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Biomedical nanoparticle carriers with combined thermal and magnetic responses

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Recommended Citation

Liu, T., Hu, S., Liu, D., Chen, S., & Chen, I. (2008). Biomedical nanoparticle carriers with combined thermal and magnetic responses. Retrieved from http://repository.upenn.edu/mse_papers/159

Postprint version. Published in *Nano Today*, Volume 4, Issue 1, February 2009, Pages 52-65. Publisher URL: http://www.sciencedirect.com/science/journal/17480132

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Biomedical nanoparticle carriers with combined thermal and magnetic responses

Abstract

Several biocompatible polymers are capable of large responses to small temperature changes around 37°C. In water, their responses include shrinkage and swelling as well as transitions in wettability. These properties have been harnessed for biomedical applications such as tissue engineering scaffolds and drug delivery carriers. A soft material/hard material hybrid in which a magnetic metal or oxide is embedded in a temperature-responsive polymer matrix can combine the thermal sensitivity with magnetic signatures. Importantly, nanosizing such construct brings about new desirable features of extremely fast thermal response time, small magnetic hysteresis and enhanced magnetic susceptibility. Remote magnetic maneuvering and heating of the hybrid nanocolloids makes possible such applications as high-throughput enzyme separation and cell screening. Robust drug release on demand may also be obtained using these colloids and nanoparticle-derived thin film devices of combined thermal magnetic sensitivity.

Keywords

nanoparticles, biomedical, thermal response, magnetic response, drug delivery

Comments

Postprint version. Published in *Nano Today*, Volume 4, Issue 1, February 2009, Pages 52-65. Publisher URL: http://www.sciencedirect.com/science/journal/17480132

Nano Today (2008) xxx, xxx-xxx



REVIEW

Biomedical nanoparticle carriers with combined thermal and magnetic responses

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- Received 18 September 2008; received in revised form 13 October 2008; accepted 13 October 2008 9

sensitivity.

KEYWORDS

- Nanoparticles; 11
- Biomedical; 12 13
- Thermal response; 14
- Magnetic response; 15

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- Drug delivery
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Contents

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ature changes around 37 °C. In water, their responses include shrinkage and swelling as well

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23 Introduction

Smart materials responsive to multiple environmental stim-24 uli are of interest to biotechnology because of possible 25 applications such as delivery carriers, separation plat-26 forms and environment sensors. Since body temperature 27 is nearly constant, a small temperature excursion about 28 it provides an environmental stimulus to be exploited. 29 Temperature-responsive soft materials used in conjunction 30 with localized heating (e.g., via hyperthermia) are there-31 fore prime candidates for biomedical applications [1]. Other 32 stimuli such as pH, glucose, stress or strain, and electro-33 magnetic fields can be combined with thermal stimulus to 34 create a multi-stimuli-responsive system. Here we focus on 35 36 magnetic stimulus which can be applied remotely. One possible application of magnetically and thermally responsive 37 smart nanomaterials is illustrated in Fig. 1 that pertains to 38 remotely controlled drug delivery. 39

Since none of the soft materials suitable for biomedi-40 cal applications is magnetic, a soft-hard hybrid construct is 41 required to combine magnetic and thermal sensitivities. The 42 soft temperature-responsive materials of choice are those 43 that form hydrogel [2], which is a three-dimensional net-44 work of polymer that retains its structure while being water 45 absorbent; i.e., it swells, but does not dissolve, in water. 46 Common biomedical uses of hydrogels include soft contact 47 lenses made of silicone or polyacrylamide and medical elec-48 trodes made of polyethylene oxide. In some hydrogels, it is 49

possible to couple water absorption and network deformation to a temperature-stimulated phase transition, so the temperature response may be manifested as a large change in the shape, rigidity, water content or hydrophobicity of the gel. The hard magnetic material of choice is iron oxide, which is relatively safe for biomedical applications and can be readily synthesized in a form of small particles to be embedded into the soft material. Iron oxide can be attracted to a magnet. Moreover, using a high-frequency field remote magnetic heating of iron oxide becomes possible thereby converting a magnetic stimulus to a thermal stimulus.

Nanotechnology offers several advantages to these materials. Nanoparticles of iron oxide do not have multiple domains found in larger magnets; the unit-cell spins of the entire nanoparticle line up and act as a single "super" spin that aligns more perfectly with the applied field giving rise to a higher magnetic susceptibility. This "superparamagnetism" unique to nanoparticles provides a stronger magnetic response than bulk magnetism. Meanwhile, breathing water in a temperature-responsive hydrogel is easier for nanoparticles because of shorter transport distance, so their response to a temperature stimulus is much faster than that of a bulk hydrogel. In addition, smaller hybrid particles form more stable colloids and they circulate better in the body; at the same time they can more easily penetrate and accumulate in the leaky, defective architecture of growing, vascularizing tumors [3,4]. Nanosized iron oxide and polymer particles can also be more readily



Figure 1 Two drug release mechanisms under magnetic heating. Gentle magnetic heating causes temperature-responsive polymer to shrink, squeezing drug out from the nanoparticle. Intense magnetic heating additionally ruptures the nanoparticle, triggering a burst-like drug release.

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digested in the body through biodegradation and clearance 77 [5]. On the other hand, the stability of the nanoparticle 78 construct and its cargo against chemical dissolution and 79 degradation may be questionable. Moreover, the magnetic 80 force on nanoparticle is very small because of small mass. In 81 the following we will discuss the current status and under-82 standing of the nanoscale hybrid systems which have been 83 developed to exploit these thermal and magnetic responses 84 for biomedical applications. 85

Temperature-responsive polymers

Like all materials polymers manifest thermodynamic struc-87 tural transitions along with associated physical or chemical 88 responses. These changes are categorized by the phase dia-80 grams. Polymers, however, are unique in that their solutions 90 may thermodynamically separate into two distinct phases at 91 92 high temperatures, whereas in other materials such phase separations usually occur at low temperatures. Of special 93 interest for biomedical applications is the behavior of a poly-94 mer-water solution which is stable below a so-called lower 95 critical solution temperature (LCST), above which the solu-96 tion partitions into two phases: water and a polymer-rich 97 phase. This is in contrast to the phase separation below 98 an upper critical solution temperature (UCST) that is more 99 commonly encountered in non-polymer systems. Such LCST 100 exists for both homopolymers and block copolymers. Some 101 common ones are listed in Table 1. 102

Among the homopolymers that exhibit LCST, the most 103 studied is poly(N-isopropylacrylamide) (poly(NIPPAm) or 104 PNIPPAm) [6] (Fig. 2a) in which the LCST behavior repre-105 sents a coil-to-globule transition in the shape of a hydrated 106 polymer chain [7]. At low temperature, the chain solubilizes 107 water which keeps the chain extended. At higher temper-108 ature, the lost entropy of the ordered water around the 109 chain becomes energetically costly, so the water leaves 110 for the bulk and the coil collapses under the hydropho-111 bic force between polymer segments. Slightly crosslinked 112 NIPPAm is therefore a thermally responsive hydrogel that 113 shrinks above the LCST by rejecting water from the polymer 114



Figure 2 Chemical formula of two polymers that exhibit LCST. (a) PNIPAAm homopolymer and (b) PEO_PPO_PEO triblock copolymer.

network. Poly(*N*-vinylcaprolactam) (PVCL) is another extensively studied homopolymer with a similar LCST behavior

[8]. Among block copolymers, the most studied are the poly(ethylene oxide)-poly(propylene oxide)-poly(ethylene oxide) (PEO-PPO-PEO) triblock copolymers [9] (Fig. 2b). PEO, also known as PEG, is frequently present as a biocompatible hydrophilic coating on nanoparticles to improve their in vivo circulation [10]; PPO, on the other hand, is more hydrophobic. Commercially known as Pluronics[®] (BASF) or poloxamers[®] (ICI) this amphiphilic polymer is a non-ionic surfactant because within each chain the PEO blocks and the PPO blocks can self-segregate into hydrophilic and hydrophobic domains, respectively. Above the LCST, interchain aggregation also occurs, forming alternating PEO and PPO layers arranged into micelles (with a hydrophobic PPO core and a hydrophilic PEO shell), cylinders, lamellas or other supramolecular structures [11]. In this sense, the LCST also represents the critical micellization temperature (CMT) [12–13]. Stabilized supramolecular structures of PEO_PPO_PEO (via chemical crosslinking, physical entanglement with another interpenetrating polymer network, or adsorption to a water/oil interface) undergo a volumetric transition at the LCST due to water solubilization/rejection in the PPO layer. Moreover, at higher concentrations swollen micelles may gel reflecting an ordering tendency akin to colloidal crystallization which maximizes the free volume, hence entropy, around individual micelles. Some PEO_PPO_PEO polymers listed in Table 1 have an LCST close to the physiological temperature (37 °C).

Natural biopolymers generally exhibit multiple structural transitions at increasing temperatures, some causing large shape changes. For example, a single strand polypeptide can reversibly transform from a helix to a coil above a characteristic temperature, and two helical strands of complementary DNA reversibly dissociate when heated above the "melting" temperature. Such changes of secondary and tertiary structures of natural biopolymers have a profound effect on their biological functionalities. The helix-to-coil transition is not the LCST type, however, unlike the coil-to-globule transition in PNIPPAm. This is because the conformation change from helix to coil [14] is mainly controlled by hydrogen bonding between amino acids (base pairs) and is relatively immune to the entropy-dominated influences of solubilization and hydrophobicity. So the UCST here is essentially the "melting" temperature of the hydrogen bond (between a carbonyl oxygen and an amine hydrogen). Synthetic block copolypeptides containing hydrophobic and hydrophilic blocks have also been synthesized to exploit their thermal responses. Hydrophobic blocks in these diblock and triblock copolypeptides typically appear as α -helices or β -sheets, whereas random coils serve as the hydrophilic blocks. However, unlike PEO_PPO-PEO block copolymers that form micelles, lamellas or other ordered supramolecular structures, the aggregation of hydrophobic blocks in these copolypeptides commonly leads to long range gelation forming an ''amorphous'' hydrogel instead [15,16]. For example, between two helices of the ''leucine zipper'' type the aggregation takes the form of side-wise lineup of the two helices, providing physical (as opposed to chemical) crosslinks for the gel [17]. The thermal behavior of these hydrogels is again the non-LCST type since they "melt" at

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Homopolymers		Modified copolymers		Pluronic [®] series and similar triblock copolymers		Natural polymers ^a	
Materials	LCST (°C)	Materials	LCST (°C)	Materials	CMT (°C)	Materials	T _{gel−sol} (°C) ^a
Poly(<i>N</i> -isopropylacrylamide), PNIPAAm [71]	30-34	Poly(NIPAAm-co-AAm) [1,21]	35 <mark>-55</mark>	L64 [12]	24–45	Gelatin/collagen [48,49]	~40
Poly(N-vinylcaprolactam), PVCL [2,71,74]	25–50	Poly(NIPAAm-co- <i>N-</i> tBAAm) [1]	<30	P65 [12]	26-49	Polysaccharides [2,86]	30—50
Poly(vinyl methyl ether), PVME [71]	37		30—39	F68 [12]	27–53	Natural polymo	ers ^b
Poly(N,N-diethylacrylamide), PDEAAm [56,71]	25 <mark>-34</mark>	PNIPAAm-CA-PCL [67]	37–38	P84/P85 [12]	19—47	Materials	T _{sol−gel} ^b (°C)
Poly(methacrylic acid), PMAA [2]	~75	PNIPAAm-b-PMMA/PBMA [79,31]	32—35	F88 [12]	22–53	Methylcellulose, MC <mark>[2]</mark>	~80
Poly(vinyl methyl oxazolidone), PVMO [2]	~65	P(NIPAAm-co-SMA) [80]	~40	P103/P104/P105 [12]	18-32	Hydroxypropylcellulose, HPC <mark>[2]</mark>	~55
poly(dimethylaminoethyl methacrylate), PDMAEMA [75]	~50	Poly(NIPAAm-co-DMAAm)	32–44	F108 [12]	21–41	Polyphosphazene derivatives [2]	33—100
poly(N-(L)-(1-hydroxymethyl) propylmethacrylamide)[76]	~30	Poly(NIPAAm)-PL(G)A [68,69]	34-50	P123 [12]	13–26	Elastin-like polypepti	des (ELPs)
Poly(silamine) [2]	~37	poly(NIPAAm-co-HPMAm) series_[82]	10—50	F127 [12]	20–36	Materials	LCST (°C)
Poly(siloxyethylene glycol) [2]	10—60	PUA-b-PNIPAAm [83]	~31	PEO—PLA—PEO [60]	19–32	Poly(GVGVP) [71,74]	28–30
Poly(vinyl alcohol), PVA [2]	~125	Peptide-modified P(NIPAAm-co-AAc) [84]	~34	PEO-PHA-PEO [85]	22–45	Poly(GVG(50% Val-30% Gly-20% Ala)P) [21,74]	40-42
Poly(vinyl pyrrolidone), PVP [2]	~160	PVCL-g-PTHF [2]	35—50	PEO—PEA—PEO [85]	14–44	Poly(GVG(6% Val-50% Gly-44% Ala)P) [21]	67

^a Most natural polymers form a gel phase below $T_{gel=sol}$. At high temperatures, they have a random coil configuration forming a sol. At low temperature, renaturation to the triple helical conformation in gelatin and the double helical conformation in polysaccharides drives the formation of physical junctions, causing gelation. ^b Some natural biopolymers (e.g., cellulose) undergo reverse thermogelation (gelation at elevated temperature from a sol state at low temperature) at $T_{sol=gel}$.

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high temperatures by breaking loose the crosslinks. Similar non-LCST behavior is found in natural hydrogels and some examples are listed in Table 1. When gelatin is cooled below the gelation temperature, random coils of polypeptides self-assemble into triple-helices of the collagen structure, providing crosslinks [18]. In this case, both hydrogen bonding and hydróphobic aggregation contribute to gelation.

Since any protein solution eventually precipitates at 184 sufficiently high temperatures, hydrophobic collapse of 185 the polypeptide backbone must be ultimately inevitable. 186 Indeed, linear polypeptides made of monomers of a single 187 amino acid species have a well defined collapse temperature 188 which rises with the hydrophilicity of the respective amino 189 acid: 24°C for valine, 40°C for proline, 45°C for alanine 190 and 55 °C for glycine [19]. Therefore, by combining different 191 amino acids, it is possible to design linear homopolypeptides 102 that hydrophobically collapse near the physiological tem-193 perature. These so-called ''elastin-like polypeptides'' (ELP) 104 behave like PNIPPAm. For example, the LCST of an ELP made 195 of Val-Pro-Gly-Val-Gly repeats is 26 °C [19], which is raised to 42 °C by randomly substituting 50% val, 30% Gly and 20% Ala 197 for the second valine in the repeats. Such ELP may be suit-198 able for temperature-responsive drug delivery applications 199 [20,21]. 200

It is clear from the above discussion that the phase 201 transitions and the associated property changes of the 202 temperature-responsive polymers are fundamentally sen-203 sitive to the chemical and structural features of their 204 building blocks as well as their surrounding [1]. This 205 is unavoidable because the LCST transition reflects a 206 delicate balance between solubilization and hydrophobic 207 collapse, which involve electrochemical equilibrium and 208 electrostatic/electrodynamic interactions. These influence-200 exerting features start with the primary structure of the 210 polymer, including the hydrophilicity/hydrophobicity of the 211 monomers and their arrangement (e.g., random copolymer versus block copolymer). They also extend to the secondary 213 structure; for example, whether the hydrophobic block is a 214 random coil, α -helix or β -sheet makes a difference [15–16]. 215 Moreover, the chemistry and physical properties of the 216 modifications to the polymer and its environment, includ-217 ing crosslinking agents, intentionally incorporated additives 218





such drugs and imaging agents or unintentionally incorporated additives such as absorbed serum proteins, and the aqueous environment it is in (pH, salt concentration and dielectric constant), can all have a profound effect. Lastly, the molecular weight and polydispersity of the polymer are obviously important parameters as well. These factors should be taken into account in the design of any materials package involving temperature-responsive polymers.

Temperature-responsive nanocolloids

Although temperature-responsive polymers may be directly conjugated with drugs and used as such, a preferred form for controlled drug delivery entails the colloidal state in which the therapeutic substance is encapsulated inside the suspended nanoparticles [4]. Nanocolloids based on temperature-responsive polymers must remain stable in physiological electrolytes such as phosphate buffered solution (PBS) and serum. The typical size range of stable colloids prepared from common temperature-responsive polymers is shown in Fig. 3. Some examples of polymerbased temperature-responsive colloidal particles are given in Table 2.

Being an amphiphilic surfactant, PEO-PEO readily forms oil-in-water micelles with a PPO core and a PEO corona. Using double emulsion (water-in-oil-in-water) techniques (e.g., Fig. 4), one can also form PEO_PPO-PEO vesicles (liposomes or nanocapsules) with a shell made of a bilayer membrane that has hydrophilic, PEO-rich outer

Table 2 Volume changes and transition temperatures of colloidal particles made of temperature-responsive polymers. Volume change is generally larger for the Pluronic[®] series than for the PNIPAAm series. It also increases in the order of nanoparticles, microspheres/beads and nanocapsules.

Materials	Volume changes (%)	Transition temperature (°C)
PNIPAAm/iron oxide Beads [87] ^a	~85	~35
PNIPAAm microsphere [88]	~83	~35
Au/Boltorn H ₄₀ -NIPAAm nanoparticle [89]	~64	~32
Pluronic [®] F127/iron oxide nanoparticles [90]	~78	20–25
Pluronic [®] F127 nanocapsules [91]	~97	~26
Pluronic [®] F127/heparin nanocapsules [22]	~99	~25
Pluronic [®] F127/poly(ethylenimine) nanocapsules [92]	92—97	~21
Au/Pluronic [®] F127 core-shell nanocapsules [93]	~96	~18
Pluronic [®] F127/PEG nanocapsules [94]	~89	~23
Pluronic [®] F68 nanocapsules [91]	~98	~40
Pluronic [®] F68/iron oxide nanocapsules [91]	~94	~40
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Please cite this article in press as: T.-Y. Liu, et al., Nano Today (2008), doi:10.1016/j.nantod.2008.10.011

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faces [22]. These colloids dilate below the LCST and shrink above the LCST, with a radius ratio typically ranging from 2 to 5 (Fig. 5). Post-formation crosslinking adds stability to the colloids without substantially affecting their thermal responses. The core of the PEO_PPO_PEO micelle can incorporate hydrophobic substance such as drug, as can the shell of the bilayer nanocapsule; meanwhile the core of the bilayer nanocapsule can be loaded with hydrophilic substance as illustrated in Fig. 4.

PNIPPAm is a homopolymer and does not self-assemble 255 into micelles. However, latex-like colloids which exhibit 256 volumetric responses to temperature changes can be pre-257 pared starting with NIPPAm monomers and proceeding with 258 polymerization under emulsifying conditions that limit the 259 reactions within emulsion micro-reactors. The product is 260 often referred to as microgel [23,24] which may actually 261 reach the nanosize (less than, say, 300 nm) for PNIPPAm 262 [25] and PVCL [26]. More generally, PNIPPAm may be 263 modified in two ways to become sufficiently amphiphilic, 264 hence capable of self-assembly into nanocolloids [1]. First, 265 when the NIPPAm blocks copolymerize with blocks that 266 are more hydrophobic, the block copolymer self-assembles 267 into micelles with a hydrophobic core and a PNIPPAm-rich 268 corona. Conversely, when more hydrophilic pendants are 269 added to NIPPAm, micelles form above the LCST with a 270 PNIPPAm core and a hydrophilic corona; the micelles can 271 then be crosslinked to maintain stability below the LCST. 272 273 Triblock copolymer with both a hydrophobic end block and a hydrophilic end block can also be prepared [27]. A simi-274 lar approach may be applied to form ELP colloids [20]. The 275 above colloids also undergo volumetric transitions with a 276 typical radius ratio ranging from 2 to 4, while their cores 277 can again incorporate hydrophobic drugs. 278

The volume reduction of the colloid is obviously accompanied by water rejection. Accordingly, bulk or shell diffusivity may change significantly. In the case of hydrogel, there



Temperature-responsive tra Figure 5 diameter reduction above the LCST. F6 having a shell made of a bilayer of the PEO_PPO_PEO triblock copolymer known as Pluronic[®] F68. Its structure is similar to that illustrated in Fig. 4(b). F68-EDC refers to similar nanocapsules in which the outer PEO'shell is crosslinked by gelatin, which in turn is crosslinked by EDC. Its structure is similar to that illustrated in Fig. 4(c). F68-10 refers to fully crosslinked nanocapsules that additionally contain iron oxide nanoparticles in the core as illustrated in Fig. 4(d). The LCST may be identified with the inflection point of the size-temperature curve. The LCST is lower in F68-EDC and F68-IO mostly because the additive (NPC, see caption of Fig. 4), which reacts with PEO to render it crosslinkable, is less hydrophilic than PEO. Crosslinking constrains swelling at low temperature, so F68-EDC is smaller than F68 below the LCST. Filling the core with iron oxide nanoparticles further reduces shrinkage above the LCST.

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is evidence of a "dry skin" forming above the LCST that 282 decreases the diffusivity [28,29]. Pronounced changes in 283 surface properties are also experienced by some colloids. On colloids that have a PNIPPAm or ELP corona the surface switches from being hydrophilic to being hydrophobic as the LCST is exceeded, causing colloid to aggregate or even precipitate from the water solution [30]. The hydrophobic nanoparticles in the aggregate actually experience an addi-289 tional squeeze caused by the inter-particle adhesion and 290 osmotic pressure [30]. Such hydrophobic colloids have a strong tendency to adhere to the living cells. These changes 292 do not occur on PEO_PPO_PEO colloids which have a PEO corona that is always hydrophilic.

Magnetic-core/shell 295

A magnetic-core or shell as a part of the colloidal nanopar-206 ticle offers three opportunities: the magnetic colloid can 297 be attracted to the region of a high magnetic field H, it 298 can experience an internal stress as non-uniform distor-299 tion arises from magnetic forces, and it can be heated by 300 a non-contact magnetic field. The attracting field can be 301 either DC or AC since the magnetic body force is the gra-302 dient of the magnetic internal energy density $1/2\chi\mu_0 H^2$, 303 where χ is susceptibility and μ_0 is the permeability of vac-304 uum. Therefore, high-susceptibility material is favored for 305 magnetic localization. On the other hand, the heating field 306 is always AC typically in the radio-frequency (RF) range, 307 10⁴ to 10⁵ Hz. Since an AC field can generate an eddy cur-308 rent, induction heating is always feasible for any conductor, 309 but it becomes more efficient for a magnetic material in 310 which magnetic hysteresis causes additional energy dissipation. To maximize the sum of eddy current (Joule) heating 312 and magnetic heating, a relatively high electrical resistivity 313 and large magnetic coercivity (mainly due to the resistance 314 to domain wall movement) is therefore favored. However, 315 nanomagnets suitable for nanocolloids are superparamag-316 netic [31], i.e., it is a single-domain ferromagnet free to 317 switch following a quasi-static field without apparent coer-318 civity. So there is little coercivity contribution and whatever 319 energy dissipation must come from some sort of internal 320 or boundary "friction" (see below) which does not pre-321 vent switching but nevertheless drags the magnetic moment 322 letting it lag the AC field. In a linear-response medium, 323 the Debye theory describes this lag in terms of a relax-324 ation time τ [32]. It then follows that maximal dissipation 325 occurs when τ^{-1} is commensurate with the frequency f, i.e., $2\pi f\tau \sim 1$, because when $2\pi f\tau \ll 1$ there is no lag and when 327 $2\pi f\tau \gg 1$ the moment stops to respond. Therefore, effec-328 tive heating obtains by tuning the frequency to the range of 329 $2\pi f\tau \sim 1$; under this condition more heat can be generated 330 by driving the field harder (higher H) and faster (higher f). 331 Lastly, magnetic distortion can be caused by either a DC or 332 AC field as long as the frequency is not much higher than 333 the resonance frequency. There is little knowledge of the 334 magneto-mechanical resonance of colloidal nanoparticles 335 although typical experiments utilizing magnetic distortion 336 are conducted with a frequency much less than 10^3 Hz, a 337 condition unlikely to contribute to much heating. 338 339

Among magnetic metals Co is perhaps the only material suitable for the magnetic-core or shell; Fe oxidizes too easily

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at the nanosize and Ni is toxic to the body. Among magnetic oxides iron oxide (IO) is preferred. Iron oxide takes the form of magnetite (Fe₃O₄) or maghemite (γ -Fe₂O₃), both having the structure of spinel although γ -Fe₂O₃ is a highly defected spinel containing many vacancies in the sublattices of both Fe^{3+} and O^{2-} . Maghemite IO is clinically used as a contrast agent for magnetic resonance imaging (MRI) because it causes a (dipolar-type) field inhomogeneity which accelerates the spin-spin relaxation/decoherence in its surrounding [33]. The use of other ferrites, such as magnetic spinels with other 3d transition metals partially substituting for Fe [34,35] and haxaferrites such as BaFe₁₂O₁₉ [36], is not advised because of increased complexity for synthesis and uncertain profile of toxicity.

Since all the above oxides are insulators, only Co may benefit from eddy current heating. However, no report exists for incorporating Co into nanosized temperature-responsive polymer colloid (Co-containing micelles made of other block copolymers have been reported [37]). The strategy to incorporate IO into the core of a temperature-responsive polymer colloid varies according to the nature of the core. In an agueous solution, IO nanoparticles readily form from Fe(II) and Fe(III) salts at ambient or near ambient temperatures. After purification and recovery, the redispersed IO in an aqueous solution may be used as one part of the feedstock in the double-emulsion procedure to form the hydrophilic core of a PEO_PPO_PEO colloid (Fig. 4(a,b)). Alternatively, internal precipitation in the hydrophilic core which contains a Fe(II)/Fe(III) solution may be triggered by a pH increase after the formation of the colloid (Fig. 4(e)). For hydrophobic cores, hydrophobic IO nanoparticles need to be first synthesized, which typically involves high temperature precipitation in a long-chain alcohol such as oleic acid [38,39]. The oily IO can then be used in the emulsion procedure to enter the hydrophobic core. Since the procedure to grow spherical oily IO nanoparticles of a narrow size distribution from 3 to 20 nm (Fig. 6(a-d)) is rather well developed, it may also be used to prepare hydrophilic IO if it is modified with an additional step to introduce a hydrophilic outer coat using ligand exchange, physical adsorption or chemical conjugation [40,41]. Magnetic-shells containing IO are also possible. Since most shells of temperature-responsive polymer colloids are hydrophilic, magnetic-shells are synthesized using hydrophilic IO. This is typically achieved by either adsorption of IO nanoparticles or precipitation from aqueous Fe precursors [26,42]. Using IO nanoparticles as seeds to initiate polymerization, other magnetic-core/polymer-shell nanocolloids can also be synthesized as reviewed by Schmidt [31].

Under magnetic heating the temperature of the magnetic nanocolloid solution gradually rises reaching a steady state of several to several tens of degrees of centigrade higher. At this temperature, the heat input from the magnetic nanoparticles equals the heat loss at the external boundary (the container, fixtures, surfaces). What is informative of magnetic dissipation is the initial heating rate, typically of the order of 0.1-1 °C/s for colloids containing IO nanoparticles. Since the energy input of the solution is entirely from the energy input of the magnetic nanoparticles, the initial heating rate of the nanoparticle should be precisely C_W/C_MV_M times that of the (water) solution. Here C_W and C_M are the volumetric specific heat of water

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Morphologies, revealed by transmission electron microscopy, of iron oxide nanoparticles prepared from an oil-based Figure 6 solution. (a and b) are solid particles and (c and d) are hollow ones. The as-prepared nanoparticles are single crystals according to lattice imaging (b) and (d). (e) After magnetic heating, some hollow nanoparticles ruptured into pieces no longer in registry with each other, as indicated by markers.

and the magnetic material, respectively, and $V_{\rm M}$ is the vol-403 ume fraction of the magnetic material in the solution. Since 404 $C_W/C_M\,{\sim}\,1$ for IO and V_M is of the order of $10^{-3},$ the initial 405 heating rate experienced by the IO nanoparticle must be of 406 the order of 10^2 to 10^3 °C/s. The steady state temperature of the IO nanoparticle depends on the heat exchange mechanisms between IO and the surrounding, which are currently unknown. However, microscopy evidence presented in Fig. 7 410 for IO nanoparticles in the core of a PEO_PPO_PEO colloid 411 after RF heating suggests a rather high temperature of pos-412 sibly several hundred degrees of centigrade. Clearly, very 413 efficient "frictional" heating has been achieved. Magneti-414 cally caused fracture of hollow IO nanoparticles is also seen 415 in Fig. 6(e), and similar transmission electron microscopy 416

observations of magnetic-heat-rupture have been reported for silica nanoparticles coated with an (single crystalline) IO shell [43].

Assuming magnetic heating involves isolated, independent nanoparticles only, in an RF field friction arises in and around a magnetic particle from two sources [44]. First, particle may tumble causing frictional heating at the particle-water interface. The relaxation time $\tau_{\rm B}$ for this mode can be estimated as the time required for Brownian motion over a characteristic distance of the order of one particle diameter. From Stoke-Einstein equation and viscous drag on a spherical particle, one can estimate $\tau_{\rm B} = 3\eta V/kT$, where η is the viscosity at the interface, V is the particle volume and kT has its usual meaning. Brownian relaxation may not be



Transmission electron micrographs of F68-10 nanocapsules (see caption of Fig. 5) that show (a) uneven shrinkage after Figure 7 exposure to 45 °C, above the LCST. After magnetic heating, some nanocapsules ruptured (b), other coarsened into irregular shaped ones (c).

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Biomedical nanoparticle carriers

responsible for the frictional heating of IO seen in Fig. 7, 431 though, because the heat from this mechanism should be 432 about equally shared between the nanoparticle and water so 433 it is unlikely for IO alone to reach a very high temperature. 121 Friction may also arise from spin rotation without crystal-435 lattice rotation. The relaxation time τ_N for this mode (Neel relaxation) is the reciprocal of the spin flipping rate which 437 is of the order of $v_{\rm D} \exp(-\frac{KV}{kT})$. Here $v_{\rm D}$ is the Debye fre-438 quency of the order of 10^{12} /s and KV is the energy barrier for 439 coherent spin flipping which may be of a magnetocrystalline 440 or shape origin. Most IO nanoparticles of several nanometers 441 in size are superparamagnetic with a blocking temperature 442 typically around 50 K or lower. At the blocking temperature, 443 $\tau_{\rm N}$ should be of the order of 10^{-2} to 10^2 s, so we estimate 444 $\tau_{\rm N}$ to be of the order of 10⁻¹⁰ s at room temperature. This 445 would make Neel relaxation too fast to add to any significant 446 friction in a RF field. However, the IO nanoparticles in Fig. 7 447 came from internal precipitation at the ambient tempera-448 ture, so they are not perfect and most likely contain a high 449 concentration of crystalline defects. Such defects may not 450 significantly affect the blocking temperature and the super-451 paramagnetic characteristics measured at low frequency, 452 but they can greatly increase the friction against spin flip-453 ping thus causing lattice heating. This seems to be the most 454 likely magnetic heating mechanism for the IO nanoparticles 455 in Fig. 7. 456

457 Biomedical applications

Magnetically and thermally responsive nanocolloids may find 458 applications in medicine and biotechnology such as drug 459 delivery and enzyme immobilization/separation. Magnetic body force can align or relocate the colloid and magnetic 461 dissipation provides a means of remote heating. Temper-462 ature excursions can trigger a change in the size, water 463 content, diffusivity, surface properties and hydrogen bond-464 ing of the colloid. Although all of these individual effects 465 have been separately illustrated in numerous studies, there 466 are very few biomedically relevant reports that demonstrate 467 the combined magnetic and thermal actions in the nanocol-468 loid setting-bulk hydrogels and large (µm to mm) latex 469 particles are excluded. In the following, we summarize these 470 studies and comment on the pertinent mechanisms. 471

472 Magnetic heating of UCST colloids

Conventional synthetic polymers may experience increased 473 diffusivity and water content when magnetically heated 474 above the UCST, which may accelerate the release 475 of trapped drug or a model dye. This was reported 476 by Schmidt and coworkers for an IO-core-containing 477 poly(ε -caprolactone) (PCL) nanocolloid loaded with a solva-478 tochromic dye; PCL exhibits an UCST of 35 °C when dispersed 479 in dimethyl sulfoxide as in this study [45,46]. A more inter-480 esting study concerns a biopolymer with hydrogen bonding 481 that melts above the UCST; magnetic heating then causes 482 the release of hydrogen-bonded drug. This was demon-483 strated by Derfus et al. using IO nanoparticles to which single 484 strand DNA was grafted: the DNA binds a dye-labeled com-485 plement below the UCST, then releases it above the UCST 486 487 at the implanted site in a mouse tumor model [47]. Melting

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Figure 8 Cumulative release of a model drug (vitamin B_{12}) from F68–10 nanocapsules (see caption of Fig. 5) at various temperatures. The rapid increase from 37 to 45 °C is mostly due to nanocapsule shrinkage from 90 to 45 nm (see Fig. 5). The much faster burst-like release during magnetic heating is due to rupture of the nanocapsule (see Fig. 7).

hydrogen-bonding in a bulk gel magnetically heated above the UCST has been used to increase the diffusivity, hence drug release from IO-containing collagen [48] and gelatin [49]. Extension to microgels of submicrometer sizes is in principle feasible but not yet reported.

Magnetic heating of LCST colloids

Magnetic heating of the NIPPAm colloid above the LCST induces aggregation and size shrinkage. Wakamatsu et al. [50] applied the first effect to IO-core/PNIPPAm-shell nanoparticles to trigger their entrapment in a column packed with hydrophobic beads. We have applied the second effect to PEO_PEO nanoparticles to squeeze out a hydrophilic drug from the core (the preparation method is shown in Fig. 4, size isotherm in Fig. 5, reconstructed magnetic-cores in Fig. 7, and the release mechanism in Fig. 1). This is the first example of utilizing magnetic heating and size shrinkage to control drug release from a nanocolloid. The profile of drug release rates shown in Fig. 8 is very favorable: very slow at 4°C and 25°C, modest at 37°C (below the LCST), much faster at 45 °C (above the LCST) and bursting upon magnetic heating. Compared to an earlier example of µm-sized colloid (NIPPAm with [O) [51], the ratio of release under magnetic heating to that of 25 °C is at least a factor of 100 higher in this nanocolloid. A further comparison to other examples of magnetically triggered drug release (with or without a temperature-responsive polymer) is shown in Table 3.

Magnetic separation of LCST/UCST colloids

Magnetic support particles have been investigated for a long 516 517 time as a separation platform in biotechnology. Nanometer 517

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Table 3 Half-life (t_{50}) for drug release (typically a small molecule) from particles with and without magnetic heating. t_{50} is the time to reach $M_t/M_{\infty} = 0.5$, where M_t/M_{∞} is the amount released at time *t* normalized by the total amount of drug contained (see Fig. 8). Much faster release with magnetic heating.

Materials	Half-time $(t_{50})^a$	Released molecules	
	Without magnetic heating	With magnetic heating	
Pluronic [®] F68/iron oxide nanocapsules [91]	42 h (37 °C)/5 h (45 °C)	5 min	Vitamin B ₁₂
Pluronic [®] F127/iron oxide nanoparticles [90]	18h (15°C)/3h (45°C)	5 min	Doxorubicin
Fe ₃ O ₄ /PAH capsules [95] ^a	15h (25°C)	30 min	FITC_dextran
Silica/iron oxide nanospheres [43]ª	>20 days (25 °C)	3 min	Fluorescent dye
Silica/iron oxide nanospheres [96] ^a	>10 days (25 °C)	15 min	Ibuprofen
Ethylene-vinyl acetate with embedded magnetic sphere [97] ^b	Not measured (>40 days)	10 times shorter ^b	Bovine serum albumin

^a Without a temperature-responsive polymer.

^b 10 mm × 10 mm × 2 mm, not a temperature-responsive polymer, no magnetic heating, faster response due to magnetic distortion.

sized colloids can reduce fouling, but magnetic separation 518 becomes much more difficult because of the smaller mag-519 netic force in comparison to colloidal forces that favor 520 suspension and Brownian motion. Colloids made of LCST 521 polymers aggregate above the LCST, so they experience a 522 much larger magnetic force and smaller colloidal forces, 523 524 thus allowing easy separation by a relatively low field. This 525 has been demonstrated by Kondo and Fukuda [52] by heating IO-containing PNIPPAm colloids (150-250 nm) above 32 °C to 526 separate immobilized enzymes on the nanoparticles. The 527 dispersion-to-flocculation transition at the LCST was also 528 utilized by the same group [53] to achieve magnetically 529 aided affinity selection of target cells from phage display 530 libraries. Similarly, magnetic separation of UCST colloids can 531 be practiced below the UCST as illustrated by Kaiser [54] for 532 IO-containing polystyrene nanoparticles. In the latter case, 533 a hydrophobic solution (cyclohexane in this study) must be 534 used. 535

536 Magnetic directing of LCST colloids

In vivo localization of nanomagnetic particles is feasible 537 according to the study of Deng et al. [55] who localized IO-538 containing PNIPPAm nanoparticles (300-500 nm) to liver in 539 a rabbit using a DC magnetic field; without a field accu-540 mulation in other organs (lung, spleen, kidney and heart) 541 was observed. The colloid was initially placed below the 542 LCST to access the swollen state to soak up doxorubicin, 543 a hydrophilic drug for cancer treatment, although in vivo 544 demonstration of drug release was not performed in this 545 study apparently because the LCST (32-37 °C) is no higher 546 than the body temperature. In principle, AC magnetic heat-547 ing (hyperthermia) can also provide a localization effect 548 for LCST colloids since above the LCST the colloid will pre-549 cipitate with a tendency to adhere to the cells. Localized 550 hyperthermia was proposed as a targeting tool to direct 551 drug-loaded LCST colloids to tumors which are warmer 552 553 $(\sim 42 \,^{\circ}\text{C})$ than the rest of the body [30], but this idea has not been demonstrated for magnetic colloids.

Membranes of magnetically and thermally responsive colloids

Various magnetic hydrogels not unlike those previously mentioned [48-49] have been studied but one serious shortcoming of the macroscopic gels is their slow response time, which scales with the size to the second power reflecting the diffusion limit of water transport [56]. This can be overcome if the macroscopic construct is itself made of nanoparticles of temperature-responsive hydrogel. Since the diffusion time of nanoparticle is very short, the response of the construct is also very fast despite its macroscopic dimension. Indeed, the nanoparticles can even be embedded in another gel without affecting the response time as long as water exchange in and out of the nanoparticles can proceed locally. One such construct with a magnetic signature is a membrane made by gelling nanoparticles or by depositing nanoparticle colloids. For example, Csetneki et al. reported a membrane made of nanoparticles with an IO-containing polystyrene core which is coated with PNIPPAm [57]. The membrane was endowed with a special microstructure by applying a magnetic field during gelation (below the LCST) with poly(vinyl alcohol) crosslinking: the magnetic nanoparticles are lined up into necklace strings due to dipole-dipole interactions. Above the LCST, shrunk nanoparticles disrupt the microstructure causing a rapid increase in permeability as demonstrated by bovine serum albumin penetration. Using spin coating, we have fabricated a 50-µm film of IO-containing PEO_PPO_PEO nanoparticles on a silicon substrate to demonstrate magnetically actuated rapid dye release from this device (Fig. 9). Micro-implant devices constructed in a similar way may be used for magnetically controlled drug delivery.

In vivo delivery

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Hyperthermia via magnetically heating of IO has been studied in mice and human cadavers to treat breast tumors, 586

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Figure 9 A device prototype made of iron-oxide-containing nanocapsules. Drug-containing F68,-10 nanocapsules (see caption of Fig. 5) were spin coated onto a Si substrate to form a 60-μm thick film. Another PEO_PPO_PEO triblock polymer (Plurónic® F127) solution with a LCST of 22 °C was used in the spin-coating solution as a ''binder'' gel. After spin coating, a gelatin coating was introduced to crosslink the PEO shell of the nanocapsule. Magnetic heating triggers drug release from F68-10. A similar implant device may be used for controlled drug release.

showing tumor shrinkage and nuclear degenerations in 590 heated malignant cells [58]. A maximum temperature ele-591 vation ΔT up to 88°C was reported. A recent study 592 demonstrated deep cranial thermotherapy using magnetic 593 heating of aminosilane-coated IO applied to human glioblas-594 toma multiforme patients who also received MRI and 595 computed tomography (CT) for evaluation [59]. At a ΔT of 596 5-12 °C, patients reported no discomfort. For drug release, 597 we already mentioned (see Magnetic heating of UCST col-598 loids section) the study of fluorophore (a model drug) release 500 from magnetically heated IO that was pre-implanted into 600 a mouse tumor model [47]. Clinical use of dextran-coated 601 IO as a MRI contrast agent has also been a well-established 602 modality for liver imaging [33]. 603

In the above applications IO colloids were delivered by direct injection to the target sites. In recent years, in vivo animal studies have been used to demonstrate the possibility of targeted delivery and imaging of IO with tethered targeting moieties; for example, folate ligand has been tethered to the dextran coating of IO via a linker to target tumor xenografts that overexpress folate receptors [60]. In theory, if the self-directed IO colloids are well localized to the targeted tumor site, they can also be magnetically heated to treat tumor, but this has not

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been demonstrated in vivo. Indeed, although multifunc-614 tional nanoparticles capable of targeted delivery of imaging 615 agents and drugs is a much discussed concept, its in vivo 616 demonstration for magnetic colloids is so far rare; we know 617 of none for temperature-responsive magnetic colloids. In a 618 recent review of application of nanotechnology in cancer 619 therapy and imaging [61], only one was cited for simultane-620 ous targeted delivery of drug and imaging agent: it delivers 621 to targeted tumor cells small interfering ribonucleic acids 622 (siRNA) that are covalently tethered to the dextran coat-623 ing of [0 [62]. This study did not utilize magnetic heating, 624 magnetic directing or thermal sensitivity. Recently, Yang et 625 al. used core-shell magnetic nanoparticles (core containing $MnFe_2O_4$, a spinel ferrite and doxorubicin, an anticancer 627 drug) tethered with a breast-cancer-targeting antibody 628 (human epidermal growth factor receptor 2 (HER2)) to 629 simultaneously detect and treat cancer xenografts in mouse 630 models [63]. Although this study used an amphiphilic block 631 copolymer of poly(p,L-lactide-co-glycolide) (PLGA) and PEG 632 for the shell, which is not temperature-responsive, it should be possible to replaced PLGA-PEG by PEO-PPO-PEO or a PNIPPA copolymer. Using such a construct, functionalities of 635 magnetic heating, magnetic directing and thermal sensitiv-636 ity can in principle be incorporated into nanocolloid systems 637

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638 for self-directed simultaneous detection and treatment of 639 diseases.

640 Designing nanoscale systems

We begin this section with a few comments on the drug 641 release mechanisms in magnetically heated LCST colloids. 642 Although a generic increase in diffusivity at higher tempera-643 ture may play a minor role, the dominant mechanisms are all 644 related to structural changes due to the LCST transition and 645 magnetic field/heating. Clearly, the volumetric shrinkage 646 provides a potentially powerful driving force for drug release 647 from the core. Effective actuation requires core shrinkage, 648 which is easier for a soft core than for a hard core [64]. 649 However, volumetric shrinkage cannot account for the mag-650 netically triggered burst-like release in Fig. 8, which is much 651 faster than that achieved by heating to 45 °C (above the 652 LCST) alone. The burst-like release is most likely due to the 653 severe disruption of the IO core by magnetic heating. Other 654 structural changes in the pore structure of the shell may also 655 play a role. The changes may be caused by a thermal distor-656 tion akin to the one associated with a heated heterogeneous 657 network structure: some regions expand while others con-658 tract. Magnetic forces may also cause a structural disruption 659 of the shell when $2\pi f \tau_B \ll 1$, as shown in a low frequency 660 (300 Hz) study on magnetically triggered on off permeabil-661 ity switch across a polyelectrolyte shell surrounding a Co/Au 662 core of $5 \mu m$ [65]. Force-directed structural movement is 663 probably not important in the RF frequency range because, 664 to effect shell distortion, $2\pi f \tau_{\rm B} \ll 1$ must be satisfied for 665 a particle of the size of the colloidal particle-a condition 666 unlikely to be met. 667

We have already emphasized the importance of the 668 LCST/UCST temperature, the structural transitions and the 669 magnetic constituent of the nanocolloid that is responsive 670 to both magnetic and temperature stimuli. For in vivo drug 671 delivery, these temperatures should be a few degrees of 672 centigrade above the physiological temperature, and prefer-673 ably there is a large change in size and surface functionality. 674 Tuning the transition temperature must be tackled at the 675 system level, since as mentioned before the transition tem-676 perature is sensitive to all chemical and physical aspects 677 of the constituents of the polymer and its surrounding. A 678 soft core is preferred to effect core shrinkage [64]. Actu-679 ation will be more effective if the transition temperature 680 and the magnetic response are sharp. This requires a pre-681 cise control of the composition and microstructure including 682 a narrow distribution of the molecular weight of the polymer 683 and of the size of the IO nanoparticles. Efficiency of mag-684 netic heating is probably sensitive to the defect chemistry 685 of the IO, its control and characterization at the nanoscale 686 presenting a challenge. Cost, synthetic ease and scalability 687 for mass production are important and mostly dependent on 688 the chemistry and processes selected. 689

A successful system design should also address other issues of material chemistry and physics. First, safety and biocompatibility demand rigorous screening to eliminate any toxic chemical in the composition of the polymer and the process residue. A particularly complicated issue is colloidal and drug stability. Structural integrity of the nanocolloid obviously calls for substantial stability of the

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constituent polymer during storage and circulation, which may be improved by crosslinking. Nanocolloids tend to have longer circulation half-lives, but to help escape the fate of rapid clearance by macrophages or the reticuloendothelial system surface hydrophilic tethers of PEG or dextran (a polysaccharide) are beneficial [66]. Tethers may also reduce the absorption of serum proteins, thus avoiding enzymatic attack at the same time. Meanwhile, biodegradability of the temperature-responsive polymer would be desirable which may be introduced by incorporating biodegradable blocks or oligomers such as PCL [67], polylactic acid (PLA) [68] and PLGA [69], including their copolymers (in the PEO-PPO-PEO triblock copolymers, they should substitute for the PPO block) [60]. Concerning drug targeting, hydrophilic tethers mentioned above will mask the transition to hydrophobicity above the LCST of PNIPPAm and PVCL, so temperaturetriggered aggregation and cell adhesion is no longer possible. In this regard, moieties for receptor or ligand bonding to enable targeted delivery is a desirable functionality that can be attached to the nanoparticles via suitable surface tethers [61]. Another important issue is the trigger for drug release. Although a long residence time after localization at the target site may sometimes be enough for delivering drug, a more efficient scheme is to utilize a device that allows for nanoparticle internalization (e.g., via receptor-mediated endocytosis) [70] and drug release (e.g., via an acid-labile linkage that is broken in the lowpH environment of endosomes) [71,72]. Lastly, drug loading is dictated by the physical chemistry of the polymer and the drug during fabrication, so a condition which simultaneously allows for polymer reaction (including self-assembly) and drug incorporation need to be found [64]. Since these aspects will again impact the transition temperature and transition characteristics, a system engineering approach must be adopted to find a satisfactory solution for this nanotechnology.

Finally, injection of particulate substance (liposomes, micelles and other natural or synthetic particles) in the submicron size range may elicit allergic reactions such as cardiovascular, respiratory and cutaneous symptoms, including death [73]. Typically, such reactions are most severe upon initial exposure, and the frequency of particulate allergy in the 5–45% range seems to be much higher than that of classical anaphylactic reactions to drugs (for example, penicillin allergy occurs in <2%). Interestingly, the trigger dose of hypersensitivity reactions in mouse models is two orders of magnitude higher than that in reactive man, so many animal studies may not fq = l the threat of possible allergic reactions (interestingly, big models appear to exhibit a similar trigger dose as reactive man). Therefore, designing safe nanoparticle delivery systems for in vivo applications may pose the most serious though least considered challenge.

Acknowledgements

This work was supported by the National Science Council of the Republic of China, Taiwan under contract No. NSC96-2627-B-009-006 and NSC96-2113-M009-027-MY2, and by the US National Science Foundation under grant No. DMR-05-20020 (MRSEC).

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Please cite this article in press as: T.-Y. Liu, et al., Nano Today (2008), doi:10.1016/j.nantod.2008.10.011

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