Protein-ligand binding measurements using fluorescence polarization

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- Fluorescence polarization assay used to probe protein-ligand interactions
- Binding affinities are consistent with activity assays
- Technique sensitivity allows analysis of a wide range of concentrations

Introduction

Human Lys-specific Histone Demethylase 1 (LSD1) promotes the demethylation of mono- or di-methyl-Lys4 on histone H31: as chromatin remodelling enzyme, LSD1 is found in complex with several partners, one of which being the REST Co-repressor 1 (CoREST1). We used the fluorescence polarization (FP) as a biophysical tool to analyse the binding properties of LSD1-CoREST1 (LC1) hetero-dimer to H3-derived peptides (Fig. 1), that have been conjugated at the C-terminal with the fluorescent label TAMRA (5-Carboxytetramethylrhodamine).

Materials & Methods

- black, 384-well, low flange microplates from Corning, UK
- TAMRA-conjugated peptides provided by NKI Protein Facility (Amsterdam)
- LC1 protein was recombinantly produced, according to reference 1
- Multi-mode microplate reader from BMG LABTECH, Germany

Experimental setup

1 nM labelled peptides were mixed to assay buffer: 15 mM KH2PO4 pH 7.2, 5% glycerol, 1 mg/ml BSA. Mix was then distributed equally across the plate. For each experiment, triplicates were prepared: starting concentration of protein LC1 was 4 μM, and then this was serially diluted 1:1 using the mix in the other wells. For competitive assays, constant enzyme concentration (around Kd) were mixed with 1 nM labelled ligand in the same assay buffer as above. Competitors were then titrated in using serial 1:1 dilutions as described for direct binding assay. The typical concentration range started from 10 μM.

CLARIOstar® instrument settings

Detection Mode: Fluorescence Polarization, Endpoint
No. of flashes: 50
Excitation: 540-20
Emission: 590-20
Target temperature: 25°C

Focus and Gain adjustments for both channels were set on a reference well containing just labelled ligand. It was chosen to set the adjusted polarization to a reference value of 35 mP. Typical adjustment values for gain were between 1600 and 2000.

Results & Discussion

We first devised a direct binding experiment using three different H3-derived peptides, all previously described as substrates and/ or inhibitors of LC1 complex. On a relative scale, results obtained are consistent with activity measured before by the working group. The difference in the peptides consisted in different lengths and/or different amino-acid compositions as listed below:

meH3 (pep01) = H3 N-terminal tail, methylated on Lys4, 21 residues
H3K4M (pep02) = H3 N-terminal tail, baring the mutate residue in position 4, 21 residues

Fig. 1: Assay principle for monitoring protein-ligand interaction using fluorescence polarization.

(A) H3 labeled peptides in solution show a rapid rotation that causes dispersion in different direction of the incident polarized light: the resulting emitted polarization signal is low. [B] Upon binding to LC1, the high-molecular weight complex results in a slow rotation of the fluorescent molecule leading to high polarization values. [C,D] Non-tagged peptides, in orange, compete for LC1 active site. As competitor concentration increases, more freely labelled peptide remains in solution. This results in a low FP value, comparable to results in (A).
SN12 [pep03] = Transcription factor SNAIL, an H3 analogue, consisting of 12 residues (N-terminal)

As shown in figure 2, all three peptides do bind to LC1.

Acquired data was analysed using Mars Software and graphically edited in GraphPad. Association constants were calculated modifying the equation for fluorescent anisotropy [see equation 1] using constant label concentration as constrain.

\[ B = \frac{L_t + K_t + R_t - \sqrt{(L_t + K_t + R_t)^2 - 4L_tR_t}}{2} \]

Eq. 1: Fitting for Fluorescence Anisotropy. \( B \) = binding, \( L_t \) = total ligand concentration, \( R_t \) = total "receptor" concentration.

To further assay the binding of histone tail derivate peptides and to confirm association parameters, we proceeded using a competition approach: for this set of experiments a new test mix was prepared, as described above. We titrated in different potential competitors, using the same enzyme concentration, which depended on \( K_d \) values obtained from direct binding assays. As shown below (Fig. 3), we were able to assess affinity of different potential binders using a different approach but still in fluorescence polarization.

Table 1 enlists all molecules used in both direct and competitive assays with the resulting affinities values [\( K_d \)].

<table>
<thead>
<tr>
<th>Ligand</th>
<th>( K_d )</th>
<th>Type of Assay</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pep01</td>
<td>107.3 nM ± 7.6 nM</td>
<td>Direct Binding</td>
</tr>
<tr>
<td>Pep02</td>
<td>76.4 nM ± 10.4 nM</td>
<td>Direct Binding</td>
</tr>
<tr>
<td>Pep03</td>
<td>81.9 nM ± 16.7 nM</td>
<td>Direct Binding</td>
</tr>
<tr>
<td>Pep A</td>
<td>147.1 nM ± 35.4 nM</td>
<td>Competition</td>
</tr>
<tr>
<td>Pep B</td>
<td>316.6 nM ± 59.9 nM</td>
<td>Competition</td>
</tr>
</tbody>
</table>

Fluorescence Polarization has proven to be a reliable technique to probe protein-ligand interaction in real-time and in solution. Both, direct and competitive assays, are useful tools to analyse binding properties. Next to the CLARIOstar very similar results were also obtained on the PHERAstar\textsuperscript{®} FS microplate reader.