Microencapsulation of self-healing agents containing a fluorescent dye

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Abstract. Two different self-healing agent candidates, *endo*-dicyclopentadiene (*endo*-DCPD) and 5-ethylidene-2-norbornene (ENB), containing a fluorescent dye surrounded by a melamine–urea–formaldehyde (MUF) shell were microencapsulated by *in-situ* polymerization and the resulting microcapsules were characterized in this work. The microcapsules showed a narrow size distribution with a spherical shape and rough outer and smooth inner surfaces for both healing agent systems. Shell thicknesses of the microcapsules were ~880±80 nm for *endo*-DCPD and ~620±60 nm for ENB. The incorporation of a fluorescent dye as tracer into self-healing agents did not disturb the formation of microcapsules. The release of self-healing liquid into the induced crack from ruptured microcapsules in an epoxy coating layer was observed using a fluorescence microscopy. The use of a fluorescent dye is very effective in the observation of a damage site.

Keywords: coatings, self-healing, microencapsulation, fluorescent dye

1. Introduction

A lot of research on self-healing technology is underway for microcapsule-based coating systems over the last decade [1–6]. In this technique, microcapsules rupture upon damage inducing cracks in the coating layer, releasing their encapsulated liquid healing agent into the crack planes. The organic and inorganic self-healing materials, surrounded by urea-formaldehyde (UF) thermosetting shell were microencapsulated, and the microcapsules were embedded into different polymer coating materials, showing protective ability on steel plate (e.g., effective inhibition from metal corrosion and rusting) [1, 2]. A norbornene-based ring opening metathesis polymerization (ROMP) healing agent, *endo*-dicyclopentadiene (*endo*-DCPD), as self healing agent was also microencapsulated with UF shell [7–12]. UF capsules filled with *endo*-DCPD were developed in nano-size, which is important in fabricating thin coating applications [3]. Recently, self-healing agents, 5-ethylidene-2-norbornene (ENB) and ENB with crosslinkers, showing a faster ROMP rate at lower catalyst loadings, were microencapsulated with a ternary melamine–urea–formaldehyde (MUF) shell in this laboratory [11, 12]. The fabrication process is relatively very simple with no external control of pH, and produced a narrow size distribution of capsules with no debris. These microcapsules also had a rough outer surface and exhibited significantly higher thermal stability and less core material permeability when compared to UF capsules. Small changes in the core material during encapsulation process (e.g., going from one core material to a closely related core material), the manufacturing procedure (e.g., order of input), the ratio of the reactants and additives (e.g., typically the ratio between the main compo-
nents of shell forming reagents), and the reaction conditions (e.g., reacting temperature and time) often lead to significant changes in the formation of acceptable microcapsules.

One of the key aspects for achieving a successful microcapsule-based self-healing system lies with production of adequate microcapsules to be embedded. Thus, microcapsules must be carefully engineered in order to possess requisite strength during handling and processing, long shelf life during storage, and excellent adhesion with the cured polymer matrix [13]. It is also critical that the liquid healing agent within microcapsules is effectively delivered into the crack planes after damage in order to provide efficient recovery of properties. Therefore, it is beneficial to track the release of healing agent through invisible cracks from incorporated microcapsules for the development of self-healing. The addition of tracers such as X-ray dye [14] and UV fluorescent material [15] for hollow fiber reinforced composites and red dye [1] for microcapsule containing coatings into the self-healing agents have been reported for visual observations of damages. In this study, microcapsules were synthesized for two different self-healing agents, endo-DCPD and ENB, containing a fluorescent dye surrounded by MUF shell. A particle size analyzer (PSA), a thermogravimetric analyzer (TGA) and a scanning electron microscope (SEM) were used to investigate particle size/distribution and thermal resistance of the microcapsules and to observe morphology of the capsules, respectively. The MUF microcapsules containing a fluorescent dye were dispersed in an epoxy coating layer and a fluorescence microscope (FM) was used to track the transport of liquid healing agent through cracks after inducing the damage on the epoxy coating.

2. Experimental
2.1. Preparation of microcapsules and coatings
Two healing agent candidates, endo-DCPD (with 95% endo-isomer, Acros Chemical Co., Belgium) and ENB (Sigma–Aldrich, USA) as core material, were microencapsulated by in-situ polymerization of melamine (M) (Sigma–Aldrich, USA), urea (U) (Sigma–Aldrich, USA) and formaldehyde solution, 37 wt% in H₂O (F) (Sigma–Aldrich, USA) to produce the MUF polymer shell in an aqueous solution. Sodium lauryl sulfate (SLS, Junsei, Japan) was used as emulsifier, and poly(vinyl alcohol) (PVA, degree of polymerization = 500, degree of hydrolysis = 99.0 mol%, Junsei, Japan) as stabilizer. Figure 1 shows the microencapsulation process in this work. A fluorescent dye (derivative of 4,4’-diamino-2,2’-stilbenedisulfonic acid, Hwasung Chemical Co., Ltd., Korea) of 0.05 g/L was dissolved in self-healing agents at room temperature for 10 min before encapsulation. Also, SLS (0.5 wt%) and PVA (6.3 wt%) aqueous solutions were prepared by heating at 70°C for 20 min and 2 h, respectively. MF prepolymer solution was obtained from a mixture of 3.81 g melamine and 6.89 g 37wt% formaldehyde aqueous solution with 70 mL distilled water by heating at 70°C for 25 min until it becomes clear. Urea (0.61 g) was dissolved in 30 mL distilled water at RT in a 250 mL reaction beaker. Subsequently, the MF solution, 30 mL SLS solution, and 30 mL PVA solution were added into the reaction beaker, after raising the agitating speed to 300 rpm. Prior to slowly adding 30 mL of core healing agent (endo-DCPD or ENB + fluorescent dye) into the beaker, the agitation speed was increased to 500 rpm, leading to the formation of small droplets of the core materials. This agitation step was allowed to continue for 10 min at RT to generate the stabilized emulsion, before the temperature was raised to the nominal reaction temperature of 85°C for 40 min and maintained for 320 min under continuous agitation. The microcapsule slurry formed after the isothermal reaction was decanted on filter paper (Advantec no. 2, Toyo Roshi Kaisha Ltd., Japan). The microcapsules were separated into individual capsules easily by hand-shaking after rinsing and drying. The microcapsules produced were mechanically dispersed into diglycidyl ether of bisphenol-A
DGEBA, equivalent weight =188 g·eq⁻¹, Kukdo Chem., Korea) epoxy coatings cured with diethylentriamine (DETA, equivalent weight=190 g·eq⁻¹, Kukdo Chem., Korea) at room temperature for 24 hours. Table 1 has chemicals used for microencapsulation and epoxy coatings.

### 2.2. Characterization of microcapsules

The thermal stability of the microcapsules was examined with a thermogravimetric analyzer (TGA, Auto-TGA Q500, TA Instruments, USA) upon heating from room temperature (RT) to 500°C for ENB- and 600°C for endo-DCPD-microcapsules at a scanning rate of 10°C/min in a nitrogen atmosphere. The size and size distribution of the microcapsules were obtained with a particle size analyzer (PSA, Mastersizer 2000, Malvern Instrument, UK). The capsule surface and shell wall thickness were analyzed using a scanning electronic microscope (SEM, JSM-6380, Jeol, Japan) after the microcapsules were spread on an adhesive tape, punctured using a razor blade, and heated on a hotplate at 150°C for 12 hours to ensure that the core material completely evaporated.

### 2.3. Damage observations

DGEBA epoxy resin was mixed with 5 wt% of Grubbs catalyst using a mechanical stirrer for 10 min at 500 rpm, followed by adding DETA curing agent in an equivalent weight and mixing for additional 10 min. 5 wt% of microcapsules was added to the epoxy solution and mixed at a slower stirring speed of 150 rpm for 10 min. Air bubbles were removed under vacuum for 10 min. The mixed epoxy resin was then poured into a mold (50 mm × 5.3 mm × 0.2 mm) made by polyurethane, and then cured at RT for 24 h and at 50°C for 4 h. A fluorescent microscope (Axiovert 40 CFL, Zeiss, Germany) was employed to observe microcapsules before damage and the transport of healing agent through cracks after damage to the cured coating layer given by hand. The excitation wavelengths used for the observation in this work were λ =350, 480, and 546 nm.

### 3. Results and discussion

#### 3.1. Thermogravimetric analysis

Figure 2 shows the weight loss curves of endo-DCPD- and ENB-microcapsules containing a fluorescent dye and the MUF capsule shell without the core material. As shown in the figure, there are gradual and similar decreases down to ~70% in weight for both endo-DCPD- and ENB-microcapsules up to ~300°C above which there are sudden weight drops for ENB-microcapsules as reported in the previous work [11]. For endo-DCPD-microcapsules, the sudden drop occurred at ~430°C, much higher than that of ENB-microcapsules. The dramatic weight loss in a particular temperature of the microcapsules is due to a sudden release of the healing agent. As the temperature increases, the microcapsules become weaker and burst beyond the critical level at a certain temperature by increasing...
the internal pressure of the microcapsules and the collapse of the weak shells [11]. Notice the sudden loss of MUF shell mass at ~420°C, indicating thermal degradation, which would lead to a dramatic decrease in mechanical strength of the shell. The gradual weight loss below the temperature at which the sudden weight loss takes place may be attributed to the diffusion and evaporation of self-healing agent through walls of microcapsules. The thicker capsule shell thickness of endo-DCPD-microcapsules leads to the dramatic weight loss temperature higher than ENB-microcapsules, as will be shown in the SEM image.

3.2. Particle size analysis
Figure 3 shows the distribution of the microcapsules using a particle size analyzer. Average diameters of endo-DCPD- and ENB-microcapsules are found to be ~80 and ~52 µm, respectively. A relatively higher viscosity of endo-DCPD than that of ENB produces larger self-healing agent droplets, leading to larger endo-DCPD-microcapsules than ENB-microcapsules in average diameter. The particle size can readily be adjusted by rpm of propeller during dispersion of healing agents in water. The shoulder in the range of 1–10 µ in this figure may be due to the particles formed by self-coagulation of the wall materials.

3.3. SEM observations
Scanning electron microscope images for microcapsules filled with endo-DCPD and ENB were respectively shown in Figures 4 and 5. As noticed from the images in both figures, a perfect sphere
with similar inner and outer surfaces was produced. There is no debris for both self-healing agents. Looking at the surface is somewhat rough external surface and smooth inner wall. The rough outer surface of microcapsules is thought to contribute to improving the adhesion between the host matrix and capsules by increasing contact area with the matrix material. The thicknesses of the shells were found to be ~880±80 and ~620±60 nm from SEM images of endo-DCPD- and ENB-microcapsules, respectively.

3.4. Fluorescence microscope observations

In this study, the morphology of microcapsules and the release of the self-healing agent from microcapsules were observed after inducing cracks on an epoxy coating layer dispersed with 5 wt% of endo-DCPD-microcapsules by means of a fluorescence microscope. Figure 6 contains fluorescent microscope images (a), (b), and (c) before damaging and (d), (e), and (f) after damaging taken at different excitation wavelengths of 350, 480, and 546 nm, respectively. The vivid images could be viewed for all excitation wavelengths in different emission colors; blue at 350 nm, green at 480 nm, and red at 546 nm. Note that the image without a fluorescent dye showed completely black in color. The images showed a spherical shape of microcapsules as observed by SEM. It is interesting to notice that the capsule shell is very bright, indicating the incorporation of fluorescent dye into the shell material. Note that the dye was mixed with self-healing agent, followed by dispersion in water and then in-situ polymerization of shell materials reaction at higher temperature. This is presumably due to the migration of dye to shell forming material during the initial stage of reaction. The images (d), (e), and (f) in Figure 6 for cracked coating layer embedded with 5 wt% of Grubbs catalyst and 5 wt% of microcapsules show a bright line starting from microcapsules. This is considered to be the trace of transport of self-healing agent between crack planes released from a microcapsule. The brightest fluorescence was observed in the vicinity of the microcapsules where the release of self-healing agent starts to the cracks because the larger amount healing agent is spread to the exit of crack. Also, it is shown that the propagating crack runs around one of small microcapsule at left-top in the image. The black spots shown in the images (d), (e), and (f) are believed to come from the Grubbs catalyst in the matrix. In this work, the addition of a fluorescent dye to self-healing agent enables to observe the track of the self-healing agent release into cracks from microcapsules at three different UV wavelengths. This means that the addition of fluorescent dye to self-healing agent may be useful in studying the details of cracks which are crucial in a microcapsule-based self-healing methodology.

4. Conclusions

In this study, microcapsules containing self-healing agents, endo-DCPD and ENB, and a fluorescent
dye surrounded by the MUF shell were successfully synthesized and analyzed to develop a self-healing coating system. Thermogravimetric analysis showed that endo-DCPD-microcapsules have a significantly higher thermal stability than ENB-microcapsules. From scanning electron microscope observations, both endo-DCPD- and ENB-microcapsules have similar morphologies in shape and inner/outer surfaces but different shell thicknesses. The use of a fluorescent dye gave rise to the clear visual images of microcapsules and cracks filled the self-healing agent from ruptured microcapsules after damage at different excited wavelengths.

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References

Figure 6. Fluorescent microscope images for a coating layer dispersed with endo-DCPD microcapsules containing a fluorescent dye (before damage, left) and a fluorescent dye in the presence of Grubbs catalyst (after damage, right) at different excited wavelengths; (a) and (d) at $\lambda = 350$ nm, (b) and (e) $\lambda = 480$ nm, (c) and (f) $\lambda = 546$ nm


