PAC-FragmentDEL – photoactivated covalent capture of DNA-encoded fragments for hit discovery†

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We describe a novel approach for screening fragments against a protein that combines the sensitivity of DNA-encoded library technology with the ability of fragments to explore what will bind. Each of the members of the library consists of a fragment which is linked to a photoactivatable diazirine moiety. Split and pool synthesis combines each fragment with a set of linkers with the version of the library reported here containing some 70k different compounds, each with an individual DNA code. Incubation of the library with a protein sample is followed by photoactivation, washing and subsequent PCR and sequencing which allows the individual fragment hits to be identified. We illustrate how the approach allows successful hit fragment identification using only microgram quantities of material for two targets. PAK4 is a kinase for which conventional fragment screening has generated many advance leads. The as yet undrugged target, 2-epimerase, presents a more challenging active site for identification of hit compounds. In both cases, PAC-FragmentDEL identified fragments validated as hits by ligand-observed NMR measurements and crystal structure determination of off-DNA sample binding to the proteins.

Introduction

The screening of small compounds (typically <16 heavy atoms) or fragments is now established as an effective approach for identification of hit compounds in drug discovery.1,2 Fragment hits have been developed into lead compounds3 and clinical candidates4 for many therapeutic targets, with 6 approved fragment-derived drugs now treating patients. There are two essential features of the approach: (1) low molecular weight compounds are sufficiently complex to make detectable interactions with a portion of a binding site but unlike larger compounds (as screened in high throughput screens) are not sterically prevented from binding5 and (2) most screens generate a number of fragment hits, providing choice for identifying suitable hit and lead compounds. Fragments that bind with affinities ($K_D$) as low as 5 mM have successfully been optimised to clinical candidates, and such weak binding is usually detected in a screen against a target using a biophysical approach such as surface-plasmon resonance (SPR), ligand-observed (LO-) or protein-observed (PO-) NMR, thermal shift analysis or X-ray crystallography.6

Fragments bind to chemically attractive sites on a protein. A recent development for chemical biology is the use of photoactivated fragments to identify and characterise such sites. This was initially demonstrated by the Cravatt group7 where a small number (14) of fragments were each prepared linked to diazirine and an alkyne handle (Fig. 1(a)). The
fragments were incubated with cells, photoactivated and click chemistry used to isolate the proteins which had been covalently labelled. GSK extended the idea with an approach they termed PhaBits, where a larger number (567) of fragments were similarly linked to diazirine and alkyne and incubated with an isolated protein. In both cases, identification of the protein or binding site used mass spectrometry, which constrains the number of fragments that can be screened and the amount of protein target required.

DNA-encoded library (DEL) technology emerged 30 years ago and has been developed as an effective method for hit identification, where successive synthesis using split and pool combinatorial methods builds a library of many millions (billions) of molecules, each of which is labelled with a specific DNA tag. The usual process is to incubate a DEL with the target that can be immobilised onto a bead, then separate the target, wash to remove low affinity or non-binding compounds and then use PCR and sequencing to identify the hits. Provided the DEL hits are retained on the protein during the wash process, the approach provides great sensitivity, allowing many millions of molecules to be screened rapidly against only small amounts of protein material. In this way, DEL screening has successfully identified hit compounds that have been progressed to a number of clinical candidates. The stringency of the wash process in DEL screening typically requires a KD for binding of compound to target of less than tens of μM, so non-covalent DELs are unlikely to find many single fragment hits for most targets. There have been reports of DELs constructed with a fragment on each strand of DNA to exploit avidity where the two fragments bind in adjacent sites or for a dynamic combinatorial approach where fragments react with each other on the surface of the protein. These approaches enrich ligands on specific targets by either crosslinking of DELs on a target or affinity enhancement.

Here we describe and demonstrate PAC-FragmentDEL, an approach where photoactivated covalent capture of DNA-encoded fragments can identify hits binding to a target. There have been reports of DELs of non-fragment sized molecules incorporating an electrophile, where forming a covalent bond with a protein requires reaction with an available cysteine side chain. In contrast, photoactivation generates a carbene which rapidly inserts into nearby bonds. The PAC-FragmentDEL approach is demonstrated with two examples. The serine/threonine protein kinase PAK4 (p21 activated kinase 4) was chosen as a positive control, where a larger number (567) of fragments were similarly linked to diazirine and alkyne and incubated with an isolated protein. In both cases, identification of the protein or binding site used mass spectrometry, which constrains the number of fragments that can be screened and the amount of protein target required.

Results and discussion

Library design and synthesis

Each molecule in a PAC-FragmentDEL library is composed of three parts: DNA encoding system, linker, and fragment (Fig. 1(b)), where the DNA encoding system records the chemical matter installed at a specific step. The linker consists of a fixed linker containing the diazirine and an extension of variable length of 0 to 13 atoms between diazirine and fragment to probe different depths of the binding site. The work reported here used two types of library as summarised in Fig. 2. Type 1 consists of sub-library 1, made from 2634 amines and sub-library 2 from 4783 fragments containing amine capping reagents (acids, aldehydes, sulfonyl chlorides), where such functional groups are required to be compatible with DEL chemistry. The Type 2 library consists of sub-libraries 3 and 4, where DEL chemistry was used to merge two smaller fragments, but without an extension linker.

The same headpiece-conjugated diazirine linker is present in all library members. The library was synthesized in a “split & pool” fashion in two cycles. The synthesis of the Type 1 sub-libraries is outlined in Scheme S1. For sub-library 1, the conjugated diazirine linker was split into 6 wells, each ligated to a unique code, followed by installation of the amino ester extension linker shown in Table 1. Upon completion of the reaction, as monitored by LCMS, the 6 wells were pooled and purified; hydrolysis of ester to free carboxylic acid afforded the cycle 1 mixture. This cycle 1 mixture was split into 2636 wells (2634 fragments and 2 nulls) and each was encoded with unique code 2, followed by installation of a fragment by amide condensation. The final step was ligation with the library ID DNA and purification. Sub-library 2 was synthesised in a similar way, with reactions differing according to the nature of the 5 extension linkers (Table 1) and electrophilic handles on the fragments. This resulted in 15 804 members of sub-library 1 and 23 905 members of sub-library 2.

![Fig. 2. Design of the PAC-FragmentDEL libraries. The same DNA and photoactivation group is present in all sub-libraries. The DNA encodes the library and the groups added in cycles 1 and 2.](image-url)
The Markush structure of the Type 2 sub-libraries 3 and 4 is also shown in Fig. 2 which were synthesised as outlined in Scheme S2† in a similar fashion to the Type 1 library but without an extension linker. Fig. S1† is a summary of the properties of the fragment component of the compounds in the two types of library. Although these properties will be affected by the DNA code, diazirine and linker, the main recognition moiety of the Type 1 libraries is well within the usual property space of most fragment libraries. Most Type 1 fragments contain less than 18 heavy atoms, a LogP less than 2, less than 3 hydrogen bond acceptors and 2 hydrogen bond donors. The two cycles of synthesis of Type 2 sub-libraries mostly generates fragments with between 18 to 26 heavy atoms, MW of 260 to 350, slightly lower a LogP and small increase in the number of hydrogen bonding groups. Although this larger molecular weight means the coverage of potential chemical space is not as great for these Type 2 compounds compared to the smaller Type 1 library compounds,22 the larger compounds may be more suitable for screening against binding sites which will bind with lower ligand efficiency.23 For many of the compounds in the Type 2 library, the cycle 1 fragment forms a structured linker, positioning the cycle 2 fragment for binding. This construction of the overall PAC-FragmentDEL library means that the same fragment is present a number of times with different extended linkers in Type 1 sub-libraries and in some cases is also present in the Type 2 sub-libraries.

**Performing a PAC-FragmentDEL screen**

The process for performing a screen is outlined schematically in Fig. 3. The combined Type 1 and Type 2 PAC-FragmentDEL (notionally 10⁷ copy number per individual DEL molecule, estimated by qPCR) is incubated with 250 pmol of the his-tagged protein for 1 h at room temperature and then exposed to UV light of 365 nm for 10 minutes on ice. The sample is then incubated with Ni-magnetic beads and washed 10 times with buffer to remove non-binding PAC-FragmentDEL members and then heat washed for 10 minutes with buffer at 75 °C and once at 95 °C to denature the protein and remove non-covalent binders. An equivalent sample is taken through the process without exposure to UV light. Two further pairs (with and without UV light exposure) are also prepared similarly; one pair contains 10 μM of a competitor compound and the other pair are control samples with blank beads (i.e. no protein).

The DNA remaining in the prepared samples is then subjected to PCR amplification, sequenced and the number of sequence counts for each fragment in each sample recorded. In brief, the analysis consists of subtraction of signal from the controls to remove fragments that have bound strongly without UV irradiation to either beads or protein or to remove fragments which show higher sequence count in the presence of the competitor (either artefacts or compounds that bind more strongly when the competitor is bound to the protein). The selected fragments are those that show enrichment compared to controls and with enhanced sequence count in the absence of the competitor. Off-DNA samples of the selected fragments (see Fig. S3† for the mapping between fragment and corresponding PAC-FragmentDEL member) are directly available for Type 1 or require synthesis for Type 2 library members.

**Initial proof of concept experiment with PAK4**

An initial set of experiments were performed on the enzyme PAK4 for individual PAC-FragmentDEL compounds encoding fragments known to bind to PAK4: 1 (with null linker) and 2 (with linker 7 from Table 1) where the $K_D$ measured by SPR

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**Table 1** Extension linkers used in construction of Type 1 libraries

| Item | Structure of extension linkers | Used in      |
|------|--------------------------------|--------------|
| 1    | Null                           | Sub-library 1|
| 2    | Sub-library 1                  | Sub-library 1|
| 3    | Sub-library 1                  | Sub-library 1|
| 4    | Sub-library 1                  | Sub-library 1|
| 5    | Sub-library 1                  | Sub-library 1|
| 6    | Sub-library 2                  | Sub-library 2|
| 7    | Sub-library 2                  | Sub-library 2|
| 8    | Sub-library 2                  | Sub-library 2|
| 9    | Sub-library 2                  | Sub-library 2|
| 10   | Sub-library 2                  | Sub-library 2|
| 11   | Sub-library 2                  | Sub-library 2|

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**Fig. 3** The PAC-FragmentDEL process. Typically a total of 3 pairs of samples (+/− UV exposure) are prepared with the library incubated with blank beads, and for the target +/− competitor.


for the isolated fragments was measured as 510 \( \mu \text{M} \) and 300 \( \mu \text{M} \) respectively. The results of incubation with the isolated PAC-FragmentDEL members (Fig. 4) demonstrate significant enrichment of the conjugated fragments binding to bead-immobilised PAK4 compared to the blank bead control when exposed to UV light.

**PAC-FragmentDEL screen for PAK4 and 2-epimerase**

For PAK4, the screen was performed in the presence and absence of 10 \( \mu \text{M} \) of compound 3 which binds to the ATP-binding site of PAK4 with a \( K_D \) measured by SPR of 50 nM. Analysis of the sequence counts obtained for the competitive hits showed that none of these PAC-FragmentDEL compounds bound to the beads (Fig. 5(a)), but some bound weakly to PAK4 in the absence of UV irradiation. In total, 301 fragments were identified that bound competitively, some with different linkers (hence more than 301 dots in the PAK4 plots in Fig. 5). 11 of the fragments were chosen for further validation based on signal strength and structural diversity. The 301 hits contained several different linkers and at least in the case of PAK4, there was no preference for a specific linker.

For 2-epimerase, there is a reported inhibitor (Epimerox\(^{24}\)) which in our hands did not show binding by isothermal titration calorimetry (ITC). We therefore generated compound 4 (manuscript in preparation) which binds to 2-epimerase with a \( K_D \) of 100 nM as measured by ITC and was competitive with UDP-GlcNAc as confirmed by LO-NMR (see Fig. S2†). The PAC-FragmentDEL screen for 2-epimerase was performed in the presence and absence of 4 and as can be seen in Fig. 5(c), the sequence count of the PAC-FragmentDEL hits is low. The sequence count for a particular library member is affected by a number of factors, in particular the success of individual reactions and thus number of compound copies. However, the overall low sequence count for 2-epimerase probably reflects weak affinity for this binding site and only 21 competitive fragments could be identified, 9 of which were selected for experiments to validate binding.

**Hit validation by NMR and X-ray crystallography**

The fragments within the selected hits from the two PAC-FragmentDEL screens were sourced or synthesised. These isolated off-DNA fragments were then validated as binding by measurement of competitive LO-NMR spectra and by determination of crystal structures of off-DNA compounds bound to the protein. All of the hits were confirmed as binding by LO-NMR, crystal structures were obtained for 10 of the 11 PAK4 hits and 5 of the 9 2-epimerase hits. The hits for which data is presented are shown in Fig. 6.

A representative set of spectra for binding to PAK4 is shown for 5 in Fig. 7, where the changes in the NMR spectra in the presence and absence of staurosporine are consistent with binding to the ATP-binding site. Fig. 8 shows detail of the crystal structures of the ATP-binding site of PAK4 with each of 5, 6 and 7 bound. Each of these hits are similar to sub-structures in known kinase hinge binding compounds, but none have been reported previously as binding to PAK4.
As explained earlier, each individual fragment can be present in both Type 1 and Type 2 PAC-FragmentDELs. 5 was a hit several times with different extended linkers in Type 1, 6 was a hit only in Type 2 library and 7 was found from both Type 1 and Type 2 libraries.

Fig. 9(a) shows the crystal structure of 2-epimerase (PDB code: 3BEO) with UDP and UDP-GlcNAc bound. A comparison of the molecular surface of this structure (Fig. 9(b)) with the structure with the hits 8 and 9 (Fig. 9(c) and (d)) emphasises the conformational change to an active form associated with substrate and cofactor binding. The fragment hits both bind in the UDP cofactor site, with the enzyme in an open, deactivated conformation. Fragment 8 was from Type 1 and 9 from Type 2 library.

Perspectives and conclusions

There are several advantages of the PAC-FragmentDEL approach. Only a small amount of tagged protein (250 pmol per sample) is required, and it is relatively rapid to identify hits, with selection from the DEL library and sequencing and analysis taking only a few weeks to perform. The solubility of the fragments that are incorporated into the library can be lower than required for most fragment screening approaches. Typically, biophysical screens (such as NMR or SPR) require aqueous solubility for the fragment in the high 100 s of μM to enable detection of weak binding. Conjugation with DNA removes this solubility requirement, at least for the PAC-FragmentDEL screen, broadening the chemical space of fragments that can be included in the library. Finally, confirmation of hits depends on which library they are from. The fragments that are conjugated in the Type 1 sub-libraries are mostly readily available; for Type 2, synthesis of off-DNA samples is required.

In the examples described here, the selection of PAC-FragmentDEL hits is by comparison of the hits in the presence and absence of a competitor ligand that binds to
the site of interest. If such a competitor is not available (and it could be a peptide or protein partner), then an alternate approach would be to block the active site by mutation(s) and compare the hits with those obtained for the wild-type protein. Similarly, comparative screens of orthologous proteins could be used to identify selective fragments.

There are many interesting features of the PAC-FragmentDEL approach which are yet to be investigated, such as which types of targets and binding sites are appropriate for the approach? The examples described here demonstrate a low false-positive rate for the fragments selected for characterisation, but will this hold for all targets? The false negative rate (which is related to the cut-offs applied in hit analysis) is of less interest – the priority in early hit discovery is to find some chemically tractable hits for progression and as with any screening, it is accepted that compounds will be missed. Another area for analysis is the impact on whether a particular fragment is found to bind of different extension linkers in Type 1 sub-libraries or the coupling of two fragments in Type 2 libraries. We have also not investigated scale up of individual PAC-FragmentDEL hits for incubation with the target protein to characterise by mass spectrometry or crystallography, where the covalent bond is formed on photoactivation. The carbene formed by photoactivation inserts into proximal H–hetearotom and H–C bonds. A limited analysis of the hits in the PhABits study showed covalent bonding to acidic side chains. It is likely that the high temperature washing step in our protocol would hydrolyse at least a proportion of any ester bonds formed but high temperature washing step in our protocol would covalent bonding to acidic side chains. It is likely that the groove is well established as a highly druggable protein kinase class of proteins for which the ATP binding site is masked by a large amount of non-specific attachment from hits that are not affected by competitor.

In conclusion, the PAC-FragmentDEL approach successfully identified fragments that were validated as binding to the two targets presented here. PAK4 is from the protein kinase class of proteins for which the ATP binding groove is well established as a highly druggable site for small molecules to bind. This target was selected as a demonstrator for the approach, with known fragments included in the screen as a positive control. 2-Epimerase has a more open and polar active site which is less ligandable and for which no small molecules have been reported to bind. The hit rate from the PAC-FragmentDEL screen confirmed the different ligandability of the two proteins and selected hits were validated as binding in solution (NMR) and by crystallography.

**Experimental**

**Protein production**

**PAK4.** A gene fragment encoding PAK4(300-591) was amplified using primers with BamHI and XhoI sites and cloned into a pGEX-6P1 vector. The L301A mutation was then generated by site-directed mutagenesis. The protein was expressed in *E. coli* BL21(DE3)pLysS cells by inducing with 1.0 mM IPTG at OD600 = 0.4, grown overnight at 18 °C. The protein was purified on a GSTrap column (Cytiva) and the GST tag cleaved with PreScission protease (Cytiva). The protein was further purified using a 26/600 Superdex 75 pg column (Cytiva) in 50 mM Tris [pH 7.5], 150 mM NaCl, 10 mM DTT.

**2-Epimerase.** A clone codon optimised for expression in *E. coli* for *B. anthracis* UDP-N-acetylgalactosamine 2-epimerase (1-371) was generated by gene synthesis (GenScript HK). Digestion of the gene and subsequent ligation into a modified pET41a vector gave an N-terminal His tag with enterokinase cleavage site. Site-directed mutagenesis (Agilent) was used to generate a N370A mutant. The protein was expressed in *E. coli* BL21(DE3)pLysS cells by inducing with 1.0 mM IPTG at OD600 = 0.7 for 16 hours at 18 °C. The protein was purified using a HisTrap HP column (Cytiva) and the His tag cleaved with enterokinase (NEB). A negative affinity step was followed by further purification using a 16/600 Superdex 75 pg column (Cytiva) in 50 mM Tris pH 7.5, 50 mM NaCl, 10% glycerol, 2 mM DTT.

**Synthesis of PAC-FragmentDEL and fragments that are not commercially available**

See ESI.†

**Screening, sequencing, and decoding**

Selection was carried out using a KingFisher Duo Prime Purification System (ThermoFisher) in a 96 well plate. 2.5 μM PAK4 (or 2-epimerase) was incubated with a mixture of the Type 1 and Type 2 sub-libraries at room temperature for 60 min in 100 μL selection buffer (25 mM HEPES, 150 mM NaCl, 0.01% Triton X-100, 10 mM imidazole, 0.3 mg ml⁻¹ ssDNA, pH 7.4). The protein–PAC-FragmentDEL mixture was incubated on ice and subjected to UV irradiation at 365 nm for 10 min. The control without UV treatment was included in parallel. Protein was immobilized on Ni-Charged Magbeads at RT for 30 min. The weak or non-specific binders were removed by 1 min wash in 500 μL selection buffer 10 times, followed by two 10 min heat washes at 75 °C and one at 95 °C in 50 mM HEPES, 300 mM NaCl, pH 7.4 buffer to remove the non-covalent binders. The DNA in the sample was quantified by qPCR and samples amplified by PCR and sequenced on an Illumina NovaSeq 6000.

The selection output was quantified by qPCR using PowerUp SYBR Green Master Mix (Thermo, A25778) and amplified by PCR using Q5 Hot Start High-Fidelity 2X Master Mix (NEB, M0494L). PCR cycle was determined by qPCR result. The resulting PCR amplicons were purified by using QIAGEN-MinElute PCR Purification Kit (QIAGEN, 28006) and quantified using a Qubit DNA High-Sensitivity Kit (Invitrogen, 32854) before library construction and sequencing. The library preparation was performed by Nextflex Rapid DNA-Seq Kit (BI00 Scientific, 5144-08) following the manufacturer’s operational manual.
Specifically, 10 ng DNA was used as input. DNA library was constructed through end-repair, dA-tailing, adaptor-ligation and 13 cycles PCR amplification followed a final purification by AMPure XP beads (Beckman, A63882). Subsequently, the resulting libraries were denatured and neutralized and subjected to 116 bp single-end sequencing on an Illumina NovaSeq platform ([Illumina, USA] by Chengdu HitGen Pharmaceuticals Inc., China.

After decoding, the DNA sequences enriched in selection were translated into chemical structures and visualized by DataWarrior files (Openmolecules). The enrichment of fragments can be determined by the sequence count which is the copy number of the corresponding fragment’s DNA barcode. The difference between groups was compared by enrichment ratio which is calculated by sequence count in different selection conditions, and background subtraction was performed by removing high signals that appeared in the blank control.

**Ligand-observed NMR (LO-NMR) validation of competitive compound binding**

1D $^1$H, STD$^{29}$ the water ligand observed via gradient spectroscopy (waterLOGSY)$^{30}$ and relaxation-filtered$^{31}$ spectra were acquired at 298 K using excitation sculpting to suppress the solvent peak on a 600 MHz Bruker Avance III spectrometer fitted with a cryoprobe. Samples contained 10 μM protein with either 100 μM or 500 μM compound in 20 mM Tris pH 7.5 (PAK4) or 20 mM Tris pH 7.5, 50 mM NaCl (2-epimerase). Competition was determined by the addition of 10 mM ligand for a period of 16 h at 19°C. The crystals were then flash-frozen in liquid N2.

**ITC to measure $K_d$ for binding of 4 to 2-epimerase**

ITC measurements were performed using an iTC200 instrument (Microcal, GE Healthcare). The feedback mode was ‘low’ with reference power setting of 4 μCals$^{-1}$. The cell was stirred at 1000 rpm and thermostated at 25 °C. All experiments were performed using the dialysis buffer (see below) with 1% DMSO (v/v). Both experiments were conducted with 6 μM ligand in the cell and 60 μM protein in the syringe. The experiments were conducted with 12 injections of 3.05 μL and 240 s spacing. The first ‘waste’ injection of 0.5 μL was discarded in each case. All data were fitted to a one site model using the provided software.

Protein (7.2 mg mL$^{-1}$) was dialysed with stirring for four hours 20 °C, in 25 mM HEPES pH 7.4, 150 mM NaCl. Upon recovery the protein was filtered (0.22 μM spin filter). The protein concentration was determined by UV absorbance spectroscopy at 280 nM using an extinction coefficient of 17 420. The dialysis buffer was degassed, this was then used for subsequent preparation of protein and ligand solutions for the titration experiments. For each experiment the ligand was freshly prepared from a 20 mM DMSO stock solution.

**X-ray crystal structures of protein–ligand complexes**

Apo crystals of PAK4. Human PAK4 L310A mutant protein was concentrated to 15 mg mL$^{-1}$ and submitted to crystallization screening experiments. Initial crystals grew overnight in 0.1 M Tris pH 8.5 and 0.8 M Sodium tartrate using the hanging-drop vapor-diffusion method at 19 °C. Larger crystals were subsequently obtained using streak seeding with seeds created from these initial crystal stocks.

PAK4 protein–ligand complexes. Crystals were prepared by soaking apo crystals in mother liquor containing 10–20 mM ligand for a period of 16 h at 19 °C. Prior to data collection, all crystals were transferred to a cryoprotectant consisting of 0.1 M Tris pH 8.5, 0.8 M sodium tartrate and 20% glycerol. The crystals were then flash-frozen in liquid N$_2$.

Apo crystals of 2-epimerase. *Bacillus anthracis* 2-epimerase protein was concentrated to 15 mg mL$^{-1}$ and submitted to crystallization screening experiments. The best crystals grew in 2–3 days from 200 mM ammonium sulphate, 50 mM Bis-Tris pH 5.6 and 28% PEG 3350 using the hanging-drop vapor-diffusion method at 19 °C. Crystals were subsequently generated overnight using streak seeding with seeds created from these initial crystal stocks.

2-Epimerase protein–ligand complexes. Crystals were prepared by soaking apo crystals in mother liquor containing 10 mM ligand for a period of 16 h at 19 °C. Prior to data collection, the crystals were flash-frozen in liquid N$_2$. No cryoprotectant was needed.
Data collection for both proteins. Diffraction data of the soaked crystals were collected at 100 K on beamline i03 at Diamond Light Source (UK). Data processing was carried out with XDS. Data collection statistics are summarized in ESI† Table S1.

Structure determination for both proteins. Crystallographic data was processed, and structures determined and refined within the CCP4 package using AceDRG to build models of each compound. The structures were determined by molecular replacement using published structures (PDBcode 2BVA for PAK4 and 3BE0 for 2-epimerase) to calculate model phases and subsequently refined using REFMAC5. Interactive graphical model building was carried out with COOT. In all structures the respective ligands were clearly defined by the initial electron density maps. See Table S1† for data collection and refinement statistics.

Data deposition
The refined crystal structures of 5, 6 and 7 bound to PAK4 and of 8 and 9 bound to 2-epimerase are deposited in the PDB with codes 8AHH, 8AHI, 8AHG, 8AHE and 8AHF respectively.

Author contributions
GL, JW, HM, and HL designed the PAC-FragmentDEL approach and wrote the initial draft of the manuscript, CS and CD performed the chemistry of DEL, QC contributed on DEL screening, Jian Liu performed the sequencing, XC and LZ analysed the DEL screening data, PT resynthesized the hits. REH oversaw the initial concept of the application of PAC-FragmentDEL to these targets and wrote the final version of the manuscript. JBM designed the details of the experiments on these targets and made ITC measurements and analysed the data. NM made SPR measurements. LMB conducted all the crystallographic structure determination and RH and HS conducted all NMR experiments. WF, JS and NH produced the protein samples. BAM and Jin Li oversaw the PAC-FragmentDEL design and manuscript writing.

Conflicts of interest
There are no conflicts to declare. All authors are employees of either Vernalis (R&D) Ltd or HitGen who funded the research.

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