Review

Benzil Photoperoxidations in Polymer Films and Crosslinking by the Resultant Benzoyl Peroxides in Polystyrene and Other Polymers

Ivan Lukáč 1,*, Branislav Husár 1,†, Martin Danko 1 and Richard G. Weiss 2

Abstract: Benzil (BZ) can be converted almost quantitatively to benzoyl peroxide (BP) in aerated polymer films upon irradiation at >400 nm (i.e., the long-wavelength edge of the \( n \rightarrow \pi^* \) absorption band of BZ, where BP does not absorb). Here, we summarize results for the photoperoxidation of BZ structures with molecular oxygen, principally in glassy polymer matrices. Some of the polymers are doped directly with BZ or its derivatives, and others, contain covalently attached BZ pendant groups from which BP groups are derived. While the decomposition of low-molecular-weight BP doped into polymer films (such as those of polystyrene (PS)) results in a net decrease in polymer molecular weight, thermal decomposition of pendant BP groups is an efficient method for chain crosslinking. Crosslinking of PS films doped with a molecule containing two covalently linked BZ or BP groups proceeds in a similar fashion. Free radicals from the covalently attached BP allow grafting of new monomers, as well. Additionally, the use of radiation filtered through masks has been used to create patterns of polymers on solid surfaces. Crosslinking of photodegradable poly(phenyl vinyl ketone) with BP structures obtained by photoperoxidation of BZ structures for the preparation of photodegradable polymer networks is described as well. In sum, the use of BZ and BP and their derivatives offers simple and convenient routes for modifying polymer chains and, especially, for crosslinking them. Specific applications of each use and process are provided. Although applications with PS are featured here, the methodologies described are amenable to a wide variety of other polymers.

Keywords: benzil; photoperoxidation; benzoyl peroxide; polymeric benzoyl peroxide; crosslinking; polystyrene

1. Introduction

Photon-induced transformations of 1,2-diphenylethanedione (benzil, BZ) and related compounds into benzoyl peroxide (BP) in glassy polymer matrices are the basis for this review. It will primarily treat photoperoxidations induced by molecular oxygen of BZ, compounds containing two BZ structures dissolved in polymers, and BZ groups covalently appended to polymers. The courses of the subsequent thermal or photochemical decompositions of the resultant BP photoproducts and their influence on the polymer properties are also described. They lead to a variety of crosslinks in the polymers. Polystyrene (PS) is featured here as a polymer for investigation.

A common method for synthesizing crosslinked PS involves copolymerization of styrene and a monomer with two or more vinyl groups. However, this method is not readily feasible for the preparation of crosslinked thin polystyrene films due to chemical, environmental and other considerations that accompany polymerization. Such films are...
prepared usually by the removal of solvents from solutions on substrates. Although less frequently used, post-polymerization crosslinking of PS offers much greater flexibility and specificity, especially in those cases which require radiation to form the needed crosslinks (e.g., the transformation of polymeric materials, such as photoresists, into micron- and submicron-sized structures for microelectronics [1,2]). Currently, the most commonly employed photoresists involve reactions that create photo-acids or photo-bases [3] in situ [4,5].

Selective surface modification and functionalization processes that involve the introduction of crosslinks have become increasingly important today to produce chemical or biosensors, especially for classical resist applications in the fields of interface engineering and nanotechnology [6,7]. Photo- or thermo-active agents to achieve these crosslinks have included a variety of reagents (e.g., nitrenes, carbenes, excited triplet states of carbonyl-containing compounds or sensitized decomposition of a perester) that can be covalently linked to a polymer or added as a low-molecular dopant containing two active groups that can react directly with a polymeric substrate to form a network [1]. In this regard, the photoperoxidation of BZ structures in the air (by molecular oxygen) is well suited to produce crosslinks within thin layers of polymers.

Additionally, there are clear advantages to linking the crosslinking agent directly to a polymer over adding a non-covalently linked dopant. Principal among these are the higher efficiency of crosslinking and the inability of the active units to separate into microphases over time, thereby reducing the efficiency of crosslinking and the distribution of the crosslinks; well-defined layers, patterns and images may be obtained. However, in all cases, the crosslinking agent must be compatible with and appropriate for the type of polymer being employed.

There are many specific examples of how these issues have been addressed. For instance, self-crosslinkable polymers have been prepared by post-modification of PS with reactive groups such as maleimide [8] or iodide [9] or by post-modification of poly(4-chloromethyl styrene) with light-sensitive structures such as benzophenone, stilbazolium, fluorenone, carbazole and sulphonyl azide [10]. Self-crosslinkable PS can also be prepared by copolymerization of styrene and a monomer containing a reactive functional group (e.g., azide [11] and phenylindene [12]). The photosensitized cleavage of peresters in copolymers containing p-vinylbenzophenone-p'-tert-butyl perbenzoate units has been found to be reasonably efficient [13,14]. Additionally, high crosslinking efficiency in surface layers of nanometer-range thicknesses has been induced by the thermal coupling reaction of two pendant benzocyclobutene structures [15] in melts and by azide-alkyne [16] groups.

Low-molecular-weight crosslinking agents such as very reactive difunctional Friedel-Crafts reagents [17] have been used in bulk with PS, and bisazides [18] have been doped into synthetic polymers for crosslinking their films. Perfluorophenylazides, which produce very high insertion yields, have been used to introduce functional groups, especially into synthetic polymers [7]. Additionally, a highly efficient negative photoresist based on poly(vinyl phenol) has been developed using bis(perfluorophenyl azides) as the crosslinking agent [19]. A major advantage of low-molecular-weight reagents is the possibility to use one dopant for crosslinking a variety of polymers. Photoperoxidation and crosslinking by a low-molecular-weight molecule containing two 1,2-dicarbonyl groups, doped into PS films, has been shown to be a very efficient crosslinking method (vide infra). BZ-containing dopants are more soluble than the corresponding BP-containing dopants in many polymers. As a result, the BZ reagents can usually effect higher concentrations and more even distributions of crosslinks than can the BP ones. Moreover, the more thermally stable BZ groups can be heated, shaped, extruded, blended, etc. in polymer processing steps prior to the thermal initiation of crosslinking in which the thermally sensitive BP initiators are generated in situ.

Diacylperoxides and many dialkyazo compounds are exploited frequently in polymer chemistry as polymerization initiators and crosslinkers due to their ease of thermal decomposition to form radicals. Although there is ample literature on polymeric azo radical
initiators, [20] the formation and subsequent use of polymeric diacylperoxides has not been reported previously and is summarized here.

2. Photochemistry of Benzil in Solution in the Absence of Molecular Oxygen

Spectroscopic and Temporal Information about Excited BZ States

The conformations of BZ excited states have been investigated extensively. Correlations among the excited state dynamics, emission properties and conformations of BZ have been reviewed in connection with the development of room-temperature phosphorescent materials [21]. Here, the focus will be on the properties of excited states of BZ dissolved in polymers or in solid solutions.

The rotational angles between the carbonyl groups of BZ differ in their ground and excited states. In the ground state, the groups are in a nonplanar (skewed) conformation with a twist angle of about 72°. Both first excited singlet and lowest triplet states have s-trans planar conformations (i.e., the −CO−CO− dihedral angle is ~180°) [22]. Therefore, both fluorescence and phosphorescence from BZ are dependent on external constraints imposed by the environment. Based upon the fact that the absorption maxima of BZ in toluene [23] and in a PS matrix [24] are both at 390 nm, we infer that the BZ ground state conformations are skewed to the same degree in the two media. The phosphorescence maximum of BZ in various solvents at 77 K (solids) is at ca. 522 nm (\(E_T = 229 \text{ kJ mol}^{-1}\)) and originates from a nonplanar (skewed cisoid) conformation; at room temperature in solution, BZ phosphorescence emanates from a planar, relaxed transoid conformation at ca. 565 nm (\(E_T = 212 \text{ kJ mol}^{-1}\)) in various solvents as well as in BZ/p-tert-butylcalix [6] and calix [8] arenes complexes [22]. The maximum of the phosphorescence from unsubstituted BZ in a glassy PMMA film (prepared by solution casting) is at 531 nm (\(E_T = 225 \text{ kJ mol}^{-1}\)) at 77 K and at 544 nm (\(E_T = 219 \text{ kJ mol}^{-1}\)) at 296 K [25]. This value indicates that the emitting species has an average −CO−CO− dihedral angle that is not completely relaxed; it is skewed with an angle greater than 72° but smaller than 180°.

Phosphorescence at 77 and 300 K from a PMMA block prepared by polymerization of methyl methacrylate in the presence of dissolved BZ exhibited two maxima that were ascribed to matrix-imposed, nearly skewed (525 nm) and trans planar (565 nm) geometries [26]. The phosphorescence maxima of BZ and its derivatives in polymer films of PS, PMMA, polyvinylchloride, polypropylene and polyethylene or BZ structures appended to the main chains in a homopolymer or copolymer have maxima at ca. 520–550 nm and are 10–20 nm lower at 77 K than at 296 K; the phosphorescence lifetimes were 3.2–5.5 ms at 77 K and 0.1–0.5 ms at 296 K [25,27,28].

Phosphorescence quenching in PS containing chain-ends labeled with BZ groups has been examined in degassed cyclohexane as a function of concentrations of added PS with and without anthracene end labels [29]. The phosphorescence lifetime was 85–90 µs at 298 K with a BZ group concentration of 10^{-5} \text{ mol dm}^{-3}, and it was independent of PS molecular weight (determined by viscometry) over a 7000–170,000 range. The inter-chain quenching did not follow a single exponential decay when BZ concentrations were \(\leq 10^{-4} \text{ mol dm}^{-3}\); double-exponential phosphorescence decay is proposed to result from absorption of light by benzil at two different times during the phosphorescence decay measurement. Experiments with the small molecule, azulene, as the quencher, indicated that although the diffusion coefficients for the PS-labeled anthracene and azulene are different, they are comparable.

BZ is an industrially important member of the class of molecules with 1,2-dicarboxyl functionality. The solution-phase photochemistry of 1,2-dicarboxyl compounds [30–33], as well as BZ itself, has been reviewed [34]. In this regard, BZ acts as a “photopromoter” in site-selective sp^3 C−H fluorination of terpenoid derivatives, and the photochemistry of BZ and other reagents to effect photo-fluorinations has been reviewed recently [35].

The lowest energy triplet state of BZ is formed almost quantitatively by the intersystem crossing of the initially excited singlet states. The symmetric Norrish type I cleavage (i.e., the C−C bond between the carbonyl groups) does not occur, at least as a principal
reaction pathway when the lowest \( n \to \pi^* \) transition band is irradiated in solution under an inert atmosphere \[36\]. It results in abstraction by the lowest energy triplet state of \( \text{BZ} \) when a good H-atom donor (RH) is present (Scheme 1). Subsequent combination and disproportionation reactions of the ketyl radical and radical formed from the hydrogen donor are responsible for the eventual photoproducts.

Scheme 1. Hydrogen abstraction by the excited triplet state of \( \text{BZ} \) from an H-donor, RH.

Additionally, recombination of two polymeric pendant ketyl radicals has been postulated to be responsible for the photodisproportionation of copolymer-bearing \( \text{BZ} \) structures in deaerated THF solution \[25\]. The lowest triplet states and ketyl radicals were detected by transient absorption measurements in this system.

When \( \text{BZ} \) is irradiated in the presence of amines, an exciplex can be formed. If the amine contains an \( \alpha \) C–H bond, the exciplex can collapse to an aminyl radical and the \( \text{BZ} \) ketyl radical \[37\]. Hydrogen abstraction is facilitated by tertiary amines and is an important photoinitiation process for light-induced radical polymerizations \[38–43\]. The use of 1,2-diketones as visible photoinitiators for the photodisproportionation of waterborne latex paints \[44\] and photoinitiation activity of water-soluble copolymers with pendant \( \text{BZ} \) chromophores has been reported as well \[45\].

A novel photoreaction of \( \text{BZ} \) has been found on a silica gel surface \[46\] and in the confined cavity of coordination compounds \[47\]. The formation of bibenzofuranone (BBF), furanone (FRN) and its hydroperoxide (FRNO) proceed by intramolecular trapping of an excited ketone carbonyl oxygen by a phenyl group.

Even under the anaerobic conditions used, it is likely that a trace amount of oxygen remained on the silica gel, and the formation of FRNO could not be avoided completely. In both types of media, the formation of an intermediate radical (FRNR) has been suggested. The emission spectra of 1,2-diketones have been measured, and their efficiency as initiators of photodegradation processes has been investigated in solid polymer matrices \[28\].

3. Photooxidation of Benzil in Solution with Molecular Oxygen

Molecular oxygen plays an important role in \( \text{BZ} \) photochemistry. In the absence of oxygen and H-atom donors (e.g., in deaerated benzene solutions), \( \text{BZ} \) is almost photochemically inert \[48,49\]. However, it is highly reactive in the presence of oxygen \[48–50\]. Thus, photooxidation of \( \text{BZ} \) in oxygenated benzene leads to several products: phenyl benzoate, benzoic acid, biphenyl and a small amount of \( \text{BP} \) \[48,49\]. Later, a considerable amount of peroxybenzoic acid was also identified \[51\]. When an aerated benzene solution of styrene copolymer with pendant \( \text{BZ} \) groups was irradiated at \( \lambda = 366 \) nm, low-molecular-weight cleavage photoproducts, as well as \( \text{BZ} \) fragments bound to the copolymer, were found \[25\].

Photooxidation of \( \text{BZ} \) has been studied extensively also as a participant in the sensitized photoepoxidation of olefins by molecular oxygen; \( \text{BZ} \) is the light-absorbing unit \[50–54\].
In addition, the mechanism of photooxidation of 1,2-diketones has been studied in the presence of olefins, focusing on the interaction of the triplet states of 1,2-diketones with O₂. Two types of reactions were found to occur competitively:

1. The formation of singlet oxygen (¹O₂) by energy transfer to ground-state triplet molecular oxygen (³O₂) [55] (note that ¹O₂ does not oxidize 1,2-dicarbonyl groups [49]);
2. Addition of ³O₂ to the triplet states of diketones, producing 1,4-biradicals.

The existence of such 1,4-biradicals has been assumed [25,48–51], but direct evidence (by laser flash photolysis) has been demonstrated only in the case of 2,2′-thenil (Scheme 2) [52]. The reaction of ground-state triplet molecular oxygen with the triplet state of BZ to form the 1,4-biradical is a spin-allowed process and is aided by the biradicaloid character of n→π⁺ triplet states of carbonyl compounds [50]. The addition of molecular oxygen to mono carbonyl and 1,2-dicarbonyl compounds and formation of 1,4-biradical and singlet oxygen species has been theoretically treated and reviewed, as well [56]. Once formed, the 1,4-biradical oxygen-adduct of BZ can decompose readily to benzoylperoxy and benzoyl radicals (Scheme 3) [50].

![Scheme 2](image)

**Scheme 2.** Photooxidation of 2,2′-thenil in the presence of oxygen: Triplet-oxygen adduct formation, its decomposition and possible uptake of molecular oxygen by a thiophene-2-carbonyl radical.

![Scheme 3](image)

**Scheme 3.** Photooxidation of BZ in aerated benzene solution: triplet-oxygen adduct formation, its scission and possible uptake of molecular oxygen by a benzoyl radical, followed by a competition between cage recombination (BP formation) and cage escape (other photoproducts) that incorporate benzoyl and benzoyleperoxy radicals.

Acylperoxy radicals are responsible for the epoxidation of olefins [50]. The mode of scission and 1,4-biradical and formation of two acylperoxy radicals is supported experimentally by the stoichiometry of the epoxidation reaction [51,57] ¹⁸O labeling experiments [51,58], the high yields of peroxybenzoic acid in the absence of reactive olefins and its low chemical yield in the presence of an olefin [51].
If the benzoyl and benzoylperoxy radicals remain in the same cage, they may combine as shown in Scheme 3 to produce BP. In solutions of low viscosity, cage-escape can dominate, and the other photoproducts may be formed (Scheme 3). The product distribution is similar to that from the thermal decomposition of BP [49].

However, despite extensive research, direct experimental evidence for the 1,4-biradical has not been obtained [52]. The investigations of the photoepoxidation of olefins [50–58] have concentrated on the role of benzoylperoxy radicals (inferring the existence of the 1,4-biradical species as a precursor). Although the same products are formed when BP is reacted thermally in benzene and from irradiation (>290 nm) of BZ in aerated benzene, the BP ground-state was shown not to be the key intermediate for product formation [49]. BZ is not an efficient photosensitizer for the decomposition of BP and, therefore, spin-allowed insertion of the ground triplet state of oxygen into the triplet state of BZ by radiation not absorbed by BP (>330 nm) would yield BP as the major product. Thus, after cage escape, the benzoyl radical reacts with molecular oxygen, resulting in two benzoylperoxy radicals.

4. Photoperoxidation of Benzil in Glassy Polymer Matrices in the Presence of Molecular Oxygen

BZ can be converted almost quantitatively to BP in aerated PS or poly(methyl methacrylate) (PMMA) films upon irradiation at >370 nm (i.e., the long-wavelength edge of the n→π* absorption band, where BP does not absorb) (Scheme 4) [24].

Scheme 4. Formation of BP in aerated glassy polymer matrices.

The photo-transformation in PS or PMMA films was followed spectroscopically as a function of conversion of BZ molecules by UV (i.e., changes in the π→π* and n→π* absorption band intensities) and by FTIR [24] (i.e., changes in the carbonyl absorption bands in the 1600–1800 cm⁻¹ region (Figure 1)). The presence of BP as virtually the sole photoprodut was confirmed by comparing the isolated product with an authentic sample [24]. The formation and then decomposition of pendant BP groups in a copolymer was studied by chemiluminescence [59].

Figure 1. Changes in FTIR spectra of BZ doped in a PS film during irradiation (>400 nm) at room temperature in air. The spectrum of PS has been subtracted from each spectrum for clarity.

The photoperoxidation process has also been studied upon irradiation (>400 nm) of BZ at ambient temperatures in aerated matrices of PS, PMMA, bisphenol A polycarbonate
(PC), bisphenol A polysulfone (PSF) and poly(vinyl chloride) (PVC) [60]. The decrease in absorbance from the 1,2-dicarbonyl group of BZ (at 1650–1700 cm\(^{-1}\)) and the increase in absorbance of the BP carbonyl group (at 1750–1800 cm\(^{-1}\)) were followed by FTIR spectroscopy in these cases. The relative rates of reaction are sensitive to the nature of the glassy polymer matrices and decrease in the order: PS > PC > PVC ~ PSF > PMMA (Table 1). The rate of BZ consumption in the polymers depends on several factors, including the concentration of dissolved molecular oxygen, its rate of diffusion and how the excited states of BZ interact with the walls of their polymer cages.

Table 1. Equilibrium molecular oxygen concentrations (\(^3\)O\(_2\)), corresponding coefficients of diffusion (D) and pseudo-first-order rate constants for BZ consumption (k) for irradiation of BZ in various polymer matrices at ambient temperature in air.

| Polymer Matrix | (\(^3\)O\(_2\)) (mol dm\(^{-3}\)) | D (cm\(^2\) s\(^{-1}\)) | k (h\(^{-1}\)) |
|---------------|-------------------------------|-------------------|--------------|
| PS            | 1.64 × 10\(^{-3}\)           | 1.10 × 10\(^{-7}\) | 2.35 × 10\(^{-2}\) |
| PC            | 4.78 × 10\(^{-3}\)           | 0.21 × 10\(^{-7}\) | 1.51 × 10\(^{-2}\) |
| PVC           | 0.28 × 10\(^{-3}\)           | -                 | 1.11 × 10\(^{-2}\) |
| PSF           | 2.27 × 10\(^{-3}\)           | 0.44 × 10\(^{-7}\) | 1.10 × 10\(^{-2}\) |
| PMMA          | 0.29 × 10\(^{-3}\)           | 0.38 × 10\(^{-7}\) | 0.69 × 10\(^{-2}\) |

The need to maintain rigid walls for the host cages if high yields of BP are to be obtained is indicated by two sets of experiments. Irradiation of BZ in an aerated, molded polyethylene film at ambient temperatures (i.e., above the glass transition temperature \(T_g\)) led to different chemistry, and the photoproduct(s) could not be identified. A part of the greater complexity may be due to an inhomogeneous distribution of guest molecules in the polyethylene films. Similarly, when the BZ-doped glassy films contained residual solvent, their irradiation led to increased absorbance at 1720 cm\(^{-1}\) and 1690 cm\(^{-1}\), probably from phenyl benzoate and benzoic acid, respectively. Thus, rigid cages that do not permit facile escape or large conformational changes of intermediates are required to affect almost quantitative photoperoxidation of BZ to BP.

It is interesting to note the differences in emission spectra of BZ in various polymers. In PE, where BP is not formed from irradiation of BZ at room temperature, no emission (phosphorescence) was observed; in glassy polymer matrices, intensive emission was observed. The difference was attributed to different extents of oxygen quenching in polycrystalline PE and glassy PMMA [28]. Note that molecular oxygen is much more soluble in solid PS than in solid PMMA.

Glassy polymer films containing BZ were also irradiated in the absence of molecular oxygen. Although the disappearance of BZ still occurs under these conditions, its rate is \(>7\times\) slower in PS and \(>4\times\) slower in PMMA than in the presence of oxygen. The photoproducts from BZ in the deaerated PS films were not determined. In the evacuated PMMA films, a low-intensity hydroxyl absorption (3600–3300 cm\(^{-1}\)) in the FTIR spectrum was detected after irradiation. It is probably a consequence of the abstraction of a hydrogen atom from PMMA by BZ triplets and subsequent combination of the radicals; some BZ-like moieties were covalently attached to the polymer chains during irradiation. These results demonstrate that despite the slow diffusion and limited concentration of molecular oxygen in the glassy polymer matrices, the (bimolecular) photooxidation of BZ, producing BP, is much more efficient than the (pseudo-first-order) photoreduction. Because BZ conversion is slow on the time scale for the diffusion of molecular oxygen, the concentration of molecular oxygen remains near its saturation values throughout the irradiations (Table 1).

We reiterate that the photooxidation of BZ in a glassy polymer matrix results in almost quantitative formation of BP. It is reasonable to assume that the same mechanisms leading to the addition of molecular oxygen to the triplet state of BZ (and, thus, the production of the 1,4-biradical) occur in solution and in a glassy polymer matrix. The separation in the mechanistic pathways probably occurs when the 1,4-biradical cleaves into a benzyloperoxy-benzoyl radical pair. At that stage, the radical pair can diffuse from the cage of its origin in...
solution, allowing the radicals to react independently as doublets; in the glassy matrix, the
radicals of the pair are constrained to remain for a long period in proximity, and favoring
their recombination as BP, a stable product at room temperature (Scheme 3).

As mentioned, a mechanism for BP formation based on the direct formation of perox-
ide from a 1,4-biradical has been suggested. In this connection, it is worth mentioning that
in this solution-phase study, the radiation employed (mostly λ > 320 nm) permitted higher
energy states to be accessed than for photoperoxidation of BZ in glassy polymer matrices
where the radiation wavelength was >370 nm. Thus, in solution, another hypothesis is that
vibrationally excited BP is produced from the direct insertion of molecular oxygen into the
carbonyl–carbonyl bond of the triplet state of BZ [49].

The excited triplet state of BZ can sensitize the formation of the 1∆g state of singlet
oxygen [61]. In that regard, singlet oxygen reactions with organic molecules in a glassy PS
matrix have been investigated [62]. For molecules that react efficiently with singlet oxygen,
rate constants in the glass are significantly smaller than in liquid toluene. However, we
suggest that some molecules that react inefficiently with singlet oxygen may become more
reactive in a glass than in solution; examples may be “self-sensitized” reactions (i.e., such
as that for BZ) in which the substrate, itself, produces the singlet oxygen.

As noted, the conformational twist angle about the –CO–CO– bond influences the
triplet state energy of BZ as well as the degree of conjugation of the carbonyl groups with
their adjacent phenyl moieties [63]. Therefore, the conformation and energy of the lowest
excited triplet state may also influence the reactivity of BZ and diacylperoxide formation.

Additionally, the effect of the photochemical transformation of 1,2-dicarbonyl com-
pounds on the molecular characteristics and the rate of the subsequent poly(ε-caprolactone)
and poly(lactide) hydrolysis was studied in poly(ε-caprolactone) and poly(lactide) that had
been doped with BZ or 1,4-bisbenzil. In addition to photoreduction of BZ and 1,4-bisbenzil,
transformation to the corresponding peroxides was observed [64,65].

Camphorquinone (CQ), a 1,2-dicarboxyl compound that lacks the phenyl auxochromes
of BZ, has also been irradiated in glassy PS matrices in the absence and presence of molec-
ular oxygen [66,67]. Interestingly, no diacylperoxide was detected upon irradiation of the
aerated films even though the relative rate of camphorquinone consumption increased
significantly with higher oxygen concentrations. In glassy PS, the photochemically induced
loss of CQ was accompanied by scission of the polymer chains. The degree of polymer
degradation was highest in the absence of molecular oxygen (under an argon atmosphere)
and lowest in an oxygen atmosphere. In the absence of molecular oxygen, the electronically
excited molecules of CQ abstract H atoms from the backbones of PS chains, leading to
subsequent chain scission. The degradation of low-density polyethylene in the presence of
cobalt stearate and BZ was also examined [68].

5. Polymers Bearing Covalently Attached 1,2-Dicarbonyl Groups

BP groups that are covalently attached to polymer chains have been formed by
irradiation of films of copolymers containing BZ pendant groups. Preparation of the
copolymers was accomplished by free-radical initiated polymerization of mixtures of
monomers containing a BZ moiety (VBZ, BZMA, PCOCO and BZS) and another monomer,
such as methyl methacrylate, styrene or phenyl vinyl ketone [69–73].

\[
\text{CQ}
\]
The copolymer of BZS and styrene was also successfully prepared by nitroxide-mediated polymerization [72,73]. Although research to date has focused on copolymers of styrene and PMMA, a copolymer of VBZ and phenyl vinyl ketone has been prepared and irradiated as well (vide supra) [74].

Successful copolymerization, defined here as random copolymerization of the monomers and their incorporation into the eventual chains in a distribution that is nearly the same as the monomer feed composition, requires that the reactivity of the monomer components be nearly the same. As indicated by comparisons of the monomer mixtures and copolymer compositions in Table 2, many of the BZ-containing monomers are much more reactive than styrene.

Table 2. Compositions of BZ-bearing monomer and styrene mixtures and their copolymers.

| BZ Monomer | Conversion wt% | mol% of BZ Group |
|------------|----------------|------------------|
| BZMA [69]  | 30             | 1.7              |
| PCOCO [70] | 18             | 3.8              |
| VBZ [71]   | 52             | 2.4              |
| BZS [72,73]| 28             | 1.2              |
| BZS [72,73]| 19             | 4.0              |

The small difference between the concentrations of BZS in the monomer mixtures and styrene and in the copolymer demonstrates that these two monomers have very similar reactivities toward polymerization and that the BZ groups are distributed randomly along the polymer chains. The reactivity of 4-substituted styrene monomers is known to increase with increasing electron-withdrawing character of the substituent [75]. The 4-CH$_2$OR styrene substituent of BZS is slightly more electron-donating than hydrogen [72,73]. Therefore, BZS is slightly less reactive than styrene, and VBZ (whose 1,2-dicarbonyl substituent is highly electron-withdrawing) is more reactive than styrene [71]. Additionally, it is known that the reactivity of a methacryloyl monomer (BZMA) [69] and of a vinyl carbonyl monomer (PCOCO) [70] are higher than that of styrene.

Polymers bearing BZ units in the main chain (I and II) or as pendant side groups [76] (III) (Figure 2) have also been synthesized via polycondensation and free radical copolymerization reactions (Figure 2 and Scheme 5).

Films of these polymers under nitrogen atmospheres have been crosslinked by UV radiation. Their potential application as negative resists has been examined [76]. Crosslinking efficiency (and, therefore, sensitivity in lithography) depends upon the radiation type and power: 311 nm from an Hg-lamp and 308 nm from both steady-state and pulsed excimer lasers have been employed. Increasing the energy of the 308 nm laser pulses from 100 to 400 J m$^{-2}$ resulted in a 4–6 fold increase of the lithographic sensitivity. The increase is attributed to the splitting of the C–C bond between the carbonyl groups (and formation of benzoyl radicals) that occurs during two-photon absorptions; changes in the radiation intensity at the low powers provided by the steady-state irradiations did not result in increased quantum efficiencies for crosslinking. The chemistry responsible for photocrosslinking achieved by scission of the 1,2-dicarbonyl groups is not entirely under-
stood. At low radiation intensities, the crosslinking mechanism is through a combination of a BZ ketyl macroradical with a macroradical of a neighboring chain.

**Figure 2.** Structures of copolymers: poly(oxyhexyleneoxy-4,4′-benzylene) (I), poly(oxy-4,4′-benzyleneoxysebacoyl) (II) and poly(phenyl methacrylate-co-4-methacryloyloxy-4′-methoxybenzil) (III).

**Scheme 5.** Photochemical transformation of BZ pendant groups into BP groups in styrene copolymers.

Copolyamides containing BZ units in the main chain (copolymide poly[iminohexamethylenimino-4,4′-(1,2-dioxoethane-1,2-diyl)dibenzoyleo-iminohexamethylene-iminoadipoyl] COPAIM) were also prepared from adipoyl chloride, 4,4′-dichloroformylbenzil and hexamethylenediamine [77,78]. Their photochemical reactions in solution and in films occurred in different ways. UV irradiation in solution led to polymer degradation; the BZ unit in the copolyamide split into two benzoyl radicals, which then reacted with solvent. UV irradiation of the copolyamide film caused the BZ units to split into two benzoyl radicals (as in the solution irradiation), but the benzoyl radicals abstract hydrogen atoms from the polyamide in this case. The coupling of the resulting macroradicals results in gelation. Gelation also depends on $T_g$ of the polymers. Those with $T_g$ higher than room temperature did not form a gel because of decreased recombination of macroradicals. The number of formed crosslinks was found to be equal to the number of bond scissions.
A copolymer of 10-methacryloyloxy camphorquinone (MCQ) and styrene has been synthesized as well [79]. Upon irradiation, no camphordiacetyl peroxide was formed. The efficiency of crosslinking is lower than in the case of the benzil analog, BZS/S. MCQ has also been used to prepare the homopolymer, copolymers and terpolymers with butyl methacrylate and/or 2-(dimethylamino)ethyl methacrylate. The ability of the CQ pendant groups in these polymers to act as photoinitiators has been examined in the photocuring of the hexanediol diacrylate/butyl acrylate mixture using irradiation with $\lambda > 400$ nm [80]. The fastest photocuring rate was found for the initiating system consisting of poly(MCQ-co-butyl methacrylate) and low-molecular-weight tertiary amine.

Polybenzil was prepared from polybenzoin in an analogous manner and was shown to be a photosensitizer [81]. Proton exchange membranes comprised of covalently crosslinked sulfonated poly(arylene ether sulfone)s with quinoxaline cross-linkages were prepared via cyclocondensation reactions between 3,3'-diaminobenzidine and benzil groups attached to the polymer chains [82].

5.1. Photoperoxidation of Pendant Benzil Groups in Glassy Polymer Matrices in the Presence of Molecular Oxygen

As with the films doped with BZ molecules, the progress of the photoinduced reactions of the copolymers, as self-supporting styrene copolymer films (20–200 µm thickness) could be followed by FTIR spectroscopy (Scheme 5, Figure 3). Especially in styrene copolymers with VBZ and BZS, the transformation of BZ to BP and subsequent transformation of BP structures can be followed easily by FTIR spectroscopy. In styrene copolymers of PCOCO and BZMA, carbonyl and ester group FTIR absorption bands occur between those of BZ and BP groups. However, side products, when formed, are difficult to detect by FTIR due to overlapping absorption bands. Although in copolymers of VBZ and styrene, the BP groups are formed almost quantitatively, a small amount of side products was detected in the copolymer of BZS. In BZS/S, for example, the decrease in IR absorbance from the 1,2-dicarbonyl part of BZ at 1650–1700 cm$^{-1}$ and the concurrent increase in absorbance from the BP carbonyl at 1750–1800 cm$^{-1}$ have been recorded as a function of irradiation time (Figure 3). The depletion of BZ and formation of BP structures in the 1150–1250 cm$^{-1}$ region can be used to follow the structural changes, as well [72,73]. UV–vis absorption spectra in the region of the diketone $n\rightarrow\pi^*$ absorption band at 390 nm [72,73] and the formation of peroxides by $^{13}$C NMR spectroscopy [72,73] were also monitored during photoperoxidations. In fact, several spectroscopic techniques can be used to follow the course of the reaction.

Figure 3. Changes in FTIR spectra of a BZS/S copolymer film during irradiation ($\lambda = 436$ nm) at room temperature in air. The absorbance bands of PS have been subtracted for clarity.
The irradiated films become partially insoluble in organic solvents as a result of the crosslinking from **BZ** peroxidation and subsequent **BP** photo-decomposition. Even irradiation of vacuum-dried **BZS/S** film at the red edge of the **BZ** \( n \rightarrow \pi^* \) absorption band \((\lambda = 436 \text{ nm})\) at room temperature led to partial crosslinking \([72,73]\). It would be interesting to prepare peroxide-containing polymers that retain their solubility after photoperoxidation of the **BZ** groups. However, it has not been possible to avoid at least some crosslinking in the photoperoxidation when irradiations are continued to nearly complete conversion. The origin of this crosslinking has been discussed \([72,73]\).

### 5.2. Dependence on Wavelength of Irradiation

The photoperoxidation selectivity of polymers bearing **BZ** structures at different wavelengths of irradiation and pulse energies has been discussed above \([71,76]\). In the presence of molecular oxygen, irradiation at >400 nm of **VBZ/S** copolymer films converts dicarbonyl groups to peroxy moieties nearly quantitatively (Figure 3); at shorter wavelengths of irradiation (366, 313 or 254 nm), the **BP** groups that are formed can undergo subsequent photolysis involving cleavage of their peroxy bonds and net crosslinking of polymer chains \([71]\). The ratio of the relative rate constants for **BZ** oxidation and subsequent **BP** decomposition upon absorption by a second photon decrease with lower wavelengths (i.e., higher energies) of irradiation.

Although formation of **BP** groups was also observed during irradiation of the **VBZ/S** copolymer film at 366 nm (Figure 4), FTIR spectra indicate that the peroxides were slowly converted to esters (N.B., peaks at 1700–1750 cm\(^{-1}\)) and benzoic acid (peaks at 1670–1700 cm\(^{-1}\)) \([71]\). The intensities of the **BP**-related bands increase to 90% conversion of the **BZ** groups (full lines in Figure 4) and then decrease upon further irradiation (dotted lines in Figure 4).

**Figure 4.** Changes in FTIR spectra of a **VBZ/S** copolymer film during irradiation \((\lambda = 366 \text{ nm})\) at room temperature in air. The absorbances from PS have been subtracted from each spectrum for clarity.

This experiment also shows that decomposition of the peroxides is principally due to excitations within the tail of the low energy absorption band and not to sensitization (energy transfer) by the remaining **BZ** groups. Although a sensitization mechanism was operative in the case of some homopolymers \([71]\), it cannot account for the reaction of the **BP** groups here because the concentration of 1,2-dione groups is at high at low conversions and low at high conversions (where the concentration of the **BP** groups becomes larger). Thus, the average separation between the remaining **BZ** and most of the **BP** groups is large, and the **BZ** groups are more distant in the copolymer than in homopolymers, making
Dexter energy transfer unlikely. Additionally, the lack of appropriate spectra overlap makes Förster energy transfer very difficult. Consistent with this interpretation, irradiation of a VBZ/S copolymer film at 313 nm led similarly to the in situ transformation of BZ to BP groups.

No absorption attributable to BP groups could be discerned in FTIR spectra of a VBZ/S copolymer film irradiated with a low-pressure mercury arc (i.e., principally at 254 nm) [71]. At this wavelength, it appears that transformation of BZ to BP groups is still the major reaction pathway, but photoinduced decomposition of BP groups proceeds rapidly after their formation. Radiation at 254 nm is absorbed strongly (aromatic $\pi \rightarrow \pi^*$ transitions), and the very high optical densities result in very little penetration beyond the irradiated surfaces (to a depth of about 2.4 $\mu$m [71]; interior portions of the film are not exposed to the photon fluxes).

The largest dependence of the photochemistry on radiation wavelength was observed for polymers containing bichromophoric (BZ and carbonyl) PCOCO structures [70]. In the copolymer with styrene (PCOCO/S), carbonylated BP groups are formed efficiently upon irradiation at $\lambda > 400$ nm, as has been described for the simpler BZ derivatives (Scheme 5). Subsequently, these BP groups can be cleaved photochemically or thermally. When irradiated at 366 nm, only a very small yield of BP groups was indicated by FTIR spectroscopy at 1760–1800 cm$^{-1}$ (Figure 5). As before, the transformation of BZ to BP (Scheme 5) is probably the principal initial reaction pathway, but photolysis of the peroxides is very rapid. During photoperoxidation, absorption at 366 nm by the 1,2-dicarbonyl groups is reduced so that the fraction of radiation absorbed by the keto groups is increased. Subsequently, energy transfer from the keto groups to the BP groups leads to the loss of the latter (Figure 5). FTIR spectra indicate that the expected products from thermolysis of the BP groups, esters of benzoic acid (1710–1760 cm$^{-1}$) and benzoic acid and carbonyl-containing moieties (1650–1710 cm$^{-1}$), are present. Similar photo-sensitization of the decomposition of peroxy esters has been reported by Neckers and co-workers [13,14] in a polymer containing benzophenone-peroxy ester bichromophoric groups.

![Figure 5. Changes in FTIR spectra of PCOCO/S copolymer film during irradiation ($\lambda = 366$ nm) at room temperature in air. The spectral bands of PS have been subtracted from each for clarity.](image)

5.3. Thermal Decomposition of Pendant BP Structures and Crosslinking of Styrene Copolymers

An example of how the thermolysis of BP structures, formed by irradiation of a BZS/S copolymer [72] film in the air in this case, can be followed by FTIR spectroscopy is shown in Figure 6. Similar changes in the spectra of BP pendant groups in other copolymer films have been observed [69,70]. The loss of absorbance by BP groups (at 1750–1800 cm$^{-1}$) and formation of new species—carbonyl-stretching bands of aromatic esters at 1710–1750 cm$^{-1}$ and of benzoic acid derivatives at 1660–1700 cm$^{-1}$—are clearly seen.
Figure 6. Temporal evolution of FTIR spectra upon thermolysis of BP groups in a preirradiated BZS/S copolymer at 91 °C. The spectral bands of PS have been subtracted from each for clarity.

It is much easier to follow the loss of BZ groups upon irradiation and the formation and loss of BP groups upon irradiation and thermolysis by FTIR in BZS/S and VBZ/S copolymers than in the analogous methacrylate and vinyl carbonyl polymers (BZMA/S and PCOCO/S) (Scheme 5). The latter contains additional ester or other carbonyl functional groups that absorb in the same infrared region where the major changes from conversions of BZ and BP occur.

Using these spectra, first-order rate constants for thermolyses of BP groups at 91 °C were calculated (Table 3) [69–72]. For alkoxy-substituted BP structures (photoperoxidized BZMA and BZS), the estimated rate constants are higher than for the BP with a less electron-donating alkyl (photoperoxidized VBZ) or electron-accepting acyl substituent (photoperoxidized PCOCO). The rate constant for photoperoxidized BZS (0.20 h⁻¹) is a bit lower than that obtained at the same temperature for the equally 4-substituted BP structure (with the same BZ-type photoperoxidized group; 0.26 h⁻¹), which was bound to a methacryloy moiety [69] for reasons mentioned above [71].

| BP Source   | k₁ (h⁻¹) * |
|-------------|------------|
| PCOCO/S [70]| 0.08       |
| VBZ/S [71]  | 0.08       |
| BZMA/S [69] | 0.26       |
| BZS/S [72]  | 0.20       |

* From linear fits with R² > 0.999.

Why are the rates for the loss of BP groups from photoperoxidized VBZ and PCOCO the same? Some insights come from the product distributions observed during thermolyses of benzene solutions of BPs with a para –OCH₃, –C₆H₅, –Cl or –NO₂ substituent on one of the phenyl rings, under an inert atmosphere and in the absence [83] or presence [84] of BF₃ as a Lewis acid catalyst. The thermolysis of the nitro-substituted BP yielded more product attached to the solvent than the other substituted BPs. Of the molecules investigated, the methoxy-substituted BP reacted fastest at one temperature. The same relative ordering of rate constants was found for the 4-substituted BP pendant groups in the glassy polymers: the alkoxy derivatives are fastest, and those with an electron-withdrawing substituent (such as nitro) are the slowest. Additionally, low-molecular-weight molecules with BP groups doped into PS films react more rapidly than the analogous groups attached to a
polymer backbone [70]. This kinetic effect has been explained as an increased possibility of inductive decomposition of dopant peroxides in polymer matrices that can form aggregates; aggregation is not possible for peroxides covalently attached to polymer backbone whose distribution of BZ precursor groups is random.

After thermolysis of covalently attached BP groups, the polymer films are completely insoluble in organic solvents due to the extensive crosslinking which is induced by reactions of the intermediate radicals. For the BP copolymers with styrene [69–73] a crosslinking proceeds as is shown in Scheme 6.

**Scheme 6.** Crosslinking induced by decomposition of BP groups in photoperoxidized styrene copolymers with initial pendant BZ groups.

The reaction in Scheme 6 is based on the homolytic cleavage of the O–O bond of BP groups and formation of two benzoyloxy radicals. The addition of benzoyloxy radicals attached to one polymer chain to aromatic rings on a vicinal polymer chain leads to crosslinking. Slow loss of CO$_2$ from benzoyloxy radicals may lead to aryl radicals as well, which can affect analogous crosslinks [85,86]. At 91 °C in a perchlorinated low-viscosity solvent, the rates of decarboxylation of a benzoyloxy [85] and a 4-methoxybenzoyloxy radical [86]. (a model for the counter-pair radical from corresponding peroxides from BZMA and BZS) are calculated to be $3 \times 10^7$ and $6.5 \times 10^6$ s$^{-1}$, respectively. The addition of the other fragment benzoyloxy radicals that are not a part of a polymer chain to a nearby polymer chain does not result in a crosslink or perceptibly change the average molecular weight of the polymer. In addition, the abstraction of a hydrogen atom from the PS backbone favors chain scissions [69].

The addition to an aromatic ring proceeds in two steps: the first is very fast, reversible complexation of a benzoyloxy radical with aromatic ring formation [49,87,88]; the second requires the loss of a hydrogen atom from the carbon atom of the aromatic ring where the addition occurred and leads to a covalent bond between a carbon atom on the aromatic ring and an oxygen atom of benzoyloxy. Despite the rapid reversion of the initial addition complex, thermolyses of BP molecules in PS films lead to the final addition of the benzoyloxy radical [89] (Scheme 6). The physical and spectroscopic properties of heated or irradiated polymers with BP groups provide strong evidence that the addition occurs efficiently there as well [69–73,90]. The other radical fragment from the pendant BP groups, the low-molecular-weight benzoyloxy moiety, can remove the H-atom (and form benzoic acid) if the glassy matrix forces it to remain in the vicinity of its macroradical partner. The large quantity of benzoic acid detected by FTIR spectra after heating the polymer films supports the occurrence of this process. The role of dissolved molecular oxygen in the polymer matrix is unknown and is worthy of investigation.

From melt rheology using Winter–Chambon criterion, it was determined that at least two crosslinking species (BP pending groups) per macromolecular chain are necessary to reach the gel point [91].
6. Photoperoxidation of and Crosslinking by a Low-Molecular-Weight Molecule Containing Two 1,2-dicarbonyl Groups Doped into Polystyrene Films

A method for crosslinking styrene-based polymer films using visible light and a low-molecular-weight additive containing two 1,2-dicarbonyl groups is shown below. The methodology follows the approaches described previously—photoperoxidation followed by decomposition of the generated benzoyl peroxide groups using either heat or light. The method is the subject of a current patent application [92].

Of the compounds in Scheme 7 for PS crosslinking, the one with X = covalent bond and Y = H (1,4-bisbenzil, BZBZ) was studied most thoroughly [93]. The temporal evolution of the FTIR spectra of a PS/BZBZ (20:1 w/w) film irradiated at >400 nm in air in Figure 7 showed a decrease of dicarbonyl absorption (1675 cm\(^{-1}\)) and an increase of peroxide absorption (1770 and 1793 cm\(^{-1}\)) that were attributed to benzoyl peroxide thermal decomposition products in polystyrene, benzoic acid and aromatic benzoates [89]. Thus, the formation of photoperoxidized PS was postulated (Scheme 8). Crosslinking of photoperoxidized PS proceeds in a manner similar to that for peroxides formed in styrene copolymers bearing BZ side structures [69–73].

To better understand the crosslinking process, the degree of crosslinking and contribution of the individual steps (photochemical and thermal) leading to it must be estimated quantitatively. In that regard, BP-O-BP facilitates estimation of the portion of the PS crosslinking from thermal decomposition of bisperoxides and, thus, provides insights into the overall mechanism (Scheme 9) [94]. The proposed mechanism for PS crosslinking in the thermal decomposition of bisperoxides is based on a correlation of crosslink density (i.e., the crosslink concentration at junction points) and the low concentrations of BP-O-BP. The crosslink density was estimated by swelling. A small amount of polymer degradation also occurs. Although the photochemical conversion of BZ-O-BZ to BP-O-BP proceeded only to about 30%, it is included in Scheme 9.

After decomposition of the BP moieties in BP-O-BP, two types of acyloxy radicals are formed (products A1 and A2) progressively. It is assumed that each acyloxy radical can be linked to a phenyl ring of PS or it can abstract a hydrogen atom (Scheme 9). Abstraction leads to the formation of benzoic acid (low-molecular-weight product B1) and ethers of benzoic acid (low-molecular-weight products B2). The addition of A1 or A2 to a phenyl ring of PS yields PS benzoate (functionalized PS C1) and PS benzoate bearing the benzoic acid moiety (functionalized PS C2). However, only the addition of both acyloxy radicals of A2 to the PS aromatic ring results in crosslinking (junction point C3 in Scheme 9). Although two BP-O-BP are required statistically to form one polymeric BP group, four BP-O-BP are required to make one crosslink (junction point) in PS.

As predicted from the hypothesized mechanism, FTIR spectra of PS crosslinked with BP-O-BP and prepared in situ from peroxycation of BZ-O-BZ are similar [94]. However, the estimated crosslink density from the complete reaction of in situ prepared BP-O-BP from added BZ-O-BZ is ca. 1/3 of that from BP-O-BP synthesized and added to PS. These crosslink densities only correspond to the content of the BP-O-BP formed photochemically. Although features of the FTIR spectra of BZ-O-BZ in PS films after irradiation indicated the presence of BP groups and exhibited changes similar to those observed after thermal decomposition of BP-O-BP in PS (N.B., the presence of IR peaks typical of benzoate and benzoic acid), the degree of crosslinking was rather low! It was concluded that during its formation, BP-O-BP also underwent decomposition, yielding products that do not contribute to crosslinking but absorb in the same wavelength region as the added and decomposed BP-O-BP.
Scheme 7. Photoperoxidation and decomposition of a low-molecular-weight dopant containing two 1,2-dicarbonyl groups for crosslinking chains of PS in films.

Figure 7. Temporal evolution of FTIR spectra of a PS/BZBZ (20:1 w/w) film irradiated at $\lambda > 400$ nm in air.
Scheme 8. Proposed course of photoperoxidation (λ > 400 nm) in air and crosslinking by BZBZ dissolved in a PS film.
Scheme 9. Simplified mechanism of PS crosslinking from reaction of BP-O-BP, either added or generated photochemically from BZ-O-BZ. The values in parentheses represent probabilities of emerging products and PS functionalities. The value (4) for BZ-O-BZ is a theoretical limit only.

7. Examples of the Application of Benzil Photoperoxidations

The feasibility of employing VBZ/S as a negative photoresist (i.e., to form non-erasable images) has been demonstrated [71]. A spin-coated 0.5 μm thick film on a silicon wafer was irradiated through a mask at 254 nm. The irradiated part became highly crosslinked, and the unirradiated parts could be dissolved in isobutyl methyl ketone to develop the image. The pattern after irradiation contained features comparable in size to the smallest apertures of the mask.

The advantages of using pre-crosslinked positive resists have been discussed [95]. Irradiation (>400 nm) in the air of a copolymer of phenyl vinyl ketone (PVK) with VBZ (PVK/VBZ) containing 1.5 wt% of VBZ structural units [74] in the film (for the photoperoxidation step), followed by thermolysis of the resulting pendant BP groups, leads to crosslinking. Subsequent irradiation at 366 nm of the crosslinked polymer causes cleavage of the PVK chain between junction points of the polymer network through Norrish type II reactions (Scheme 10) [96–99]. Thus, the PVK/VBZ copolymer represents a novel type of photoresist based on polymer network de-crosslinking. The process involves three steps: (1) photo-generation of peroxide-containing moieties; (2) crosslinking via thermolysis of the peroxides; and (3) subsequent photoinduced de-crosslinking of the polymer network. This material provides positive-tone images after UV exposure (>330 nm) and development in an organic medium, such as isopropyl methyl ketone.
Scheme 10. Cleavage of PS chains via Norrish type II photoreactions.

Decomposition of photogenerated BP groups can be utilized for surface modification by grafting [100]. Thermal decomposition of BP groups of a photoperoxidized BZMA/S was successfully utilized for grafting by immersing the film in a boiling 15% aqueous solution of methacrylic acid. Additionally, photosensitized decomposition (λ = 366 nm) of BP groups of a photoperoxidized PCOCO/S film was used as a graft by immersing it in acrylic acid. Some examples of applications of the photoperoxidation of pendant copolymer BZ groups are presented in Table 4.

Table 4. Some examples of applications of photoperoxidation of pendant copolymer BZ groups.

| Copolymer       | Application                                      | [Ref.] |
|-----------------|--------------------------------------------------|--------|
| VBZ/S           | Negative photoresist (254 nm)                    | [71]   |
| PVK/VBZ         | Positive photoresist based on polymer network    |        |
|                 | de-crosslinking                                  | [74]   |
| BZMA/S          | Thermal grafting of methacrylic acid            | [100]  |
| PCOCO/S         | Photografting (366 nm) of methacrylic acid      | [100]  |
| BZMA/MMA        | Copolymer grafting on polyethylene surfaces     | [100]  |

8. Conclusions

The photochemistry of BZ in solution leads to various products whose nature depends on the presence or absence of molecular oxygen, hydrogen-atom donors, the wavelength of irradiation, temperature and reactant concentration.

Upon irradiation at >370 nm, BZ, either doped or covalently bound to a polymer matrix, can be converted to BP almost quantitatively. The extent of conversion depends on the nature of the substituents on the BZ structure. The presence of side products can be detected conveniently by FTIR spectroscopy in frequency regions between where the diagnostic peaks of the starting BZ and the BP peroxide products appear.

Cleavage of photochemically generated BP groups bound to a polymer backbone leads to extensive crosslinking by the addition of intermediate benzoyloxy macroradicals to aromatic rings of PS or other aromatic polymers.

The proposed mechanism of the PS crosslinking in the thermal decomposition of compounds containing two BP groups (BP-O-BP) is based on correlations of crosslink density (concentration of crosslinks, concentration of junction points) and concentrations of BP-O-BP (at low concentrations of BP-O-BP). From these correlations, one crosslink requires four BP-O-BP structures. The yield of acyloxy radicals from thermal scission of BP-O-BP is quantitative. Each acyloxy radical can add to an aromatic ring or accept a hydrogen atom from a benzoyloxy radical–phenyl ring addition complex with equal probability. For the peroxides generated from compounds containing two BZ structures (BZ-O-BZ) in a polymer film, the crosslink density depends on the yield of the photooxidized product containing two BP groups (N.B., BP-O-BP).

Because of the high reactivity of benzoyloxy radicals, the approaches described here should be amenable to crosslinking a wide variety of other polymers besides PS. Crosslinkers with two covalently attached BZ groups offer two important practical advantages over
those containing two covalently attached BP structures: (1) they are safer to handle; (2) they are usually more soluble in polymers.

However, crosslinking with a BZ-type crosslinker can be achieved in two steps by irradiating initially (at >370 nm) in the presence of molecular oxygen followed by thermal cleavage of the as-produced BP compounds to generate benzoyloxy radicals. Alternatively, crosslinking can occur in one step by irradiation at shorter wavelengths, where BP groups are formed and decomposed sequentially.

Self-supported PS films of thickness in the nm range cannot be prepared for reasons dealing with the mechanical properties of the polymer. Nano-thickness layers of PS film surfaces can be modified by crosslinking using BZ structures bound to styrene copolymer chains and using doping agents containing two BZ groups. Crosslinking by BP groups formed photochemically by irradiation that does not overlap with the BP absorptions can be initiated thermally. BP groups prepared from BZ groups in a VBZ/S copolymer irradiated at 254 nm are simultaneously decomposed with the same light, crosslinking only layers within ~2.4 µm of the surface due to strong absorption of the PS phenyl rings [71]. This depth is significantly greater than the nm range.

Clearly, the methods of crosslinking described here offer advantages over other published methods for the modification of surfaces and polymer films. Those advantages are described by several examples within this text.

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