALKWQIQG	83	63	$1.9 \times m$	1.08 Å (7Q1R)
apCC-Di-A	GQLEQELA	ALDQEIA	AAEQELA	ALDWQIQG	15	–	–	–
apCC-Di-B	GQLKQRRR	ALKQRIA	ALKQRRR	ALKWQIQG	23	–	–	–
apCC-Di-AB	apCC-Di-A+ apCC-Di-B				93	69	$0.9 \times d$	1.68 Å (7Q1T)

Helicity, percentage α helix determined by circular dichroism from the MRE₂₂₂; T_M , midpoint of the thermal unfolding transition. AUC, molecular weight measured by analytical ultracentrifugation relative to expected masses for a monomer (*m*) or heterodimer (*d*); XRD, resolution of X-ray crystallography data, and Protein Data Bank deposition ID in brackets.

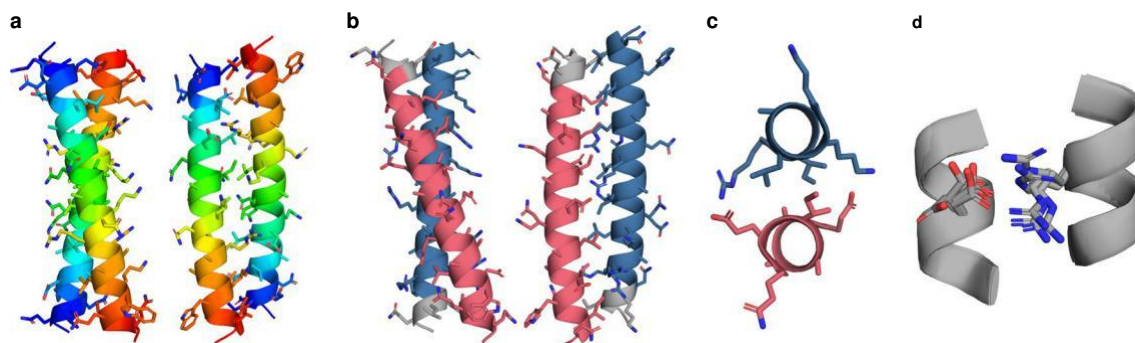


Fig. 2 | X-ray crystal structures reveal antiparallel CC as designed. **a,b**, Orthogonal views of X-ray crystal structures of apCC-Di (**a**) and apCC-Di-AB (**b**). In **b** the individual chains are apCC-Di-A (red) and apCC-Di-B (blue). The N-terminal turns are coloured grey. **c**, A single heptad from apCC-Di-AB viewed along the superhelical axis showing the definitive knobs-into-holes packing observed in CCs. **d**, Overlay of designed Arg–Asp interactions from the experimental apCC-Di and apCC-Di-AB structures. Arg–N₁ to Asp–O_δ distances of ≤ 3.5 Å were taken as indicating salt-bridge formation.

apCC-Di and apCC-Di-AB revealed antiparallel helical dimers as designed (Fig. 2a,b and Extended Data Figs. 1 and 2). Moreover, the program SOCKET2 (ref. 45) only identified knobs-into-holes interactions, the structural signature of CCs, within the designed antiparallel interfaces (Fig. 2c and Extended Data Figs. 3–5); and the introduced Arg–Asp pairs made salt bridges in three of four cases (Fig. 2d).

Next, we sought to confirm that the folded states were antiparallel in solution. First, guided by the X-ray crystal structures, we introduced the fluorescent 4-cyano-L-phenylalanine (4CF) and a fluorescence quencher L-selenomethionine (MSE)⁴⁶ at proximal interhelical *g* and *d* positions. This was done for apCC-Di and apCC-Di-AB, and with N-terminal 4CF and C-terminal MSE and vice versa (Supplementary Table 1, Supplementary Figs. 10–16 and Extended Data Fig. 6). As controls, we made analogous changes to the parallel CC-Di design³⁴ and introduced 4CF and MSE at the C termini of the apCC-Di-AB design. Only the control peptides fluoresced (Fig. 1e and Extended Data Fig. 7). By contrast, fluorescence was completely quenched in the antiparallel designs with juxtaposed 4CF and MSE residues (Fig. 1e).

Second and to test the designs in cells, we reconfigured a bimolecular fluorescence complementation (BiFC) assay to examine peptide assembly and orientation directly in *Escherichia coli* cells⁴⁷. Using a variant of the fluorescent protein Venus split after residue 154 to give 2 fragments, V1 and V2 (refs. 48–50), we fused one de novo peptide to the C terminus of V1. The other test peptide was fused to either the N or C terminus of V2 (Extended Data Fig. 8 and Supplementary Figs. 31–34). The hypothesis was that any fragment complementation and fluorescence should report on antiparallel or parallel peptide association, respectively; that is, antiparallel CCs should increase the fluorescence readout when fused to the C terminus of V1 and the N terminus of V2. Experimentally, this was

done by transforming *E. coli* cells with expression plasmids encoding synthetic genes for the peptide-split protein fusions, and then the interaction was monitored by direct measurement of fluorescence from the cells. The background fluorescence from V1 plus V2 alone was low (Fig. 1f and Extended Data Fig. 8). As a positive control, we tested the homomeric, parallel CC-Di sequence in the system³⁴. With CC-Di on the C terminus of V1 and the N terminus of V2 there was an approximately 1.4-fold increase in fluorescence above baseline. Whereas, with CC-Di C terminus to both fragments the signal increased by approximately 7-fold. These results are consistent with the parallel orientation of CC-Di and validate the assay. Turning to the new designs, fusing apCC-Di to the N terminus of V2 gave a larger signal and an approximately 8-fold increase in fluorescence relative to the C-terminal fusion, consistent with apCC-Di being antiparallel. Furthermore, with apCC-Di-A fused to the C terminus of V1 the better complementary fusion had apCC-Di-B at the N terminus of V2 with an approximately 4-fold increase in fluorescence over the alternative construct. Thus, both apCC-Di and apCC-Di-AB are confirmed as antiparallel dimers in solution and in cells.

apCC-Di-B delivers functional cargo to subcellular locations. With a verified antiparallel heterodimer in hand, we turned to the delivery into eukaryotic cells. We resynthesised apCC-Di-A and apCC-Di-B each with an N-terminal 5-carboxytetramethylrhodamine (TAMRA) fluorophore (Supplementary Figs. 17–19). The peptides were added separately to cultures of HeLa cells, fixed and imaged after 1 h. Only TAMRA-apCC-Di-B entered cells (Fig. 3a). Moreover, it did so at 4 °C indicating passive uptake directly into the cytosol rather than via active endocytosis. To confirm this, we pre-incubated cells with endocytosis inhibitors, MiTMAB or Dynasore, or the electron transport chain inhibitor, NaN₃, for 30 min before the addition

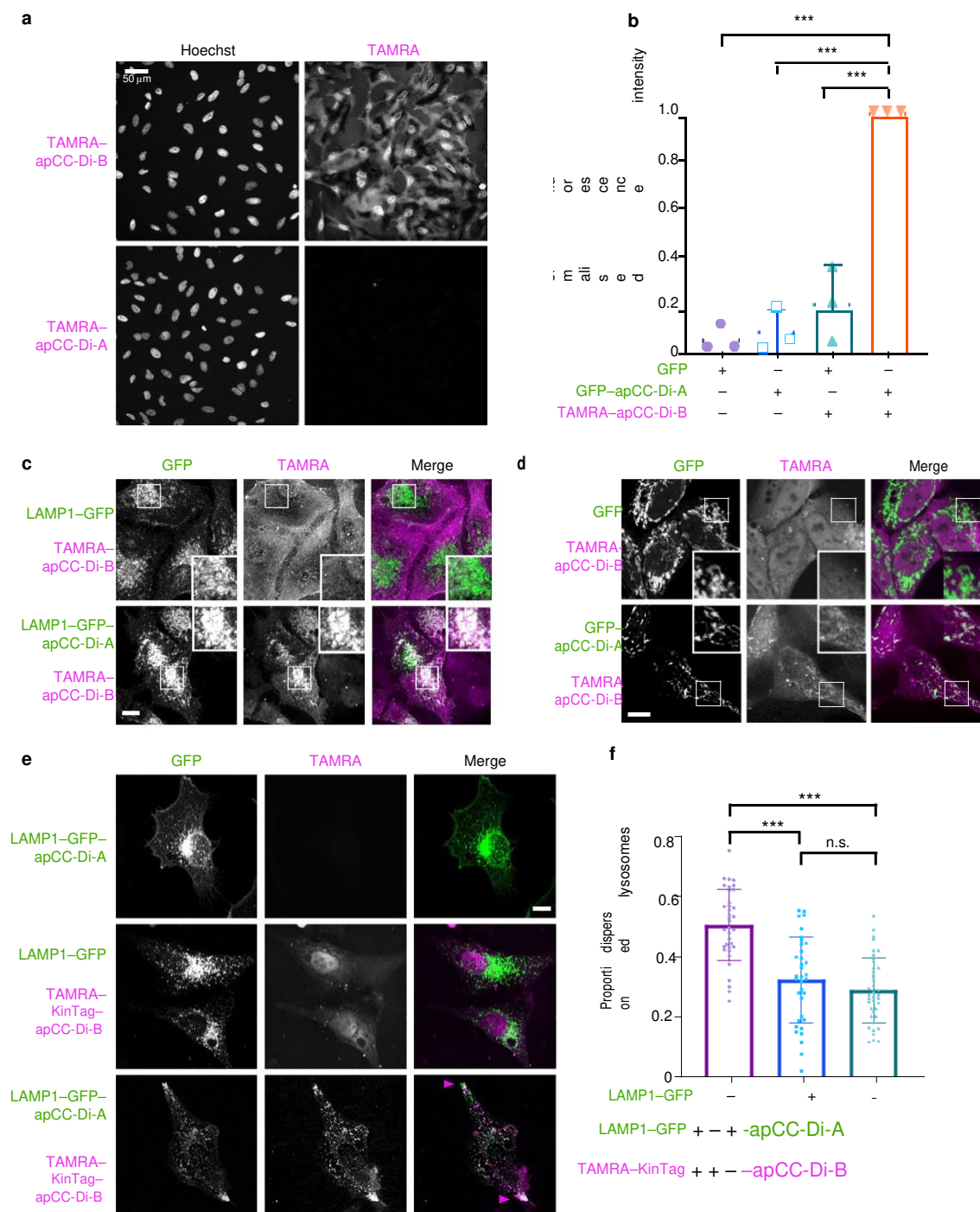


Fig. 3 | apCC-Di-B is cell penetrating and binds to apCC-Di-A in mammalian cells. **a**, HeLa cells treated with 2 μ M TAMRA-labelled apCC-Di-B (top) or apCC-Di-A (bottom) peptides for 1 h. apCC-Di-B is cell penetrating as shown by TAMRA fluorescence in the cytoplasm and nucleus; apCC-Di-A does not enter cells. Scale bar, 50 μ m. **b**, Cells transfected with GFP or GFP-apCC-Di-A and treated with TAMRA-apCC-Di-B or vehicle control were subject to GFP immunoprecipitation and quantification of TAMRA immobilised on beads measured by fluorescence of boiled bead eluant at 555 nm. Fluorescence was normalised to the signal of cells transfected with GFP-apCC-Di-A and treated with TAMRA-apCC-Di-B in three independent experiments. Data are presented as mean \pm s.d. and an unpaired *t*-test is used for statistical analysis. ****P* < 0.0001. TAMRA-apCC-Di-B binds to GFP-apCC-Di-A in cell lysate and is immobilised on beads. **c**, Cells transfected with LAMP1-GFP (top) or LAMP1-GFP-apCC-Di-A (bottom) and treated with TAMRA-apCC-Di-B. Co-localisation of GFP and TAMRA fluorescence on lysosomes indicates apCC-Di-AB dimerisation. Scale bar 20 μ m. **d**, Cells transfected with a mitochondrially targeted GFP nanobody, mitoGFP-en, and GFP (top) or GFP-apCC-Di-A (bottom) and treated with TAMRA-apCC-Di-B. Co-localisation of GFP and TAMRA fluorescence on mitochondria indicates apCC-Di-AB dimerisation. Scale bar, 20 μ m. **e**, Cells transfected with LAMP1-GFP-apCC-Di-A (top and bottom) or LAMP1-GFP (middle) and treated with TAMRA-KinTag-apCC-Di-B (middle and bottom) or vehicle control (top). KinTag is delivered to lysosomes by apCC-Di-AB dimerisation and promotes transport of lysosomes to the cell periphery, purple arrows indicate peripheral accumulations. Scale bar, 10 μ m. **f**, Quantification of lysosome dispersal from **e** by comparison of fluorescence intensity of perinuclear cluster with total LAMP1 fluorescence in a minimum of 35 cells (*n* = 36, 35, 40 cells from left to right), pooled from three independent experiments. Data are presented as mean \pm s.d. and an unpaired *t*-test is used for statistical analysis. ****P* < 0.0001; n.s., *P* = 0.2421.

of apCC-Di-B. In all three cases, peptide uptake by the cells was unaffected (Extended Data Fig. 9). Thus, apCC-Di-B is a de novo CPP consistent with the basic, arginine-rich sequence design. To determine whether apCC-Di-B taken up by mammalian cells can still bind apCC-Di-A, we added TAMRA-apCC-Di-B to HeLa cells transfected with plasmids for transient expression of green fluorescent protein (GFP) alone or of a GFP-apCC-Di-A fusion. Cells were lysed, GFP proteins were immunoprecipitated, and any associated TAMRA fluorescence signal was analysed using a microplate reader. TAMRA-apCC-Di-B signal only co-immunoprecipitated with the GFP-apCC-Di-A fusion (Fig. 3b).

Next, we asked if the apCC-Di-AB system could target subcellular organelles. For this, we constructed a fusion with the lysosomal-associated membrane protein 1, LAMP1-mGFP-apCC-Di-A, to present an exposed peptide bait on late endosomes and lysosomes⁵¹. Transfection of this gene into HeLa cells followed by exogenous treatment with TAMRA-apCC-Di-B showed striking recruitment to LAMP1 compartments (Fig. 3c). In addition, we used a mitochondrially targeted GFP nanobody (pMito-GBPen) to recruit GFP-apCC-Di-A to mitochondria⁵². Treatment of these cells with TAMRA-apCC-Di-B resulted in its localisation to mitochondria (Fig. 3d).

Finally, to determine if apCC-Di-B could deliver functional moieties that drive new PPIs in cells, we extended it with a ‘KinTag’ sequence (Supplementary Table 1 and 2). Recently, we developed this de novo sequence to bind endogenous kinesin-1 light chains and, so, provide a synthetic adapter between kinesin-1 heavy chain motors and potential cargoes⁵³. Cells expressing the LAMP1-mGFP-apCC-Di-A fusion were treated with TAMRA-KinTag-apCC-Di-B as an exogenous reagent. TAMRA and GFP fluorescence co-localised to lysosomal puncta indicating formation of the apCC-Di-AB complex (Fig. 3e), implying that the apCC-Di-B component facilitated cell entry and localisation to the intended target. Moreover, compared with controls, the lysosomes dispersed from the perinuclear region of the cell and accumulated at the cell periphery (Fig. 3e,f). Interestingly, this led to altered cell shapes and GFP- and TAMRA-positive projections (Fig. 3e). This is consistent with the KinTag component of the exogenous reagent binding to and strongly activating kinesin, which—mediated by the apCC-Di-AB bridge—transports its lysosome cargo along microtubules to the cell periphery.

Discussion

In summary, we have designed and fully characterised homo- and heteromeric antiparallel CC dimers. This includes in-cell confirmation of the antiparallel orientation using a variation on the split-Venus BiFC assay, validated with known parallel interacting CCs, which should be useful to others investigating the orientation of PPI domains in cells. Others have reported designs for antiparallel CC dimers^{42,54–62}. However, to our knowledge, apCC-Di-AB and apCC-Di are the first heteromeric and the second homomeric de novo design to be confirmed structurally to atomic resolution⁴⁰. In addition, we have shown that the basic, arginine-rich B peptide enters directly into the cytosol of mammalian cells; it is a new cell-penetrating peptide. Once in cells, the B peptide can seek and bind its acidic partner introduced as parts of synthetic genes. This can be exploited using the B peptide as an exogenous reagent for targeting intracellular proteins, complexes and organelles labelled with the complementary A peptide. Finally, the system can deliver an orthogonally functional peptide, which we demonstrate by recruiting an endogenous motor protein to relocate organelle cargoes.

Others have reported the use of CC peptides, notably natural leucine zippers, in combination with CPPs to import cargoes into cells^{63–65}. Our system is different as it combines cell penetration and partner binding within a single CC peptide motif. As we have demonstrated, this allows one half of the system to be functionalised for

both cell penetration and partner binding. In turn, this can be used to import dyes for imaging or short bioactive peptides to engage with active subcellular processes. We anticipate that this and related de novo designed peptides will add to the armoury of reagents available for identifying and following subcellular assemblies and processes in mammalian cells.

Online content

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at.

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Methods

Bioinformatic analysis. The CC database, CC+ (http://coiledcoils.chm.bris.ac.uk/ccplus/search/dynamic_interface) was filtered using the following options: redundancy, $\leq 50\%$; α helices, 2; orientation, antiparallel; partnering, any; chains, any; repeats, any; and length, > 11 residues. The SOCKET text files were downloaded for the filtered list of CCs. The knob residue identity at α and d positions (and their corresponding hole residue identity and relative heptad positions) were extracted from the SOCKET files into dictionaries for analysis. The same number (11,855) of α and d positions, and their corresponding hole residues, were extracted from 1,275 biological units. Scripts were written in Python and are available online as a Jupyter Notebook session (https://github.com/Hoecker-Lab/Designed_apDimers).

Peptide synthesis. Standard Fmoc solid-phase peptide synthesis was performed on a 0.1-mM scale using CEM Liberty Blue automated peptide synthesis apparatus with inline UV monitoring. Activation was achieved with N,N' -diisopropylcarbodiimide (DIC)/6-chloro-1-hydroxy-1H-benzotriazole (Cl-HOBt). Fmoc deprotection was performed with 20% vol/vol morpholine/ N,N -dimethylformamide (DMF) with addition of 5% formic acid to prevent aspartimide formation. Double couplings were used for β -branched residues and the subsequent amino acid. All peptides were synthesised from C to N terminus as the C-terminal amide on Rink amide resin or ChemMatrix solid support. N-terminally acetylated peptides were produced by addition of acetic anhydride (0.25 ml) and pyridine (0.3 ml) in DMF (5 ml) for 30 min at room temperature. Fluorescently labelled peptides were produced by addition of TAMRA (0.1 mM, 2 eq.), 1-[bis(dimethylamino)methylene]-1H-1,2,3-triazolo[4,5-b]pyridinium 3-oxide hexafluorophosphate (0.095 mM, 1.9 eq.) and N,N -diisopropylethylamine (0.225 mM, 4.5 eq.) in DMF (3 ml) to DMF-washed peptide resin (0.05 mM) with agitation for 3 h. Resin was washed with 20% piperidine in DMF (5 ml) for 2×30 min to remove any excess dye. All manipulations were carried out under foil to exclude light. Biotinylated peptides were produced by addition of Cl-HOBt (1 ml, 0.5 M) to a solution of biotin (0.25 mM, 5 eq.) and DIC (1 ml, 1 M) in DMF and the mixture added to DMF-washed peptide resin with agitation at room temperature for 2 h. Peptides were cleaved from the solid support by addition of trifluoroacetic acid (9.5 ml), triisopropylsilane (0.25 ml) and water (0.25 ml) for 3 h with shaking at room temperature. The cleavage solution was reduced to approximately 1 ml under a flow of nitrogen. Crude peptide was precipitated upon addition of ice-cold diethyl ether (40 ml) and recovered via centrifugation. The resulting precipitant was dissolved in 1:1 acetonitrile and water (approximately 15 ml) and lyophilised to yield crude peptide as a solid.

Peptide purification. Peptides were purified by reverse-phase high-performance liquid chromatography (HPLC) on a Phenomenex Luna C18 stationary phase column (150 \times 10 mm, 5- μ m particle size, 100- \AA pore size) using a preparative JASCO HPLC system. A linear gradient of 20–80% acetonitrile and water (with 0.1% trifluoroacetic acid) was typically applied over 30 min with gradient adjusted for each peptide to optimise separation. For acidic peptides, a linear gradient of 10–60% acetonitrile, and ammonium bicarbonate (25 mM) in water was typically applied over 30 min with gradient adjusted for each peptide to optimise separation. Where peptides were insoluble in HPLC solvents, dimethyl sulfoxide was added to improve solubility. Chromatograms were monitored at wavelengths of 220 and 280 nm. The identities of the peptides were confirmed using matrix-assisted laser desorption/ionisation–time-of-flight (MALDI–TOF) mass spectrometry using a Bruker ultrafleXtreme II instrument in reflector mode. Peptides were spotted on a ground-steel target plate using dihydroxybenzoic acid or α -cyano-4-hydroxycinnamic acid as the matrix. Masses quoted are for the monoisotopic mass as the singly protonated species. Masses were measured to 0.1% accuracy. Peptide purities were determined using a JASCO analytical HPLC system, fitted with a reverse-phase Kinetex C18 analytical column (100 \times 4.6 mm, 5- μ m particle size, 100- \AA pore size). Fractions containing pure peptide were pooled and lyophilised. Peptide stocks were dissolved in water and their concentrations determined by UV–Vis at 280 nm on a ThermoScientific Nanodrop 2000 spectrophotometer using the known extinction coefficients for tyrosine ($\epsilon_{280} = 1,280 \text{ M}^{-1} \text{ cm}^{-1}$) and tryptophan ($\epsilon_{280} = 5,690 \text{ M}^{-1} \text{ cm}^{-1}$). The total extinction coefficients for a sequence were calculated by adding the extinction coefficient for each chromophore. For TAMRA-coupled peptides concentration was calculated by measurement of UV absorbance at 555 nm ($\epsilon_{555}(\text{TAMRA}) = 85,000 \text{ mol}^{-1} \text{ cm}^{-1}$) in 50 mM sodium phosphate (pH 8).

Circular-dichroism spectroscopy. CD data were collected on a JASCO J-810 or J-815 spectropolarimeter fitted with a Peltier temperature controller (Jasco UK). Peptide samples were dissolved at 100 μ M concentration for individual peptides or 50 μ M for each peptide in mixed systems, in PBS (8.2 mM sodium phosphate, 1.8 mM potassium phosphate, 137 mM sodium chloride, 2.7 mM potassium chloride at pH 7.4). CD spectra were recorded in 1-mm path length quartz cuvettes at 20 $^{\circ}\text{C}$. The instruments were set with a scan rate of 100 nm min^{-1} , a 1-nm interval, a 1-nm bandwidth and a 1-s response time and scans are an average of 8 scans recorded for the same sample. Thermal-denaturation data were acquired

at 222 nm between 5 $^{\circ}\text{C}$ and 95 $^{\circ}\text{C}$, with settings as above and a ramping rate of 40 $^{\circ}\text{C hr}^{-1}$ and are a single recording. Baselines recorded using the same buffer, cuvette and parameters were subtracted from each dataset. The spectra were converted from ellipticities (deg) to mean residue ellipticities (MRE; $\text{deg.cm}^2.\text{dmol}^{-1}.\text{res}^{-1}$) by normalising for concentration of peptide bonds and the cell path length using the equation:

$$\text{MRE} \left(\text{deg.cm}^2.\text{dmol}^{-1}.\text{res}^{-1} \right) = \frac{\theta}{c \times l \times b}$$

Where the variable θ is the measured difference in absorbed circularly polarised light in millidegrees, c is the millimolar concentration of the specimen, l is the path-length of the cuvette in centimetres and b is the number of amide bonds in the polypeptide, for which the N-terminal acetyl bond was included but not the C-terminal amide.

Sedimentation-equilibrium analytical ultracentrifugation. Analytical ultracentrifugation (AUC) sedimentation-equilibrium experiments were conducted at 20 $^{\circ}\text{C}$ in a Beckman Optima XL-I or XL-A analytical ultracentrifuge using an An-60 Ti rotor (Beckmann Coulter). Solutions were made up in PBS at 100 μ M total peptide concentration. The experiments were run in two-channel centrepiece. The samples were centrifuged at speeds in the range 44–60 krpm and scans at each recorded speed were duplicated. Data were fitted to single, ideal species models using SEDFIT (v15.2b)/SEDPHAT, comprising a minimum of four speeds. 95% confidence limits were obtained via Monte Carlo analysis of the obtained fits.

Affinity measurements. Affinities were estimated by plotting the midpoint of the thermal unfolding transition (T_M) against the total concentration of peptide. Thermal-denaturation curves were measured between 5 and 95 $^{\circ}\text{C}$, at total peptide concentrations between 15 and 100 μ M, and the T_M extracted from the second derivative of the spectra. All measurements were performed in PBS (8.2 mM sodium phosphate dibasic, 1.8 mM potassium phosphate monobasic, 137 mM sodium chloride, 2.4 mM potassium chloride, pH 7.4).

X-ray crystallography. Sitting-drop vapour-diffusion experiments were set up in three-well IntelliPlates (Art Robbins) using a Phoenix pipetting robot (Art Robbins). Drops for the homomers were pipetted in a 3:2 ratio (peptide:reservoir solution) and the heteromers were pipetted in a 1:1 ratio. Crystals were obtained at 20 $^{\circ}\text{C}$ directly from a sparse matrix screen using the commercially available JCSG Core I-IV screens (Qiagen). Hit conditions can be found in the Supplementary Table 4. Crystals were mounted using CryoLoops (Hampton Research), transferred into a cryogenic solution made of the corresponding reservoir solution supplemented with 25% glycerol, and flash-cooled in liquid nitrogen. Diffraction data were collection at $-173 \text{ }^{\circ}\text{C}$ in 0.1 $^{\circ}$ slices on beamlines BL 14.1 and BL 14.2 at the BESSY II electron storage ring operated by the Helmholtz-Zentrum Berlin. Data were processed using DIALS (2.0.2)⁶⁸ or XDSAPP2 (2.0)⁶⁹ with XDS (Build 20200417). Resolution cut-off was selected according to the Pearson's correlation coefficient ($CC_{1/2}$) greater than 0.30, signal-to-noise ratio (I/σ) greater than 0.5, and a completeness of more than 75% in the highest resolution shell. The data were phased via molecular replacement with Phenix.phaser (2.8.3)⁷⁰ using a model poly-alanine α -helix without accounting for translational non-crystallographic symmetry. Model building as well as manual and automated refinements were done iteratively using Coot (0.9.6)⁷¹ and either Phenix (1.19.2_4158).refine⁷² or Refmac5 (ref. 73).

Fluorescence-quenching assay. Fluorescence-quenching experiments were performed following previously published protocols⁴⁶. In brief, using a Jasco Fluorimeter, peptides (100 μ M each) were excited at 240 nm and emission spectra recorded between 260 and 400 nm. All measurements were recorded in 50 mM sodium phosphate, pH 7, and are the average of five individual spectra.

Biomolecular fluorescence complementation assay. Genestrings (Eurofins Genomics) encoding individual CCs were cloned using standard restriction enzyme–ligation cloning. Both V1 (N-Venus) and V2 (C-Venus) constructs^{48,49} were amplified by PCR from the entire protein sequence and then cloned into arabinose-inducible expression cassettes in the medium copy plasmid backbones pVRC₃ and pBAD, respectively. Site-directed mutagenesis was carried out following the QuikChange II Protocol (Agilent). Full protein sequences for all constructs are given in Supplementary Table 2.

To monitor the interaction of the CCs in cells, TB28 cells (MG1655 Δ LacIZYA)⁷⁴ were transformed with the required plasmid combination as indicated. Colonies were picked in triplicate at least and overnight cultures were grown at 37 $^{\circ}\text{C}$ in M9 minimal medium with 0.25% glycerol, 0.5 mM CaCl₂, 2 mM MgSO₄, 2 μ g ml⁻¹ thiamine and 0.2% casamino acids (with 100 μ g ml⁻¹ ampicillin and 25 μ g ml⁻¹ chloramphenicol where required). The cultures were then used to inoculate 10-ml⁻¹ cultures of the same medium with the addition of 0.2% arabinose, and then were grown at 37 $^{\circ}\text{C}$ until reaching an OD₆₀₀ \approx 0.5. Five-ml samples were centrifuged at 5,000 rpm for 15 min, the supernatant removed and then resuspended in 250 μ l of PBS. Additional 1-ml samples were centrifuged at 13,000 rpm for 10 min, the supernatant removed and frozen for western

blotting at a later stage. Duplicate 100- μ l samples of each culture were placed in a black-bottomed 96-well plate, and the fluorescence read using a CLARIOstar microplate reader (BMG Labtech), using the yellow fluorescent protein (YFP) with the following settings: 497 nm excitation (15 nm bandwidth), 540 nm emission (20 nm bandwidth), 517.2 nm dichroic. YFP fluorescence (relative fluorescence units) was normalised by dividing by the OD₆₀₀.

Western blotting. Pellets were resuspended in appropriate volumes of sodium dodecyl sulfate loading buffer and dithiothreitol, boiled at 95 °C for 3 min, then 10 μ l loaded, along with 10 μ l PageRuler Plus Prestained Protein Ladder (Thermo), onto 8–15% Mini-PROTEAN Gels (Bio-rad) and run at 160 V until the loading dye reached the bottom of the gel. Proteins were then transferred onto a Immobilon-P membrane (Merck) in standard transfer buffer (1 \times Tris–glycine, 10% methanol) for 2 h at 230 mA. After this, the membranes were washed in PBS and blocked overnight in 10% milk in PBS with gentle rocking. The blots were then washed in PBS, and then incubated with a His primary antibody (Abcam) (1:5,000) in 10% milk in PBS for 2 h at room temperature with gentle rocking. After washing, the horseradish-peroxidase-conjugated secondary antibody was added (Abcam) (1:5,000) in 10% milk in PBS and incubated for 1 h. After washing, 2 ml Amersham ECL detection reagent (GE) was added to each blot and incubated for 1 min, and then placed in a cassette along with standard X-ray film for the desired interval before being placed in a developer.

Cell culture. HeLa cells were maintained in high glucose Dulbecco's modified Eagle's medium (Gibco Invitrogen) with 10% (vol/vol) fetal calf serum (Sigma-Aldrich) and 5% penicillin/streptomycin (PAA) (herein referred to as DMEM) at 37 °C and 5% CO₂. For transfection, cells were seeded in 6-well plates on fibronectin coated 13-mm coverslips at a density of 1 \times 10⁵ cells per well and incubated at 37 °C and 5% CO₂ for 16 h before transfection. Cells were transfected with 0.4 μ g DNA using Effectene transfection reagent according to the manufacturer's instructions (Qiagen). After transfection cells were incubated at 37 °C, 5% CO₂ for 16 h. Peptide treatments were in 1 ml DMEM 37 °C and 5% CO₂ for the time indicated. Cells were fixed by addition of 4% paraformaldehyde in PBS at room temperature for 10 minutes (2 ml per well for 6-well plate) and washed three times with PBS. Confocal images were collected using a Leica SP5II system with a 63 \times objective running Leica LAS X and are presented as maximum intensity projections. Figures were assembled using ImageJ in conjunction with Inkscape (0.92). Image analysis was conducted in ImageJ (1.53c) and data was processed using Graphpad Software Prism 8.0.

GFP immunoprecipitation. For GFP immunoprecipitation, HeLa cells were plated at a density of 2 \times 10⁶ per 10-cm dish 4 h before transfection. Cells were incubated on ice and washed in 2 \times 5 ml wash buffer (25 mM HEPES pH 7.4, 150 mM NaCl) followed by 1 ml lysis buffer (25 mM HEPES pH 7.4, 150 mM NaCl, 0.1% NP-40, 0.1% Triton-X 100 and protease inhibitor cocktail) for 10 min. Cells were centrifuged at 13,000g for 10 min at 4 °C and the supernatant incubated with 30 μ l of prewashed (3 \times 800 μ l wash buffer, 2,000 rpm) GFPTrap beads for 120 min at 4 °C. Fifty microliters of supernatant was retained for analysis of cell lysate. Beads were centrifuged at 2,000g for 2 min, the supernatant removed and washed (3 \times 800 μ l wash buffer, 2,000 rpm). Samples were analysed for GFP expression by gel electrophoresis and western blotting. Elution from beads was achieved by addition of 1% sodium dodecyl sulfate at 95 °C for 10 min. TAMRA fluorescence intensity measurements of eluents were performed at 555 nm on a CLARIOstar (BMG Labtech) microplate reader at room temperature.

Data processing and plotting. Unless otherwise stated, all other data were processed and plotted in Python (3.8.5) using Numpy (1.19.2) and matplotlib (3.1.3).

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The CC database CC+ is open and publicly accessible. The coordinate and structure factor files for homomer-S, apCC-Di, apCC-Di-AB_var & apCC-Di-AB have been deposited in the Protein Data Bank with accession codes 7Q1Q, 7Q1R, 7Q1S and 7Q1T, respectively. All the raw data used in this publication has been deposited in the Zenodo repository (<https://doi.org/10.5281/zenodo.6519961>).

Code availability

The scripts used for bioinformatic analysis are available from a Zenodo repository (<https://doi.org/10.5281/zenodo.6518524>).

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Author contributions

G.G.R., J.A.C. and W.M.D. contributed equally to the project. G.G.R., W.M.D., B.H. and D.N.W. conceived and developed the project idea. G.G.R. conducted the bioinformatic analysis and designed the peptides. W.M.D. and G.G.R. synthesised and purified the peptides and conducted the biophysical analysis. W.M.D. conducted and analysed the fluorescence-quenching assays. H.F.T. and N.J.S. conducted and analysed the BiFC assay. S.S. and G.G.R. conducted the X-ray crystallography. J.A.C. and M.P.D. conducted and analysed all mammalian cell experiments. G.G.R., B.H. and D.N.W. wrote the manuscript with contributions from all authors.

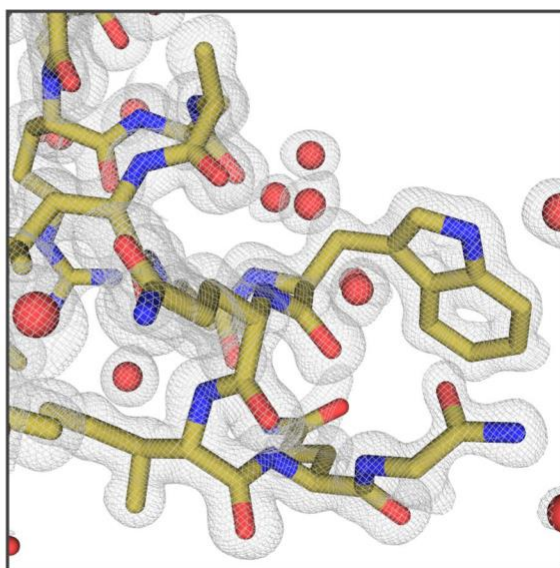
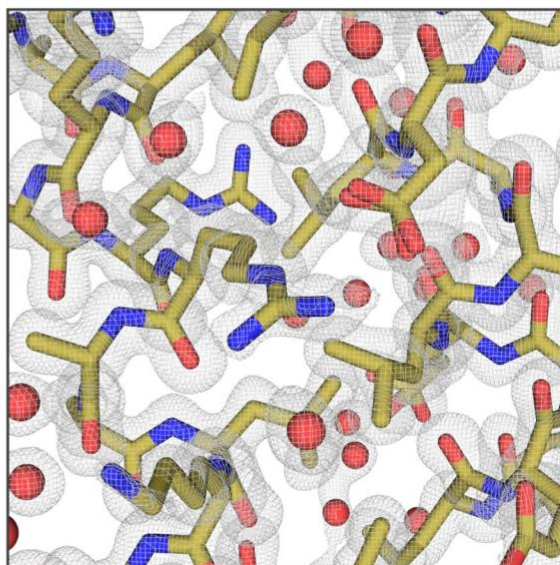
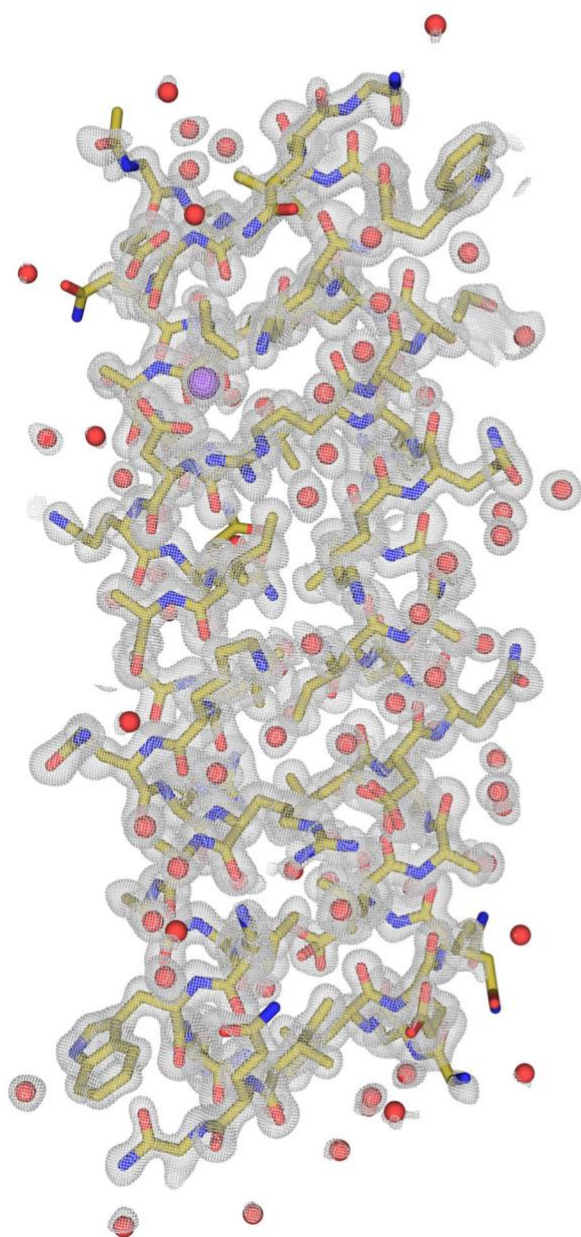
Competing interests

The authors declare no competing interests.

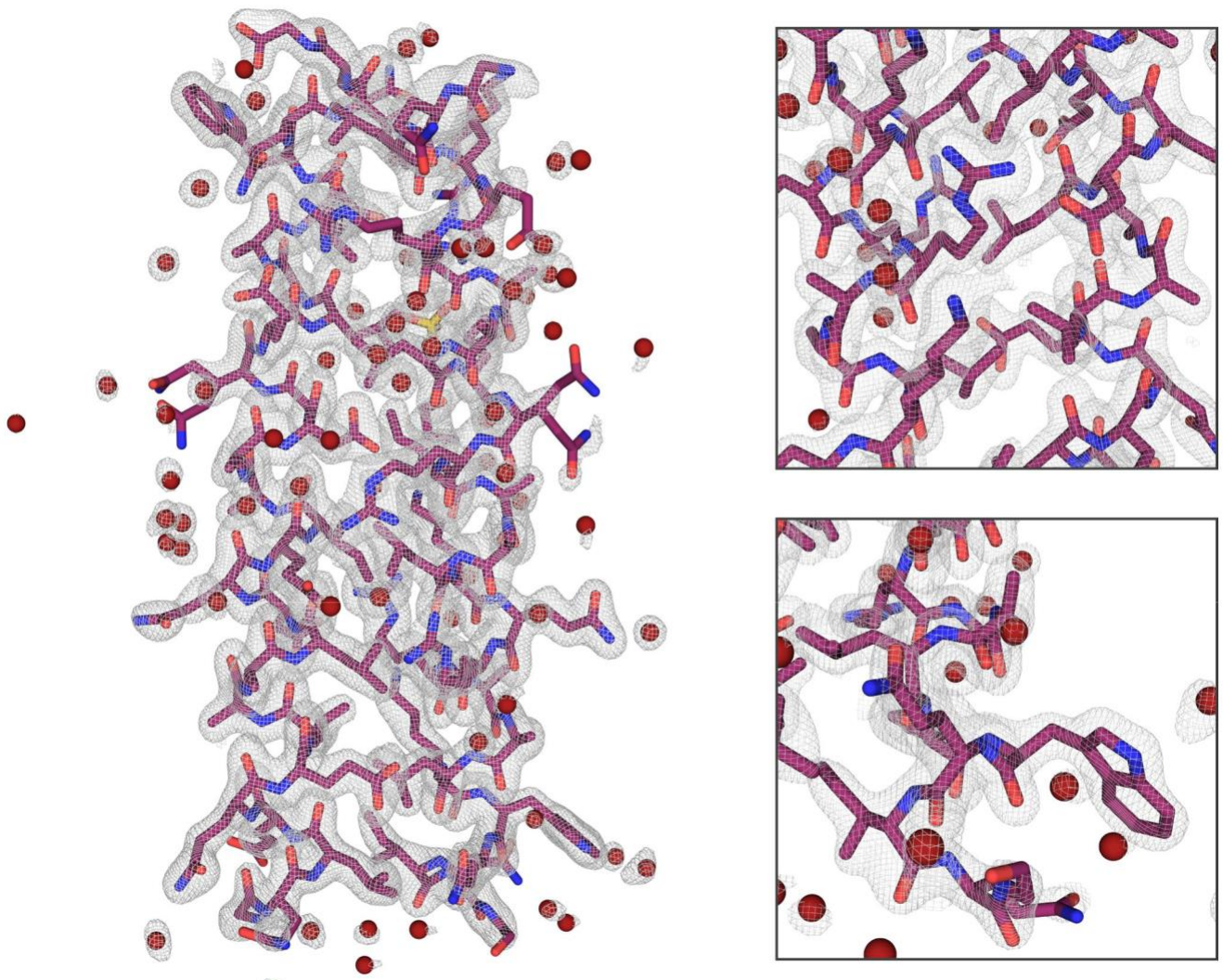
Additional information

Correspondence and requests for materials should be addressed to Birte Höcker or Derek N. Woolfson.

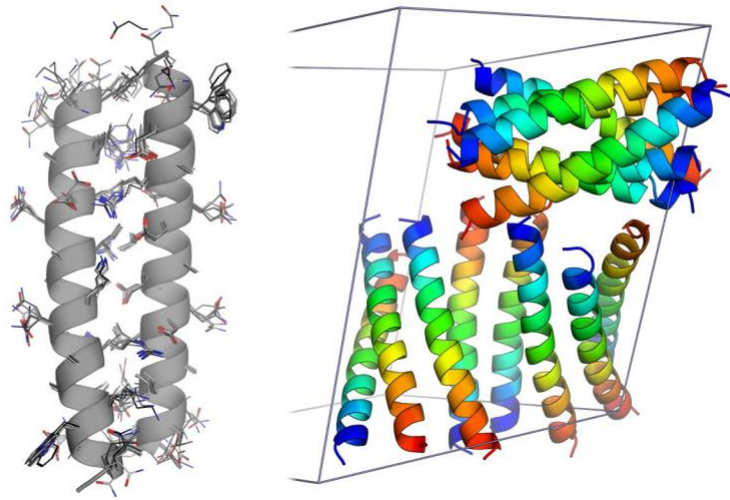
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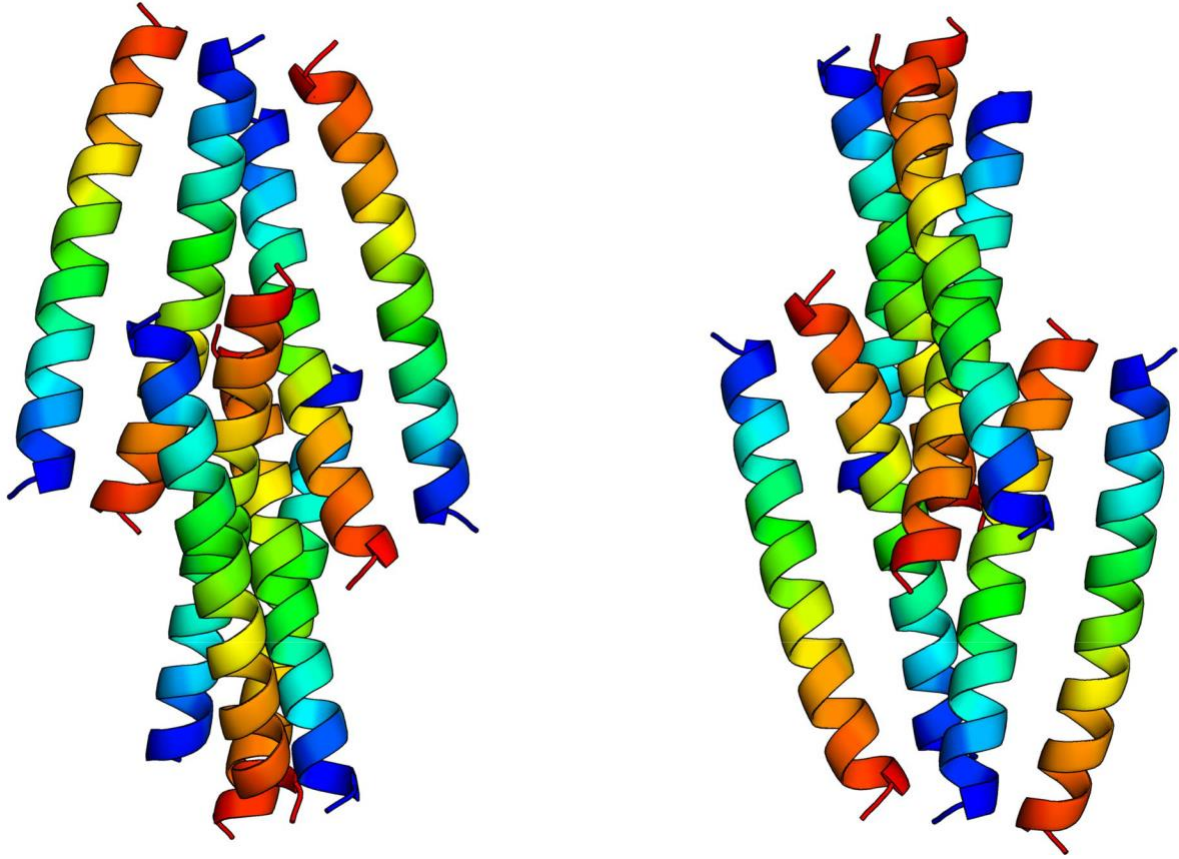
Extended Data Fig. 1 | Electron density map of apCC-Di crystal structure. Fo-Fc difference map of the apCC-Di crystal structure represented as a grey wire mesh overlaid on a stick model. The map was contoured at 1 rmsd.



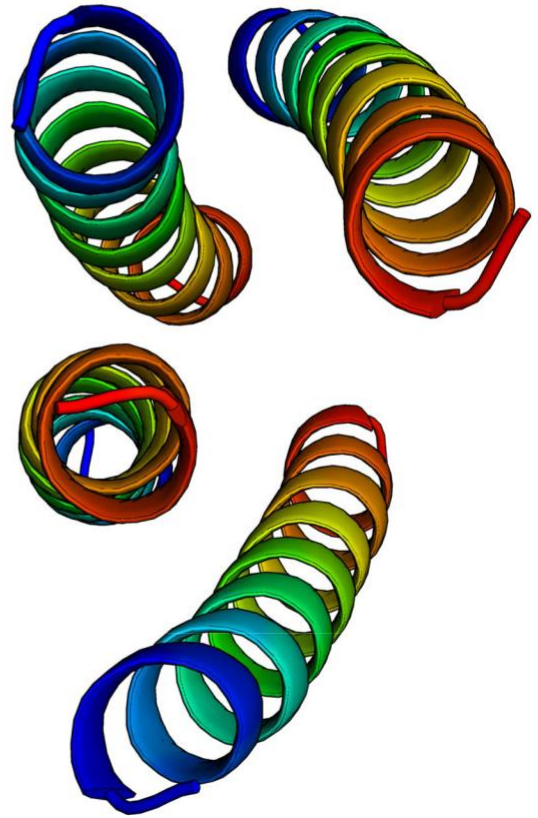
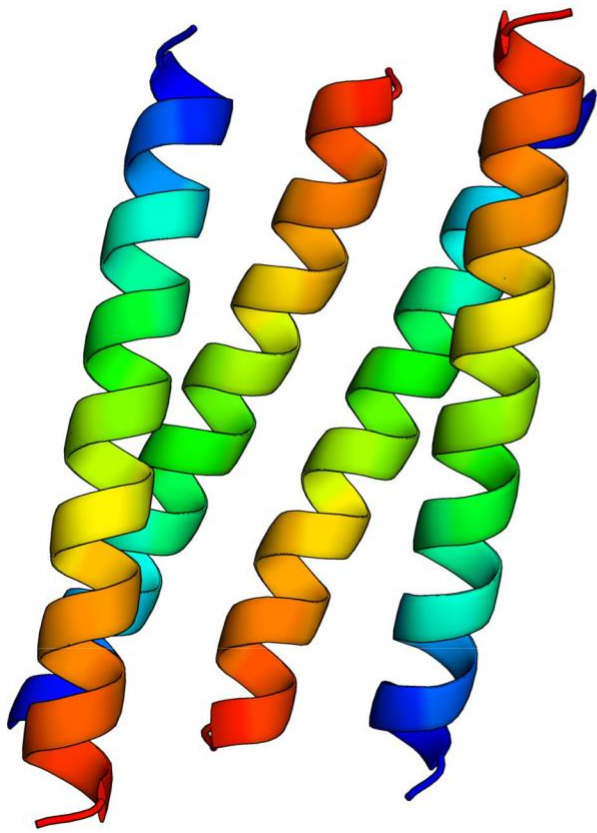
Extended Data Fig. 2 | Electron density map of apCC-Di-AB crystal structure. Fo-Fc difference map of the apCC-Di-AB crystal structure represented as a grey wire mesh overlaid on a stick model. The map was contoured at 1 rmsd.





Extended Data Fig. 3 | The asymmetric unit of apCC-Di-AB_var is composed of 14 chains. While there are minor differences between the surface-exposed residues of the seven dimer pairs, all the biological units are structurally similar (left). The high number of biological units in the asymmetric unit is a result of crystal packing. In the crystal form, apCC-Di-AB_var forms perpendicular layers of right-handed α -helical fibres (right). Between each dimer pair, the individual peptides self-associate to form KIH parallel interfaces, leading to alternating parallel and antiparallel interfaces.

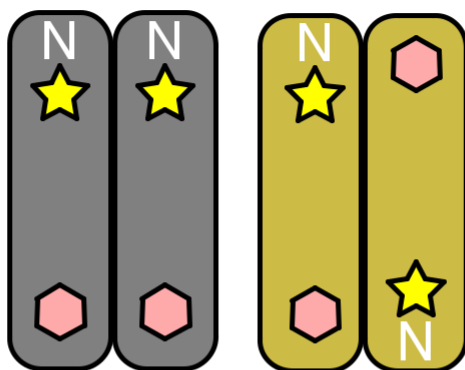


Extended Data Fig. 4 | Symmetry mates for the crystal structure of apCC-Di. Four symmetry mates are depicted above that form a repeating pattern in the crystal structure. SOCKET 2 analysis of the structure reveals four antiparallel CC dimers. There are no inter-biological unit CC interactions.

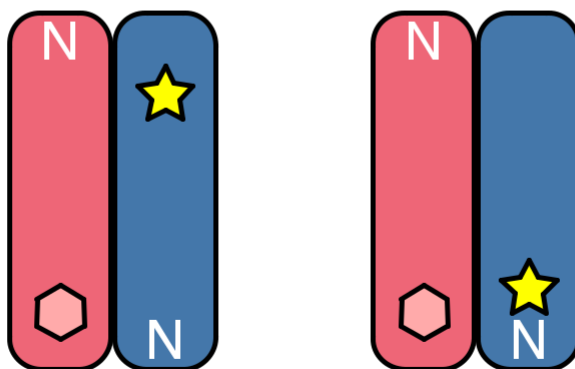


Extended Data Fig. 5 | Symmetry mates for the crystal structure of apCC-Di-AB. Two symmetry mates are depicted above that form a repeating pattern in the crystal structure. SOCKET 2 analysis of the structure reveals two antiparallel CC dimers. There are no inter-biological unit CC interactions.

 Fluorophore (4-cyano-phenylalanine)
  Quencher (Selenomethionine)

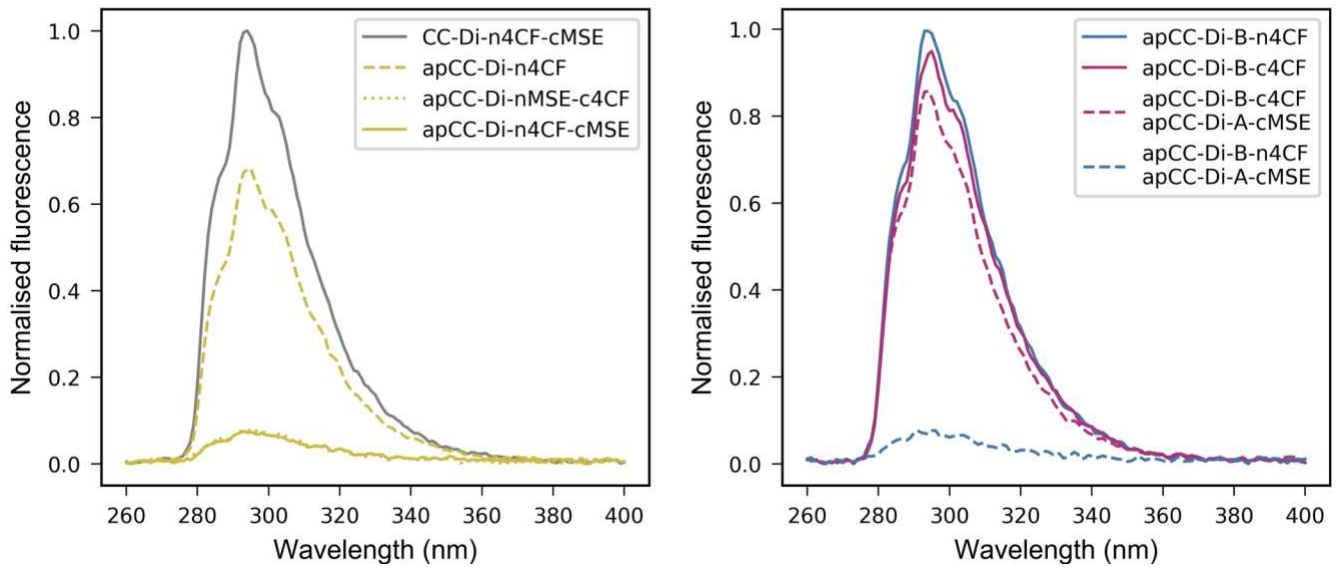


CC-Di-n4CF-cMSE apCC-Di-n4CF-cMSE

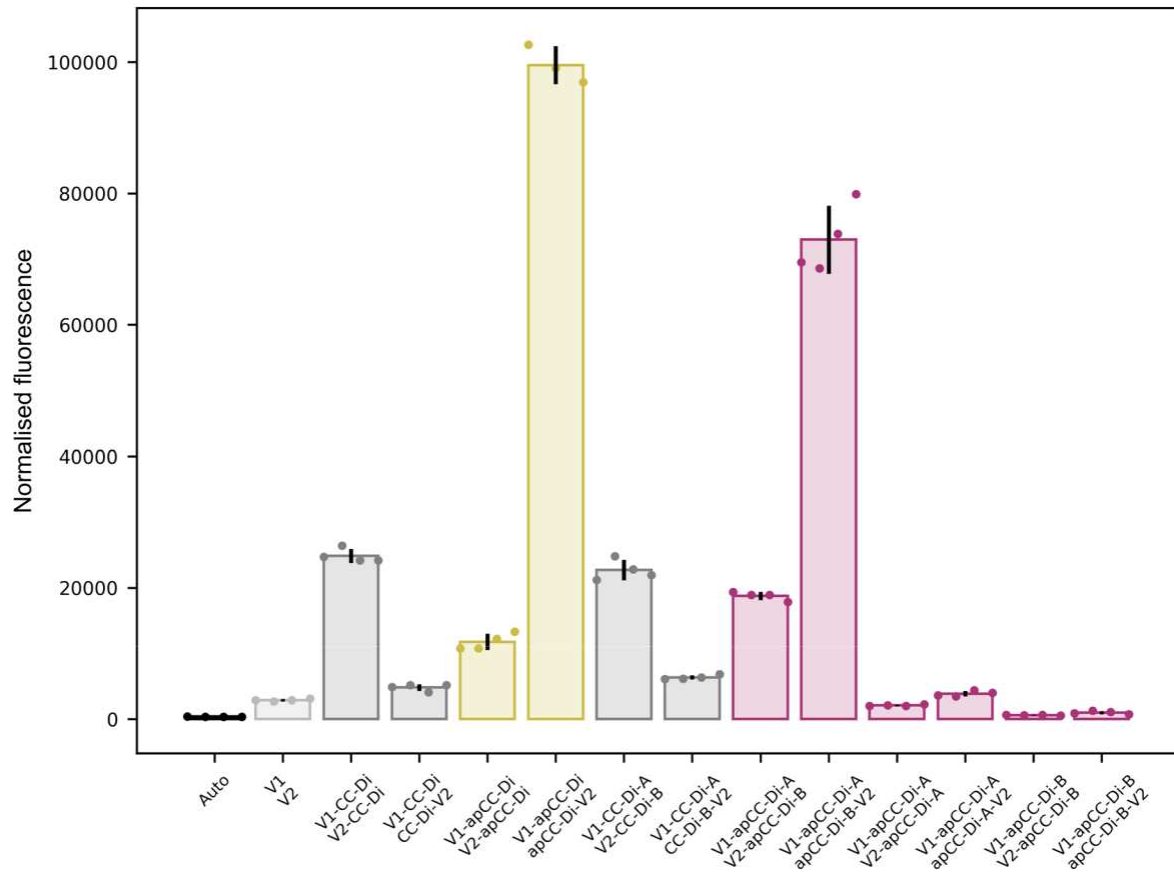


apCC-Di-A-cMSE apCC-Di-A-cMSE
 apCC-Di-B-c4CF apCC-Di-B-n4CF

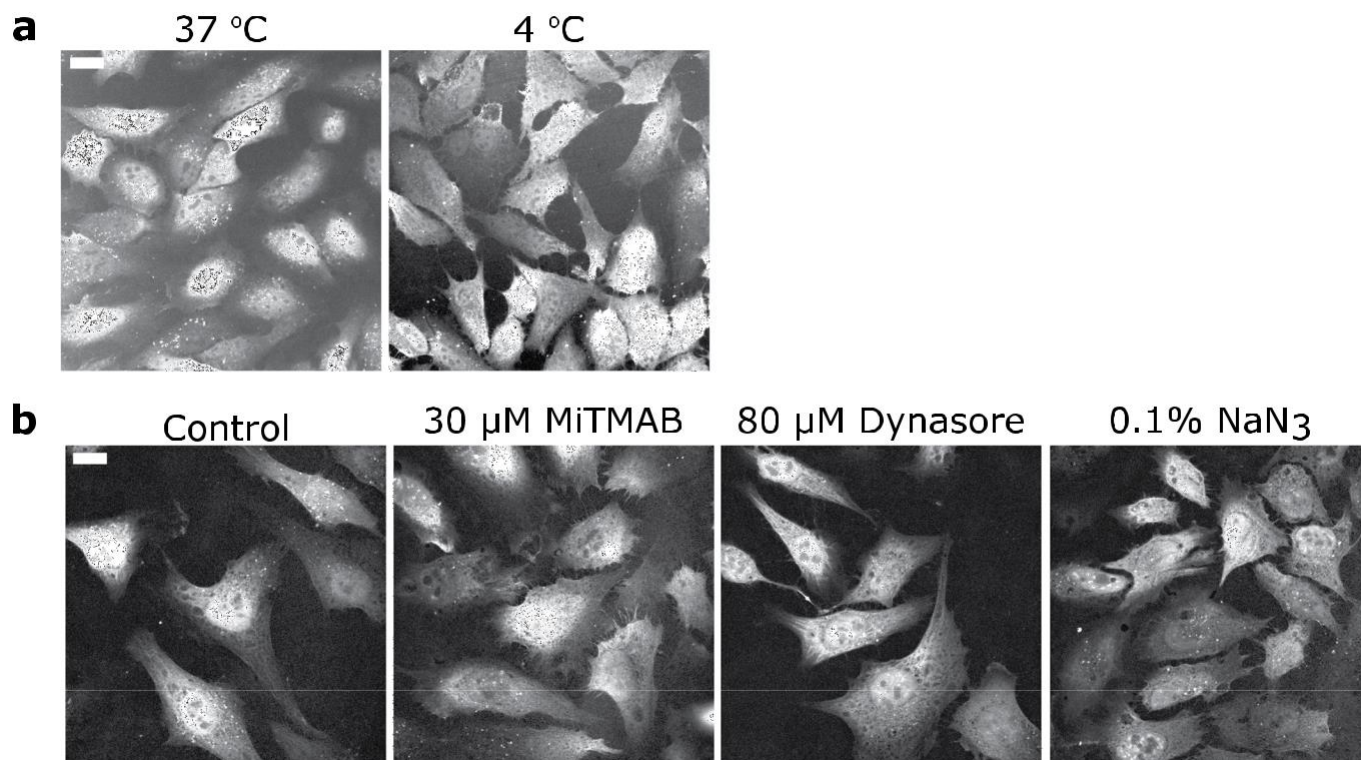
Extended Data Fig. 6 | Cartoon representation of mutation positions in designed coiled-coil (CC) dimers for the fluorescence quenching assay. For homomeric CCs, the fluorophore was incorporated near the N termini and a fluorescence quencher near the C termini. For parallel CCs (top left) this should lead to fluorescence, whereas for antiparallel CCs (top right) this should lead to quenching. For antiparallel heteromeric CCs, incorporating the fluorophore and fluorescence quencher near the same termini (bottom left) should lead to fluorescence, whereas incorporating at opposite termini should lead to quenching (bottom right).



Extended Data Fig. 7 | Extended plots for the fluorescence-quenching assay for labelled apCC-Di, apCC-Di-AB containing additional controls. (left) Fluorescence quenching assay for homomeric peptides. Controls include the parallel coiled coil CC-Di and swapping the fluorophore and the fluorescence quencher between the termini of apCC-Di. (right) Fluorescence quenching assay for the designed heteromeric peptides. The control peptides include labelling both apCC-Di-A and apCC-Di-B near the C-termini, which does not lead to quenching in an antiparallel orientation. Key: n and c indicate mutations near the N and C termini, respectively; 4CF, 4-Cyano-L-phenylalanine fluorophore; MSE, L-selenomethionine fluorescence quencher. Conditions: 100 μ M concentration of each peptide, 50 mM sodium phosphate, pH 7.



Extended Data Fig. 8 | Extended bimolecular fluorescence complementation (BiFC) assay containing controls for homomerisation of apCC-Di-A and apCC-Di-B, and controls for heteromerisation of CC-Di-A and CC-Di-B (a parallel heteromeric system). BiFC assay for apCC-Di and apCC-Di-AB. V1 and V2 are N- and C-terminal fragments of the Venus yellow fluorescent protein, where V1 represents the V1 mutant. Peptide names that precede the Venus fragment name denote fusion to the C termini of the coiled coil, for example apCC-Di-B-V2. Values are normalised according to cell density (OD_{600}) and are presented as mean values ± 1 SD from $n \geq 3$ technical replicate measurements (dots).



Extended Data Fig. 9 | Cell penetration of apCC-Di-B occurs at 4 °C and in the presence of endocytosis inhibitors. **a**, Representative confocal images of HeLa cells treated for 1 h with 2 μM TAMRA-labelled apCC-Di-B at 37 °C and 4 °C. These representative images are taken from $n = 3$ biological replicates. **b**, The same for cells at 37 °C pre-incubated with 0.1% DMSO (as a control) and with endocytosis inhibitors 30 μM MiTMAB, 80 μM Dynasore, and 0.1% NaN₃ in the DMEM before treatment with 2 μM TAMRA labelled apCC-Di-B peptide in media containing each of the inhibitors for 1 h. apCC-Di-B remains cell penetrating at 4 °C and in the presence of the inhibitors and at 37 °C, as shown by TAMRA fluorescence in the cytoplasm and nucleus. Scale bar 10 μm. These representative images are taken from $n = 3$ biological replicates.

