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## Marinopyrrole A Target Elucidation by Acyl Dye Transfer

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While a variety of methods exist,<sup>1</sup> target identification continues to pose a fundamental challenge in elucidating a natural product's mode of action.<sup>2</sup> The development of solutions to this problem is essential to advancing drugs into the discovery pipeline. The complex structural diversity that characterizes natural products complicates the adoption of routine protocols for these studies.<sup>3</sup> A systematic approach to target elucidation would both facilitate the introduction of drugs into clinical trials and challenge leads early on should undesired targets be identified.<sup>4</sup>



**Figure 1.** Application of acyl dye transfer to natural product mode of action studies. (a) A dye (L) labeled natural product (NP) probe 1 binds to target protein 2 forming complex 3. Ligation-directed acyl transfer in complex 3 results in the displacement of phenol 4 and formation of labeled protein 5. (b) Typical workflow associated with antitumor mode of action studies on a natural product. The process begins by verifying the activity of the probe from (a) in live cells. The process continues with steps that apply the probe analysis, and mass spectral based protein identification.

The process of determining the therapeutic mode of action of a natural product typically begins by developing probes bearing an affinity and/or fluorescent tag.<sup>5</sup> The aim in this exercise is to construct a probe with activity comparable to that of its parent natural product. Often a loss in activity can be circumvented by

use of bio-orthogonal approaches<sup>6</sup> (Huisgen cycloaddition reactions or Click chemistry,<sup>7</sup> Diels–Alder cycloaddition,<sup>8</sup> or olefin metathesis<sup>9</sup>), wherein the reporter is added bio-orthogonally after the natural product binds to its target. Though sufficient for cellular studies, the lack of a covalent link between the labeled natural product and its target commonly thwarts enrichment strategies (e.g., immunoprecipitation).

Cross-linking or photo-cross-linking,<sup>10</sup> a common solution to this problem, is frequently plagued by additional complications. First, the preparation of a dual tag results in an enlarged probe,<sup>11</sup> which often exhibits only a fraction of the activity of its parent natural product.<sup>2b</sup> Second, cross-linking methods are generally low yielding and unselective, increasing the complexity of downstream mass spectrometric (MS) analyses.<sup>12</sup> Finally, and perhaps most critically, the cross-linked target still bears a covalently linked natural product that is subject to xenobiotic metabolism (e.g., cytochrome P450 oxidation).<sup>13</sup> Such metabolic processing leads to the formation of multiple products, each of which is detected by MS, obscuring subsequent protein identification and binding-site mapping studies. In the present study, the interaction of a natural product with its target(s) is addressed via an elimination or displacement reaction, wherein the natural product is removed from the reaction milieu, acting as a leaving group.<sup>14</sup>





Given the use of acyl phenols in protein reactive dyes,<sup>15</sup> we examined the feasibility of this motif as a transfer agent for the immunoaffinity fluorescent (IAF) tag (Figure 1a). Though the attachment of a phenol-containing linker would enable this method to be applied to natural products in a general sense, the marinopyrroles, a family of phenolic 1,3'-bipyrroles including **6** and **7**, were selected for this study.<sup>16</sup> We chose these materials based on (1) their availability from culture (strain CNQ-418) at mg/L scale; (2) their unknown mode of action; (3) their activity in tumor cell lines (GI<sub>50</sub> values for **6** and **7** were 10  $\mu$ M in HCT-116 cells and 620 and 500 nM for **6** and **7**, respectively, in A549 breast cancer cells); and (4) their *bis*-phenol structure, previously shown to undergo clean

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acylation to **8** with acetic anhydride (Scheme 1) without a loss in activity (GI<sub>50</sub> value of 0.8  $\mu$ M for **8** in HCT-116 cells, an ~12-fold increase over **6** or **7**).<sup>16</sup>

Based on this evidence, *mono*-labeled **13** (45% yield) and *bis*labeled **14** (traces) were prepared and analyzed using a combination of MS and NMR analysis. HMBC analysis confirmed the position of the tag in *mono*-labeled **13** (see Scheme 1). The low yield of the *bis*-labeled **14** was due to its instability, rapidly converting to **13** in aqueous media with a half-life of  $t_{1/2} = 40$  min in PBS pH 7.2 at 32 °C. *Mono*-labeled **13** was much less prone to hydrolysis with a half-life of  $t_{1/2} = \sim 2$  d in PBS pH 7.2 at 32 °C.



**Figure 2.** Uptake and subcellular localization of probes **13** and **14** in HCT-116 cells. Confocal fluorescent images depicting the red fluorescence from the uptake of **6** in cells treated with (a)  $10 \,\mu$ M **6** for 1 h or (b)  $10 \,\mu$ M **6** for 6 h. A set of confocal fluorescent images showing blue (447 nm), red (692 nm), and two color mixed (mix) fluorescence in HCT-116 cells treated with (c)  $10 \,\mu$ M **13** for 6 h or (d)  $10 \,\mu$ M **14** for 6 h. Probe **14** likely hydrolyzed to **13** under these conditions. (e) Image of cells treated with  $10 \,\mu$ M **13** for 6 h and then washed, fixed, and stained for nucleic acids with Syto-60 (red) and actin with FITC-phalloidin (green). (f) Identical cells as in (e) stained for nucleic acids (red) and transfer of the IAF tag to actin (blue). Note the overlap of green in (e) with blue in (f). All cells were treated under conditions wherein control **12** was completely washed from cells (inset images). Wavelengths denote the center of emission filter. Bars denote 10  $\mu$ m.

Probes 13 and 14 mimicked the activity of marinopyrrole A (6). Probe 14, with a GI<sub>50</sub> value of 1  $\mu$ M, was ~10-fold more active than the parent natural products 6 and 7. Using two-color confocal microscopy, we compared the native red fluorescence from marinopyrrole A (6) at 692 nm (Figure 2a,b) to the blue fluorescence from the IAF tag in 13 (Figure 2d) and 14 (Figure 2c). The presence of a similar pattern of red and blue staining confirmed that probes 13 and 14 shared the same subcelluar localization in HCT-116 cells and, therefore, provided a sufficient mimic of 6. The colocalization of natural product 6 and dye-labeled probes 13 and 14 also showed that cells were not stained from the hydrolytic release of 9. Importantly, control 12 was readily washed from cells under each study (insets, Figure 2a–d).

The putative protein target of the marinopyrrole probe was initially established using an immunoprecipitation (IP) protocol.<sup>2c</sup>

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**Figure 3.** Marinopyrrole probes target actin. (a) A fluorescent gel showing cell lysate obtained from HCT-116 cells ( $10^8$  cells) treated with  $10 \ \mu$ M **13** in L1 and  $10 \ \mu$ M **14** in L2 for 12 h. (b) A Silver Blue stained SDS-PAGE gel depicting proteins immunoprecipitated from the lysate in (a) using Affigel 10 resin containing 3.5 mg/mL of an anti-IAF antibody XRI-TF35. (c) SDS-PAGE gel depicting the HCT-116 cell lysate used in (b). (d) LC/ MS/MS analysis of the IP product in (b) containing actin and actin binding proteins. (e) A fluorescent SDS-PAGE gel of 5  $\mu$ M rabbit muscle actin (protein = a, lanes L1–L10) treated with IAF tags **10–11** and probes **13–14** at 32 °C. Reactions were conducted under conditions wherein probe **13** did not label a lane marker (NativeMark, Invitrogen; protein = lm, lane LM). Fluorescent gels were imaged by excitation at 360 nm.

HCT-116 cells ( $\sim 10^8$  cells) were incubated with 10  $\mu$ M **13** for 12 h. The cells were collected, lysed, and screened for proteins that were labeled with the IAF tag. Two fluorescent bands at 40–45 kDa were apparent in the crude lysate (Figure 3a). After IP, three bands were obtained (Figure 3b). Mass spectral analysis (Figure 3d) indicated that the two lower bands were actin, while the upper band arose from actinin. The fact that actinin was observed during IP experiments in the lysate, while not fluorescent, is expected given its tight complexation with actin.

The selective binding of the probe to actin was validated with purified proteins. Samples of rabbit muscle actin (Worthington Biochemicals) were treated with **13** or **14** to yield comparable fluorescently labeled bands (lanes L4–L7, Figure 3e). Even with a 50-fold increase in concentration (lanes L1, L2, L3, L9, and L10 in Figure 3e), the nonspecific labels **10** and **11** failed to stain actin. The reactivity observed in Figure 3e was indeed due to a specific binding event as pretreatment of actin with **6** effectively blocked the uptake of dye from **14** (lane L8, Figure 3e).

Finally, mass spectral methods were applied to determine the site of dye transfer to actin. Protein from the reaction in lane L6 (Figure 3e) was trypsin digested, purified by HPLC based on the absorbance of the IAF dye at 350 nm, and subjected to LTQ-MS/S with a NanoMate direct infusion system. Evaluation of the resulting spectra using the proteomics search tool, InSpecT, to find peptides that were modified by the IAF tag (M+286 Da), resulted in the identification of the modified peptide V<sub>98</sub>APEEHPTLLTEAPLNPKANR<sub>118</sub>. Because InSpecT annotated the modification to be anywhere from K<sub>115</sub> to R<sub>118</sub>, we manually annotated the data to determine the site of localization. The manual annotation of the low-resolution data suggested that the modification could be found at K<sub>115</sub>. To further

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improve the localization of the probe, the same sample was reanalyzed by LTQ-FTICR-MS, giving rise to a mass of 861.7839 Da within 6.6 ppm of the calculated mass of the modified peptide. Furthermore, all the MS<sup>2</sup> b and y fragment ions observed by LTQ-FTICR-MS matched to within 0.02 Da enabling us to localize the site of modification to  $K_{115}$ . The unmodified peptide V<sub>98</sub>APEEHPTLLTEAPLNPK<sub>115</sub> was also identified, which indicates that this residue was only partially modified.



Figure 4. Marinopyrrole probe 13 transfers an IAF tag selectively to residue K<sub>115</sub> of actin. (a) Structure of actin (green) noting the position of the targeted K<sub>115</sub> residue (blue) and other lysine residues (red). (b) Side and (c) frontal views of the putative marinopyrrole binding pocket (aqua shaded). Images were developed using a complex between rabbit skeletal muscle actin and latrunculin A at 2.85 Å (pdb id: 1IJJ). The side chain amine of  $K_{115}$  (star) and the latrunculin A binding pocket (sphere) are noted.

Lysine 115 occupies a distinct site on actin (star, Figure 4a) that neither overlaps the ATP, gelsolin, and profilin binding sites (see Supporting Information) nor overlaps with the pocket targeted by other known natural products such as latrunculin A (dot, Figure 4a), a marine natural product that binds to the nucleotide binding cleft of actin. Structural modeling identified a channel that exists adjacent to the amine terminus of K<sub>115</sub>, suggesting a putative marinopyrrole-binding site (Figure 4b,c).

Transfer of the IAF dye from the natural product to actin was further supported by secondary analyses. First, microcalorimetry analyses verified the binding between marinopyrrole A (6) and rabbit muscle actin with a  $K_d$  of 0.12  $\pm$  0.2  $\mu$ M. The strength of this complex compares favorably to a reported value of 0.2  $\mu$ M for latrunculin A (albeit at a different binding site, see Figure 4a).<sup>17</sup> Second, using a fluorescence assay (Cytoskeleton Inc.),<sup>18</sup> 6 was shown to inhibit the polymerization of a pyrene-labeled G-actin with an IC<sub>50</sub> value of  $39.5 \pm 6.2 \,\mu\text{M}$  (see Supporting Information). Lastly, actin fibers in live HCT-116 cells treated with probe 13 were blue fluorescent (Figure 2f) from the binding of 13, as evident by costaining with FITC-phalloidin (Figure 2e).

This investigation provides definitive support for the use of a dye transfer protocol via an acyl phenol motif for mode of action studies. By transferring the dye from the natural product to its protein target, we were able to streamline cellular, target elucidation, and binding site determination studies into a single workflow (see Figure 1). Because bioactive natural products bind to proteins in selective binding pockets, the process of natural product mediated ligation may also be used to direct protein modifications site specifically. Studies are now underway to evaluate the scope of this reaction centered on the electronic nature of the phenol and its ability to transfer its payload.

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Supporting Information Available: Synthetic methods, copies of NMR spectra, protocols for cell imaging studies, IP studies and actin assays, detailed methods for the mass spectral studies, copies of original images in Figures 2-3 as well as additional references are available free of charge via the Internet at http://pubs.acs.org.

## References

- (a) Leslie, B. J.; Hergenrother, P. J. Chem. Soc. Rev. 2008, 37, 1347. (b) Wang, X.; Imber, B. S.; Schreiber, S. L. Bioconjugate Chem. 2008, 19, 585
- Recent examples: (a) Bowers, A.; West, N.; Taunton, J.; Schreiber, S. L.; Bradner, J. E.; Williams, R. M. *J. Am. Chem. Soc.* **2008**, *130*, 11219. (b) Kotake, Y.; Sagane, K.; Owa, T.; Mimori-Kiyosue, Y.; Shimizu, H.; Uesugi, (2)M.; Ishihama, Y.; Iwata, M.; Mizui, Y. Nat. Chem. Biol. 2007, 3, 570. (c) Hughes, C. C.; MacMillan, J. B.; Gaudêncio, S. P.; Fenical, W.; La Clair, J. J. Angew. Chem., Int. Ed. 2009, 48, 728.
- (3)(a) Gough, J. D.; Crews, C. M. Ernst Schering Res. Found. Workshop. 2006, 61. (b) Drahl, C.; Cravatt, B. F.; Sorensen, E. J. Angew. Chem., Int. Ed. 2005, 44, 5788.
- (4) Altmann, K. H.; et al. ChemBioChem. 2009, 10, 16.
  (5) (a) Peddibhotla, S.; Dang, Y.; Liu, J. O.; Romo, D. J. Am. Chem. Soc. 2007, 129, 12222. (b) Alexander, M. D.; et al. ChemBioChem. 2006, 7, 409
- (6) (a) Carrico, I. S.; Carlson, B. L.; Bertozzi, C. R. Nat. Chem. Biol. 2007, 3, 321. (b) Sadaghiani, A. M.; Verhelst, S. H.; Bogyo, M. Curr. Opin. Chem.
   Biol. 2007, 11, 20. (c) Heal, W. P.; Wickramasinghe, S. R.; Leatherbarrow,
   R. J.; Tate, E. W. Org. Biomol. Chem. 2008, 7, 2308. (d) Channon, K.; Bromley, E. H.; Woolfson, D. N. *Curr. Opin. Struct. Biol.* **2008**, *18*, 491. (a) Colombo, M.; Peretto, I. *Drug Discov. Today* **2008**, *13*, 677. (b) Hein,
- C. D.; Liu, X. M.; Wang, D. Pharm. Res. 2008, 25, 2216. (c) Moorhouse, A. D.; Moses, J. E. ChemMedChem. 2008, 3, 715
- (8) (a) Weisbrod, S. H.; Marx, A. Chem. Commun. 2008, 30, 5675. (b) de Araújo, A. D.; Palomo, J. M.; Cramer, J.; Seitz, O.; Alexandrov, K.; Waldmann, H. Chemistry 2006, 12, 6095.
- (a) Binder, J. B.; Raines, R. T. *Curr. Opin. Chem. Biol.* **2008**, *12*, 767. (b) Kirshenbaum, K.; Arora, P. S. *Nat. Chem. Biol.* **2008**, *4*, 527. (9)
- (10) Tanaka, Y.; Bond, M. R.; Kohler, J. J. Mol. Biosyst. 2008, 4, 473.
- (11) Guizzunti, G.; Brady, T. P.; Malhotra, V.; Theodorakis, E. A. *Bioorg. Med. Chem. Lett.* 2007, *15*, 320.
- (12) Hurst, G. B.; Lankford, T. K.; Kennel, S. J. J. Am. Soc. Mass Spectrom. 2004, 15, 832
- (a) Dekant, W. EXS 2009, 99, 57. (b) Johnson, W. W. Drug Metab. Rev. (13)2008, 40, 101.
- (14) During the preparation of this manuscript a team led by Hamachi disclosed a similar concept using ligand-directed tosylation: Tsukiji, S.; Miyagawa, M.; Takaoka, Y.; Tamura, T.; Hamachi, I. *Nat. Chem. Biol.* **2009**, *5*, 341. (b) Tsukiji, S.; Wang, H.; Miyagawa, M.; Tamura, T.; Takaoka, Y.; Hamachi, I. J. Am. Chem. Soc. 2009, 131, 9046.
- (15) Gee, K. R.; Archer, E. A.; Kang, H. C. Tetrahedron Lett. 1999, 40, 1471-1474
- (16) Hughes, C. C.; Prieto-Davó, A.; Jensen, P. R.; Fenical, W. Org. Lett. 2008, 10.629
- Coué, M.; Brenner, S. L.; Spector, I.; Korn, E. D. FEBS Lett. 1987, 213, (17)316.
- (18) (a) Zhai, L.; Zhao, P.; Panebra, A.; Guerrerio, A. L.; Khurana, S. J. Biol. Chem. 2001, 276, 36163. (b) Pollard, T. D. Anal. Biochem. 1983, 134, 406
- JA903149U