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A new safranin based three-component photoinitiating system for high resolution and low shrinkage printed parts via digital light processing†

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Additive manufacturing or 3D printing has attracted the interest of researchers in industry and academia because of its outstanding features. In this study, a new three-component photoinitiating system (PIS) consisting of safranin O (SFH<sup>+</sup>), thiol derivatives and diphenyl iodonium salt was used for the free radical photopolymerization of a diacrylate monomer (SR349) in DLP 3D printing. The photoinitiating characteristics of this PIS were evaluated and advantageously compared to those of a conventional PI (TPO) by using RT-FTIR. It is shown that the proposed PIS could be used as an efficient PIS for free radical photopolymerization. In addition, the resolution and shrinkage of printed parts in the presence of this three-component PIS were measured and compared to those printed using TPO as a photoinitiator. The resolution of printed parts was determined by using SEM and profilometry techniques. In addition, photorheometry was used to evaluate the linear shrinkage of samples. Moreover, the initiating mechanism of the three-component PIS was studied by using laser flash photolysis (LFP). A photocyclic mechanism was outlined for the three-component PIS which demonstrated this mechanism would be very beneficial for DLP 3D printing

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# Introduction

In the 1980s, three dimensional (3D) printing or additive manufacturing (AM) was first introduced as a technology for fabricating 3D objects using a computer-aided design (CAD) file.1-3 Thanks to its ability to produce complex geometries in a simple one-step process, little waste of material, and being user-friendly, the applications of this technology in different fields such as medical implants and prosthetics, 4,5 digital dentistry, 6,7 tissue engineering, 8,9 the aerospace industry, 10,11 construction12,13 and sensors14 are growing fast. It is expected that the world's 3D printing market would experience 18% growth annually and reach up to 23 billion USD by 2022.15

Vat photopolymerization is one of the most popular 3D printing techniques utilizing a light source for solidifying a photopolymer. 16,17 Digital light processing (DLP), based on vat photopolymerization method, is very similar to lithography and is also known as dynamic mask photolithography.3 Build time, oxygen sensitivity and adhesion between different layers are improved in this technique.18,19 However, vertical resolution, dimensional accuracy, and shrinkage are still the main challenges of DLP 3D manufactured parts.20-22 Vertical resolution depends on the penetration depth of incident light into liquid photopolymer. Several researches have studied and tried to solve this challenge by using different types of light absorbers and inhibitors in formulations.23,24 Zissi et al. studied the influence of adding a non-reactive absorbing chemical on the depth of cure in a 3D printing system.25 They reported that the cure depth and width could be controlled by adjusting the photoabsorber concentration. Choi et al. investigated the effect of different concentrations of Tinuvin®327, as a light absorber, on the quality of fabricated objects. It was demonstrated that not only light absorber concentration but also exposure energy could affect the resolution of complex geometries.26 Chin et al. studied the effect of a red-shifted UV absorber (Tinuvin® CarboProtect®) on cure depth and kinetics in visible light initiating resin for 3D printing. They illustrated an enhancement in the geometrical and dimensional accuracy of stereolithography process operating under visible blue light.27 However, Zabti et al. showed that the presence of light absorbers such as Tinuvin®327 in formulation increased dimensional error which could be originated from shrinkage.28

Shrinkage, as the other main challenge, was studied by several researches.<sup>29</sup> For example, Stampfl et al.<sup>30</sup> and Schoerpf et al.31 used monofunctional monomers and chain transfer agents and Chiappone et al. 32 proposed using acrylate/epoxy hybrid systems to overcome this problem to some extent.

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However, these strategies caused other problems such as deterioration of long-term mechanical properties of photo-cured systems and also the requirement of using dual photoinitiator systems (i.e., radical and cationic photoinitiators), respectively.

In recent years, visible light initiating systems have been highly regarded in order to solve some UV radiation issues such as high heat build-up, safety, and curing of thick samples. 33,34 However, there is very limited photoinitiator leading to photodissociation in the visible region.35 Therefore, using two/threecomponent photoinitiating systems (PIS), consisting of a photosensitizer and one or two coinitiator(s) have been pointed out. In these systems, the active species are usually generated via an electron/energy transfer between a photosensitizer and coinitiator.36 The photosensitizer is typically an organic dye such as safranin,37 methylene blue,38 rose bengal39 or ketone sensitizer like isopropyl thioxanthone (ITX).40 The second 3 component is an electron or/and a hydrogen donor that is usually an amine or thiol, and the third component is an electron acceptor such as onium salt.41

According to previous studies, safranin O (SFH<sup>+</sup>) can be used as a high-performance photosensitizer in two- or threecomponent photoinitiating systems. Several researchers have reported the radical generation ability of safranin in combination with different coinitiators. 42-45 Therefore, in this study, a threecomponent PIS based on safranin O was employed in digital light processing 3D printing to investigate the dual function of the dye as photosensitizer and photoabsorber. As mentioned before, the former was responsible for active species production and the latter could play a role in controlling the curing depth and improving the vertical resolution of printed parts. For this purpose, a combination of safranin O, thiol derivative and onium salt was chosen as three-component PIS in an ethoxylated diacrylate system (SR349). In addition, the effect of curing protocol of this PIS on shrinkage was investigated. The performances of this new three-component PIS regarding resolution and shrinkage were compared to those obtained using diphenyl (2,4,6-trimethylbenzoyl) phosphine oxide (TPO) as photoinitiator. Finally, the photochemical mechanisms associated with the generation of the active species in this PIS were investigated.

# **Experimental**

#### **Materials**

Safranin O (SFH<sup>+</sup>) (dye content > 85%), pentaerythritol tetrakis(3mercaptopropionate) (RSH, 95%), diphenyl (2,4,6-trimethylbenzoyl)phosphine oxide (TPO) and dimethyl sulfoxide (DMSO) were purchased from Sigma-Aldrich. Diphenyliodonium hexafluorophosphate (IOD+, 98%) was obtained from TCI. Ethoxylated (3) bisphenol A diacrylate (SR349) was obtained from Sartomer. Acetonitrile (99.97%) and isopropyl alcohol were received from Biosolve. All chemicals were used as received. Chemical structures of the main materials are shown in Table 1.

### **DLP printing system**

B9Creation V1.2 3D printer (B9Creator Company, USA) was used for printing complex geometries to study the resolution and

Table 1 Chemical structures of used materials

Name	Structure
SR349	
$SFH^{+}$	$H_3C$ $N^+$ $NH_2$ $CI^-$
RSH	HS O SH
$\mathrm{IOD}^{^{+}}$	PF <sub>6</sub>
TPO	$ \begin{array}{c c} O & H_3C \\ P - C & CH_3 \end{array} $ $ H_3C$

shrinkage properties of the printed parts. The appropriate file format of black and white images of sliced samples was produced by B9Creator software. The radiation was provided by using a modified D912HD projector (Vivitek) with a cut-off filter in the range of 380-800 nm. The radiation intensity measured at the surface of the vat was 170 mW cm<sup>-2</sup>.

The 3D printer had an X-Y resolution of 30 μm, the layer thickness was adjusted on 20 µm and each layer was exposed to radiation for 6 s, while the exposure time for the first two layers was set to be 30 s. Printed parts were submerged in 90% isopropyl alcohol for less than a minute to wash off the surface from unreacted monomers. The printed parts were air-dried at room temperature before post-curing under a medium pressure mercury lamp for about 60 seconds.

For characterization and comparing the X-Y and Z resolutions of printed samples, a 3D benchmark design was selected from Auto desk Ember website shown in Fig. 1. The minimum size printed accurately in each direction was considered as X-Y and Z resolution of the samples.

Two formulations for 3D printing were prepared, as shown in Table 2. The first one consist of SR349 monomer, which was diluted with 10 phr DMSO and three-component PIS based on SFH<sup>+</sup>/RSH/IOD<sup>+</sup>. The composition of the three-component PIS was designed according to literature and this sample was named 3KSRD. In the second sample, TPO was used as one of the most commonly used photoinitiators in photoinduced 3D

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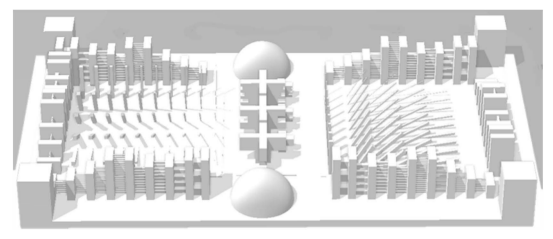


Fig. 1 3D view of benchmark design used for X-Y and Z resolution.

Table 2 Composition of the 3D printing sample formulation

Sample code	Composition
3KSRD	SR349/DMSO (10 phr)/SFH <sup>+</sup> (0.2 phr)/RSH(3 phr)/IOD <sup>+</sup> (3 phr)
1KTPO	SR349/DMSO (10 phr)/TPO(1 phr)

printing techniques. This sample was named 1KTPO as a reference to compare the performance of the new proposed PIS with that of a conventional one. In order to compare the results obtained with 1KTPO and 3KSRD, the TPO concentration was adjusted to obtain the same final conversion as 3KSRD sample. Therefore, the appropriate concentration of TPO was found to be 1 phr. Worth mentioning that the concentrations of SR349 and DMSO were kept the same to avoid any interference on the printing properties such as resolution, shrinkage and dimensional accuracy.

#### Characterization and measurements

Vertex 70 FTIR spectrophotometer (Bruker Optik GmbH, Germany) in its real-time mode was used to study the photopolymerization kinetics of formulations. The resolution of spectra was equal to 4 cm<sup>-1</sup>. Samples were irradiated by a xenon lamp (Hamamatsu L9566-04) when IR spectra were recorded. It should be mentioned that the irradiation wavelength was adjusted at  $\lambda$  < 395 nm by using a cut-off filter. Light intensity at the sample surface was set to be 230 mW cm<sup>-2</sup>. The reference sample (containing TPO) was irradiated by a 395 nm LED device (Roithner Laser Technik) which provided 100 mW cm<sup>-2</sup> intensity at the surface of the sample. The degree of C=C conversion, C, was calculated according to eqn (1):

$$C(\%) = \frac{A_0 - A_t}{A_0} \times 100 \tag{1}$$

where  $A_0$  and  $A_t$  are the area of the IR absorption band at 1637 cm<sup>-1</sup> of samples at the starting point and time t, respectively. All spectra were baseline-corrected before the integration with the software OPUS 7.0. To be sure about the reproducibility

of the results, each experiment was repeated at least three times. Finally, the rate of photopolymerization,  $R_p$ , was determined from the slope of the conversion-time profile.

Linear shrinkage measurements of samples were conducted based on Shah et al. method, 47 by using an MCR 302 rheometer from Anton Paar, which was combined with a light source accessory for the curing of samples. The experimental setup is presented in Fig. 2. The test was performed in the oscillation mode of the rheometer. The smooth aluminum plate was fixed at an initial 0.5 mm gap and a zero Newton normal force was applied to samples. The samples were irradiated in the same condition as RT-FTIR analysis. The light was directed under a glass plate by using an optical fiber, which would allow the photopolymerization of samples. The shrinkage of sample led to the gap reduction. So the variation of the gap during the photopolymerization would represent the shrinkage according to eqn (2), where the  $d_{\text{final}}$  and  $d_{\text{initial}}$  are the distance between the glass and aluminum plate at the beginning and end of photopolymerization, respectively.

Shrinkage (%) = 
$$\left(1 - \frac{d_{\text{final}}}{d_{\text{initial}}}\right) \times 100$$
 (2)

Scanning electron microscopy (SEM) was performed in order to investigate the quality of printed specimen (FEI Quanta 400 microscope, Thermo fisher scientific, DEU). The imaging was carried out under low vacuum condition.

The size and shape of printed pattern were evaluated by an optical profilometer (Altisurf 500 workstation, Altimet, FRA) equipped with a microforce sensor for solid films.

Cary 4000 UV-vis spectrophotometer (Agilent Technologies, USA) was used to determine the UV-visible spectra of samples at room temperature. Acetonitrile was used as a solvent for the preparation of samples.

Triplet-triplet absorption and photobleaching spectra were measured by using an LP900 laser flash photolysis analyzer (Edinburgh Instruments, UK). The details of the instrument, its accessory, experiment conditions and procedure were reported elsewhere.<sup>48</sup> It would be noted that the light excitation was

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performed at  $\lambda_{\rm exc} = 518$  nm because SFH<sup>+</sup> has a maximum absorption at this wavelength. The constant of quenching rate of excited states  $(k_a)$  was determined by using a well-known Stern-Volmer equation:

$$\frac{1}{\tau} = \frac{1}{\tau_0} + k_{\mathbf{q}}[\mathbf{Q}] \tag{3}$$

which  $\tau$  and  $\tau_0$  are the lifetime of triplet state in the presence and absence of quencher Q, respectively.

# Results and discussion

As mentioned before, 1KTPO sample was designed to have the same final conversion as 3KSRD system. This could be seen in Fig. 3, where the photopolymerization kinetics of the two samples were studied using RT-FTIR technique. However, it should be noted that the polymerization for 1KTPO sample was found to be four times faster than that for 3KSRD system.

#### Effect of PIS on resolution

Most of the TPO based systems contain a UV absorber which increases the absorbance and decreases the penetration depth of light. In this study, no absorber was added to the sample in order to study the intrinsic resolution obtain with the different PIS. X-Y and Z resolutions of both 3KSRD and 1KTPO samples are gathered in Table 3 as evaluated from the printed benchmark design (Fig. 4). It can be seen that 3KSRD resolution was

clearly better than that of 1KTPO in all directions. This is particularly the case in the Z direction. In that case, it seems that SFH<sup>+</sup> could successfully control the penetration depth of incident light by its light-absorbing capability. To support this justification, the penetration depth,  $D_p$ , of the light in the presence of safranin and TPO compound was calculated by using eqn (4).3

$$D_{\rm p} = \frac{1}{2.3(\varepsilon_{\rm I}[{\rm I}] + \varepsilon_{\rm A}[{\rm A}])} \tag{4}$$

where  $\varepsilon_{\rm I}$  and  $\varepsilon_{\rm A}$  are the molar extinction coefficient of initiator and absorber (when present), respectively, and [I] and [A] are the concentration of initiator and absorber, respectively. Based on the values reported for the molar extinction coefficients in the literature45,49 and keeping in mind that no absorber was used in this study, the  $D_p$  of 3KSRD sample was found to be significantly smaller than that of 1KTPO (i.e., 40 vs. 600 μm). It approves that SFH<sup>+</sup> had a dual-function as photosensitizer and photoabsorber in the system.

Table 3 Dimensional resolution measurements

Sample	X-resolution (μm)	Y-resolution (μm)	Z-resolution (μm)
1KTPO	500	500	>1000
3KSRD	100	100	100

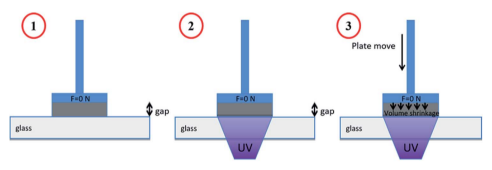


Fig. 2 Schematic representation of the shrinkage measurement apparatus.

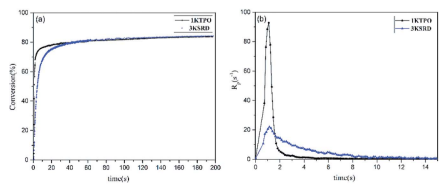
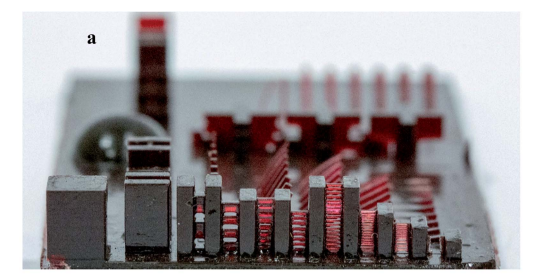


Fig. 3 Comparison of 3KSRD and 1KTPO polymerization: (a) conversion (b) rate of photopolymerization.



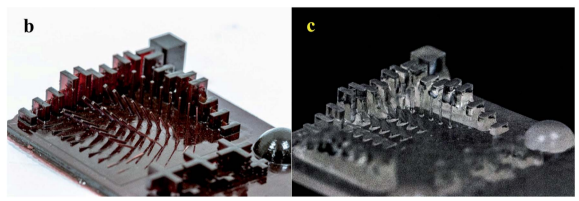


Fig. 4 Photographs of the printed benchmark design for (a) 3KSRD sample (Z-resolution), (b) 3KSRD sample and (c) 1KTPO sample.

# Effect of PIS on shrinkage

As noted earlier, shrinkage in printed parts is an important intrinsically due to the free radical photopolymerization.21,28 For this reason, the effect of new proposed PIS on the shrinkage was studied by photorheometery technique. As can be seen in Fig. 5, the linear shrinkage of 3KSRD and 1KTPO were measured to be 0.9 and 4.2%, respectively. It means that the shrinkage of printed part improved significantly when the three-component PIS was used. As the monomer and photopolymerization conditions are almost the same for both samples, different behavior of the two systems could be related to the different photoinitiating systems. In fact, as seen previously in Fig. 3, the polymerization rate of 1KTPO system was remarkably higher than that for 3KSRD system which led to a higher inhomogeneity and internal stress formation in the cured sample. It seems that this higher inhomogeneity and internal stress caused a more considerable shrinkage in 1KTPO system.

It is worth mentioning that our proposed new three-component PIS (*i.e.*, SFH<sup>+</sup>/RSH/IOD<sup>+</sup>) could provide a soft-start curing protocol which caused achieving relatively high conversion and low shrinkage in an appropriate time. In other words, 3KSRD sample with a slow rate of polymerization may

facilitate the relaxation of the polymer chain, thereby leading to a lowering of the shrinkage. <sup>50,51</sup>

Considering the high resolution and low shrinkage of 3KSRD sample, the capability of this formulation for the

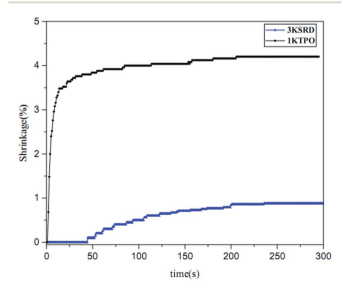
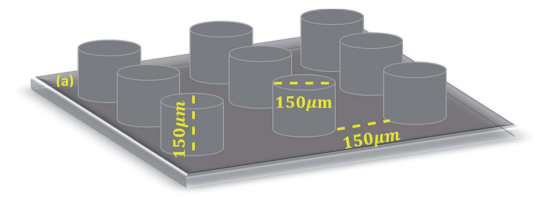


Fig. 5 Effect of PIS on shrinkage.



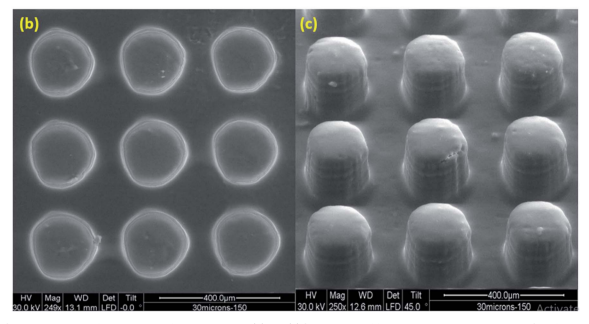


Fig. 6 (a) Schematic image of simple micro-featured pattern, (b) and (c) SEM images of fabricated microstructures by 3KSRD formulation from different angles.

printing of a simple and a complicated (i.e. Eiffel tower) micro-featured pattern was demonstrated. The simple micro-featured pattern included vertical cylinders with 150  $\mu m$  height and 150  $\mu m$  diameter, as shown in Fig. 6a. The pattern was successfully printed with reasonable fidelity. The topography and profilometry of the printed objects were shown in

Fig. 6b, c and 7, respectively. These figures confirmed clearly the dimensional accuracy and high resolution of printing system.

In addition, the photograph of a 3D printed mini-Eiffel tower was depicted in Fig. 8. As can be seen, all the details of the tower were correctly printed.

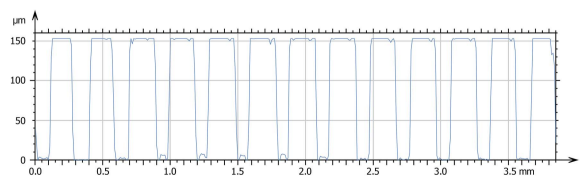


Fig. 7 Lateral profile of printed cylinder from 3KSRD sample with 150 μm in height and diameter.

### Photopolymerization kinetics

The conversion and photopolymerization rate of twocomponent (*i.e.*, 2KSR and 2KSD samples) and threecomponent PIS (*i.e.*, 3KSRD sample) were studied to understand the effect of each coinitiator on the photopolymerization kinetics. The specifications of the samples and the results are compiled in Table 4 and Fig. 9.

The results show that both two-component PIS and the three-component PIS, could initiate the free radical photopolymerization quite effectively. However, the final conversion and polymerization rate of 2KSR and 2KSD were found to be considerably lower than that of 3KSRD. It seems that the lower number of created radicals in the two-component PIS in comparison to that of three-component PIS is much lower, thereby leading to slower conversion rates. <sup>36,53</sup>

By looking more closely to the first seconds of polymerization in Fig. 9, it can be seen that there was an inhibition time (about 3.5 s) in both 2KSD samples. Indeed, aryl radicals produced by the SFH<sup>+</sup>/IOD<sup>+</sup> reaction in the 2KSD system were prone to molecular oxygen. By contrast, thiyl radicals in SFH<sup>+</sup>/RSH system are less sensitive to oxygen inhibition and, therefore, lead to lower inhibition time.<sup>48</sup>

In addition, Fig. 10 shows that the maximum rate of polymerization for 3KSRD was found to be three times higher than those for 2KSR and 2KSD samples. The higher reactivity of



Fig. 8 Photograph of a 3D printed Eiffel tower by using 3KSRD sample.

Table 4 Sample formulation, degree of conversion and polymerization rate

Sample	Composition (phr) SFH <sup>+</sup> /RSH/IOD <sup>+</sup>	Conversion (%)	Maximum rate of polymerization $(s^{-1})$
2KSD	0.2/—/3	60	5.4
2KSR	0.2/3/—	64	4.1
3KSRD	0.2/3/3	84	22.0

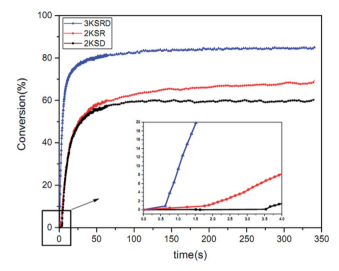


Fig. 9 Conversion vs. time for two- and three-component PIS samples.

three-component system could be explained by different mechanism as deeply studied recently. Indeed, two mechanisms can be introduced to explain these observations. Firstly, both coinitiators can react in parallel with the excited state of the photosensitizer, leading to higher production of radicals. In that case, no synergistic effect occurs and the efficiency is directly related to the excited state quenching efficiency by both coinitiators. Stocondly, sequential reactions in which one of the coinitiators reacted with the excited photoinitiator and then followed by the reaction of the second coinitiator with the photoproduct formed from the photoinitiator in the first reaction. In addition, the ground state photoinitiator was recovered through this second reaction which could absorb the light and react again, leading to a photocyclic initiating system.

## Photoinitiation mechanism

In order to study the photoinitiating mechanisms of two- and three-component PISs, laser flash photolysis (LFP) experiments were performed. At first, the UV-vis spectra of SFH $^+$  and the mixture of (SFH $^+$ /RSH/IOD $^+$ ) were measured. As can be seen in Fig. s1,† the absorption spectrum of SFH $^+$  had a maximum around 520 nm ( $\varepsilon=44~000~{\rm M}^{-1}~{\rm cm}^{-1}$ ) and both coinitiators absorbed in the wavelength less than 375 nm. It was observed that the spectrum of SFH $^+$  did not change by the addition of both coinitiators, so ground-state interactions between these

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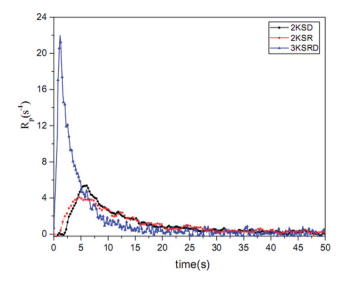


Fig. 10 Polymerization rate vs. time for two- and three-component PIS samples.

components could be discarded. In addition, it can be understood that only SFH<sup>+</sup> was excited when using visible light. On the other hand, it is well-known that the irradiation of SFH<sup>+</sup> does not initiate polymerization solely, thus the polymerization process would involve the excitation of the SFH<sup>+</sup> followed by further reactions.45

It is worth mentioning that according to the transient absorption spectrum of SFH<sup>+</sup>, there were three distinct peaks at 430, 520 and 820 nm. The peaks at 430 and 820 nm belong to triplet-triplet absorption of SFH+ (3SFH+) and the peak at 520 nm belongs to SFH<sup>+</sup> photobleached species. The absorption peaks of semi-oxidized and semi-reduced form of SFH+ appeared in 420-430 nm range.<sup>54</sup>

SFH<sup>+</sup>/IOD<sup>+</sup>. To understand the role of each component in active species production, firstly the photoinduced electron transfer (PET) reaction between <sup>3</sup>SFH<sup>+</sup> and IOD<sup>+</sup> was evaluated. The reaction of SFH<sup>+</sup> triplet state (<sup>3</sup>SFH<sup>+</sup>) with diphenyl iodonium hexafluorophosphate as a strong electron acceptor should result in semi-oxidized SFH+ ('SFH2+). The Gibbs free energy change of this reaction could be determined by using Rehm-Weller equation:

$$\Delta G_{\rm ET} = E_{\rm ox} (D) - E_{\rm red} (A) - E_{\rm T} + C$$
 (5)

where  $E_{ox}$  (D) refers to electron donor oxidation potential and  $E_{\rm red}$  (A) is electron acceptor reduction potential,  $E_{\rm T}$  is photoinitiator triplet state energy and C stands for a Coulombic term which is negligible in acetonitrile. Based on the data presented in Table s1,† the Gibbs free energy change was found to be -0.77 eV which indicates the spontaneous nature of the reaction.

Fig. 11 shows the results of LFP analysis for the reaction between <sup>3</sup>SFH<sup>+</sup> and IOD<sup>+</sup> (eqn (6) and (7)). As shown in Fig. 11a, <sup>3</sup>SFH<sup>+</sup> was found to decay faster when IOD<sup>+</sup> concentration increased. Furthermore, the formation of a permanent signal at 520 nm in Fig. 11b, implied that the concentration of SFH<sup>+</sup> decreased due to the reaction between <sup>3</sup>SFH<sup>+</sup> and IOD<sup>+</sup>.

The quenching rate constant  $k_q^{-3SFH^+/IOD^+}$  was calculated to be  $5 \times 10^5$  L mol<sup>-1</sup> s<sup>-1</sup> by using Stern-Volmer plot, which was in agreement with previous reports. 42,43 One of the products of eqn (7) (IOD) could be dissociated into an active aryl radical (Ph) and a stable molecule (eqn (8)). These results were consistent with the RT-FTIR results: as it is shown in Fig. 9, free radical polymerization of 2KSD system could reach 60% final conversion that meant the aryl radicals could efficiently initiate the polymerization.

$$SFH^{+} \stackrel{h\nu}{\to} {}^{3}SFH^{+} \tag{6}$$

$$^{3}SFH^{+} + IOD^{+} \rightarrow \cdot SFH^{2+} + IOD^{\bullet}$$
 (7)

$$IOD. \rightarrow Ph. + PhI$$
 (8)

The photobleaching signal of SFH<sup>+</sup> in the presence of enough amount of IOD+, was followed by increasing the concentration of RSH to investigate the effect of the second coinitiator on the mechanism of three-component PIS. In other words, if there would be a photocyclic reaction, the photobleaching signal of SFH+ would be returned to zero. It means that safranin dye in the ground state form was recovered by the reaction between ('SFH2+) and thiol. However, as can be seen in Fig. s2,† the photobleaching signal of the SFH<sup>+</sup> decreased when the amount of RSH was increased. This clearly indicated that no reaction occurred to recover SFH<sup>+</sup> in ground state. Furthermore,

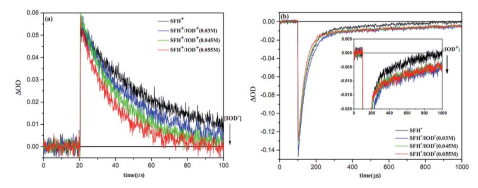


Fig. 11 (a) The decay of triplet-triplet absorption [ $^3$ SFH $^+$ ] measured at 820 nm and (b) ground state photobleaching of [SFH $^+$ ] recorded at 520 nm in the presence of various concentrations of IOD<sup>+</sup> ( $\lambda_{exc} = 518$  nm, 6–7 mJ per pulse).

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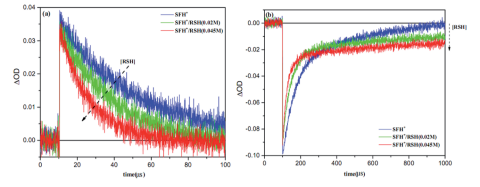


Fig. 12 (a) The decay of triplet-triplet absorption [ $^3$ SFH $^+$ ] measured at 820 nm and (b) ground state photobleaching of [SFH $^+$ ] recorded at 520 nm in the presence of various concentrations of RSH ( $\lambda_{exc} = 518$  nm, 6–7 mJ per pulse).

the decrease in the photobleaching signal of SFH<sup>+</sup> in the presence of RSH could express that there was a parallel reaction between RSH and the residual amount of unreacted <sup>3</sup>SFH<sup>+</sup>.

SFH<sup>+</sup>/RSH. One of the possible ways of producing active species in the proposed three-component PIS was the reaction between <sup>3</sup>SFH<sup>+</sup> and RSH as the primary photoreaction (eqn (9) and (10)). It seems that the neat electron transfer reaction between <sup>3</sup>SFH<sup>+</sup> and RSH could be ignored due to its positive Gibbs free energy change (+0.31 eV). On the other hand, hydrogen abstraction reaction could occur between <sup>3</sup>SFH<sup>+</sup> and RSH as a well-known hydrogen donor. 55,56 Indeed, as shown in Fig. 12a, <sup>3</sup>SFH<sup>+</sup> signal at 820 nm was effectively quenched by increasing the amounts of RSH and the quenching rate constant  $k_0^{^{3}\text{SFH}^+/\text{RSH}}$  was calculated to be  $7 \times 10^5 \text{ L mol}^{-1} \text{ s}^{-1}$ which was in the same order of magnitude for aliphatic thiols.<sup>57</sup> The photobleaching signal of SFH<sup>+</sup> in the presence of RSH is shown in Fig. 12b. Generation of a permanent signal by adding RSH, would be a confirmation for eqn (10).58,59 Thiyl radicals produced in this reaction could initiate the radical polymerization which was in agreement with RT-FTIR results of 2KSR system.

$$SFH^{+} \xrightarrow{h\nu} {}^{3}SFH^{+} \tag{9}$$

$$^{3}SFH^{+} + RSH \rightarrow SFH_{2}^{+} + RS$$
 (10)

$$SFH_2^{\bullet+} + IOD^+ \rightarrow SFH^+ + Ph\bullet + PhI$$
 (11)

As shown in Fig. 13, the photobleaching signal of SFH<sup>+</sup> in the presence of RSH came back to zero when IOD was added to the system. This feature indicated that SFH<sup>+</sup> was efficiently recovered during the reaction (eqn (11)). Consequently, it could be concluded that there was a photocyclic reaction which could produce two different radical species (i.e., RS' and Ph') which could both initiate the radical polymerization in a proper manner. Most importantly, synergistic effect of producing more active species and recovery of the photosensitizer in threecomponent PIS improved the polymerization rate and final conversion, as showed RT-FTIR results.

It is worth noting that SFH<sup>+</sup> acted as a photoabsorber in this system as well which improved the resolution of 3D printed

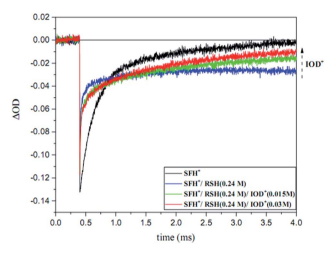


Fig. 13 Ground state photobleaching of [SFH+] recorded at 520 nm in the presence of RSH at 0.24 M and various concentration of IOD<sup>+</sup> ( $\lambda_{\rm exc}$ = 518 nm, 6-7 mJ per pulse).

articles. Therefore, the photocylic behavior of SFH<sup>+</sup> would also be beneficial from this point of view.

# Conclusions

A three-component photoinitiating system based on safranin was introduced to print complicated geometries by using digital light processing 3D printing. The following statements can be concluded from this work:

- (1) The feasibility of using SFH<sup>+</sup>/RSH/IOD<sup>+</sup> mixture as an efficient three-component PIS for free radical photopolymerization of diacrylate monomers (i.e., SR349) was demonstrated.
- (2) The resolution and shrinkage of DLP 3D printed parts in the presence of SFH<sup>+</sup>/RSH/IOD<sup>+</sup> and a conventional photoinitiator (i.e., TPO) were compared.
- (3) The resolution and shrinkage of DLP 3D printed parts improved remarkably when SFH<sup>+</sup>/RSH/IOD<sup>+</sup> was used as PIS.
- (4) The incorporation of IOD<sup>+</sup> coinitiator to the mixture of SFH<sup>+</sup> and RSH caused the regeneration of SFH<sup>+</sup> through a photocyclic reaction.

(5) The reaction mechanism of SFH<sup>+</sup>/RSH was studied by using laser flash photolysis technique for the first time.

# Conflicts of interest

There are no conflicts to declare.

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