



# A catalyst-free, temperature controlled gelation system for in-mold fabrication of microgels†

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**Anisometric microgels are prepared via thermal crosslinking using an in-mold polymerization technique. Star-shaped poly(ethylene oxide-*stat*-propylene oxide) polymers, end-modified with amine and epoxy groups, form hydrogels, of which the mechanical properties and gelation rate can be adjusted by the temperature, duration of heating, and polymer concentration. Depending on the microgel stiffness, the rod-shaped microgels self-assemble into ordered or disordered structures.**

Amine-epoxy chemistry is a long-established industrial technology, commonly applied in the fabrication of resins and adhesives. The reaction mechanism, mixing, and reaction kinetics have been extensively described before. These include the curing of epoxy resins with simple short amine linkers,<sup>1,2</sup> kinetic experiments investigating the addition reaction of primary and secondary amines in relation to their stoichiometry,<sup>3,4</sup> mixing behavior,<sup>5</sup> and water uptake of crosslinked networks.<sup>6</sup> To prepare hydrogels, the amine-epoxy click reaction was implemented for biocompatible poly(ethylene glycol) (PEG) or thermoresponsive poly(*N*-isopropylacrylamide) (pNIPAAm) based systems. In contrast to commonly applied (photo)chemical free radical acrylate polymerization or copper catalyzed azide-alkyne click chemistry, the amine-epoxy ring-opening addition works independently of an initiator or a catalyst. Macro- and microporous hydrogels were fabricated by crosslinking bifunctional PEG-epoxy with cystamine and diamino-poly( $\epsilon$ -caprolactone) using a salt as a sacrificial template.<sup>7</sup> Alternatively, PEG hydrogels containing antibacterial quaternary ammoniums were developed by combining thiol-yne and amine-epoxy click chemistry.<sup>8</sup> In the case of pNIPAAm, prepolymer backbones were functionalized with side-chain epoxy functions and coupled *via* degradable polyamidoamines to form an

injectable hydrogel system.<sup>9</sup> When epoxy functions in the pNIPAAm hydrogels were modified with proteins *via* their free amines, a positive effect on cell viability and proliferation was observed *in vitro*.<sup>10</sup> As this nucleophilic addition reaction can be induced *via* a thermal trigger, it is a facile but highly effective crosslinking mechanism that can be applied in a variety of hydrogel fabrication methods.

One type of hydrogel that has recently gained increasing attention is anisometric microgels, as they can be applied as building blocks to induce anisotropy in macroscopic materials. They have been produced *via* in-mold polymerization techniques using perfluoropolyether (PFPE) molds with low surface energy and defined cavities.<sup>11,12</sup> Commonly, (PEG)-acrylates are cross-linked into particles or microgels with varying size, shape, and geometry by photochemical curing using radical initiators in an inert atmosphere.<sup>13</sup> Functional groups that enable post-process modification with, for example, bioactive domains, were implemented by mixing side-chain functionalized acrylate moieties pre-crosslinking.<sup>12,15</sup> To fabricate soft rod-shaped microgels with polymer concentrations down to 10 wt/V%, sPEG-acrylates were dispersed in liquid inert PEG (200 g mol<sup>-1</sup>) as a non-volatile solvent.<sup>13</sup> *Via* Michael-type addition, these microgels can be functionalized with thiol-containing moieties.<sup>14</sup>

In this report, a simple and more versatile crosslinking alternative is presented to apply for in-mold polymerization. Star poly(ethylene oxide-*stat*-propylene oxide) (sPEG) polymers are functionalized with epoxy or amine groups to enable thermally induced catalyst and radical-free click chemistry as a versatile alternative to the established acrylate-based systems. The sPEG polymer backbone consists of 20% propylene oxide (PO) suppressing the characteristic crystallization of PEG at molecular weights exceeding 700 g mol<sup>-1</sup> (Fig. 1a).<sup>16</sup> Their fluid characteristics open up a broad range of potential applications, excelling those of commercially established PEG derivatives. Moreover, the multi-arm backbone of sPEGs enables fast cross-linking, while maintaining free endgroups to implement further modification. Amine-epoxy systems especially benefit from these properties as three different reactive groups can be

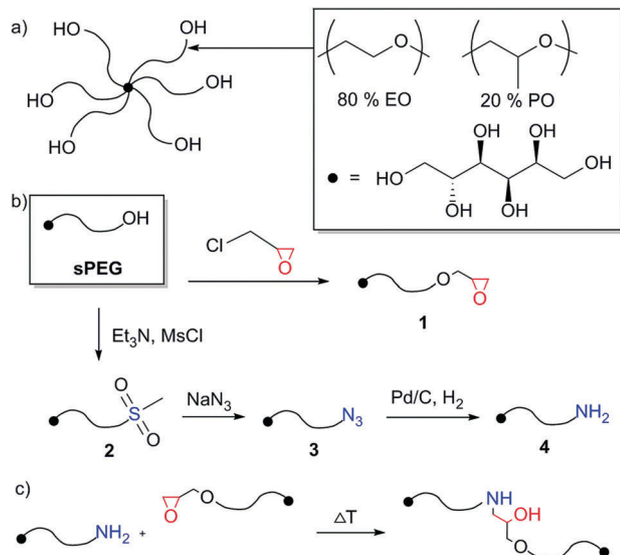
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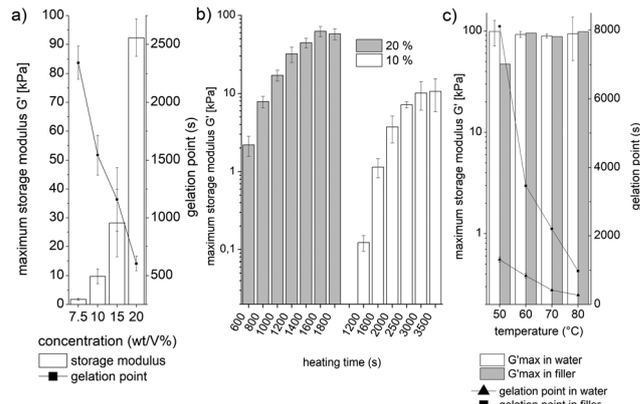


**Fig. 1** (a) The sPEG backbone contains a defined ratio of EO and PO moieties, inhibiting the crystallization at elevated molecular weight; (b) synthesis procedure for the production of sPEG-NH<sub>2</sub> and sPEG-epoxy (detailed procedures are provided in the ESI<sup>†</sup>); (c) gelation mechanism: a nucleophilic addition of amine to epoxy function.

provided: amine, epoxy, and hydroxyl moieties (Fig. 1). This allows for coupling a broad range of functions, *e.g.*, acrylates, carboxylic acids, amines, epoxy compounds (*e.g.* fluorophors<sup>17</sup>), NHS-active esters, alcohols, or even anhydrides. As a result, amine-epoxy gels can be specifically modified according to the desired application.

To obtain sPEGs with epoxy and amine endgroups, synthesis procedures are established to functionalize the liquid sPEG (Fig. 1b and ESI<sup>†</sup>). Briefly, sPEG-epoxy (**1**) is produced by the activation of sPEG with a strong inorganic base, followed by stirring with epichlorohydrin overnight. The procedure is adapted from previous reports.<sup>18</sup> sPEG-amine (**4**) is synthesised *via* a three-step reaction procedure, adapted from the literature.<sup>19</sup> The sPEG substrate is converted with triethylamine and mesylchloride leading to sPEG-mesyate (**2**). After reflux overnight with sodiumazide, compound **3** is obtained. Alternatively to an elaborate homogeneous catalysed Staudinger reduction, a heterogeneous catalysis with palladium on carbon in an 85 bar hydrogen atmosphere is performed in a high pressure autoclave reactor to reduce **3** to sPEG-amine.

The sPEG, modified with amine and epoxy endgroups, provides a system where three different parameters can be altered to control the mechanical properties, swelling behaviour, and gelation time of the hydrogels in an aqueous environment: polymer concentration, heating time, and the reaction temperature. The gelation time and mechanical properties are quantified by rheology. Prepolymer solutions with an equimolar ratio of amine to epoxy are applied at different polymer concentrations. The duration of heating and temperature are varied, while a solvent trap is installed around the sample to minimize water evaporation. First, the influence of the prepolymer concentration is investigated at 60 °C (Fig. 2a). This reveals that a minimum sPEG content of

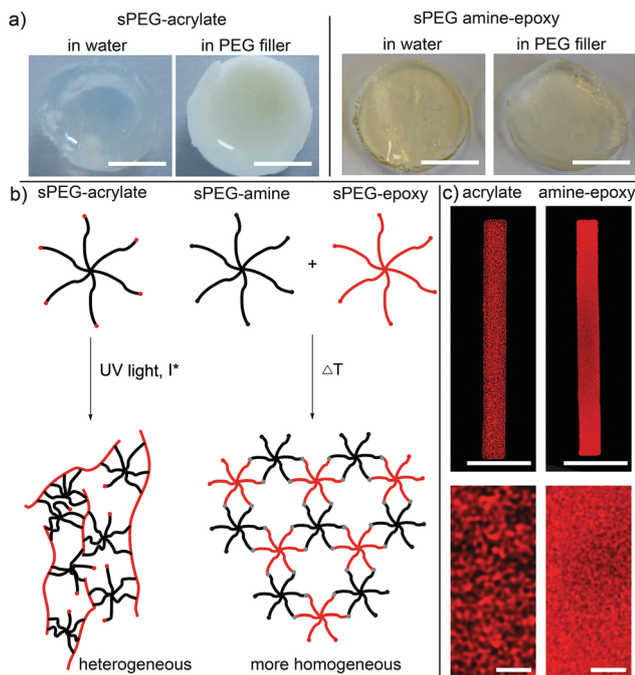


**Fig. 2** Hydrogel characterization (maximum  $G'$  and gelation point) (procedures and detailed conditions are listed in the ESI<sup>†</sup>). (a) Variation of sPEG concentration for heating at 60 °C, (b) the effect of different heating times at 60 °C for both 10 and 20 wt/V%, (c) gelation at different temperatures for both water and non-volatile PEG filler as a dispersing agent (20 wt/V% reactive sPEG); measurements are reproduced three times except for 2c with the filler due to long gelation times.

7.5 wt/V% is necessary to achieve crosslinked hydrogels, which have a maximum  $G'$  of 1.8 kPa. By increasing the polymer concentration, a range of different  $G'$  is achieved up to 92 kPa at 20 wt/V%. In addition to the increase in storage modulus, the initiation period for gelation decreases from above 2000 s to less than 600 s. Secondly, different heating times are compared to initiate and stop the gelation reaction and adjust the stiffness of the hydrogels (Fig. 2b). 10 or 20 wt/V% sPEG solutions are applied to the rheometer's heating plate at 25 °C and consequently heated to 60 °C for specific durations. In the case of 20 wt/V%, a range of gels with  $G'$  maximum of several kPa up to 62 kPa is realized as the heating time increases from 600 to 3500 s, respectively. When reducing the polymer concentration by half, the  $G'$  maximum ranges from approximately 100 Pa to 10 kPa, respectively. Thirdly, the effect of the dispersing agent on the storage modulus and gelation rate is investigated as water will be replaced by a non-volatile PEG filler for the microgel fabrication (Fig. 2c and chapter 6, ESI<sup>†</sup>). When comparing different heating temperatures, similar  $G'$  values are achieved independent of the dispersing agent or temperature. On the other hand, the gelation point is strongly related to the heat input and differs depending on the dispersing agent. In aqueous solution, a minimum gelation time of 261 s is detected at 80 °C, while for decreasing temperature, the gelation time prolongs up to 1236 s at 50 °C. In contrast, in the non-volatile PEG filler, gelation occurs four to eight times slower and the maximum  $G'$  is still not reached after 50 000 s in the case of 50 °C.

To compare hydrogels prepared with sPEG-acrylate *versus* sPEG-amine-epoxy, hydrogel discs are prepared in polydimethylsiloxane (PDMS) molds. In the case of the sPEG-amine-epoxy hydrogels, the solution is crosslinked at 60 °C for 45 minutes inside an oven. When using non-volatile PEG as the dispersing agent, the filler is successfully washed out post-gelation (Fig. S10b in the ESI<sup>†</sup>). As demonstrated before, hydrogel discs prepared with an sPEG-acrylate/linear PEG solution are, in contrast to





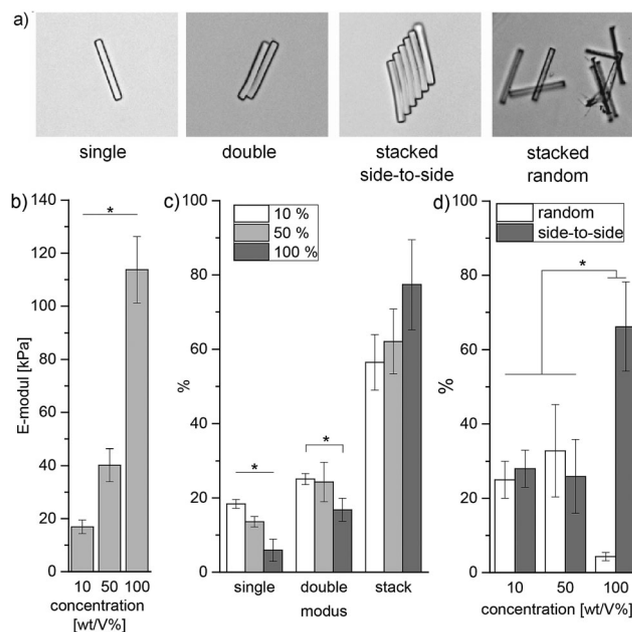
**Fig. 3** Transition from crosslinking in bulk to in-mold polymerization for the production of microgel rods. (a) Comparison of bulk-hydrogels: discs are produced in PDMS molds in either water or linear PEG ( $200 \text{ g mol}^{-1}$ ) as an inert filler, polymer concentration: 20 wt/V%, scale bar: 12 mm; (b) schematic of network formation using sPEG-acrylate or sPEG-amine-epoxy gelation systems: the acrylate produces heterogeneous networks, while the amine-epoxy networks are more ordered with more defined porosity. (c) STED-microscopy of microgels, fabricated with 20 wt/V% of the different pre-polymers and rhodamine B-acrylate, scale bar: 20  $\mu\text{m}$ ; below: magnification of porous microgel structure, scale bar: 1  $\mu\text{m}$ .

water-based gelation, white and non-transparent (Fig. 3a).<sup>13</sup> This change in optical properties indicates a reaction induced phase separation,<sup>20</sup> leading to hydrogels with a predominantly mesoporous character. In contrast, hydrogel discs fabricated with sPEG-epoxy and sPEG-amine are comparably transparent, regardless of whether the reactive polymer is dispersed in water or the inert PEG filler (Fig. 3a). This distinction in internal network structure may be explained by the different crosslinking mechanisms (Fig. 3b). Acrylate groups crosslink with the formation of linear polymer chains and distributed side functions, which connect the linear chains together to form a crosslinked network. Amine-epoxy click reactions (Fig. 1c) on the other hand result in a step-wise polymerization process, leading to a more ordered network.<sup>21</sup> The similar appearance of crosslinked hydrogels from sPEG-amine and sPEG-epoxy, when either water or the filler is used to disperse the polymer, is reflected in a similar swelling behaviour of the hydrogel discs at the same polymer concentration (Fig. S10a in the ESI†). Post-modification of the hydrogels is demonstrated by the coupling of rhodamine B-acrylate, indicating the presence of free amines after thermal crosslinking (Fig. S11 in the ESI†).

In addition to hydrogel discs, rod-shaped microgels ( $5 \times 5 \times 50 \mu\text{m}$ ) are produced *via* in-mold polymerization using the PEG filler (details in chapter 9, ESI†).<sup>13</sup> In the case of sPEG-amine

and sPEG-epoxy, the filled mold is placed in an oven at  $60 \text{ }^\circ\text{C}$  for 45 minutes. When compared to the sPEG-acrylate microgels, the thermally crosslinked microgels differ significantly in internal structure as imaged with stimulated emission depletion (STED) microscopy (Fig. 3c). To visualize the structure, rhodamine B-acrylate is coupled to the microgels by mixing the dye into the prepolymer solution before crosslinking. These results demonstrate that amine-epoxy microgels contain a more homogeneous porosity with a mesh size in the range of 1–100 nm, while the pores for the sPEG-acrylate microgels are more heterogeneous and in the micrometer range. Similar to the hydrogels, the stiffness of the microgel rods is altered by the polymer concentration and measured using atomic force microscopy (AFM) (Fig. 4b, procedure details in the ESI†). Solutions with sPEG-amine and sPEG-epoxy at 10 wt/V% result in softer rods with an elastic modulus of  $17 \pm 2.6 \text{ kPa}$ , whereas rigid and stiff microgels with moduli of  $40 \pm 6.2 \text{ kPa}$  and  $114 \pm 12 \text{ kPa}$ , respectively, are obtained with 50 and 100 wt/V% functional sPEG.

Interestingly, different self-assembly behaviours in water of the purified microgels are observed depending on the prepolymer concentration, and therefore on the stiffness of the individual microgels. Concentrated microgel solutions are compared for three prepolymer concentrations, 10, 50, and 100 wt/V%. The rods are counted and categorized by their degree of aggregation into four modes: single, double, stacked random, and stacked side-to-side. Stacks contain 3 or more microgels. A direct correlation



**Fig. 4** Analysis of microgel self-assembly: (a) stacking modes, the microgels are imaged using bright field microscopy; (b) individual microgel stiffness depending on the reactive polymer concentration, determined using AFM (procedure in the ESI†); (c) self-assembly is correlated with the reactive pre-polymer concentration used to prepare the microgels. The number of rods is counted and categorized according to the groups defined in (a); (d) comparison of random *versus* side-by-side microgel aggregation with respect to the polymer concentration. All measurements are reproduced three times.



between individual microgel stiffness and aggregation behaviour is observed when comparing the stacking modes and the rods' prepolymer content (Fig. 4b). With increasing microgel stiffness, the tendency to build aggregates rises and the amount of isolated microgels drops significantly. In the case of 10 and 50 wt/V% microgels, the stacks show similar amounts of random and ordered stacked gels, whereas for the stiffer 100 wt/V% rods, 66 ± 12% of the microgels are part of side-to-side stacks with a minimal tendency for random aggregation for only 4.3 ± 1.1% of the microgels (Fig. S10 in the ESI†). As microgels with an increased polymer concentration are more hydrophobic and behave more like solid particles, elevated attraction forces may be induced.<sup>22</sup> The interfacial free energy inside the more hydrophilic liquid is minimized *via* self-assembly of the microgels into ordered structures. Here, the high aspect ratio provided by the anisometric shape of the microgels further promotes this assembly behavior.<sup>22</sup>

In summary, a thermally induced gelation system is established, based on liquid sPEGs that crosslink *via* catalyst free amine-epoxy click chemistry. The system excels by its versatility, as the stiffness and gelation rate can be adjusted by three different parameters: polymer concentration, gelation temperature, and heating time. In addition, a broad range of possibilities are available to implement post-modifications. The thermal crosslinking mechanism facilitates many hydrogel fabrication methods, such as in-mold polymerization to fabricate soft microgels. In contrast to the UV polymerized microgels using this technique, the microgels formed *via* the click-reaction possess a more homogeneous nanometer mesh size. Depending on the individual microgel stiffness, self-assembly behaviour in an aqueous solution is altered. Therefore, the presented gelation system opens new avenues to prepare a platform of hydrogels and microgels with variable properties depending on the application.

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## Conflicts of interest

There are no conflicts to declare.

## Notes and references

- J. M. Laza, C. A. Julian, E. Larrauri, M. Rodriguez and L. M. Leon, *Polymer*, 1999, **40**, 35–45.
- F. González Garcia, B. G. Soares, M. E. Leyva and A. Z. Simões, *J. Appl. Polym. Sci.*, 2010, **117**, 2762–2770.
- J. P. Bell, *J. Polym. Sci., Part A-2*, 1970, **8**, 417–436.
- D. H. Kim and S. C. Kim, *Polym. Bull.*, 1987, **18**, 533–539.
- M. R. Vanlandingham, R. F. Eduljee and J. W. Gillespie Jr, *J. Appl. Polym. Sci.*, 1999, **71**, 699–712.
- P. O. S. B. Cursaru and M. Teodorescu, *U.P.B. Sci. Bull., Series B*, 2010, **72**, 99–114.
- Z. A. A. Hamid, A. Blencowe, B. Ozcelik, J. A. Palmer, G. W. Stevens, K. M. Abberton, W. A. Morrison, A. J. Penington and G. G. Qiao, *Biomaterials*, 2010, **31**, 6454–6467.
- C. Zhou, V. X. Truong, Y. Qu, T. Lithgow, G. Fu and J. S. Forsythe, *J. Polym. Sci., Part A: Polym. Chem.*, 2016, **54**, 656–667.
- A. K. Ekenseair, K. W. M. Boere, S. N. Tzouanas, T. N. Vo, F. K. Kasper and A. G. Mikos, *Biomacromolecules*, 2012, **13**, 1908–1915.
- M. O. Pehlivaner Kara and A. K. Ekenseair, *Biomacromolecules*, 2017, **18**, 1473–1481.
- J. P. Rolland, B. W. Maynor, L. E. Euliss, A. E. Exner, G. M. Denison and J. M. DeSimone, *J. Am. Chem. Soc.*, 2005, **127**, 10096–10100.
- M. P. Kai, A. W. Keeler, J. L. Perry, K. G. Reuter, J. C. Luft, S. K. O'Neal, W. C. Zamboni and J. M. DeSimone, *J. Controlled Release*, 2015, **204**, 70–77.
- J. C. Rose, M. Cámara-Torres, K. Rahimi, J. Köhler, M. Möller and L. De Laporte, *Nano Lett.*, 2017, **17**, 3782–3791.
- J. C. Rose, D. B. Gehlen, T. Haraszti, J. Köhler, C. J. Licht and L. De Laporte, *Biomaterials*, 2018, **163**, 128–141.
- K. P. Herlihy, J. Nunes and J. M. DeSimone, *Langmuir*, 2008, **24**, 8421–8426.
- J. Groll, J. Fiedler, E. Engelhard, T. Ameringer, S. Tugulu, H.-A. Klok, R. E. Brenner and M. Moeller, *J. Biomed. Mater. Res., Part A*, 2005, **74A**, 607–617.
- J. Yu, Z. Su, H. Xu, X. Ma, J. Yin and X. Jiang, *Polym. Chem.*, 2015, **6**, 6946–6954.
- R. M. Laine, S. G. Kim, J. Rush, R. Tamaki, E. Wong, M. Mollan, H. J. Sun and M. Lodaya, *Macromolecules*, 2004, **37**, 4525–4532.
- R. Mahou and C. Wandrey, *Polymers*, 2012, **4**, 561.
- E. Girard-Reydet, H. Sautereau, J. P. Pascault, P. Keates, P. Navard, G. Thollet and G. Vigier, *Polymer*, 1998, **39**, 2269–2279.
- V. M. Litvinov and A. A. Dias, *Macromolecules*, 2001, **34**, 4051–4060.
- Y. Du, E. Lo, S. Ali and A. Khademhosseini, *Proc. Natl. Acad. Sci. U. S. A.*, 2008, **105**, 9522.

