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Photogenerated Base in Polymer Curing and Imaging: The use of a-Keto Carbamates as Novel Photoprecursors of Amines

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Several newly developed a-keto carbamates are useful as photoprecursors of amines. These photogenerated catalysts that are designed primarily for use in chemically amplified resist materials can also be used for the curing of polymers containing base crosslinkable reactive groups. The new a-keto carbamates can undergo light triggered photocleavage both in the solid-state and in solution to give free amines. The photoactive benzoxyloxy carbonyl groups of these compounds are active with ultraviolet radiation below 400nm. The influence of various steric and electronic effects on the photoliberation of free amines from a-keto carbamates has been studied and the effect of structural modification of the aryl rings was used to probe the electronic factors of the photocyclization. The great potential of these photoactive carbamates as organic sources of photogenerated amines was demonstrated using both classical spectroscopic techniques (UV, IR and NMR) and GC-MS analysis of the photolysis products. In all cases the photocleavage reaction is near quantitative and the quantum yields have been measured over a range of wavelengths from 250-370 nm.
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Summary.
Several newly developed α-keto carbamates are useful as photoprecursors of amines. These photogenerated catalysts that are designed primarily for use in chemically amplified resist materials can also be used for the curing of polymers containing base crosslinkable reactive groups. The new α-keto carbamates can undergo light triggered photocleavage both in the solid-state and in solution to give free amines. The photoactive benzoinyloxycarbonyl groups of these compounds are active with ultraviolet radiation below 400 nm. The influence of various steric and electronic effects on the photoliberation of free amines from α-keto carbamates has been studied and the effect of structural modification of the aryl rings was used to probe the electronic factors of the photocyclization. The great potential of these photoactive carbamates as organic sources of photogenerated amines was demonstrated using both classical spectroscopic techniques (UV, IR and NMR) and GC-MS analysis of the photolysis products. In all cases the photocleavage reaction is near quantitative and the quantum yields have been measured over a range of wavelengths from 250-370 nm.


**Introduction.**

While early studies with photochemically labile protecting groups have originally been carried out with the goal of simplifying complex synthetic schemes, their use has more recently been extended to a variety of application in microlithography. A number of photodeprotection schemes that also require the controlled release of reactive species in-situ include caged biomolecule release, or combinatorial solid phase peptide synthesis. Our original work with chemically amplified resist materials has involved the in situ phototriggered formation of acid within a reactive polymer film. The now well known family of resists based on the chemistry of polymers with t-butyl carbonate (t-BOC) and t-butyl ester groups has been used commercially by IBM for the mass production of DRAM memory chips since the early 1980’s. Numerous others have used our approach in the design of analogous resist materials that also involve the photogeneration of acid in the key imaging step. In contrast, the complementary approach of base photogeneration has received little attention. Recently, we have developed a strategy for base photogeneration by masking amines and diamines as photoactive carbamates. Under the action of light, these reactive carbamates decompose to liberate the base in a reactive form. The first generation of photoprecursors of organic base are already finding application in a variety of imaging schemes requiring photogenerated base. The success of our amine photorelease strategy is further illustrated by the development of a number of base catalyzed chemically amplified resist materials.

Of all the common photolabile amino protecting groups, we found that those derived from the 3,5-dimethoxy-α,α-dimethylbenzyloxy carbonyl (Ddz) and o-nitrobenzylloxy carbonyl moieties are particularly well suited to the photogeneration of organic base. While base photogenerators derived from these photoprotecting groups have proven useful, they do suffer from some restrictions. For instance, the utility of the o-nitrobenzyl class of base photogenerator is limited by the formation of reactive o-nitrosocarbonyl compounds, which may be toxic and also may undergo deleterious side reactions. Over-irradiation products such as azo-coupled intermediates act as internal light filters and ultimately restrict the use of these base photogenerators to thin film applications. Similarly, severe anti-bleaching is a significant problem with the 3,5-dimethoxy-α,α-dimethylbenzyloxy carbonyl class of photobase generators.
It is these potential limitations of our first generation of amine photoprecursors that has prompted the development of new photochemistry for organic solid state applications. As indicated in the previous report, desyl photochemistry offers several desirable features which render it particularly attractive for base photogeneration. These features include improved photoefficiency, enhanced rate of photorelease and increased stability of the photo by-products. In this report, we describe novel applications of benzoin photochemistry to the preparation of efficient photoprecursors of organic bases: primary and secondary amines. Encouraged by preliminary results from the solution photolysis of cyclohexylamino and piperidino carbamates of 3',5'-dimethoxybenzoin\textsuperscript{14}, we focus further on the novel photorearrangement of the 3',5'-dimethoxybenzoin chromophore (Scheme 1), while the preceding report described the synthesis of a variety of carbamates based on the 3',5'-dimethoxybenzoin chromophore and its substituted analogs, we now focus on the photochemistry of these \(\alpha\)-keto carbamates. The carbamates were chosen for detailed evaluation based on 1) their synthetic accessibility and 2) their ability to contribute to structure-activity relationships pertinent to base photogeneration. Our aim is to provide a basic understanding of the issues that determine the efficiency of amine photogeneration from 3',5'-dimethoxybenzoin carbamates and substituted derivatives.

**Results and Discussion.**

1) Design Considerations

Previous experience in designing base photogenerators\textsuperscript{9}, suggests some of the most critical features to consider include: 1) favorable UV characteristics, 2) quantum efficient photodecomposition, 3) solid-state photoreactivity, 4) photorelease of the base in a reactive form, 5) stability of photo by-products 6) thermal stability, 7) ease of synthesis. This list is only a general guide because specific applications place additional custom requirements on any base photogenerator. The materials chosen for detailed study were restricted to the cyclohexylamine photogenerators shown in Scheme 2. By only selecting photoprecursors of cyclohexylamine, a direct comparison of the chromophore structure-photosensitivity relationship could be readily made. The \(\alpha\)-keto carbamates of interest are those containing the following chromophores: the parent desyl group as in (1), the 3,3',5,5'-tetramethoxybenzoin moiety as in (2), the 3,5-dimethoxybenzoin group as in (3), the 3',4,5'-trimethoxybenzoin moiety as in (4), the 3',5-dimethoxy-4-methylthiobenzoin group as in (5), 2-(3,5-
dimethoxyphenyl)-1-(2-naphthalenyl)ethanone as in (6) and 2,2-di-(3',5'-dimethoxyphenyl)-1-phenylethanone as in (7). These materials are characterized in terms of the design criteria outlined above.

2) Photochemistry of Carbamates Derived from α-Hydroxy Ketones

α-Keto carbamates absorb UV light from just below 370 nm down to the deep UV. Excitation of the carbonyl chromophore is believed to cause photocyclization to the benzo[b]furan with concomitant liberation of the amine type base (Scheme 1 and 2). Due to the nature of the α-keto chromophore, UV exposure at longer wavelengths is thought to cause photocyclization via direct n-π* excitation. However, photolysis at the higher energy end (π-π* excitation) can also result in the liberation of base although the photocyclization may proceed less efficiently as it requires some energy transfer process. A variety of the spectroscopic methods e.g. 1H NMR, IR, and UV proved useful in monitoring the photochemistry of these new photogenerators. GC and GC-MS facilitated identification of the photoproducts.

a) Solution Photosensitivity

i) Photochemistry of Unsubstituted α-Keto Carbamates

Monitoring the photolysis of benzoin cyclohexyl carbamate (1) by a number spectroscopic methods served to indicate the complexity of the photochemistry of the α-keto chromophore. A significant conversion of benzoin cyclohexyl carbamate (1) to free cyclohexylamine was observed on monitoring the photolysis by 1H NMR spectroscopy (Table 1). However, despite the high conversion, no evidence was observed for the clean photocyclization. In fact, the 1H NMR spectrum became very complicated, implying that photodecomposition of this material proceeds by a variety of competing pathways. This was further confirmed by monitoring the photolysis of benzoin carbamate (1) by infra-red spectroscopy. In this case, the keto carbonyl stretch (1699 cm⁻¹) disappears while the carbamate carbonyl stretch (1722 cm⁻¹) remains substantially intact. A weak carbonyl stretch at 1658 cm⁻¹ also appears possibly due to photooxidation. These observations are consistent with photodecomposition of benzoin carbamate (1) proceeding by a variety of mechanisms of which photocyclization to benzo[b]furan
plays only a minor role. Furthermore, a complex multicomponent mixture is observed by TLC after 350
nm photolysis of benzoin carbamate (1).

ii) Photochemistry of Aryl substituted α-Keto Carbamates

The importance of the meta activation by 3',5'-dimethoxy substituents in the photocyclization of
α-keto carbamates to benzo[b]furan is apparent in the 350 nm solution photolysis of some α-keto
carbamates as readily monitored by $^1$H NMR spectroscopy in dry, degassed acetonitrile-d$_3$. Table 1
shows examples of α-keto carbamates which undergo efficient photogeneration of cyclohexylamine on
350 nm photolysis. For all these carbamates, the integral of the carbamate cyclohexyl methine
[cyclohexyl CH, m, δ ~ 3.2-3.4 (dependent on exact structure of photosensitive carbamate)] decreased,
while the expected multiplet of the cyclohexyl methine of free cyclohexylamine appeared at δ 2.52, with
increasing 350 nm exposure in a Rayonet photochemical reactor. By ratioing the cyclohexyl methine
integral for free cyclohexylamine to the aryl resonances, the % conversion after 60 min. 350 nm exposure
was calculated for a variety of α-keto carbamates as shown in Table 1. % Conversion varied from 37-
98% depending on the chromophore structure. Besides identifying cyclohexylamine, $^1$H NMR
spectroscopy allowed the expected benzo[b]furan to be identified as the major photo by-product on 350
nm photolysis of 3',5'-dimethoxy substituted benzoin carbamates, (2), (3), (4), (5) and (6). A
characteristic singlet of the 3-benzo[b]furan proton in the photoproduct mixture appears with increasing
photolysis. The chemical shift of this proton is included in Table 1. In these systems which undergo
smooth photocyclization, the % conversion for formation of benzo[b]furan tends to parallel that for
amine generation. The cleanliness of the photocyclization of these 3,5-dimethoxy activated benzoin
carbamates is nicely illustrated by following the changes in the $^1$H NMR aryl resonances with increasing
exposure dose for the conversion of carbamate (2) to benzo[b]furan (2a) in tetrahydrofuran-d$_3$ (Figure
1). A plot of % conversion against irradiation time for the 350 nm photolysis of carbamate (2) further
demonstrates the efficiency of the benzoin photocyclization (Figure 2). The same clean photochemistry is
not apparent with other α-keto carbamates. For instance, systems lacking the 3',5'-dimethoxy substitution
pattern e.g., (1) undergo particularly complex photodecomposition.
The clean photogeneration of free cyclohexylamine from all 3',5'-dimethoxy substituted benzoin carbamates was further confirmed by both gas chromatography using coinjection and GC-MS studies (C\text{6}H\text{11}NH\text{2} M^+=99). For these cyclohexyl carbamates, (2), (3), (4), (5) and (6), GC-MS indicated the major by-products were the expected substituted benzo[b]furans (2a), (3a), (4a), (2a) and (6a) respectively. Other minor products were identified as the corresponding benzyl ketone e.g. (2b), the corresponding substituted benzil e.g. (2c), the corresponding N-acylcyclohexylamine e.g. (2d) and dicyclohexylurea (Scheme 2). Formation of these minor photoproducts can be rationalized in terms of the known benzoin photochemistry. For instance, the benzyl ketone e.g. (2b), is formed by loss of the carbamoyloxy moiety. This mode of photocleavage is common in benzoin photochemistry and tends to occur when a good leaving group is present at the α-keto position. Indeed, the photolabile phenacyl protecting group is known to cleave via this pathway\textsuperscript{15}. The benzil photoproduct e.g. (2c), is likely formed by cleavage of the oxycarbonyl bond followed by photooxidation. In the case of unsymmetrical benzoin carbamates, only the corresponding unsymmetrical benzil was identified by GC-MS. Since there was no indication of any symmetrical benzils, formation of these 1,2-ethanediones is unlikely to be the product of acyl radical recombination. Similar cleavage of oxycarbonyl bonds is known to occur in appropriately substituted benzoins. For instance, photolysis of 4,4'-dimethoxybenzoin acetate is known to form 4,4'-dimethoxybenzil as the major product\textsuperscript{16}. As for the N-acylcyclohexylamine e.g. (2d), this class of photoproduct is likely formed by combination of acyl and cyclohexylamino radicals. The acyl radicals are most likely formed by the well known Norrish Type I photocleavage. Cyclohexylamino radicals may be formed in a number of ways. For example, the carbamoyloxy radicals liberated in the formation of the benzyl ketone e.g. (2b), will likely undergo facile decarboxylation. Alternatively, cleavage of the oxycarbonyl bond to give the substituted benzil e.g. (2c), would be accompanied by formation of a N-formylcyclohexylamino radical. This species would be expected to liberate cyclohexylamino radicals via decarbonylation. Interestingly, no indication of the corresponding aldehyde from the acyl radical formed by Norrish Type 1 cleavage was available by GC-MS. This implies recombination of acyl and amino radicals is strongly favored over hydrogen abstraction. The presence of trace amounts of dicyclohexylurea can be rationalized as follows. Combination of N-formylcyclohexylamino and cyclohexylamino radicals would give dicyclohexylurea. Another possibility is
the reaction of photogenerated cyclohexylamine with cyclohexyl isocyanate. The isocyanate being formed by loss of a hydrogen atom from the N-formylcyclohexylamino radical.

Further evidence for the photochemical rearrangement of these α-keto carbamates was obtained by following the changes in the infra-red spectrum of a solution of carbamate (3) in dry acetonitrile with increasing UV exposure dose. As seen in Figure 3, 350 nm Raynet exposure of carbamate (3) caused the gradual disappearance both the keto carbonyl stretch (1700 cm\(^{-1}\)) and the carbamate carbonyl stretch (1723 cm\(^{-1}\)). This change is consistent with the proposed mode of photodecomposition (Scheme 2) in which photocyclization to liberate the free amine causes these stretches to disappear. In addition, evidence for the photogeneration of free amine was gained by monitoring the changes in the N-H stretching frequency of this substrate (3) (Figure 4). In this case, UV exposure at 350 nm resulted in the gradual depletion of the carbamate N-H stretch at 3360 cm\(^{-1}\), with concomitant appearance of stretches due to the photogenerated cyclohexylamine in the region 3500 to 3700 cm\(^{-1}\). Similarly smooth disappearance of both carbonyl stretches was observed on irradiation of carbamates (2) and (4). These findings are consistent with smooth photocyclization and concomitant amine liberation occurring in these 3',5'-dimethoxy substituted benzoin carbamates. In the case of carbamates, (5) and (6), the disappearance of both carbonyl groups was complicated by the appearance of two weak absorptions at 1675 and 1685 cm\(^{-1}\) respectively. This indicates the photocyclization of carbamates (5) and (6) is complicated by alternate side reactions open to the desyl chromophore. These include photocleavage by Norrish Type I and C-O homolytic pathways, both of which would produce carbonyl containing products. Such competing photochemistry may also explain the slightly lower yields for amine generation from carbamates (5) and (6) (Table 1).

Beyond 350 nm, the extinction coefficients of these carbamates is typically of the order 50-200 L mol\(^{-1}\).cm\(^{-1}\). Considering the low absorption of these carbamates at longer wavelengths, the efficiency of the photocyclization and concomitant amine generation on 350 nm photolysis is truly remarkable. In addition carbamates which absorb the most light at longer wavelengths are not the most efficient according to \(^1\)H NMR studies. For instance, amine photogeneration from carbamate (6) (\(\varepsilon_{(365)}\) 212 L mol\(^{-1}\).cm\(^{-1}\)) is less efficient than carbamate (2) (\(\varepsilon_{(365)}\) 45 L mol\(^{-1}\).cm\(^{-1}\)). This indicates a subtle relationship exists between photocyclization and the nature of the benzoyl chromophore.
In carbamates containing bathochromic substituents, the UV spectrum of these materials was strongly influenced out to ca. 340 nm. For example, the 4-thiomethyl substituted carbamate (5) and the 1-naphthalenyl carbamate (6) had extinction coefficients at 336 nm of 5049 and 2008 L mol⁻¹ cm⁻¹ respectively. Therefore, it is apparent that structural variation of the benzoyl chromophore does influence the long wavelength absorption. However, in these systems the absorption drops off quickly beyond 340 nm and the substituent effect is not strong enough to cause a significant bathochromic shift out to 350 nm.

The result of following the changes in the UV spectrum of a dilute solution of carbamate (2) is shown in Figure 5. Three isobestic points are observed: 251, 266 and 340 nm respectively. Overall, the change in the UV spectrum is representative of this class of α-keto carbamates and is consistent with photocyclization to form the corresponding benzo[b]furan (2a) (Scheme 2). During irradiation of carbamate (2), the maxima originally at 267 nm is replaced by an intense absorption maxima at 300 nm. due to photocyclization to the conjugated benzo[b]furan (2a). Interestingly, the absorption due to the aryl π-π* transition (250-265 nm) decreases on photocyclization. Furthermore, beyond ca.340 nm, the absorption tail of the carbonyl chromophore disappears with increasing 350 nm photolysis. This bleaching of the long wavelength n-π* transition was only observed for 3,3',5,5'-tetramethoxybenzoin carbamate (2). Other derivatives e.g. (3) and (6) only show bleaching of the deep UV π-π* absorption while carbamates (4) and (5) show antibleaching. The propensity for bleaching of both the long wavelength tail (> 340 nm) and the deep UV absorption (ca. 250-265 nm) is a particularly attractive feature of this photogenerator (2). Based on the observed photobleaching of carbamate (2), this material holds particular promise as a photocatalyst in thick film applications at both deep UV and far UV exposure wavelengths. This is in contrast to the α-nitrobenzyl derived class of photobase generators in which anti-bleaching has restricted their use in some applications. The overlay plot (Figure 5) generated on solution photolysis of α-keto carbamates is comparable to that reported by Sheehan et. al. for 3',5'-dimethoxybenzoin acetate. This is encouraging in that photocyclization to liberate amine seemingly proceeds by an analogous pathway to that used to liberate carboxylic acids (vide infra). The photolysis of 3',5'-dimethoxy substituted carbamates, (2) through (6), is remarkably clean by TLC. In these instances, the respective benzo[b]furans, (2a) through (6a) are the major products, and only trace amounts of other by-products are present.
Preparative photolysis

Preparative photolysis of cyclohexyl carbamates of both 3,3',5,5'-tetramethoxybenzoin (2) and 3',5'-dimethoxybenzoin (3) proceeded smoothly in acetonitrile. In both cases, the major products were cyclohexylamine and the appropriate substituted benzo[b]furan (2a) and (3a), respectively. The fluorescent benzo[b]furans were isolated as the major photoproduts in excellent yields (80-85%) by flash chromatography. In the case of 3,3',5,5'-tetramethoxybenzoin cyclohexyl carbamate (2), a deoxybenzoin photoproduct (2b), formed by loss of the carbamoyloxy moiety, was also isolated in 7% yield from the photolysate. Formation of such products have also been observed during GC-MS product studies (vide supra). Comparison of the extinction coefficients of the starting carbamates with the respective benzo[b]furan photoproducts indicates that the photoproducts absorb significantly less light in the 250-265 nm region. For example, the extinction coefficient of 3',5'-dimethoxybenzoin cyclohexyl carbamate (3) at 254 nm is 12341 L mol\(^{-1}\) cm\(^{-1}\), while that of 2-phenyl-5,7-dimethoxybenzofuran (3a) is 5972 L mol\(^{-1}\) cm\(^{-1}\). This difference explains the deep UV photobleaching of these photoactive carbamates.

iii) Photochemistry of 2,2'-Di-substituted \(\alpha\)-Keto Carbamates

By \(^1\)H NMR spectroscopy, the 2,2'-diaryl substituted cyclohexyl carbamate (7) was observed to photorelease a significant amount of free cyclohexylamine (Table 1). However, no evidence for the formation of benzo[b]furan (7a) was available by \(^1\)H NMR spectroscopy. By GC-MS, only cyclohexylamine, dicyclohexylurea and N-benzoylecyclohexylamine could be identified. Infra-red spectroscopy indicated that photocyclization proceeds somewhat sluggishly, with both the keto carbonyl stretch at 1700 cm\(^{-1}\) and the carbamate carbonyl stretch at 1727 cm\(^{-1}\) slowly disappearing on 350 nm Rayonet exposure. However, the rate of disappearance of the keto carbonyl stretch is faster than the carbamate carbonyl stretch and a weak absorption appears at 1685 cm\(^{-1}\). Overall, photolytic studies on the 2,2'-disubstituted \(\alpha\)-keto carbamates (7) resemble the less efficient 3',5'-dimethoxybenzoin carbamates (5) and (6). Based on these experimental observations, it seems likely that 2,2'-disubstituted \(\alpha\)-keto carbamates undergo amine photogeneration via a number of competing pathways, in which photocyclization plays only a minor role.

b) Solid-State Photosensitivity

i) Photochemistry of Unsubstituted \(\alpha\)-Keto Carbamates
Based on the results of monitoring the solution photochemistry of this class of benzoin derived base photogenerators, the solid state photodecomposition of these α-keto carbamates was investigated in limited detail. The solid state photodecomposition was conveniently monitored by irradiating thin poly(methacrylonitrile) (PMAN) films containing a few mol % benzoin carbamate (I) and following the changes in the UV spectrum. The change in the UV spectrum is inconsistent with efficient photocyclization to form the corresponding benzo[b]furan (1a) because the long wavelength n-π* transition of the carbonyl chromophore does not disappear. This observation is in agreement with the solution IR study, in which both carbonyl absorptions did not disappear during UV exposure. Although the mechanism of cyclohexylamine photogeneration may not proceed solely via photocyclization, benzoin cyclohexyl carbamate (I) has been reported to be an efficient base photogenerator for deep UV lithography. This probably stems from the ability of benzoin cyclohexyl carbamate (I) to photoliberate cyclohexylamine via alternate pathways and in such applications is likely aided by the observed bleaching at 254 nm.

ii) Photochemistry of Aryl substituted α-Keto Carbamates

The intramolecular nature of the photocyclization of 3',5'-dimethoxy substituted benzoins suggests that photogeneration of amines can be effected in the solid state. This proved to be the case with all benzoin carbamates containing this substitution pattern undergoing photodecomposition as monitored by infrared and UV spectroscopy. The solid state photoreactivity was established by coating sodium chloride and quartz substrates with 1 μm thick poly(methacrylonitrile) (PMAN) films containing approximately 5 mol % of the benzoin carbamate.

Photocyclization was monitored using infrared spectroscopy to follow the simultaneous disappearance of both keto and carbamate carbonyl absorptions with increasing UV exposure dose. Photocyclization to liberate base was found to be general in nature, and occurred on irradiation over a broad range of wavelengths ranging from deep UV to far UV. For instance, 3,3',5,5'-tetramethoxybenzoin carbamate (2) undergoes photocyclization to liberate cyclohexylamine on irradiation 254, 313, 336 and 365 nm. The observed change in the infrared spectrum on 254 nm photolysis of such films is illustrated for the photocyclization of α-keto carbamate (2) (Figure 6). There is a substantial difference in the extinction coefficients at 254 and 365 nm in these carbamates. For example, the extinction coefficient of carbamate (2) at 254 nm is 6180 L mol⁻¹ cm⁻¹, while at 365 nm it is 102 L mol⁻¹.
In a ca. 1 μm thick PMAN films containing 5 mol % of carbamate (2), this amounted to absorption of 78.8% of the light at 254 nm and only 7.1% of the light at 365 nm. However, despite the low absorption at 365 nm, the photochemistry is still relatively efficient at such long wavelengths.

The result of following the changes in the UV spectrum of thin films of PMAN containing about 5 mol % α-keto carbamate (3) on 254 nm exposure is shown in Figure 7. The deep UV maxima at 270 nm is replaced by an intense absorption maxima at 300 nm. The change in the UV spectrum is representative for this class of α-keto carbamates and is consistent with photocyclization to the conjugated benzo[b]furan taking place in the solid state. This spectral change is accompanied by bleaching of the deep UV π-π* absorption in the region 250-265 nm. Such bleaching is desirable in many applications and many of these base photogenerators hold significant promise as photocatalysts particularly in thick film applications where photobleaching at the exposure wavelength is critical.

Furthermore, the similarity between the solid state and the solution photolyses, as monitored by IR and UV spectroscopy implies the photochemistry is not appreciably altered in the solid state. This consideration is of importance in extending the utility of these base photogenerators to a variety of solid state applications.

iii) Photochemistry of 2,2'-Di-substituted α-Keto Carbamates

Based on the results of monitoring the photodecomposition of this class of α-keto carbamates in solution, the solid state photocyclization of these materials was not investigated in sufficient detail to warrant further discussion at this time.

3) Quantum Yield

In order to determine the photoefficiency of base photogeneration from α-keto carbamates, the standard equation (Eq. 1) defining quantum yield was rearranged to a more convenient form (Eq. 2). This equation expresses the photoefficiency of the benzoin photocyclization in a manner that allows the quantum yield to be evaluated experimentally.

\[
\Phi = \frac{\text{moles of } \alpha\text{-keto carbamate consumed}}{\text{einstein absorbed}} \quad \text{Equation 1}
\]
\[ \Phi = \frac{\text{starting film concentration (mol \cdot cm}^2) \times \% \text{ conversion}}{\text{radiation flux (einstins} \cdot \text{s}^{-1} \cdot \text{cm}^2) \times \text{time (s)} \times \% \text{ absorbed}} \]  

Equation 2

In the interest of evaluating the quantum yield for photocyclization over a range of common UV exposures wavelengths we were most interested in looking at \( \alpha \)-keto carbamates which possessed favorable UV characteristics in the range 250-400 nm. In this regard, 3,3',5,5'-tetramethoxybenzoin cyclohexyl carbamate (2) was considered a particularly attractive candidate because it possessed a weak but bleachable UV absorption maxima which extended beyond 350 nm. Accordingly, the solid state quantum yield for the photodecomposition of 3,3',5,5'-tetramethoxybenzoin cyclohexyl carbamate (2) was evaluated at 254, 313, 336 and 365 nm, as described in the Experimental Section. The quantum yields are shown in Table 2 and range between 0.05 and 0.07. In strict terms, these quantum yields are for the disappearance of the starting \( \alpha \)-keto carbamate (2). However, the quantum yields may be assumed to be representative of cyclohexylamine photogeneration. This assumption is considered valid because carbamate (2) has demonstrated clean generation of cyclohexylamine and benzo[b]furan, uncomplicated by side products, based on \(^1\)H NMR analysis of solution photolysates at low conversion. The quantum yields were evaluated at low % conversion (5-25%) by back extrapolating a plot of log quantum yield against number of quanta absorbed, to the point of zero quanta absorbed. Figure 8 illustrates the result of the quantum yield determination performed on 365 nm irradiation. Considering the limited sensitivity of the quantitative infrared technique for measuring such low % conversion values, the quantum yields at 254, 313, 336 and 365 nm are in excellent agreement within experimental error. The overall quantum yield may be assumed to be ca. 0.06 and is largely independent of exposure wavelength. Assuming that photocyclization only proceeds through direct excitation of the keto carbonyl group then a more significant variation in quantum yield would be expected on 254 nm (aryl \( \pi-\pi^* \) excitation) versus 365 nm (carbonyl \( n-\pi^* \) excitation) exposure. The fact that the quantum yield is not significantly affected by the exposure wavelength suggests energy transfer within the molecule is particularly efficient and that photodecomposition proceeds via different mechanisms depending on the exposure wavelength. Since GC-MS product studies on the photolysis of 3,3',5,5'-tetramethoxybenzoin cyclohexyl carbamate (2) indicate that the same products are formed on 254, 313, 336, and 365 nm photolysis, it seems likely that the photocyclization proceeds via the same excited keto moiety. This
being the case, exposure beyond 300 nm likely proceeds via direct carbonyl n-\(\pi^*\) excitation. On the other hand, exposure at 254 nm likely involves some secondary energy transfer from the excited 3,5-dimethoxyphenyl moiety to the keto carbonyl group.

The solid state quantum yield of ca. 0.06 for amine photogeneration is significantly lower than that observed for the photoliberation of acetate and phosphate in solution\textsuperscript{17,19,20}. This result can be rationalized as follows. First, reduced molecular motion in the solid state probably increases the difficulty of attaining appropriate conformations for photoreaction. Second, phosphate and acetate are significantly better leaving groups than the carbamoyloxy moiety.

In the case of benzoin phosphate, photocleavage is quenched by common triplet quenchers indicating that the reaction proceeds via the triplet manifold. On the other hand, the photocleavage of carboxylate and phosphate esters of meta methoxy activated benzoins is not quenched by naphthalene. This suggests the photocleavage in such systems proceeds via a singlet excited state or via a short lived triplet state. Based on the available mechanistic data, Givens \textit{et al.}\textsuperscript{19} have proposed an ion-pair mechanism to explain the photocyclization of desyl phosphate esters. This mechanism may be extended to rationalize the photogeneration of amines from 3',5'-dimethoxybenzoin carbamates, as shown in Scheme 3. On irradiation, n-\(\pi^*\) excitation of the carbonyl group leads to formation of the excited triplet state (8). Subsequent homolysis produces a radical pair (9) which undergoes rapid electron transfer to yield an ion pair (10). After ring closure, the liberated anion acts as a base. This base removes the bridgehead proton from (11), forming the benzo[b]furan photoproduct. The unstable carboxylated amine then loses carbon dioxide, liberating the free amine. The benzylic carbocation intermediate (10) is postulated based on the photochemical meta effect\textsuperscript{21}. In the photochemically excited state, meta methoxy groups are known to be particularly efficient in stabilization of a developing positive charge in a benzylic site. In methoxy substituted benzenes, the electron density meta to the methoxy substituents is enhanced in the photochemical excited state. In contrast, a significantly lower electron density exists at the meta sites in the ground state. Since the photocyclization pathway is the preferred mode of photocleavage for 3',5'-dimethoxybenzoinyl carbamates, it seems probable that photocyclization proceeds via a meta benzylic carbocation as shown in Scheme 3. However, since photocleavage of the related phenacyl protecting group\textsuperscript{15} is known to proceed by C-O homolysis followed by radical combination, this mechanism should also be considered. In this case, the benzoinyl radical formed by homolysis
abstracts a hydrogen atom from the solvent to give a deoxybenzoin product. Indeed, a small amount of a deoxybenzoin product (2b) was isolated from the photolysate from 3',5'-dimethoxybenzoin carbamate (2). This indicates the C-O homolysis / radical combination mechanism is only a minor photocleavage pathway. Rather, the major photocleavage pathway requires that the benzoynyl radical (9) preferentially undergoes rapid electron transfer to give an ion pair (10) which reacts as shown in Scheme 3. Similar mechanistic pathways involving competition between pure homolysis and combined homo- / heterolytic cleavage have recently been shown to be important in the photochemistry of meta methoxy benzylic esters.22

Similar mechanistic arguments may also be used to explain the enhanced photocyclization yields in meta methoxy substituted benzoynyl carboxylate esters. In contrast, the photochemistry afforded by benzoynyl phosphate seems to be unique in undergoing efficient photocyclization without meta stabilization. This indicates the nature of the leaving group as well as the structure of the benzoynyl chromophore play an integral role in determining the photocleavage mechanism of substituted benzoins.

4) Thermal Stability

For benzoyn carbamates to be potentially useful in the formulation of resist materials or imaging systems, they must be thermally stable under standard lithographic processing conditions that may include prolonged heating at temperatures upwards of 150°C. The thermal properties of these α-keto carbamates was determined by a combination of differential scanning calorimetry (DSC) and thermogravimetric analysis (TGA). As shown in Table 3, this requirement was generally fulfilled for carbamates derived from α-hydroxy ketones. This is further illustrated by the DSC curve for carbamate (2) (Figure 9), which shows an exotherm at 118°C corresponding to the melting point of the material. The decomposition temperature was determined by TGA, which shows that near quantitative weight loss occurs on heating carbamate (2) above 290°C in a nitrogen atmosphere (Figure 10). In an air atmosphere, the decomposition temperature is approximately 60°C lower for carbamate (2) (Table 3). Furthermore, it is known that many common photoacid generators exhibit decreased thermal stability in a phenolic matrix.23 In contrast, these base photogenerators retain their thermal stability in a phenolic environment. For instance, both the melting point and decomposition temperature of carbamate (2) were only lowered by 5°C on mixing with poly(vinyl phenol) as determined by DSC. Because of their thermal
stability, these materials offer a potential advantage over many common photocatalysts used in microlithography and polymer curing applications.

Conclusions.

Substituted benzoin carbamate derivatives of amines show great potential as photoprecursors of organic bases. These α-keto carbamates are cleaved by UV irradiation below 370 nm down to the deep UV both in solution and in the solid state. The substitution pattern on the aryl rings and at the 2-position was varied to determine the optimal chromophore structure. Cyclohexyl carbamates derived from 3',5'-dimethoxybenzoin and its substituted analogs were particularly photoefficient affording high yields of free cyclohexylamine and the corresponding substituted benzo[b]furan as determined by $^1$H NMR spectroscopy and GC-MS product studies. The photocleavage for this class of substituted carbamates proved particularly clean, affording minimal side products. In contrast the photochemistry of the o-nitrobenzyl class of photobase generators is particularly complex. The parent benzoinyl cyclohexyl carbamate and 2,2'-disubstituted α-keto carbamates also formed cyclohexylamine, but the photocleavage was complicated by other photoproducts besides the expected benzo[b]furan. The solid state quantum yield for the photogeneration of cyclohexylamine from the cyclohexyl carbamate of 3,3',5,5'-tetramethoxybenzoin was found to be ca.0.06 over the range 250 to 370 nm. In terms of photoefficiency, these materials are less efficient than the best o-nitrobenzyl carbamates. However, many of the 3',5'-dimethoxybenzoinyl carbamates offer a significant advantage in that they are photobleachable. The combination of clean photochemistry and photobleachable absorption makes this class of base photogenerator ideally suited to thick film applications. These materials are currently being evaluated in a number of applications requiring base photogeneration. These include polymer crosslinking for imaging and curing, and polymer modification reactions of utility in coatings, microlithography and other imaging applications.

Experimental Procedures.

General Procedures

Melting points and boiling points are uncorrected, melting points were recorded on a Gallenkamp melting point instrument. Unless stated otherwise, infrared spectra were obtained as KBr Disks using a
Nicolet FTIR/44 spectrometer. Ultraviolet-visible spectra were measured in acetonitrile solution using a Hewlett-Packard 8450 Diode Array Spectrophotometer. NMR spectra were recorded in CDCl₃ on a Bruker AF250 spectrometer using tetramethylsilane as internal standard. DSC was performed using a Perkin Elmer cell. TGA was performed on a Perkin Elmer thermobalance. Both techniques used a heating rate of 10°C/min. GC was performed on a Hewlett-Packard 5890 Series II Gas Chromatograph equipped with a 5% phenyl methyl silicone fused silica capillary column operating in split injection mode. GC-MS was performed on a Hewlett-Packard GC interfaced to a Hewlett Packard mass-selective detector. Solution photolyses were performed in a Rayonet Photochemical Reactor RPR100 equipped with 16 RPR3500A lamps. Microanalyses were performed by Desert Analytics, Tucson, AZ.

Quantum Yield Determination

a) Sample Preparation

The samples used to determine quantum yields were prepared in the following manner. Poly(methacrylonitrile) (PMAN) (11.0 wt %) was dissolved in nitromethane. The polymer solution was filtered through a 0.45µm Acrodisc-PTFE filter and diluted with nitromethane to 8.91 wt %. Approximately 5.36 mol % (relative to polymer) of 3,3′,5,5′-tetramethoxybenzoin cyclohexyl carbamate (2) was added and the solution stirred until the carbamate dissolved. The resulting solution was applied to standard silicon, sodium chloride and quartz substrates with a Headway Research Spin-coater. All films were dried at 90°C for 15 min. and then in vacuo for 12h. Film thicknesses were measured on a Tencor Alpha Step 200 and were in the range 1.1-1.3µm

b) Evaluation of Quantum Yield

In order to evaluate the quantum yield for the benzoin photocyclization, all the terms in Eq. 2 must be determined. This was achieved in the following manner. UV spectra of the sample films on quartz disks were recorded on a Hewlett Packard 8450A spectrophotometer. Thus by knowing the film absorbance and the extinction coefficient, the starting film concentration for each sample could be determined by using the Beer-Lambert law (Eqns 3-5) where A is absorbance, I₀ is the incident radiant flux, Iₗ is the transmitted radiant flux, ε is the molar extinction coefficient (L mol⁻¹ cm⁻¹), c is concentration (mol L⁻¹), and l is the path length (cm).
\[ A = \log_{10} \left( \frac{I_0}{I_t} \right) = \varepsilon c_l \quad \text{Equation 3} \]

\[ c_l = \frac{A}{\varepsilon} \quad \text{Equation 4} \]

\[ \text{starting film concentration} = \frac{c_l}{1000} = \frac{A}{1000\varepsilon} \quad \text{Equation 5} \]

The changes in the infrared absorbance spectrum of the sample films coated on sodium chloride disks were monitored by quantitative infrared spectroscopy. This was done with use of a Nicolet FTIR/44 spectrometer equipped with software for quantitative analysis. The photochemical rearrangement was followed by monitoring the disappearance of both the keto carbonyl stretch at 1700 cm\(^{-1}\) and the carbamate carbonyl stretch at 1716 cm\(^{-1}\) in relation to the polymer reference band. The polymer reference band used was the C-H stretch at 3050-2800 cm\(^{-1}\). The ratio (C=O/C-H) of the peak under area under the carbonyl absorption and the reference C-H absorption was calculated initially and then again after each successive UV exposure dose. The ratio of the amount of desyl substrate present after irradiation to the amount present initially multiplied by one hundred gives the percent desyl substrate remaining after exposure (Eq. 6).

\[ \% \text{ substrate remaining after exposure} = \frac{\text{final} \ (\text{area C} = \text{O band/area C - H band})}{\text{initial} \ (\text{area C} = \text{O band/area C - H band})} \times 100 \quad \text{Equation 6} \]

\[ \% \text{ conversion of substrate} = 100 - \% \text{ substrate remaining} \quad \text{Equation 7} \]

From this, the percent conversion caused by each exposure was found by subtraction (Eq. 7). For each exposure, two measurements of percent conversion were made on each disk. Conversion values between 5 and 25% were then used directly in the quantum yield calculation. At all exposure wavelengths, plots of % conversion against exposure time for 3,3',5,5'-tetramethoxybenzoin cyclohexyl carbamate (2) showed a near linear response within this region, thus implying that all the incident light was being absorbed by the substrate and therefore the inner filter effect was minimal at this extent of
reaction. This is illustrated for the 313 nm exposure of 3,3',5,5'-tetramethoxybenzoin cyclohexyl carbamate (2) (Figure 11).

UV exposure was performed using an OAI exposure tool in conjunction with an Oriel narrow band pass filter at the following exposure wavelengths 254, 313, 336, and 365 nm. The sample coated sodium chloride disks were placed at a distance from the lamp where the light output varied no more than 5% over the exposed region and was in the range 0.20-6.23 mW cm⁻² depending on the exposure wavelength. The radiation flux at each exposure wavelength was calculated by dividing the light intensity by the energy of one einstein of irradiation at that particular wavelength. Exposure times were obtained directly from an OAI exposure timer.

The percent absorbed is the amount of light absorbed by the sample film at that particular exposure wavelength and was calculated by setting the value for the incident light equal to 100% and subsequently rearranging the Beer-Lambert law (Eqns 8-11) where \( I_{abs} \) is the absorbed radiant flux.

\[
I_{abs} = I_o - I_t \quad \text{Equation 8}
\]

\[
I_t = I_o \times 10^{-\varepsilon cl} \quad \text{Equation 9}
\]

\[
I_{abs} = I_o \left(1 - 10^{-\varepsilon cl}\right) \quad \text{Equation 10}
\]

\[
\% \text{ absorbed} = 100 \left(1 - 10^{-A}\right) \% \quad \text{Equation 11}
\]

Thus with all the terms in Eq. 3 having been evaluated, the quantum yield at each exposure wavelength was calculated for several exposure doses. A plot of log quantum yield against the number of quanta absorbed was then used to determine the absolute quantum yield by back extrapolation to zero quanta absorbed. This method of determination corrects for any internal filtering by the benzofuran photo by-product. This procedure was used to calculate the quantum yield for photocyclization of 3,3',5,5'-tetramethoxybenzoin cyclohexyl carbamate (2) at 254, 313, 336 and 365 nm.
Preparative Photolysis

Photogeneration of Cyclohexylamine and Preparation of 2-(3,5-Dimethoxyphenyl)-5,7-dimethoxybenzo[b]furan (2a).

A pyrex flask containing a solution of carbamate (2) (0.234g, 0.512mmol) in dry, degassed acetonitrile (100ml) was irradiated for 2h in a Rayonet photolysis chamber fitted with 350 nm lamps. During this time, samples were removed and monitored by GC which indicated that cyclohexylamine was being liberated. After 2h, TLC confirmed the formation of one major photoproduct along with other trace polar photoproducts. The solvent was removed in vacuo and the residual oil purified by flash chromatography (10% ethyl acetate / 90% hexane) to give 2-(3,5-dimethoxyphenyl)-5,7-dimethoxybenzo[b]furan (2a) as a white solid (0.137g, 85%). An analytically pure sample was obtained by recrystallization from ethyl acetate-hexane.

M.p. 106-108°C (Lit.20 m.p. 110-111°C). Microanalysis: Calculated for C_{18}H_{18}O_{5} (314.32). C 68.78, H 5.77; Found C 68.58, H 5.46%. UV: $\varepsilon_{286sh}$ 24205, $\varepsilon_{300}$ 28727, $\varepsilon_{318sh}$ 19136. IR ν 1611, 1602, 1594, 1491, 1342, 1204, 1160, 1149, 1036, 824 cm$^{-1}$. $^1$H NMR δ 3.86 and 4.02 (each 3H, each s, Ar(1) 5.7-OCH$_3$), 3.88 (6H, s, Ar(2) 3.5-OCH$_3$), 6.47 (1H, t(J$_m$=2.2Hz), Ph 4-H). 6.48 (1H, d(J$_m$=2.2Hz), Bf 6-H), 6.63 (1H, d(J$_m$=2.2Hz), Bf 4-H), 6.95 (1H, s, Bf 3-H), 7.03 ppm (2H, d(J$_m$=2.2Hz), Ph 2,6-H). $^{13}$C NMR δ 55.52 (q), 56.08 (q), 55.83 (q), 94.37 (d), 97.34 (d), 100.95 (d), 102.45 (d), 102.99 (d), 130.38 (s), 132.15 (s), 139.37 (s), 145.44 (s), 156.31 (s), 156.86 (s), 161.00 (s) ppm.

Further elution of the column allowed isolation of one minor photoproduct as a yellow oil (0.011g, 7%). This material was tentatively identified as 1-(3,5-dimethoxyphenyl)-2-(3,5-dimethoxyphenyl)ethaneone (2b) by $^1$H NMR spectroscopy.

$^1$H NMR δ 3.77 and 3.83 (each 6H, each s, Ar(1) and Ar(2) 3,3',5,5'-OCH$_3$), 4.18 (2H, s, 2-CH$_2$), 6.37 (1H, t(J$_m$=2.2Hz), Ar(2) 4-H), 6.43 (2H, d(J$_m$=2.2Hz), Ar(2) 2,6-H), 6.65 (1H, t(J$_m$=2.2Hz), Ar(1) 4-H), 7.16 ppm (2H, d(J$_m$=2.2Hz), Ar(1) 2,6-H).

Photogeneration of Cyclohexylamine and Preparation of 2-phenyl-5,7-dimethoxybenzo[b]furan (3a).

A pyrex flask containing a solution of carbamate (3) (0.196g, 0.493mmol) in dry degassed acetonitrile (100ml) was irradiated for 2h in a Rayonet photolysis chamber fitted with 350nm lamps.
During this time samples were removed and monitored by GC which indicated that cyclohexylamine was being liberated. After 2h, TLC confirmed the formation of one major photoproduct along with other trace polar photoproducts. The solvent was removed in vacuo and the residual oil purified by flash chromatography (10% ethyl acetate / 90% hexane) to give 2-phenyl-5,7-dimethoxybenzo[b]furan (3a) as a white solid (0.100g, 80%). An analytically pure sample was obtained by recrystallization from hexane.

M.p. 68-69°C (Lit. 17 m.p. 89.1-89.6°C). Microanalysis: Calculated for C_{16}H_{14}O_3 (254.27). C 75.57, H 5.55; Found C 75.50, H 5.30%. UV: ε(246) 9301, ε(292sh) 25067, ε(300) 25817, ε(317sh) 15855. IR ν 1606, 1479, 1429, 1594, 1218, 1214, 1147, 824, 765, 761 cm⁻¹. ¹H NMR δ 3.87 and 4.03 (each 3H, each s, Ar(1) 5.7-OCH₃), 6.47 (1H, d(Jₘ=2.2Hz), Bf 6-H), 6.64 (1H, d(Jₘ=2.2Hz), Bf 4-H), 6.97 (1H, s, Bf 3-H), 7.35 (1H, m approximates to t, Ph 4-H), 7.43 (2H, m approximates to dt, Ph 3,5-H), 7.88 ppm (2H, m approximates to d, Ph 2,6-H). ¹³C NMR δ 55.11 (q), 55.83 (q), 94.35 (d), 97.21 (d), 101.87 (d), 124.92 (d), 128.46 (d), 128.69 (d), 130.36 (s), 130.49 (s), 139.42 (s), 145.48 (s), 156.49 (s), 156.84 (s) ppm.

Acknowledgement. Support of portions of this work done at Cornell University by the Office of Naval Research is acknowledged with thanks.
References.


Scheme 1

\[ \text{MeO} \quad \text{OCONR}_1R_2 \quad \text{MeO} \]

\[ \text{hv} \quad 250-365 \text{nm} \]

\[ \text{MeO} \quad \text{O} \quad \text{Me} \]

Where \( R_1 = H, R_2 = \text{C}_6\text{H}_{11} \),
\( R_1, R_2 = \text{-(CH}_2\text{)}_5 \).
Scheme 2

Where others includes:

Where:
- \( R_1 = R_2 = H, R_3 = 4-OMe \) (1)
- \( R_1 = H, R_2 = 3,5-OMe, R_3 = H \) (2)
- \( R_1 = 4-OMe, R_2 = 3,5-OMe, R_3 = H \) (3)
- \( R_1 = 4-SMe, R_2 = 3,5-OMe, R_3 = H \) (4)
- \( R_1 = 3,4-C_6H_4(2-Naphthyl), R_2 = 3,5-OMe, R_3 = 3,5-DiOMePh \) (5)
- \( R_1 = H, R_2 = 3,5-OMe, R_3 = 3,5-DiOMePh \) (6)
Scheme 3

1) hν
2) ISC

Homolysis

Ring Closure

SET
<table>
<thead>
<tr>
<th>Carbamate</th>
<th>% Conversion&lt;sup&gt;a&lt;/sup&gt;</th>
<th>δ(3-Bf-H)&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benzoin (1)</td>
<td>75</td>
<td>Not Observed</td>
</tr>
<tr>
<td>3',5',5'-TetraMeO (2)</td>
<td>87</td>
<td>7.14</td>
</tr>
<tr>
<td>3',5'-DiMeO (3)</td>
<td>93</td>
<td>7.12</td>
</tr>
<tr>
<td>3',4,5'-DiMeO (4)</td>
<td>98</td>
<td>6.96</td>
</tr>
<tr>
<td>3',5'-DiMeO-4-MeS (5)</td>
<td>37</td>
<td>7.07</td>
</tr>
<tr>
<td>3',5'-DiMeO-2-Naphth (6)</td>
<td>62</td>
<td>7.25</td>
</tr>
<tr>
<td>2,2-Di-(3',5'-MeO) (7)</td>
<td>62</td>
<td>Not Observed</td>
</tr>
</tbody>
</table>

<sup>a</sup> % Conversion calculated from the integral ratio of the photogenerated cyclohexylamine methine (CH, m, δ 2.52) and the aromatic resonances after 60min 350nm Rayonet exposure.

<sup>b</sup> Chemical shift for resonance of 3-H in benzofuran photoproduct in MeCN-d3 quoted ppm downfield from TMS.
TABLE 2

Quantum Yields\textsuperscript{a} For The Photocyclization Of 3,3',5,5'-Tetramethoxybenzoin Carbamate (2) (With Concomitant Liberation of Cyclohexylamine)

<table>
<thead>
<tr>
<th>$\Phi_{254\text{nm}}$</th>
<th>$\Phi_{313\text{nm}}$</th>
<th>$\Phi_{336\text{nm}}$</th>
<th>$\Phi_{365\text{nm}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.053</td>
<td>0.071</td>
<td>0.051</td>
<td>0.065</td>
</tr>
</tbody>
</table>

\textsuperscript{a}Quantum yield evaluated from a plot of log $\Phi$ versus the number of quanta absorbed, by back extrapolation to zero quanta absorbed, from data taken at conversion values unaffected by the inner filter effect.
<table>
<thead>
<tr>
<th>Carbamate</th>
<th>Melting Point</th>
<th>Decomposition Temperature&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benzoin (1)</td>
<td>87-88°C</td>
<td>193°C</td>
</tr>
<tr>
<td>3,3',5,5'-TetraMeO (2)</td>
<td>116-118°C</td>
<td>233°C</td>
</tr>
<tr>
<td>3',5'-DiMeO (3)</td>
<td>149-151°C</td>
<td>203°C</td>
</tr>
<tr>
<td>3',4, 5'-DiMeO (4)</td>
<td>105-107°C</td>
<td>218°C</td>
</tr>
<tr>
<td>3',5'-DiMeO-4-MeS (5)</td>
<td>152-154°C</td>
<td>252°C</td>
</tr>
<tr>
<td>3',5'-DiMeO-2-Naphth (6)</td>
<td>161-162°C</td>
<td>230°C</td>
</tr>
<tr>
<td>2,2-Di-(3',5'-MeO) (7)</td>
<td>170-172°C</td>
<td>231°C</td>
</tr>
</tbody>
</table>

<sup>a</sup> Decomposition temperature was taken as the temperature of 5% weight loss as determined by Thermogravimetric Analysis (in air).
Figure Titles

Figure 1: Change in $^1$H NMR aromatic resonances of carbamate (2) as a 38.6 mM solution in THF-d8 with increasing 350 nm photolysis in Rayonet.

a) Prior to exposure.
b) After 15min exposure.
c) After 30min exposure.
d) After 60min exposure.
e) After 90min exposure.

Figure 2: Plot of % conversion against exposure time for carbamate (2) as a 38.6 mM solution in THF-d8 on 350 nm exposure in Rayonet.

Figure 3: Change in carbonyl region of carbamate (3) as a 11.8 mM solution in MeCN with increasing 350 nm photolysis in Rayonet.

a) Prior to exposure.
b) After 5min exposure.
c) After 10min exposure.
d) After 15min exposure.
e) After 20min exposure.
f) After 30min exposure.
g) After 45min exposure.
h) After 60min exposure.
i) After 120min exposure.

Figure 4: Change in N-H stretching region of carbamate (3) as a 11.8 mM solution in MeCN with increasing 350 nm photolysis in Rayonet.

a) Prior to exposure.
b) After 5min exposure.
c) After 10min exposure.
d) After 15min exposure.
e) After 30min exposure.
f) After 60min exposure.
g) After 120min exposure.
h) After 240min exposure.

Figure 5: Change in UV spectrum of carbamate (2) as a 6.164x10$^{-5}$M solution in MeCN with increasing 350 nm photolysis.

a) Prior to exposure.
b) After 15s exposure.
c) After 30s exposure.
d) After 45s exposure.
e) After 75s exposure.
f) After 120s exposure.
g) After 180s exposure.
h) After 300s exposure.
i) After 450s exposure.
Figure 6: Change in IR spectrum of carbamate (2) (5.36 mol%) in a 1.265µm thick PMAN film with increasing 254 nm exposure dose.

a) Prior to exposure.
b) After 50mJcm⁻² exposure.
c) After 100mJcm⁻² exposure.
d) After 200mJcm⁻² exposure.
e) After 300mJcm⁻² exposure.
f) After 400mJcm⁻² exposure.
g) After 500mJcm⁻² exposure.
h) After 750mJcm⁻² exposure.
i) After 1.0Jcm⁻² exposure.

Figure 7: Change in UV spectrum of carbamate (3) (4.67 mol%) in a 1.375µm thick PMAN film with increasing 254 nm exposure dose.

a) Prior to exposure.
b) After 30mJcm⁻² exposure.
c) After 75mJcm⁻² exposure.
d) After 125mJcm⁻² exposure.
e) After 200mJcm⁻² exposure.
f) After 300mJcm⁻² exposure.
g) After 500mJcm⁻² exposure.
h) After 1.0Jcm⁻² exposure.

Figure 8: Plot of log quantum yield against number of quanta absorbed for 365 nm exposure of carbamate (2) (5.36 mol%) in a 1.265µm thick PMAN film.

Figure 9: Differential scanning calorimetry trace for carbamate (2).

Figure 10: Thermogravimetric analysis trace for carbamate (2).

Figure 11: Plot of % conversion against exposure dose for 313 nm exposure of carbamate (2) (5.36 mol%) in a 1.265µm thick PMAN film.
Fig 2

% Conversion

Time (s) x10^3
The graph shows the % transmittance of samples exposed to increasing laser doses with a 1.41 micron thick PAM matrix. The samples were exposed to 100mj/sqcm with a dose at 254nm.
Fig 7

JFC1/11/25A1-4.67MOL7 in 1.375UM PMAN FILM, 254NM EXPOSURE

a) Prior 60 exposure
b) After 30mS/cm² 254nm exposure
c) " 75 " " "
d) " 125 " " "
e) " 200 " " "
f) " 300 " " "
g) " 500 " " "
h) " 15 " " "

WAVELENGTH (nm)
Curve 1: TGA

Sample Weight: 2539 mg

JFC1646/67A1

% Weight (Wt %)

Temperature (°C)

0.0 10.0 20.0 30.0 40.0 50.0 60.0 70.0 80.0 90.0 100.0

0.0 100.0

Time

TEMP1: 30.0 °C TIME1: 0.0 min RATE1: 10.0 °C/min
TEMP2: 400.0 °C TIME2: 0.0 min RATE2: 10.0 °C/min
TEMP3: 900.0 °C TIME3: 0.0 min

JFC

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7 Series Thermal Analysis System
Mon Jun 6 20:09:09 1994