Analysis of Amyloid Nanostructures Using Photo-cross-linking: In Situ Comparison of Three Widely Used Photo-cross-linkers

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Supporting Information

ABSTRACT: Photoinduced cross-linking (PIC) has become a powerful tool in chemical biology for the identification and mapping of stable or transient interactions between biomacromolecules and their (unknown) ligands. However, the value of PIC for in vitro and in vivo structural proteomics can be realized only if cross-linking reports accurately on biomacromolecule secondary, tertiary, and quaternary structures with residue-specific resolution. Progress in this area requires rigorous and comparative studies of PIC reagents, but despite widespread use of PIC, these have rarely been performed. The use of PIC to report reliably on noncovalent structure is therefore limited, and its potentials have yet to be fully realized. In the present study, we compared the abilities of three probes, phenyl trifluoromethyl diazirine (TFMD), benzophenone (BP), and phenylazide (PA), to record structural information within a biomolecular complex. For this purpose, we employed a self-assembled amyloid-like peptide nanostructure as a tightly and specifically packed model environment in which to photolyze the reagents. Information about PIC products was gathered using mass spectrometry and ion mobility spectrometry, and the data were interpreted using a mechanism-oriented approach. While all three PIC groups appeared to generate information within the packed peptide environment, the data highlight technical limitations of BP and PA. On the other hand, TFMD displayed accuracy and generated straightforward results. Thus TFMD, with its robust and rapid photochemistry, was shown to be an ideal probe for cross-linking of peptide nanostructures. The implications of our findings for detailed analyses of complex systems, including those that are transiently populated, are discussed.

Photoinduced covalent cross-linking (PIC) represents a powerful approach with which to probe biological systems and processes. At the level of interaction networks, cross-linking enables (unknown) interacting partners to be captured, for example, proteins that bind functionalized peptides, proteins that bind functionalized small molecules, small molecules that bind to functionalized proteins, proteins that bind to functionalized proteins, and functionalized proteins that oligomerize. Additionally, at the level of domains, residues, or even atoms, PIC can also provide information on secondary or tertiary structure. The use of PIC to interrogate complex biological systems, structures, and networks, both in vitro and in vivo, has increased in recent years, based on advances in site-specific incorporation of PIC groups into proteins (using total synthesis, semi-synthesis, or incorporation by the use of modified codons) combined with enhancements in analytical methodology, specifically mass spectrometry (MS), that enable the products of cross-linking to be identified uniquely in residue-specific detail.

Benzophenone (BP), phenylazide (PA) and trifluoromethyl diazirine (TFMD) are three commonly used PIC groups, to which various advantages and disadvantages have been ascribed. Diazirines are capable of rapid, indiscriminate reactions but tend to be synthetically and/or commercially less accessible. BP is capable of high-yielding PIC, but long irradiation times are often required and its excitation is reversible. PA generates a highly reactive nitrone that rearranges to an intermediate with strong reactivity preferences. Given their commercial availability, PA and BP have been employed widely in PIC studies of protein interactions, despite misgivings about their suitability for extracting molecular-level information. For BP the caveats include the potential for cross-linking chemistry to be determined by diffusion rather than supramolecular structure, while for PA cross-links can be templated by covalent preference rather than noncovalent structure. The $K_d$ (affinity) and dynamics of different protein–protein/ligand or peptide interactions vary widely; hence different PIC reagents may exhibit differences in reactivity, product distribution, and yield depending on the system. In selecting appropriate PIC reagents, each reagent therefore needs to be compared, but surprisingly, despite the widespread use of cross-linking in the analysis of protein systems, comparative studies of these three PIC groups are rare.

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linking efficiency and/or yield\textsuperscript{21–23} but to our knowledge, a comparison of the utility of these reagents for garnering per residue\textsuperscript{20} information on (supra)molecular structure has not been performed to date, leaving open the question as to the optimal reagent for analysis of protein interactions in structural detail.\textsuperscript{24} This requires consideration of the underlying mechanistic cross-linking chemistry and any reactive preferences\textsuperscript{25} unique to the PIC group in question. Such studies are critical if PIC is to be used to address modern challenges in proteomics, biotechnology, and structural molecular biology, which require the (conventional) ability of PIC to map pairwise interactions, married with a structural rationale for such interactions.

Recently, Schofield and co-workers reported a comparative study in which five PIC groups were compared on the basis of cross-linker reactivity (kinetics and yield) and, using proteolytic digestion, their abilities to report on the interaction of a ligand with human 2-oxoglutarate oxygenase.\textsuperscript{26} The authors observed that different PIC groups generate different cross-links and concluded that PIC group selection should be empirically guided or that such studies should employ a selection of different PIC groups. Our group\textsuperscript{27,28} has previously employed diazirines to study supramolecular organization in amyloid-like peptide nanostructures using ion mobility spectrometry (IMS)\textsuperscript{29,30} coupled with tandem mass spectrometry (MS/MS) to identify the cross-links formed in residue-specific detail. These studies showed that individual cross-links within peptide nanostructures can furnish information on conformation and intermolecular association within highly ordered amyloid-like assemblies of a model peptide (A\textsubscript{β\textsuperscript{16–22}}).\textsuperscript{27,28} Simultaneously, Sinz and co-workers used diazirines to determine a β-turn preference for a peptide in solution.\textsuperscript{16} Given the rigid, close packing of individual peptides within amyloid-like nanostructures of A\textsubscript{β\textsuperscript{16–22}}, these structures form an ideal system with which to systematically compare cross-links obtained for BP, PA, and diazirine (TFMD) within the same self-assembled structure. Here we use a unique A\textsubscript{β\textsuperscript{16–22}} supramolecular structure (the fibrillar form resulting from aggregation at pH 7.0)\textsuperscript{34} to illustrate the advantages and disadvantages of different PIC reagents (BP, PA, and TFMD) that should be considered when designing experiments for structural analysis of protein architectures and interacting surfaces. The results revealed potential technical limitations of PA and BP, while highlighting the favorable properties of TFMD. TFMD photolyzed as readily as PA and generated a product distribution that was easy to interpret. Importantly, TFMD formed only peptide–peptide intermolecular cross-links in the presence of supramolecular structure and light. The implications of these results for the study of biomolecular interactions using PIC are discussed, acknowledging the favorable properties of diazirines\textsuperscript{1} and the availability of technologies with which to incorporate them into biomolecules using chemical synthesis\textsuperscript{31} or genetic methods,\textsuperscript{1,2,32} notably TFMD-phenylalanine\textsuperscript{34} (employed in the present study). Taken together, our results highlight the potential of TFMD as a reagent with which to pursue in vitro and in vivo structural proteomics and complex systems research using PIC.

\section*{RESULTS AND DISCUSSION}

\textbf{Synthesis and Nanostructure Characterization.} We selected A\textsubscript{β\textsuperscript{16–22}} amyloid-like nanofibrils that form readily at pH 7.0 in vitro\textsuperscript{35} as the scaffold for comparison of the utility of different PIC reagents, since this stable, closely packed, and robust structure provides an ideal environment in which to photolyze and compare all three PIC groups. A\textsubscript{β\textsuperscript{16–22}} is also a good model on account of its synthetic accessibility,\textsuperscript{35,36} its ability to self-assemble into different but uniquely defined nanostructures independent of amino acid substitutions, and its amenability to analysis by high-performance liquid chromatography (HPLC)\textsuperscript{35,36} and/or MS.\textsuperscript{28} Also of interest is its role as a mimic of the biologically important A\textsubscript{β} peptides involved in Alzheimer’s disease.\textsuperscript{31,37,38} Assembly of A\textsubscript{β\textsuperscript{16–22}} is sensitive to variation in primary structure, but it is rare that destabilizing modifications abrogate aggregation.\textsuperscript{35,36,39} Rather, amino acid substitutions lead to subtle effects (e.g., on morphology or aggregation propensity of this peptide).\textsuperscript{31,35,36,40}

Once incorporated within A\textsubscript{β\textsuperscript{16–22}} nanostructures, each PIC functionality was photolyzed using standard laboratory apparatus (6-W illuminator with 254- and 365-nm lamps). Products formed within the supramolecular structure (i.e., through supramolecular templating) were compared with those obtained in the absence of supramolecular structure (i.e., as a monomer in solution). Previously, A\textsubscript{β\textsuperscript{16–22}} modified with a TFMD group at Phe-20 (A\textsubscript{β\textsuperscript{16–22}}-TFMD\textsubscript{20}) was shown to display self-assembly behavior similar to that of wild-type (WT) A\textsubscript{β\textsuperscript{16–22}}.\textsuperscript{27} Presently, A\textsubscript{β\textsuperscript{16–22}} analogues with azido and benzoyl modifications at Phe-20 (termed A\textsubscript{β\textsuperscript{16–22}}-PA\textsubscript{20} and A\textsubscript{β\textsuperscript{16–22}}-BP\textsubscript{20}, respectively) were prepared via Fmoc solid-phase peptide synthesis (Figure 1, Supplementary Tables ESI 1 and 2 and Supplementary Figures ESI 1–5).

\begin{figure}[h]
\centering
\includegraphics[width=0.8\textwidth]{figure1.png}
\caption{Structures of A\textsubscript{β\textsuperscript{16–22}} and photoreactive analogues.}
\end{figure}

Peptides were incubated under aggregation-promoting conditions (0.4 mM peptide, aqueous solution, buffered at pH 7.0, 15 days, 4 °C, quiescent). Sedimentation-HPLC\textsuperscript{43} revealed that all peptides aggregated substantially (mole fraction >75%, see Supplementary Table ESI 3 and Supplementary Figures ESI 7–10 for details). Mixed samples containing WT A\textsubscript{β\textsuperscript{16–22}} spiked with photoreactive analogues (ratio 4:1, WT:modified) also yielded aggregates in which the modified peptide co-assembled with the WT peptide (see below). All aggregates were analyzed further using negative-stain transmission electron microscopy (TEM) (Figure 2 and Supplementary Table ESI 4), allowing higher-order supramolecular structure\textsuperscript{44} (e.g., fibrils versus nanotubes) and sample homogeneity to be evaluated. All assemblies were fibrous, indicating that no gross perturbations to the A\textsubscript{β\textsuperscript{16–22}} supramolecular structure had resulted from the substitutions made to primary structure.

\textbf{Cross-linking and Analysis of Monomeric Peptides.} To study PIC in the absence of supramolecular structure, we irradiated peptides that had been diluted into hexafluoroiso-
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Figure 2. TEM of nanostructures formed upon incubation of Aβ16–22 and its photoreactive analogues at 4 °C for 15 days. (a–c) Fibers formed from stock solutions of Aβ16–22, Aβ16–22–PA20, or Aβ16–22–BP20, respectively. (d–f) Fibers formed when a stock solution of Aβ16–22 was spiked (ratio 4:1) with Aβ16–22–TFMD20, Aβ16–22–PA20 or Aβ16–22–BP20 respectively (scale bars = 100 nm).

propanol (HFIP), a solvent in which the peptides are monomeric.27 Molecularly dissolved peptides were irradiated (254 or 365 nm) for varying amounts of time (typically 0, 5, or 60 min) and analyzed using liquid chromatography (LC) interfaced to an ion trap mass spectrometer. The choice of wavelength was guided by experiment (see UV–vis spectra in Supplementary Figures ESI 5 and ESI 6) and, where necessary, by relevant information from the literature. The results of all photolysis experiments are summarized in Table 1. WT Aβ16–22 did not photolyze at either wavelength on the time scale of a typical photolysis reaction (Supplementary Figure ESI 11). Irradiation of Aβ16–22–TFMD20 at 365 nm generated a distribution of adducts as described previously.27 Presently, it was noteworthy that a significant quantity of the diazirine remained at 5 min, but that this had been consumed by 60 min. The dominant photolysis product was assigned as a hexafluoropropyl ether (m/z 1142; change in mass, Δm = +140), formed via insertion of singlet carbene and/or solvolysis of photoisomerised Aβ16–22–TFMD20. Other major products were assigned as a water adduct (m/z 992; Δm = −10) and cyclized monomer (m/z 974; Δm = −28) (Supplementary Figure ESI 12). The cyclized monomer was identified as a macrolactone, as evidenced by the formation of an aminolysis product (m/z 1033; Δm = +59) upon reaction with n-propylamine.

Photolysis of Aβ16–22–PA20 at 254 nm (i.e., near its λmax) occurred on a time scale similar to that of Aβ16–22–TFMD20 at 365 nm but generated more products (Supplementary Figure ESI 13). As with Aβ16–22–TFMD20, the major product (m/z 1075; Δm = +140) was an HFIP adduct. This product could be one of three isomeric structures: an O-substituted hydroxylamine (formed via a singlet nitrene), a hemiaminal (via a triplet nitrene),45 or a substituted azezine (formed via a 1,2-didehydroazepine). A second major ion (m/z 939) was assigned as an oxidation product corresponding to a monomer having lost N2 and gained two oxygen atoms (putative nitro compound). Significantly, in contrast to the results obtained for Aβ16–22–TFMD20, covalent dimerization was observed. The clearest evidence was m/z 992, which indicates dimerization through one intermolecular cross-link with an additional HFIP insertion. Clearly, formation of dimeric products in HFIP (i.e., a solvent that inhibits molecular association) is cause for concern,

Table 1. Summary of Results for All Photolysis Experiments

| phase         | λ (nm) | t (min) | Aβ16–22 | Aβ16–22–PA20 | Aβ16–22–BP20 |
|---------------|--------|---------|---------|--------------|--------------|
| solution      | 254    | 5       | ND<sup>7</sup> | ND           | no change<sup>b</sup> |
|               |        | 60      | no change | ND           | ND           |
|               | 365    | 5       | ND       | partial photolysis; appearance of HFIP adduct and other insertion products | partial photolysis; appearance of HFIP adduct and other insertion products |
|               |        | 60      | no change | complete photolysis; HFIP adduct is the dominant product | photolysis nearing completion; HFIP adduct is the dominant product; homodimer observed |
| homoaggregate | 254    | 60      | ND       | complete photolysis; putative oxidation products dominate | no change |
|               | 365    | 60      | no change | complete photolysis; peptide insertion products dominate | complete photolysis; putative oxidation products dominate |
| heteroaggregate| 254    | 60      | NA<sup>a</sup> | ND           | no change |
|               | 365    | 60      | NA       | heterodimer observed | no change |

<sup>a</sup>ND = no data. <sup>b</sup>All changes reported are relative to a dark control. <sup>a</sup>NA = not applicable.

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because it suggests PA can form intermolecular cross-links independently of supramolecular templating.

For Aβ16–22-BP20, we were unable to observe the carbonyl n→n* transition (for which λ_{max} typically falls in the range 330–350\text{nm}) that generates the reactive intermediate (BP triplet). This absence is explained by some unusual solvent effects, which were the subject of a recent report by Lewandowska-Andralojc et al.\textsuperscript{47} HFIP induces a blue shift of...
the weak n→π* absorption, causing it to become hidden under the
tail of the strong aromatic π→π* absorption (see
Supplementary Figure ESI 5). Although the authors were
unable to determine the precise position of λ_{\text{trans}} when the HFIP
content was above 80% (v/v), they were still able to observe a
proportion of BP’s n→π* triplet spectroscopically when the
sample was irradiated at 340 nm. On this basis, our 365-nm UV
source should have been effective in generating the reactive BP
triplet (noting this is routinely the case).6 However, when the
experiment was performed, no changes were observed in the
sample, even after 60 min of irradiation (Supplementary Figure
ESI 14). A brief (5 min) irradiation at 254 nm was also
ineffective in initiating cross-linking (see Supplementary Figure
ESI 15).

Cross-linking and Analysis of Aggregated Peptides.
Subsequent experiments focused on the PIC of peptide
aggregates. Each aggregate was irradiated, isolated by
centrifugation, and then disaggregated in HFIP as detailed
previously.27 Analysis of photolysis products was conducted
using LC–MS, and where signals needed to be resolved further,
using the separate capabilities of IMS–MS on a Waters
Synapt HDMS mass spectrometer. The results obtained for
homopolymeric Aβ_{16-22}-TFMD_{20} aggregates (i.e., samples
comprising only one peptide monomer) assembled at 4 °C
were indistinguishable from those reported previously (Supple-
mentary Figure ESI 16).27 The major photolysis products were
peptide insertion products (m/z 974; Δm = −28), a water
adduct (m/z 992; Δm = −10), a water-quenched covalent
dimer (doubly charged m/z 984; Δm = −28 for the first
monomer and −10 for the second), and an HFIP adduct (m/z
1142; Δm = +140). We previously identified the HFIP adduct
as resulting from isomerization of TFMD to the diaoisomer
which in the presence of mildly acidic HFIP gives a reactive
carboxylation that is quenched by HFIP.27 Note that m/z 984
had not appeared when HPLC-purified Aβ_{16-22}-TFMD_{20}
was photolyzed in HFIP (i.e., no intermolecular cross-links formed
in the absence of supramolecular structure). Photolysis of
heteropolymetric Aβ_{16-22}-Aβ_{16-22}-TFMD_{20} aggregates (i.e.,
samples comprising WT and modified monomers in a 4:1 ratio)
generated a product distribution (Figure 3) containing a
covalent heterodimer (doubly charged m/z 935; Δm = −28).
Previous MS/MS analyses27 of the various photolysis products
enabled cross-link positions to be elucidated, including
intramolecular cross-linking between TFMD-Phe-20 and Glu-
22 (identified by post-PIC aminolysis) and an intermolecular
cross-link from TFMD-Phe-20 to Lys-16, which was consistent
with an in-register antiparallel β-sheet structure.

The product distribution obtained upon photolysis of
heteropolymeric Aβ_{16-22}-PA_{20} aggregates (Figure 4a) was
more complicated than that obtained for Aβ_{16-22}-TFMD_{20}.
The products were not well resolved by LC–MS, so the sample
was reanalyzed using IMS–MS. The IMS–MS 3D driftscope
plot (Figure 4b) reveals a distribution of monomers, dimers,
and trimers (n = 1, z = 1; n = 2, z = 2; and n = 3, z = 3; where n
is the number of monomers from which the species is
composed and z is the ion charge) with the same m/z (m/z
∼908; cf. data obtained when Aβ_{16-22}-PA_{20} had been photo-
lyzed in HFIP solution). As in diazirine-mediated PIC, this
distribution is characteristic of intermolecular and/or intram-
olecular cross-linking with loss of 1 × N_{2} per monomer
(Figure 4c). Interestingly, a HFIP adduct (m/z 1075) also
appeared in the present experiment; this must have formed by
solvolysis of a photolysis product during the HFIP
disaggregation step of analysis. To examine PA cross-linking
in more detail, we interrogated the structure of the cyclized
monomer (single charged m/z 908) using IMS–MS/MS. Just
as for cyclized Aβ_{16-22}-TFMD_{20}, a discontinuous series of b
ions resulting from single amide cleavages (Roepstorff–
Fohlman–Bienzmann nomenclature)48,49 indicated that an intram-
olecular cross-link had formed between the PIC functionality
(PA) and Glu-22 (Figure 4d). This cross-link is consistent with
the peptide adopting a β-strand conformation. Other major
species resolved by IMS–MS were oxidation products that had
lost one molecule of nitrogen per monomer (Δm = −28) and
gained one or two oxygen atoms (Δm = +16 or +32,
respectively). As a result, PA gives rise to substantially more
complex reaction products compared with those of TFMD.

The observed product distributions are consistent with the
work of others, where products arising from the photolysis of
PA in aerated media have been characterized.50,51 In terms of
PIC methodology, a key consideration is whether the oxidation
products react further to form cross-links. The literature is
conflicting: in a report by Waddell and co-workers, nitro
derivatives of PA were considered benign products that do not
react photochemically or otherwise,51 whereas in a PIC study
by Escher and Schwzyer, an inhibitor containing 4-nitro-L-
phenylalanine was unexpectedly observed to cross-link photo-
chemically to α-chymotrypsin.52 Additionally, where PIC
mechanisms are concerned, it is well-known that triplet
phenylnitrene can dimerize via an azo cross-link, and that
azo cross-links are also possible. The extent to which these
reactions should contribute to cross-linking in an aerated
medium is unclear from the literature, with reported yields
being variable.50,51,53

Irradiation of the mixed aggregate (Aβ_{16-22}-Aβ_{16-22}-PA_{20})
yielded a product distribution similar to that from irradiation of
Aβ_{16-22}-PA_{20} alone (Supplementary Figure ESI 17), except
that a covalent heterodimer (doubly charged m/z 901) was also
formed. Due to the added complexity of the spectra and the
fact that dimers were obtained by cross-linking of monomeric
Aβ_{16-22}-PA_{20} in HFIP (see above), these products were not
analyzed further.

Irradiation of homopolymeric Aβ_{16-22}-BP_{20} aggregates for 60
min at 365 nm did not cause changes to the sample (Figure
5 and Supplementary Figure ESI 18), and accordingly, nor did
irradiation of aggregated Aβ_{16-22}-Aβ_{16-22}-BP_{20}. As with the
findings described earlier (i.e., no reactivity upon irradiation
in the presence of HFIP), the results observed could arise for
a number of reasons; it could be that insufficient BP triplet
was generated or, alternatively, that an excited state was formed
but was unable to react. Analysis of the control sample revealed
putative intermolecular cross-links formed in the dark
(Supplementary Figure ESI 18). Specifically, a weak signal
centered on m/z 989 (i.e., ∼m/z units less than the m/z of
monomeric Aβ_{16-22}-BP_{20}) indicated dimerization with loss of
water, which was confirmed using IMS–MS (Figure 5a). There
are two explanations: (i) intermolecular cross-linking occurred
via BP triplet with subsequent acid-catalyzed dehydration of the
cross-link,54 or (ii) cross-linking occurred via condensation of
ground-state BP with the Lys ε-amine of a second monomer.
Given that cross-linking did not take place during irradiation,
the latter explanation (i.e., imine formation in the dark)
is plausible, especially considering that imine formation by 4-
benzoyl-L-phenylalanine has been observed by others.55 To
investigate whether the observed imine cross-links were
specific, the supernatant of a centrifuged assembly mixture

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was examined for evidence of $m/z$ 990. If imine formation is templated within an aggregate, cross-linked units should not appear in the solution phase. The fact that the supernatant did not contain $m/z$ 990 suggests imine formation was templated. This dimer also appeared for the solubilized $\text{A}_{16-22}\cdot\text{BP}_{20}$ aggregate, alongside a heterodimer, in which the BP group of $\text{A}_{16-22}\cdot\text{BP}_{20}$ had apparently condensed with the Lys ε-amine of $\text{A}_{16-22}$ (Figure 5b). The imine dimers of $\text{A}_{16-22}\cdot\text{BP}_{20}$ and of $\text{A}_{16-22}\cdot\text{TFMD}_{20-22}$-BP$_{20}$ are structurally analogous to the covalent heterodimer formed from $\text{A}_{16-22}\cdot\text{A}_{16-22}\cdot\text{TFMD}_{20-20}$ which was found to contain a cross-link at position 20 of $\text{A}_{16-22}\cdot\text{TFMD}_{20}$ and Lys-16 of $\text{A}_{16-22}$.

**Conclusions.** In summary, we have used amidoyl-like nanostructures of $\text{A}_{16-22}$ to evaluate the effects of TFMD, PA, and BP groups on peptide aggregation, irrespective of the substitution introduced, and the ability of these different crosslinkers to report on their supramolecular environments via PIC. None of TFMD, PA, or BP impaired the ability of $\text{A}_{16-22}$ to form nanostructures when incorporated at residue Phe-20, and in all cases self-assembled fibrillar structures with similar morphology were formed. Major differences were observed, however, in the PIC experiments. TFMD photolyzed readily, generating a simple distribution of products, for which detailed analysis was straightforward and interpretation of supramolecular structure was possible, although care should be taken to account for products resulting from decomposition of TFMDs linear diazoisomer under acidic conditions, as reported previously. Importantly, TFMD generated only intermolecular peptide–peptide cross-links in the presence of supramolecular structure. PA also yielded structural information via the formation of inter- or intramolecular cross-links. However, control experiments revealed that cross-linking was not determined solely by supramolecular templating. Finally, BP did not form photolysis products under the conditions employed but did form intermolecular imine cross-links in the dark. Thus, in this model system, PA and BP demonstrated the potential to generate inaccuracies, while TFMD more closely resembled an ‘ideal’ PIC reagent. We note that modified tRNA/tRNA synthetase pairs have been developed for the incorporation of diazirine-based amino acids using recombinant expression, while the synthesis of diazirine-modified amino acids (Phe, Trp, Leu, Met, Lys, and Pro) is readily achievable, ensuring that diazirines can be incorporated into any peptide or protein of choice. Furthermore, the carbene resulting from TFMD excitation, which has a picosecond lifetime, represents the most reactive of the PIC groups in common use, opening the door to mechanistic studies on dynamic complex systems.

More broadly, PIC is most powerful where it can be used to provide molecular level information, either within a supramolecular structure or between interacting protein partners of complexes, be these either long-lived or transient. The approach employed here, making use of state-of-the-art MS techniques, permits a large number of PIC products to be analyzed and allows cross-linking sites to be uniquely identified. These can then be interpreted in the context of chemical reactivity and noncovalent structure. The peptide nanostructure used in this work highlights the ability of PIC using TMFD-Phe to report on molecular recognition, even for structures that involve dense packing of side chains. By photolyzing PIC reagents in the presence of a high concentration of imine cross-link products. However, factors such as cross-linker yield and rate will also be relevant; in this work the PIC reagent as one of the seven amino acids in the sequence represents around 20% of the mass of the sample for homopolymeric aggregates and around 5% of the mass of the sample for heteropolymeric aggregates. When a protein system is used (200–400 amino acids), the mass of cross-linker will be proportionally much lower, making it harder to study cross-links by MS and/or by gel electrophoresis. Thus, a cross-linker optimized to capture the interacting partner (e.g., slower rate of cross-linking but high yield) might be preferred to one that is optimized to capture structural information (e.g., indiscriminate reactivity capturing key noncovalent contacts, but also other species due to the dynamics of the interaction under study). Further, preferred chemical reactivity of the PIC group may not bias the outcome of a PIC experiment, and in some cases, such reactivity may even be desired (e.g., where a binding site contains an amino acid with which the PIC group preferentially reacts). Thus, empirically guided PIC selection or use of multiple PIC groups simultaneously will mitigate against erroneous structural interpretation of a cross-linking study. PIC analysis of biomolecular complexes remains a challenging endeavor, but
one that can reward the investigator with otherwise inaccessible information. We anticipate that the findings reported herein should aid the choice of reagent, the interpretation of information, and therefore the study of complex dynamic biomolecular interactions and systems using PIC.

**METHODS**

Peptides were prepared by automated solid-phase peptide synthesis using a CEM Liberty peptide synthesizer operated without microwave irradiation and purified by reverse phase HPLC. Nanostructures were prepared by dilution of a 20 mM peptide stock solution in DMSO with buffer to 0.4 mM final concentration, followed by incubation at ambient temperature (4 °C), in the dark, without agitation, for 15 days. Nanostructures were characterized using TEM with uranyl acetate staining. Photolysis reactions were carried out at ambient temperature using a 6 W ultraviolet lamp (254 or 365 nm) and analyzed after 5 and/or 60 min. For solution-phase studies, the peptide stock solution was diluted with HFIP and irradiated immediately. For experiments with nanostructures, aliquots of the assembly mixture were gently homogenized prior to irradiation. Irradiated nanostructures were then isolated by centrifugation and treated with HFIP to effect dissolution prior to LC–MS, IMS–MS, and IMS–MS/MS analyses. For full experimental details, see the Supporting Information.

**ASSOCIATED CONTENT**

1. Supporting Information

Synthetic methods and characterization, methods for assembly and characterization of nanostructures, and additional MS analyses and methodology. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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