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derived carbon particles to reinforce photo-cured shape memory composites†

Surface modification and characterization of waste

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Carbon fillers have been a source of inspiration to accommodate a range of surface chemistries for different applications. In this study different surface chemistries have been compared for shape memory effect on polymeric composites. Sugar industry waste (fly ash) has been utilized to prepare carbon particles named FCB. Surface modification of FCB has been done in two steps, oxidation and thiolation, respectively. In the first step, different reagents have been used to anchor the surface of FCB with oxygenated functionalities. In the second step, oxygenated FCB has been treated with a thiolating agent to covalently link thio groups on its surface. Polymeric composites have been photo cured with both types of particles, separately. A thermal actuation study has been carried out to check the shape recovery behavior of the composites. A guick shape recovery has been observed for thiolated FCB composites, due to thio linkages in the polymeric network. Samples have been characterized by scanning electron microscopy (SEM), attenuated total reflectance (ATR), dynamic light scattering (DLS), thermal gravimetric analysis (TGA), pH, conductivity, acid content particle dispersion, and composite gel content.

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1 Introduction

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Shape memory materials (SMMs) have the ability to retrieve their original shape in the presence of a particular stimulus. Researchers found a huge number of artificial materials for shape recovery under specific stimuli, including shape memory alloys,1 amorphous polymers,2 ceramics3 and shape memory gel.4 SMMs are being used in various industries i.e., textiles,5 pharmaceuticals,6 gas and oil,7 commercial products8 and aerospace.9 Epoxy matrices for SMMs present a vital range of encouraging properties i.e. convenient processing, low cost and light weight, to fabricate micro-devices,10 deployable structures,11 actuators,12 biomedical equipment13 and origami structures.14 The shape memory of epoxy, as per its strain, varies with regularity of interlinked hard and flexible segments. Strain in shape memory epoxies can be increased by two approaches. The first involves intense application temperature during deformation, and the second involves designed material nature.

One of the reported designed materials in epoxy for SMMs is carbon reinforcement.¹⁵ Surface chemistry of carbon particles, modified with different modifiers, affect their interactions with epoxy monomer¹⁶ as well as augment the mechanical properties,17 elastic and fracture toughness, thermal18 and electrical conductivities.19 Functionalization of carbon materials has been reported for reduced shape recovery time in epoxy composites relating to type of functionality introduced.20-22

The main focus of this research is to incorporate waste derived carbon particles into epoxy matrix for synthesis of SMMs. Moreover, carbon based epoxy composites are usually cured through heat,20 which is a source of pollution. But this research work has prepared functionalized carbon-epoxy composites through a unique eco-friendly photochemical approach.23 This photo-curing approach not only has minimized pollution risks but also has distinctive nature in energy saving24 and reduced curing time of composites.25 Composites prepared through this approach has been observed with high curing conversion and increased thermo-mechanical properties.23

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Materials

Fly ash (sugar industry waste) has been used to obtain filler particles (FCB). Potassium dichromate (BDH laboratory), hydrochloric acid (37% Sigma), bismithtrichloride (Sigma),

Both of these methods produce irregularity in epoxy composites to enhance shape memory properties.

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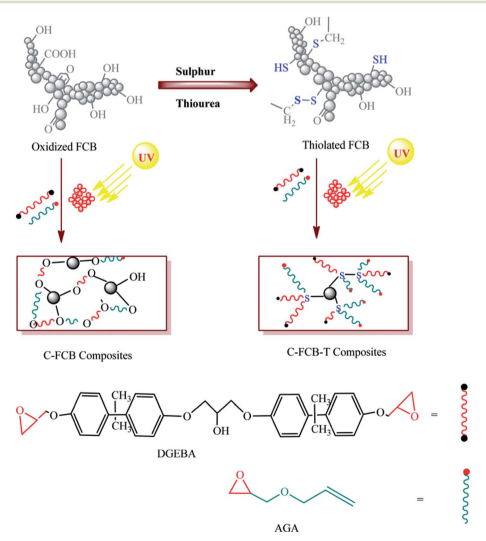
$$\begin{array}{c|c} CH_3 & CH_2 &$$

Fig. 1 Epoxies: DGEBA (a), AGE (b); PI (c).

hydrogen peroxide (35% solution), phenyl hydrazinium (Sigma), chloroaceticacid (Sigma), potassium chlorate (Sigma), sodium azide (Sigma), sulfur (Sigma) and thio-urea (Sigma) have been utilized as surface modifiers. Diglycidyl ether bisphenol A (DGEBA, Sigma), allyl glycidyl ether (AGE, \geq 99%, Sigma) have been utilized as monomers (Fig. 1a and b). Triarylsulphoniumhexaflouroantimonate salt (50 wt% propylene carbonate, Sigma) has been used as photo-initiator (PI) see Fig. 1c.

3 Methods

FCB was prepared by thermal treatment of fly ash mixed with reagents (as per Table S1†) in nitrogen atmosphere from r.t. to 500 °C with a ramp rate of 10 °C min $^{-1}$. Thiolation of prepared FCB particles (as per Table S2†) was carried out in nitrogen atmosphere from r.t. to 165 °C with a ramp rate of 10 °C min $^{-1}$. The obtained mixture was ground to powder.



Scheme 1 Preparation of filler and composites.

Table 1 Sample formulation for composite synthesis

Sample ID	DGEBA : AGE	Filler 1 wt%	PI : monomer
	(mole ratio)	of monomer	(mole ratio)
Pristine epoxy C-FCB C-FCB1-9 C-FCB-T C-FCB-T1-T9	1:2	Nil FCB FCB1-9 FCB-TP FCB-T1-T9	0.001:1

3.1 Composites synthesis

Composite samples were prepared, as per Scheme 1 and Table 1, through photo curing approach.²³ Weighed amount of both monomers with 1 wt% of functionalized filler were mixed by using magnetic stirrer for 25–30 minutes. PI was irradiated, with UV light (6 W) for one minute and mixed with filler-epoxy mixture.

3.2 Characterization

FCB particles were characterized by different instrumental and chemical analyses, including: FTIR/ATR (ALPHA-P Broker) spectra with absorbance range of 4000–500 cm⁻¹; SEM (NTRC; at 20 kV with ETD detector); DLS (Nanosizer, ZS90 Melvern), TGA (SKZ-1060) and acid contents using equation (i) through direct Boehm's titration.²⁶

$$n_{\text{CSf}} = [\text{B}] \times V_{\text{B}} - [\text{HCl}]V_{\text{HCl}} \times V_{\text{B}}/V_{\text{a}}$$
 (i)

For zeta potential (ζ , ZP) analysis 0.005 g of FCB particles were dispersed in 10 mL distilled water and ultrasonically mixed for 10 min. ζ values exposed the charge present at the surface of particles. For conductivity and pH test 0.1 g of each sample was dispersed in 10 mL distilled water, stirred for 30 min and pH/conductivity were taken at room temperature.

Composites were characterized by FTIR/ATR (ALPHA-P Broker) spectra with absorbance range of 4000–500 cm⁻¹, SEM (NTRC; at 20 kV with ETD detector); DSC and TGA (SKZ-1060), gel contents according to ASTM D2765-84, by using eqn (ii) and (iii)

$$\%Extract = (W_s - W_d)/W_s \times 100$$
 (ii)

$$\%$$
Gel content = $100 - \%$ extract (iii)

 $W_{\rm s}=$ weight of specimen being tested, $W_{\rm d}=$ weight of dried gel. ATR degree of cure (DC_{ATR}) was estimated by formula (iv), where A_0 and A_t denoted uncured and cured epoxy peak area (926 cm⁻¹), respectively.

%
$$DC_{ATR} = (A_0 - A_t/A_0) \times 100$$
 (iv)

Curing conversion through thermal-DSC (TA Q2000 DSC) was calculated by eqn (v).

% DC_{DSC} =
$$(1 - H_{DSC}/H_{Tot}) \times 100$$
 (v)

Shape recovery behavior of prepared composites was studied by indirect heating above glass transition temperature (T_g) *i.e.* 85–90 °C, which generated soft domains in samples, triggering chain mobility. Then samples were cooled and molded into desired temporary shape. Shape recovery time was found by placing deformed samples in pre-heated glass chamber. This heating released the stored elastic energy and recovery of original shape was observed by using eqn (vi)

$$\%SR = l_R/l_D \times 100$$
 (vi)

% SR = percentage shape recovery; $l_{\rm D}$ = length of total deformed shape; $l_{\rm R}$ = length of recovered shape.

4 Results and discussion

FCB particles have been prepared by thermal treatment of sugar industry waste. Different oxygen containing moieties (hydroxyl, carboxylic acid, phenolic and lactic acid) were introduced onto the surface of particles through different modifiers. The second phase of modification was carried out to introduce thio offshoots on oxidized FCB particles. Both oxidized and thiolated particles were used as filler in epoxy matrix to prepare shape memory composites.

FTIR/ATR absorbance analysis (500–4000 cm⁻¹) showed that after treatment FCB-1, 5 and 6 (Fig. 2a) have peaks at around 3300 cm⁻¹ verifying hydroxyl group generation. Characteristic peak at 1600 cm⁻¹ in all samples suggest presence of carbonyl group.²⁷ A further confirmation of these functionalities has

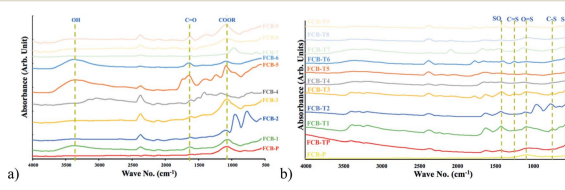


Fig. 2 ATR analysis of (a) oxidized, (b) thiolated FCB particles.

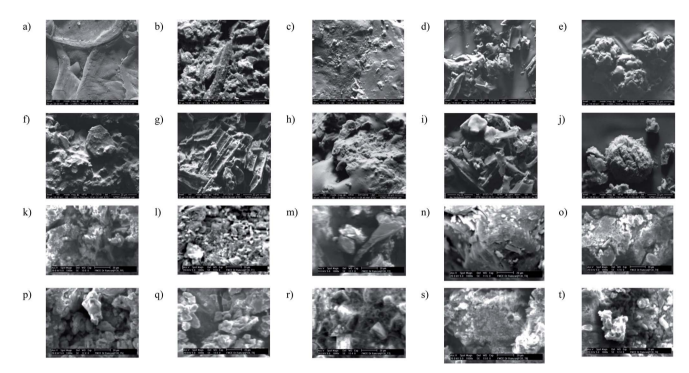


Fig. 3 SEM images of oxidized (a) FCB, (b-j) FCB 1-9, (k) FCB-TP, (l-t) FCB T1-T9

been done through Bohem titration (Table 3). In thiolated particles (Fig. 2b) prominent peaks in FCB-T1, T2, T3 and T6 at 560 cm⁻¹ (S–S), 640 cm⁻¹(C–S), 1030 cm⁻¹ (S=O), 1250 cm⁻¹ (C=S) and 1340 cm⁻¹ (sulfate/sulfones) confirmed effectiveness of thiolation treatment.^{27,28} Sulfur functionalities particularly disulfide and sulfones have been reported for shape memory response.²⁹ A distinctive peak has been observed at 914 cm⁻¹ in two samples *i.e.* FCB-2 and FCB-7, treated with potassium dichromate and potassium chlorate. Both of these modifiers have been reported for oxirane manufacturing.^{30,31}

TGA (25–800 °C) of oxidized FCB showed better thermal stability than thiolated FCB (Fig. S1a and b†). This might be attributed to the fact that during preparation, oxidized FCB particles developed compact agglomerated structures whereas thiolated particles grew amorphous morphologies. All oxidized FCB particles showed 20% weight loss at 800 °C, except FCB-5 which showed 40% weight loss. In contrast to that all thiolated particles started decomposing at 300 °C and showed 60–80% weight loss at 800 °C, except FCB-T1 which showed maximum weight loss at 340 °C and FCB-T2 which showed minimum weight loss at 800 °C (Fig. S1b†). These findings indicate that thiolated samples have better thermal dissipation behavior, which proved to be a strength in thermally actuated shape memory behavior.

SEM analysis of oxidized and thiolated samples (Fig. 3) has been recorded to observe morphological changes in FCB structure. FCB has quite a smooth surface, but after modification all samples (Fig. 3b-j) presented rough surfaces. Harsh oxidative treatment has transformed larger smooth particles into smaller itched surface structures. No doubt, particle size is an essential component for its dispersive behavior in solvent or

media, yet, surface chemistry is inevitably vital in this context. FCB-4 (Fig. 3e) has shown excellent dispersion in commonly used solvents *i.e.* acetone, ethyl acetate and dichloromethane. The sample has acidic pH (4.2), higher conductivity (23.265 mS cm $^{-1}$) and ZP (-20.1 mV) than FCB, indicating effective surface modification for long lasting dispersion.

Table 2 Zeta potential, pH and conductivity of oxidized and thiolated FCB particles

		Ծ (mS cm ⁻¹)		
Sample ID	ζ (mV)	Melvern zeta	Conductometer	pН
FCB	-02.5	0.101	00.865	9.9
FCB-1	-22.0	1.090	27.700	7.8
FCB-2	-18.4	0.640	56.052	7.3
FCB-3	-22.7	0.245	01.775	3.4
FCB-4	-20.1	0.664	23.265	4.2
FCB-5	3.36	0.661	06.283	4.8
FCB-6	39.0	0.402	06.152	5.0
FCB-7	-36.2	0.911	23.578	8.5
FCB-8	-30.9	0.422	07.180	6.8
FCB-9	-19.2	0.277	06.142	6.7
FCB-TP	03.7	0.007	00.754	9.9
FCB-T1	13.9	0.823	17.092	8.3
FCB-T2	02.7	0.420	09.953	7.5
FCB-T3	10.4	0.0731	00.610	8.2
FCB-T4	22.4	0.134	09.247	4.8
FCB-T5	-18.5	0.037	03.765	5.6
FCB-T6	-21.5	0.252	02.559	6.3
FCB-T7	25.7	0.044	27.592	6.7
FCB-T8	-10.7	0.057	02.170	6.2
FCB-T9	-16.3	0.050	01.537	6.1

Table 3 Acid contents of unmodified and modified particles

Sample ID	Carboxylic content $(\mu \text{mol } g^{-1})$	Lactonic content $(\mu \mathrm{mol}~\mathrm{g}^{-1})$	Phenolic content $(\mu \text{mol g}^{-1})$	Total acid content $(\mu \text{mol } g^{-1})$
FCB	0.01	0.04	0.54	0.59
FCB-1	0.175	0.46	1.44	2.075
FCB-2	0.35	0.3	1.425	2.075
FCB-3	0.25	0.125	1.525	1.9
FCB-4	0.25	0.125	1.675	2.05
FCB-5	0.225	0.35	1.625	2.2
FCB-6	0.325	0.125	1.6	2.05
FCB-7	0.15	0.1	1.5	1.75
FCB-8	0.266	0.084	1.575	1.925
FCB-9	0.275	0.075	1.625	1.975
FCB-TP	00	0.44	0.44	0.88
FCB-T-1	00	0.28	0.6	0.88
FCB-T-2	00	0.55	0.43	0.98
FCB-T-3	00	0.65	0.31	0.96
FCB-T-4	0.04	0.36	0.52	0.92
FCB-T-5	00	0.36	0.54	0.9
FCB-T-6	00	0.32	0.56	0.88
FCB-T-7	00	0.28	0.67	0.95
FCB-T-8	00	0.37	0.65	1.02
FCB-T-9	00	0.4	0.44	0.84

Upon thiolation, a visible change observed is shiny white coating on particles, although slight morphological alterations have also been observed (Fig. 3k-t). The surface coated with thio moieties have also been confirmed by FTIR analysis. Surface covered with S-S, S=O and C=S have a major role in shape memory designing, by developing soft region of polymeric assembly.

A range of ZP for prepared samples indicates qualitative and quantitative diversity of charges present on particles' surfaces (Table 2). Oxidative treatment of unmodified FCB enhanced ZP values, then thiolation comparatively decreased ZP of samples. An augmentation of ZP at the first modification step signposts incorporation of charged bodies on FCB surface, then decline in ZP in the second modification step indicates substitution of

charges with other functionalities. Moreover, in both treatments a reduction in pH of samples has been observed. Oxidation process dropped high pH, causing samples to be more acidic, consequently strengthening cationic curing of epoxy through activated monomer (AM) mechanism. However, on the other side, hydrogen bonding probability increased causing a delay in shape recovery of oxidized FCB composites.

A diverse range in surface chemistry of carbon particles (carboxyl, phenolic, lactonic *etc.*) helps to control composite properties especially through inter–intra collaborations. An estimation of surface oxy groups of carbon materials has been made through Boehm titration.^{26,32,33} FCB-5, among oxidized FCB particles, has been observed with the highest amount of acidic contents, likewise FCB-T8 has been observed with

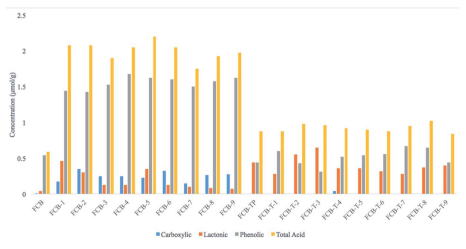


Fig. 4 Acid contents of unmodified and modified particles.

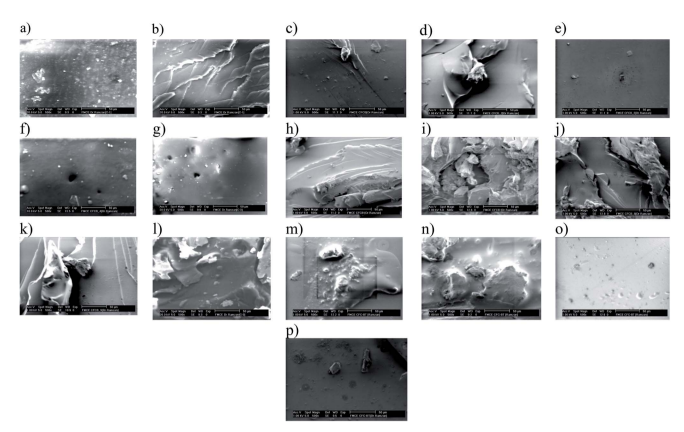


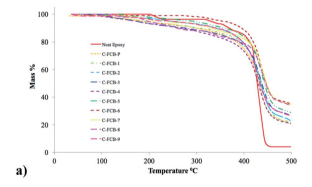
Fig. 5 SEM images of (a) Neat epoxy, (b) CFCB, (c-k) CFCB-1-9, (l) CFCB-TP, (m-o) CFCB-T1-T3, (p) CBCF-T6.

maximum acidic contents in thiolated samples (Table 3 and Fig. 4). Both these samples may have a trend to establish hydrogen bonded crosslinking ultimately producing rigid polymeric structure, with the least shape recovery behavior. Overall oxidized particles have been found with greater acidic contents, which have also been confirmed through FTIR/ATR analysis and ZP values.

Uniform dispersion of particles in solvent is essential to avoid agglomeration and subsequent improvement in composite properties. Unmodified and modified FCB particles have been dispersed in a range of solvents (acetone, ethyl acetate, dichloromethane and water) to check their dispersion stability (Fig. S2–S5†).

Oxidized particles showed excellent dispersion in water (Fig. S2a†), whereas only thiolated particle FCB-T1 showed comparable dispersion in water. The reason for oxidized samples could be high concentration of hydroxyl and carbonyl functionalities supporting dispersion, whereas for thiolated sample reason could be higher ratio of ionic bonds, originated from bismuth trichloride.

Dispersion analysis of prepared samples has also been done in a range of solvents, and different responses have been found. In acetone FCB-3, FCB-4 & FCB-T4 showed good dispersion (Fig. S3a and b†). In ethyl acetate FCB-4 and FCB-T1 showed excellent dispersion (Fig. S4a and b†). In DCM, FCB-3, FCB-4 and FCB-T1 gave better dispersion (Fig. S5a and b†). The observations clearly suggest that a variety in surface chemistry



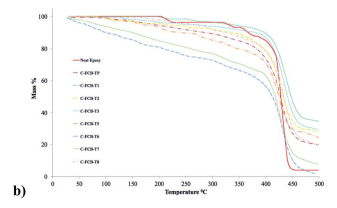


Fig. 6 TGA of (a) neat epoxy, unmodified and oxygenated FCB composites, (b) thiolated FCB composites.

Table 4 Shape recovery measurements of composites

	Deformed shapes				Trans	Dogovowy		
Sample ID	2 s	5 s	10 s	15 s	20 s	Trans Temp. (°C)	Recovery time (s)	Recovered shape
C-FCB-2	3	3		_		85	20	-
C-FCB-5	8	3	3	-	3		35	
C-FCB-6	2	2)	-	_		25	
C-FCB-7	8	8	8	5	<u>_</u>		55	
C-FCB-8	8	2	<	<	-		55	
C-FCB-T-1	C.	6			-		20	
C-FCB-T-2	0						20	
C-FCB-T-3	U	1	-	-			20	
C-FCB-T-6	3	×	-				10	

and morphology can support an array of inter-intra connections between particle-media.

After complete analyses and understanding of surface chemistry, prepared oxidized and thiolated particles were incorporated in epoxy matrix to design shape recovery composites. A comparison has been made between pristine epoxy, unmodified FCB composites and modified FCB composites. SEM images (Fig. 5b-p) evidenced dispersion uniformity of filler and matrix in all samples, irrespective of filler surface chemistry. This dispersion evenness might be attributed to UV curing approach, which links a separate phase of active center generation from PI, with sufficient homogenization period for filler and matrix.

Prepared composites were characterized for thermal stability by TGA in temperature range of 25–500 $^{\circ}$ C (Fig. 6 and Table S3†). It was observed that neat epoxy showed better thermal stability before 400 $^{\circ}$ C and the least thermal stability after 400 $^{\circ}$ C, which might be attributed to the fact that neat epoxy made a compact and thermally stable polymer till 400 $^{\circ}$ C which afterwards started disintegrating. On the contrary, composites showed quick weight loss till 400 $^{\circ}$ C, which indicates that oxy surface functionalities of fillers helped combustion process and

thermal damage to filler (carbon natured) subsequently destroyed polymer. After 400 $^{\circ}$ C, composite samples got stabilized by heat dissipation behavior of residual filler.

Curing conversion of neat epoxy and composite samples has been determined to verify crosslinking in polymeric network (Table S4 and Fig. S6†). It has been observed that incorporation of filler enhanced curing conversion, representing a strong filler–matrix interaction. Data revealed that composites having functionalized filler particles showed higher conversion than composite with unmodified filler.

Prepared composites have been tested for shape recovery by using indirect heating. Here the epoxy systems containing FCB particles as filler have been subjected to deformation and checked for shape recovery time at $T_{\rm g}$. Thiolated filler composites showed faster shape recovery than oxidized filler composites (Tables 4, S5 and Fig. S7†). The reason of difference in thermally actuated shape recovery response lies in surface functionalities developed during modification protocols. In oxidized composites oxygenated functionalities played dual role, *i.e.* hydroxyl functionalities not only contributed in epoxy curing through well-known activated monomer mechanism, but also developed hydrogen bonding at the same time.

Activated monomer mechanism develops C–O–C bond that adds to soft region of polymeric assembly, whereas hydrogen bonding develops linkages that reduce degrees of freedom which results in hard region of polymeric composites. On the contrary, in thiolated composites, thioether, disulfide, carbonyl sulfide and sulfone linkages all take responsibility for constructing soft region of polymer, hence support shape recovery upon thermal actuation. All these linkages are much more elastic than the oxygenated groups and helps in fast shape recovery.

Shape recovery measurements of composites are presented in Fig. S7.† Neat epoxy was prepared as standard to analyze the trend of shape recovery in matrix, and it was observed to have 63 s time to recover 100% of original shape. Out of oxidized samples C-FCB-2 showed minimum time (i.e. 20 s) to 100% shape recovery. The reason behind is elastic connections of FCB-2 with polymeric matrix. FTIR of the said sample showed the least hydroxyl species and a peak at 914 cm⁻¹ confirming oxirane like structure. Both these factors support the fact that FCB-2 (hard region) got incorporated into epoxy matrix through glycidyl linkage (soft region) which provided the support to quick shape recovery. The sample had neutral pH (7.3) which supports implemented UV curing procedure. FTIR of sample FCB-7 also showed the least hydroxyl content and oxirane peak at 914 cm⁻¹, but the shape recovery time is 2.5 times greater. The reason is justified by sample's basic pH (8.5) which decelerates implemented UV curing procedure, hence decreases the elastic connections between FCB-7 and epoxy matrix.

Thiolated particles imparted quick shape recovery behavior in composites. Samples C-FCB-T6 showed the least shape recovery time (10 s) in all prepared samples. Particle FCB-T6 was prepared from FCB-6 having acidic pH and high concentration of hydroxyl and carbonyl group, which anchored sulfur as thioether and carbonyl sulfide. Thioether itself has elasticity, and carbonyl sulfide developed thioether linkage as per proposed Scheme S1.†

Electropositive carbon got attacked by oxirane oxygen, then electronegative sulfur attacked another oxirane carbon, to make a link between epoxy monomers, during propagation.

5 Conclusion

Modification of filler particles with different modifiers, have resulted in specific surface functionalities, which has been confirmed through chemical and instrumental analyses. Oxidation seeded oxygenated functionalities that contributed hydrogen bonding in different components of composites, whereas thiolation introduced thioether, disulfide, carbonyl sulfide and sulfones onto particles' surface for thiolated bridges between different components of composites. Instrumental analyses have confirmed thiolated FCB with smaller morphology and better thermal dissipation. This added on to the shape recovery phenomenon in thiolated composites. Prepared particles have shown a range of pH and zeta potential values, labelling their effective dispersion and interaction capability with a range of solvents/media. Curing conversion of composites, determined by gel contents, ATR and DSC, have

shown thiolated composites with the highest values in comparison to oxygenated composites and neat matrix. Thiolated composites have shown very quick shape recovery within 10–20 s, whereas oxidized composites have shown it in 20–55 s and neat epoxy in 63 s. Structure activity study has shown the role of thioether surface functionalities in quick shape recovery, making them a promising candidate to be further explored for micro-devices, deployable structures, actuators, biomedical equipment and origami structures.

Data availability

The raw data has already been provided in the article to reproduce the findings.

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Conflicts of interest

The authors declare that they have no conflict of interest.

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