



Contents lists available at ScienceDirect

Polymer

journal homepage: www.elsevier.com/locate/polymer

Additive manufacturing with a flex activated mechanophore for nondestructive assessment of mechanochemical reactivity in complex object geometries

Bo Cao ^a, Nicholas Boechler ^b, Andrew J. Boydston ^{a,*}^a Department of Chemistry, University of Washington, Seattle, WA 98195, United States^b Department of Mechanical and Aerospace Engineering, University of California, San Diego, La Jolla, CA 92093, United States

ARTICLE INFO

Article history:

Received 27 February 2018

Received in revised form

24 April 2018

Accepted 10 May 2018

Available online xxx

Keywords:

Additive manufacturing

Mechanochemistry

Vat photopolymerization

Lattice structures

Microstructured materials

ABSTRACT

We used digital light processing additive manufacturing (DLP-AM) to produce mechanochemically responsive test specimens from custom photoresin formulations, wherein designer, flex activated mechanophores enable quantitative assessment of the total mechanophore activation in the specimen. The manufactured object geometries included an octet truss unit cell, a gyroid lattice, and an “8D cubic lattice”. The mechanophore activation in each test specimen was measured as a function of uniaxial compressive strain applied to the structure. Full shape recovery after compression was exhibited in all cases. These proof-of-concept results signify the potential to use flex activated mechanophore for nondestructive, quantitative volumetric assessment of mechanochemistry in test specimens with complex geometries. Additionally, the integration of DLP-AM with flex activated mechanophore build materials enabled the creation of customizable, three-dimensional mechanochemically responsive parts that exhibit small molecule release without undergoing irreversible deformation or fracture.

© 2018 Elsevier Ltd. All rights reserved.

1. Introduction

Mechanophores are molecular moieties that respond chemically to external mechanical stimuli [1]. In materials that contain mechanophores, macroscopic strains are translated into molecular-scale deformations at the mechanophore, which augment the potential energy surface toward specific chemical reactivity. To date, mechanophores have been demonstrated in polymers and network materials to achieve behaviors such as color change (mechanochromism) [2–4], enhanced luminescence [5], fluorescence [6], catalysis [7], self-reinforcement [8], polymer backbone structural reconfiguration [9], and small molecule release [10]. Besides molecularly designing new mechanophores that undergo force-induced bond cleavage [11], cycloreversion [12], or electrocyclic ring opening [13], efforts have been made to bridge functional mechanophores with engineering applications. For example, Zhao and coworkers fabricated electro-mechano-chemically responsive elastomeric display systems that were able to generate fluorescent patterns by coupling the mechanochemical isomerization of

spiropyran with macroscopic shape deformation caused by applied electric fields [14]. In a demonstration of autonomously self-reinforcing materials, Craig and coworkers explored polymeric materials that contained *gem*-dibromocyclopropane mechanophores, wherein spontaneous nucleophilic substitution resulted in crosslinking and strengthening of the material under otherwise destructive forces [8].

As polymer mechanochemistry continues to develop as an avenue for designing stimuli-responsive materials, one can begin to consider opportunities at the interfaces of mechanochemistry and manufacturing. One of the most rapidly developing manufacturing techniques is additive manufacturing (AM), commonly referred to as “3D printing,” which has enabled the production of complex three-dimensional (3D) structures with relative ease [15]. Recently, the fabrication of rapidly customizable mechanochromic devices was achieved through the integration of custom mechanochromic filaments with melt material extrusion AM [16,17]. These reports signified a potentially exciting opportunity to pair molecular-level strain sensitivity with custom macroscopic object geometries. As an early step toward realizing the potential of AM with mechanochemically responsive build materials, we sought a mechanophore that could quantitatively “report” activation throughout a

* Corresponding author.

E-mail address: ajb1515@uw.edu (A.J. Boydston).

geometrically complex object.

In many cases, mechanophore activation is assessed through spectroscopic means, which involves either fabrication of test samples that are specific to the method of analysis, or processing of the bulk material in a way that destroys the test specimen (e.g., dissolving the material for solution-based analyses). Mechano-chromic responses can help address this challenge, but quantitative assessment is often limited to smooth, flat surfaces that are optically accessible. For instance, lattice and microstructured materials would each present formidable challenge to comprehensive assessment of mechanochromism. We hypothesized that mechanophores capable of releasing extractable, small molecules could provide a means toward quantifying mechanophore reactivity throughout the entire volume of a test specimen without requiring dissolution, digestion, or otherwise destroying the specimen. Flex activated mechanophores, such as oxanorbornadiene (OND), are potentially good candidates for such applications [18]. The OND mechanophore was previously demonstrated in bulk thermoplastics and polyurethane elastomers to release a small molecule (benzyl furfuryl ether) that could be extracted and quantified after mechanical activation (Fig. 1) [10,18]. One notable caveat is that the efficiency of extraction would be dependent upon the swelling characteristics of the bulk material and the dimensions of the test specimen. In this study, we present the first demonstration, and characterization, of 3D structures with complex object geometries fabricated via AM, which incorporate a flex activated mechanophore that enables the quantitative assessment of mechanophore activation within mechanically reversible regimes.

2. Results and discussion

We first selected an appropriate AM method based on the intended target geometries of our test specimens (e.g. lattice materials) and the inherent reactivity of the OND mechanophore. The desire to have overhangs and void spaces within lattice designs prompted the use of vat photopolymerization [19]. Moreover, the general thermal instability of OND substrates was likely to preclude melt material extrusion. As a subset of vat photopolymerization, we applied digital light processing additive manufacturing (DLP-AM) in our studies to enable efficient printing with visible light [20], therefore also avoiding photochemical degradation of the OND that

occurs upon exposure to UV light [21]. Previous success with customizable photoresins for DLP-AM, including the production of elastomeric and graded materials, encouraged the possibility of incorporating an appropriately functionalized OND mechanophore [22,23]. The stability of the OND moiety within the DLP-AM process was examined by irradiating an OND-diol in deuterated chloroform (CDCl_3) solution with white light from the projector used during the AM process. After 2 h of exposure to the projector light, the NMR spectrum (Figure S1) showed no changes in comparison with the pristine OND-diol. This result suggested that the AM conditions would not cause photochemical degradation of the OND moiety.

To incorporate oxanorbornadiene chemically into crosslinked polymer networks during the DLP-AM process, we prepared an oxanorbornadiene-dimethacrylate (OND-DMA) as previously described [10]. This structure enables crosslinking during photocuring with the mechanophore positioned within the bridging network segment, such that the mechanical strain can be transduced into chemical potential. The complete photoresin was comprised of 98.52 wt % of 2-hydroxyethyl acrylate (HEA), 0.49 wt % of Irgacure 819 as photoinitiator, and 0.99 wt % of OND-DMA. Additionally, 0.01 wt % of Nile Red was added to the resin as a light absorber to decrease the outgrowth during AM. The DLP-AM setup is diagrammed in Figure S2 and the schematic of the layer-by-layer printing process is shown in Figure S3, illustrating the direct formation of the mechanophore-containing polymer networks from liquid resins.

To investigate the printability of the resin and characterize the mechanical properties of the printed material, dogbone tensile test specimens (dimensions: 63.5 mm \times 9.53 mm \times 3.20 mm) were printed and evaluated via uniaxial tension tests. We first investigated the effects of layer cure time on the printed material's mechanical properties by the measuring the load as a function of monotonically increasing strain until the point of specimen fracture. As presented in Fig. 2a, the stress-strain curves of test specimens printed with layer cure times ranging from 5 s to 11 s showed similar mechanical behavior. The tests specimens also demonstrated a significant elongation at break, with strains ranging from 156% to 188% (Fig. 2b, elongation measured by tracking two marks made within the specimen's gauge region using a video extensometer). The gel fraction of a representative disc-shaped sample was determined by performing Soxhlet extraction (detailed

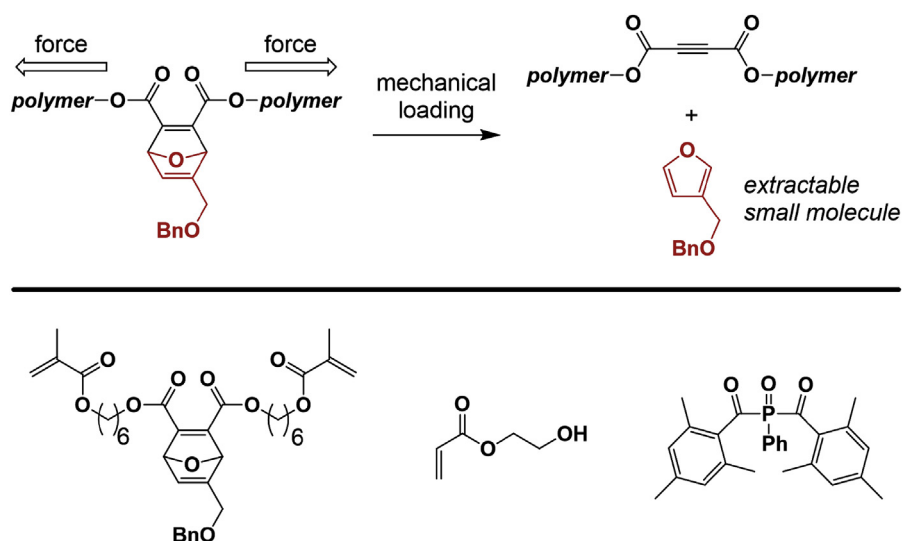


Fig. 1. (top) Generalized depiction of mechanochemical flex activation of an oxanorbornadiene under mechanical load. (bottom) Chemical structures of components of the photoresin formulation used in this study; left-to-right: OND mechanophore, 2-hydroxyethyl acrylate (HEA), Irgacure 819.

Download English Version:

<https://daneshyari.com/en/article/10154933>

Download Persian Version:

<https://daneshyari.com/article/10154933>

[Daneshyari.com](https://daneshyari.com)