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14 ABSTRA	Т							
Our final rer	ort for this prop	ram summerize	s how we (Harvard)	have or	eatly impro	wed	our understanding of	
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15 SURIEC	L TEBWS							
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Report Title

Programmable Matter - Final Report

ABSTRACT

Our final report for this program summerizes how we (Harvard) have greatly improved our understanding of assembly using magnetic levitation, and mechanical agitation. Additionally, we report on how we have developed several types of soft-actuators that change compliance and shape based on pneumatic pressurization.

The Nuzzo group (University of Illinois, Urbana-Champaign) describe a new form of stimuli-responsive polymeric actuator—a chemically-driven, mechano-polymer—that provides a new approach to effecting efficient, multidimensional and potentially "smart" programmable movements.

The Grzybowski group (Northwestern) report on how they employ precisely controlled, to a micrometer scale, external magnetic fields to direct assembly of nonmagnetic and magnetic colloidal systems into colloidal crystals or core-shell assemblies.

Enter List of papers submitted or published that acknowledge ARO support from the start of the project to the date of this printing. List the papers, including journal references, in the following categories:

(a) Papers published in peer-reviewed journals (N/A for none)

<u>Received</u> 2012/01/03 1I 8	Paper Madhav Mani, George Whitesides, Michael Brenner, Filip Ilievski. Self-assembly of magnetically interacting cubes by a turbulent fluid flow, Physical Review E, (01 2011): 0. doi: 10.1103/PhysRevE.83.017301
2012/01/03 1 7	Filip Ilievski, Aaron D. Mazzeo, Robert F. Shepherd, Xin Chen, George M. Whitesides. Soft Robotics for Chemists, Angewandte Chemie International Edition, (02 2011): 0. doi: 10.1002/anie.201006464
2012/01/03 1/ 6	Sindy K.Y. Tang, Ratmir Derda, Aaron D. Mazzeo, George M. Whitesides. Reconfigurable Self-Assembly of Mesoscale Optical Components at a Liquid-Liquid Interface, Advanced Materials, (06 2011): 0. doi: 10.1002/adma.201100067
2012/01/03 1/ 5	Katherine A. Mirica, Filip Ilievski, Audrey K. Ellerbee, Sergey S. Shevkoplyas, George M. Whitesides. Using Magnetic Levitation for Three Dimensional Self-Assembly, Advanced Materials, (09 2011): 0. doi: 10.1002/adma.201101917

TOTAL: 4

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(b) Papers published in non-peer-reviewed journals (N/A for none)

Received

<u>Paper</u>

TOTAL:

Number of Papers published in non peer-reviewed journals:

(c) Presentations

Number of Presentations: 0.00

Non Peer-Reviewed Conference Proceeding publications (other than abstracts):

Received

Paper

TOTAL:

Peer-Reviewed Conference Proceeding publications (other than abstracts):

Received

Paper

TOTAL:

Number of Peer-Reviewed Conference Proceeding publications (other than abstracts):

(d) Manuscripts

Received 2012/01/19 1! 3	<u>Paper</u> Sindy K.Y. Tang, Ratmir Derda, Aaron D. Mazzeo, George M. Whitesides. Reconfigurable Self-Assembly of Mesoscale OpticalComponents at a Liquid–Liquid Interface, Advanced Materials ()				
2012/01/19 1: 1	Filip Ilievski, Madhav Mani, George M. Whitesides, Michael P. Brenner. Self-Assembly of Magnetically Interacting Cubes by a Turbulent Fluid Flow, Physical Review E ()				
2012/01/19 1: 4	George M. Whitesides, Katherine Mirica, Audrey Ellerbee, Sergey S. Shevkoplyas. Using Magnetic Levitation for Three Dimensional Self-Assembly, Advanced Materials ()				
TOTAL: 3					
Number of Manusc	ripts:				
	Books				
Received	Paper				
TOTAL:					
	Patents Submitted				
Quality Control for G.M. Whitesides, S	Plastic Injection, Trieard, A.K. Ellerbee PCT/US/11/62399 filed 11/29/11				
Soft Robotics for Cl G.M. Whitesides, Z. A.Stokes PCT/US11/61720 fi	nemists, . Nie, R.F. Shepherd, F. Ilievski, X.Chen, R. Martinez, W.J. Choi, A.D. Mazzeo, S.Morin, S.W. Kwok, led 11/21/11				
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On-Demand and Re Z. Suo, D.J. Moone	On-Demand and Reversible Drug Release by External Cue Z. Suo, D.J. Mooney, N.D. Huebsch, X, Zhao				
	Patents Awarded				

None during reporting period

Awards

	Graduate Students			
ſ	NAME	PERCENT_SUPPORTED	Discipline	
	Gregory Morrison	0.25		
	FTE Equivalent:	0.25		
	Total Number:	1		
	Names of Post Doctorates			
ſ	NAME	PERCENT SUPPORTED		
	Ramses Martinez	0.25		
	Robert Shepherd	0.25		
	Filip Ilievski	1.00		
	Adam Stokes	0.75		
	Manza Atkinson	0.50		
	Ahmet Faik Demirors	1.00		
	FTE Equivalent:	3.75		

Names of Faculty Supported

6

NAME	PERCENT SUPPORTED	National Academy Member
George M. Whitesides	0.01	Yes
Zhigang Suo	0.10	
Michael Brenner	0.10	No
L. Mahadevan	0.00	
Ralph Nuzzo	0.10	
Bartosz Grzybowski	0.10	
FTE Equivalent:	0.41	
Total Number:	6	

Names of Under Graduate students supported

NAME

PERCENT SUPPORTED

FTE Equivalent:

Total Number:

Total Number:

Student Metrics

section only applies to graduating undergraduates supported by this agreement in this reporting period	
The number of undergraduates funded by this agreement who graduated during this period: 0.	.00
science, mathematics, engineering, or technology fields: 0.0	00
 number of undergraduates funded by your agreement who graduated during this period and will continue to pursue a graduate or Ph.D. degree in science, mathematics, engineering, or technology fields: 	00
Number of graduating undergraduates who achieved a 3.5 GPA to 4.0 (4.0 max scale): 0.0	00
Number of graduating undergraduates funded by a DoD funded Center of Excellence grant for	
Education, Research and Engineering: 0.0	00
The number of undergraduates funded by your agreement who graduated during this period and intend to work for the Department of Defense 0.0	00
he number of undergraduates funded by your agreement who graduated during this period and will receive	
scholarships or fellowships for further studies in science, mathematics, engineering or technology fields: 0.0	00

NAME

Total Number:

Names of personnel receiving PHDs

NAME

Total Number:

	Names of other research staff	
NAME	PERCENT_SUPPORTED	
T.J. Martin	0.40	
Tracie Smart	0.15	
FTE Equivalent:	0.55	
Total Number:	2	

Sub Contractors (DD882)

1 a. Northwestern University Evanston Campus 1 b. Office for Sponsored Research Northwestern University IL Evanston 602081110 Sub Contractor Numbers (c): Patent Clause Number (d-1): Patent Date (d-2): Work Description (e): The Grzybowski lab specializes in experimental and theoretically complex systems. Their work will Sub Contract Award Date (f-1): 5/15/2008 12:00:00AM Sub Contract Est Completion Date(f-2): 10/30/2011 12:00:00AM 1 a. Northwestern University Evanston Campus 1 b. Research and Sponsored Programs 633 Clark Street Evanston IL 602081110 Sub Contractor Numbers (c): Patent Clause Number (d-1): Patent Date (d-2): Work Description (e): The Grzybowski lab specializes in experimental and theoretically complex systems. Their work will Sub Contract Award Date (f-1): 5/15/2008 12:00:00AM Sub Contract Est Completion Date(f-2): 10/30/2011 12:00:00AM 1 a. University of Illinois - Urbana - Champaign 1 b. Office of Sponsored Programs 1901 S. First Street Champaign IL 61820 Sub Contractor Numbers (c): Patent Clause Number (d-1): Patent Date (d-2): Work Description (e): The Nuzzo lab specializes in designing, developing, and analyzing systems of components. Their re Sub Contract Award Date (f-1): 5/15/2008 12:00:00AM Sub Contract Est Completion Date(f-2): 10/30/2011 12:00:00AM 1 a. University of Illinois - Urbana - Champaign 1 b. Grants & Contracts Office Office of Business & Financial Services IL. 618207406 Champaign Sub Contractor Numbers (c): Patent Clause Number (d-1): Patent Date (d-2):

Work Description (e): The Nuzzo lab specializes in designing, developing, and analyzing systems of components. Their rc

Sub Contract Award Date (f-1): 5/15/2008 12:00:00AM

Sub Contract Est Completion Date(f-2): 10/30/2011 12:00:00AM

Inventions (DD882)

5 MagLev Self Assembly

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Patent Fil	ed in Foreign Countries? (5d-2)	Ν		
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5a:	George M. Whitesides			
5f-1a:	President and Fellows of Harvard	l College		
5f-c:	1350 Massachusetts Ave			
	Cambridge	MA	02138	
5a:	Sergey S. Shevkoplyas			
5f-1a:	President and Fellows of Harvard	l College		
5f-c:	12 Oxford Street			
	Cambridge	MA	02138	
5a:	Audrey Ellerbee			
5f-1a:	President and Fellows of Harvard	l College		
5f-c:	12 Oxford Street			
	Cambridge	MA	02138	
5a:	Filip Ilievski			
5f-1a:	President and Fellows of Harvard	l College		
5f-c:	12 Oxford Street			
	Cambridge	MA	02138	

5 Method for Measuring "Closness" in a network

Patent Filed in US? (5d-1) Y Patent Filed in Foreign Countries? (5d-2	2) N			
Was the assignment forwarded to the contracting officer? (5e) Foreign Countries of application (5g-2):				
5a: Gregory Charles Morrison				
5f-1a: President and Fellows of Har	vard College			
5f-c: 29 Oxford Street				
Cambridge	MA	02138		
5a: Lakshminarayanan Mahadeva	an			
5f-1a: President and Fellows of Harv	vard College			
5f-c: 1350 Massachusetts Avenue				
Cambridge	MA	02138		

5 On-demand and Reversible Drug Release by External Cue

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5a:	Zhigang Suo	,	, , ,	1
5f-1a:	President and Fellows of Harvard	College		
5f-c:	1350 Massachusetts Avenue	C		
	Cambridge	MA	02138	
5a:	David J Mooney			
5f-1a:	President and Fellows of Harvard	College		
5f-c:	29 Oxford Street			
	Cambridge	MA	02138	
5a:	Nathaniel D. Huebsch			
5f-1a:	President and Fellows of Harvard	College		
5f-c:	29 Oxford Street			
	Cambridge	MA	02138	
5a:	Xuanhe Zhao			
5f-1a:	President and Fellows of Harvard	College		
5f-c:	29 Oxford Street			
	Cambridge	MA	02138	
5 Quar	ity Control for Plastic Injection			
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5a:	Audrey Ellerbee			
5f-1a:	President and Fellows of Harvard	College		
5f-c:	12 Oxford Street			
	Cambridge	MA	02138	
5a:	George M. Whitesides			
5f-1a:	President and Fellows of Harvard	College		
5f-c:	1350 Massachusetts Ave			
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President and Fellows of Harvard C	College		
12 Oxford Street			
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Aaron Mazzeo			
President and Fellows of Harvard C	College		
12 Oxford Street			
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Stephen Morin			
President and Fellows of Harvard C	College		
12 Oxford Street			
Cambridge	MA	02138	
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Cambridge MA 02138

5a:	Robert Foster Shepherd				
5f-1a:	President and Fellows of Harvard College				
5f-c:	12 Oxford Street				
	Cambridge	MA	02138		
5a:	Zhihong Nie				
5f-1a:	President and Fellows of Harvard Co	ollege			
5f-c:	12 Oxford Street				
	Cambridge	MA	02138		
5a:	George M. Whitesides				
5f-1a:	President and Fellows of Harvard Co	ollege			
5f-c:	1350 Massachusetts Avenue				
	Cambridge	MA	02138		
5a:	Ramses Martinez				
5f-1a:	President and Fellows of Harvard Co	ollege			
5f-c:	12 Oxford Street				
	Cambridge	MA	02138		
5a:	Adam Stokes				
5f-1a:	President and Fellows of Harvard Co	ollege			
5f-c:	12 Oxford Street				
	Cambridge	MA	02138		

Scientific Progress

Please see attached.

Technology Transfer







Programmable Matter W911NF-08-1-0143 Final Report 8/1/11 – 10/31/11

DARPA/DSO ATTN: BAA 10-65 Dr. Gill Pratt 3701 North Fairfax Drive Arlington VA 22203-1714

Technical POC: Dr. Gill Pratt, DARPA/DSO Submission: Gill.pratt@darpa.mil

Submission from:

George M. Whitesides

Department of Chemistry and Chemical Biology Harvard University Cambridge, MA 02138 gwhitesides@gmwgroup.harvard.edu Tel: 617 495 9430 Fax: 617 495 9857 *Laboratory Manager* T.J. Martin tjmartin@gmwgroup.harvard.edu Tel: 617 495 9432 Fax: 617 495 9857

OUTLINE

1. HARVARD UNIVERSITY (WHITESIDES GROUP)

This program has greatly improved our understanding of assembly using magnetic levitation, and mechanical agitation. Additionally, we have developed several types of soft-actuators that change compliance and shape based on pneumatic pressurization. From our new understanding, we have published four papers (1-4), and filed two patents. In this final report, we summarize our key findings.

1.1. PROGRAMMING THE SHAPE OF SELF-ASSEMBLED OBJECTS VIA MAGNETIC FIELDS

1.1.1. Introduction

In our prior Programmable Matter reports, we described the levitation and three-dimensional (3D) selfassembly (SA) of mm-sized diamagnetic objects, suspended in a paramagnetic liquid, in a non-uniform magnetic field. The method employed a device that consists of two permanent magnets oriented with like poles facing each other (anti-Helmholtz configuration), and a container filled with paramagnetic medium. We assembled objects in 3D by suspending them in paramagnetic medium, and positioning the container between the two magnets. The final positioning of these objects in the liquid and their self-assembly is directed by gravitational forces, magnetic forces, and steric interactions (physical contact) among and between the objects and the container.

In order to program the shape of the assembled objects, we adjust the shape of the magnetic field in which the levitation is performed, the average density of the levitating objects, as well as the density distribution of each component. By controlling these parameters, we have assembled discrete objects into hierarchical structures and then changed the shapes of the structures in-situ.



Fig. 1.1.2 (top, left) Magnetic field shape to align (top, right) two mirrors that guide a laser through (bottom, left-right) a green filter, red filter, and IR filter.(4)

1.1.2. Field mediated self-assembly

By applying permanent magnets of equal strength, in an anti-Helmholtz configuration, a gradient is established, with the largest field at the magnetic poles and the lowest field at the centre of the poles. Depending on the magnetic susceptibility of an object, it will position its centre of mass at different positions within this field gradient, w.r.t. the paramagnetic fluid in which it is suspended.

We have demonstrated the utility of this phenomenon by magnetically levitating mirrors and optical filters and then allowing them to self assemble. The result is a simple device that selectively filters an incident laser beam (Fig. 1.1.2) (4). We also demonstrated the utility of magnetic levitation to self assemble soft and hard millimetre scale components.

1.1.3. Field and template mediated self-assembly



Figure 1.1.3 (top, left) Magnetic field shape to align (top, right) two mirrors that guide a laser through (bottom, left-right) a green filter, red filter, and IR filter.

Using only the magnetic field gradient for self-assembly does have some limitations – particularly, it does not currently provide a route for organization within a plane of equal field strength. To circumvent this limitation, we have incorporated levitating templates to direct the assembly of the levitating components within the field gradient (3). Fig. 1.1.3, a demonstrates templated assembly of magnetically levitated beads into an FCC packing.

In addition to templating assembly from the bottom up, we also directed the assembly internally to a template (Fig. 1.1.3, b) and externally from a template (Fig. 1.1.3, c).

1.2. Self assembly using mechanical agitation

1.2.1. Introduction

Figure 1.2.2 Photographs of Delrin (white) and Torlon (dark) spheres on an aluminum dish after agitation at for 90 sec and 150 seconds respectively. The lattice structure changed from predominately hexagonal (red) to predominately square (blue).

1.2.2. Energy dependent assembly

We studied the behavior of beads under different agitation conditions this led to the formation of various lattices depending on the rates of charging. When one type of bead charged quicker than the other the first structure formed was predominantly hexagonal. This structure could change to square when the number of both types of beads was equal and the energy was high enough to allow phase transition.

1.2.3. Finite size effects

We studied the influence of the dish size on the lattice formation of beads with a constant size. We found the structure was open with four to five beads of one type surrounding the beads of the other type when the size of the beads was large (d_{bead} ~ 0.1 d_{dish}) compared to the size of the dish. When the size of the beads was small We continued to develop systems in which selfassembly is driven by electrostatic forces between particles that acquire electrical charge triboelectrically. The particles that we used in these studies were polymer spheres and cylinders which either charged positively, negatively, or remained neutral when shaken together inside containers with flat horizontal bottoms. We also started the development of systems that self-assemble without attractive directional interactions.



Figure 1.2.3 Photographs of Delrin (white) and Torlon (dark) spheres (d=3.18mm) in a 2:1 ratio on 40mm and 100mm aluminum dishes. In the large dish, beads formed close packed hexagonal crystals and some unordered areas. In the small dish, beads do not pack closely

 $(d_{bead} \sim 0.0003 d_{dish})$ compared to the size of the dish, hexagonal structures formed. Towards the rim of the dish the hexagonal structures adopted more open conformations to follow the curvature of the dish.

1.2.4. Assembly in mixed systems

We studied the assembly of objects with different shapes, e.g. cubes and beads. The combination of these two objects in a 1:1 number ratio led to the formation of square structures independent of the shape of the objects; the main difference being that we needed higher agitation energies for cubes than for beads due to their inability to roll. When we changed the number ratio to 2:1 we formed unordered hexagonal structures if beads were in excess and square structures when cubes were in excess.

1.3. Soft actuators and Robotic gripper



Figure 1.3.2 (a) mould used to replicate the actuator in (b). (c) mould with 90° channels(2)

1.3.1. Introduction

The shape and compliance of moulded elastomer networks can change upon pneumatic pressurization. We have developed two classes of soft actuators, based on pneumatically actuated elastomeric networks. We have then used one of these classes to assemble a soft robotic gripper (2).

1.3.2.Bending Actuator

When a typical balloon inflates, it does so isotropically. By fixing one side of the balloon, and inflating, it will curl around the fixed side. In this program, we used soft lithography to fabricate inflatable channels in ultrasoft elastomers (pneunets) and then we programmed their inflation to cause bending (Fig. 1.3.2a, b). Previously, soft lithography relied on replicating silicone from moulds patterned using photo-lithography; these

moulds are usually simple 2D extrusions. In this program, we used a 3D printer to fabricate moulds with topography – a new development in soft lithography. By varying the position and angle of the channels on the mould, we were able to change the mode of bending of the replicated pneu-net from in plane (Fig. 1.3c, d) to a helical out of plane bending (Fig. 1.3d, e).

1.3.3. Reverse Curl



Figure 1.3.3. (a) Low stiffness material is stretched and then high stiffness material is chemically bonded to the surface of the stretched material. Upon relaxation of the strain, the low stiffness material attempts to return to its original dimensions by buckling the high stiffness material. (b) The rest state of the strain patterned composite is a coiled strip that (c) extends upon pressurization. We have also fabricated an *extending actuator* which begins in a coiled state and extends/straightens upon pressurization. We used our 3D printing device to form a negative mould of a pneumatic channel and mould this channel using low stiffness Ecoflex ©. This soft channel is then uniaxially strained and the

strain state is permanently patterned into the pneumatic channel by bonding a high stiffness material, such as paper, to one side of the channel, Figure a. Figure b is the result of this patterning using Ecoflex© as the channel material and paper as the strain fixing material, clearly the paper buckles as the channel relaxes some of the initial strain. Figure c is the result of pressurizing the channel in Figure b. As the microchannel stretches to reach the degree of strain encountered when the

paper was affixed, the buckling disappears and the microchannel extends. Possible uses of this soft actuation mode are chameleon tongue like grabbers, frog-like jumpers, and unfurling of sheets like sails.

1.3.4.Gripper

We arranged the bending actuators from section 1.3.2 into a gripping device (Fig. 1.3.4). Essentially, six pneu-nets, similar to those in Fig. 1.3.2c, d, where arranged at 60° to one-another in a planar array. We connected all six pneu-nets to a single pressure source–when we applied pressure, the gripper curled around an object. (2)

The compliant nature of the gripper allows it to wrap around irregularly shaped objects; additionally, the low pressures required to actuate the gripper allows it to handle delicate objects: We used this gripper to pick up an uncooked chicken egg and a live mouse without damaging them and without complex sensory feedback.



Figure 1.3.4. (left, top to bottom) A gripper picks up an uncooked chicken egg without breaking it. (right, top to bottom) A larger gripper picks up a live, anesthetized mouse without hurting it.

1.3.5. Reverse Gripper

Using a similar concept to the gripper based on bending actuation, we arranged the reverse curling actuators from **section 1.3.3 to create a "reverse" gripper. This gripper uses** pressure to actuate, but maintains grip in the rest state – without the need to continue applying pressure (Fig. 1.3.5).

We have used this gripper to pick up a variety of irregular objects (Fig. 1.3.5, e-g).



Figure 1.3.5. Process to fabricate a reverse gripper by bonding a prestretched Ecoflex body against paper. (e-g) Gripping different shapes: a frame, an sphere and a plastic cup.

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2. UNIVERSITY OF ILLINOIS, URBANA-CHAMPAIGN (NUZZO GROUP)

2.1. DRIVEN, LIGHT PROGRAMMABLE HYDROGEL ACTUATOR

We describe a new form of stimuli-responsive polymeric actuator—a chemically-driven, mechanopolymer—that provides a new **approach to effecting efficient, multidimensional and potentially "smart"** programmable movements. Prior work has provided numerous methods to trigger mechanical responses in organic actuators including, as examples: pH¹; electric field²; ion concentration³; and chemical or biological reagents^{4,5}. Among the varying types of external stimuli that have been explored, light driven mechanics have provided one of the most efficient and direct methods to achieve a programmable form of mechanical actuation⁶. This study focuses on eliciting a different form of light triggered mechanochemical actuation, one that elicits photochemically programmed mechanical responses that in turn might be harnessed to drive hierarchical forms of assembly/motion that are in fact dissipatively driven.

In this model system (one demonstrating a proof of principles for this class of phase change material), we use visible light (v ca. 450 nm) to trigger a peroxide coupled water splitting reaction that produces protons, which subsequently induce movements in pH responsive hydrogels. A schematic illustration of the light induced hydrogel response system is shown in Fig. 1a. Ruthenium (II)-trisbipyridine complexes have been shown to be an effective photosensitizing agent for this coupled sequence of reactions^{7,8}. The photostimulated conversion of persulfate at pH ~5 is particularly effective. This value is also appropriate for constructing an acid-modifiable pH sensitive hydrogel. For this demonstration we choose a polymer gel based on a polyacrylic acid backbone, because it affords transition pH value of exactly 5, thus matching the activity profile of the ruthenium trisbipyridine complex. In the secondary couple, metal oxide nanoparticles are used as the functional catalyst in the water-splitting reaction. We used in this case IrO2 for reasons of its well-known, and highly efficient, catalytic activity and stability under oxidizing conditions^{9,10}.

The chemistry used to construct the gel is a standard photo initiated polymerization under irradiation at 365 UV light. Acrylic acid (monomer, 175 mg, 2.4mM), methacrylate modified Ru(II)-tribipyridine (photosensitizer monomer, 1mg, 1.5x10⁻³mM) and N,N-methylenebis acrylamide (crosslinker, 5.00 mg, 0.03 mM) were dissolved in 1mL dimethyl sulfoxide (DMSO):H₂O=1:1 (v/v), then adding in DAROCUR 1173 (photoinitiator, 30 μ L,1.8x10⁻⁴ mM) to form the pre-gel solution. This chemistry allows facile prototyping of actuating objects using projection masks to shape the optic field. The gel is immersed in an IrO2 nanoparticle suspension overnight before application. This treatment strongly (essentially irreversibly) entrains the oxide nanoparticles in the gel, presumably via an associative binding to the positively charged Ru(II) segments in the polymer network (the IrO2 nanoparticles are stabilized against aggregation by coordination to negatively charged citrate groups during its synthesis).¹¹ In a prototypical demonstration, the composite gel is immersed in an air saturated water solution that has a pH value

equaling 5.6(Fig. 1b-1), since the concentration of CO2 is about $3.5x10^{-4}$ atm. Under light exposure in the presence of the reagent Na₂S₂O₈ ($3.3x10^{-4}$ M), a catalytic water splitting reaction is initiated that oxidizes H₂O, liberating oxygen and a correlated flux of protons. It is the latter elicit a subsequent localized decrease in pH. The latter change in pH value induces a swelling in the pH sensitive hydrogel (Fig. 1b-2)^{12,13}. Terminating the irradiation shuts down the water splitting reaction. The pH of the solution begins to recover, as a result—by consuming HCO₃⁻ in the system bath, the protons produced through water-splitting reaction can be partially neutralized. As the pH value increases again, the dimensions of the pH sensitive gel are partially restored (Fig. 1b-3). These are behaviors that operate in what is essentially a weak buffer limit.

These mechanics were harnessed to demonstrate an improved design-a double-layered hydrogel actuator, which can effect several cycles of reversible shape changes, ones that are self-sustained by contact with the ambient environment and fully controllable in phase by light. In the last report, we suggested a prototype of a promising integration strategy for an actuator, by combining 2 layers of hydrogels as a bimorph: (a) Ru(II) trisbipyridine monomer segment incorporated poly acrylic acid (Ru-PAA), which has the same formula as described above; (b) poly-hydroxyethyl methacrylate (PHEMA), which is photopolymerized from a mixture of deionized water (solvent, 1.5 g, 83 mmol), 2-hydroxyethyl methacrylate (monomer, 3.0 g, 23 mmol), ethylene glycoldimethacrylate (crosslinker, 60 mg, 0.3 mmol) (1.1 wt.%), and 2,2-dimethoxy-2-phenyl-acetophenone (photoinitiator, 195.0 mg, 0.76 mmol)¹⁴. In the new design the two pre-gel phases are set side by side and photopolymerized in a single layer gel, as shown in Fig. 2a. The pre-gel solutions as currently formulated scatter the incidented light so irradiance levels through the gel are not even. This feature embeds an intrinsic strain in the polymerized bimorph, generating a spontaneous curvature due to the gradient change in crosslinking density vertically, which is further discontinuous at the material boundary in the curved structure. The light driven water splitting reaction leads the Ru-PAA end to deswell, which reduces the swelling ratio differences between the top and bottom regions of the gel, thus relaxing its curvature. The PHEMA domain has no phase change capacity for this reason does not undergo any obvious change in volume. It is however moved in translation by the dynamics of the Ru-PAA gel (Fig. 2b). After the system was moved to the dark state, the gel begins to recover (Fig. 2c). The resulting asymmetric swell and deswell in between two ends of a single layer gel creates a walking motion, as shown in the plot (Fig. 2d) of tracking the position of the gel.

As currently formulated, the light programmable system does not fully recover the initial (pre-light) pH state (and thus bimorph strain), requiring as a result changing the solution environment to restore the initial curvature. Even though the protons can be continuously produced stably in this pH range, the consumption of the protons is solely dependent upon the following reaction (absent other reagents):

$$H_2CO_3 \leftrightarrow H^+ + HCO_3^-$$
 pKa=6.35 (eq.1)

The amount of bicarbonate is fixed by the concentration of initially CO₂ saturated solution. Given that the initial pH value equals 5.5, the new pH value reached at equilibrium post irradiance is 5.16. To fully optimize the actuator in a multi-cycle context will require an addition of a specific buffering system that does not impede the formation of the local pH gradient needed to drive the gel phase change response yet still effectively recovers an initial (more basic) swollen state in the PAA gel segments. Solid carbonates and other proton neutralizing additives are currently being examined to this end.

2.2. Stretchable electrodes embedded with a pH-responsive

POLYMERIC ACTUATOR

Previous work¹⁵ has demonstrated that, by performing the oxygen reduction reaction (ORR) in a microfluidic ambient, the electrochemical process can produce remarkable pH gradients in between working electrode (WE) and counter electrode (CE). The magnitude of these gradients is sufficient to induce a gradient change in the volume of essentially any pH responsive gel. We have developed a new system that allows us to exploit such gradients in a new form of polymeric actuator, one embedding stretchable electrodes of a microelectrochemical reactor into the pH responsive polymer network.

The fabrication of programmable hydrogel devices mainly includes fabricating stretchable electronic meshes on a Si or glass substrate, followed by transferring them onto a carrier substrate, and finally embedded the array into a responsive hydrogel matrix through photo patterning. The fabrication starts from depositing a sacrificial layer of PMMA on Si substrate in order to provide a means to release the electrode mesh. A thin polyimide layer is then deposited, followed by the steps to construct the active electrical components, in this case the pH actuating electrodes. Another polyimide layer is then deposited to sandwich the electrical components (Fig. 3a). The meshes are then completed by dry etching in an oxygen plasma. The sacrificial PMMA layer is then stripped in acetone, and the electrode meshes transferred onto a carrier substrate through several steps of transfer printing using a PDMS stamp and thermal release tape (Fig. 3b). After connecting with a cable for wiring out the bus connections, the electronic meshes, after being transferred onto a glass substrate (Fig. 3c), are incorporated within the hydrogel. To do so, a prepared monomer solution is poured into a container where the electronic meshes are then floated over it (Fig. 3d). The hydrogel prepolymer is then photopolymerized using a mask to align the electrical array with the polymer shape's design. In order to sandwich the electronic mesh into the receiving hydrogel, a second hydrogel prepolymer solution is then added and polymerized in registry through the same mask. The polymerized hydrogel with embedded array is then immersed into water to remove residual monomers, solvents, and initiator.

Similar to the previous bimorph design, we used photolithography to pattern the gel with a gradient crosslink motif. The top layer in this case is poly acrylamide (PAAm) and the bottom layer poly acrylic acid (PAA). In 1mL of reactants, the pre-gel solution of PAAm contains acrylamide (monomer, 175 mg, 2.4mM), **N,N'**-methylenebis acrylamide(crosslinker, 20 mg, 0.1 mM) and Darocur 1173(photoinitiator, 30

 μ L,1.8x10⁻⁴ mM). The PAA layer of the bimorph incorporates acrylic acid (monomer, 175 mg, 2.4mM), N, N'-methylenebis acrylamide (crosslinker, 20 mg, 0.03mM), methacryloxyethyl thiocarbamoyl rhodamine B (photo curable dye, 1mg, 1.5x10⁻³mM), and Darocur 1173 (photoinitiator, 30 μ L, 1.8x10⁻⁴ mM) in 1 mL pre-gel solution. We used a lesser amount of crosslinkerer in this PAA gel is to maximize the swelling difference of the two layers, The pink colored rhodamine dye in PAA serves to differentiate the two layers (Fig. 3e).

The electrodes, once integrated in the bimorph gel, are immersed at a 0.1M KCl solution with pH=4, in order to achieve maximum strain change (the pKa of acrylic acid is 4.35). The electrochemical ORR is performed by applying a -1.1 V potential to the WE vs. a Ag pseudo reference. As the ORR proceeds, the WE performs the reaction:

$$O_2 + 4H^+ + 4e^- = 2H_2O$$
 (eq.2)

This partial reaction locally increases the pH value in the vicinity of the WE by consuming protons, leading the PAA layer to swell and induce curvature (Fig. 3f). We are working at these early stages of this project to enhance both the magnitude and reversibility of the pH responsiveness of the hydrogel strain mediated through the ORR. We believe this type of hybrid design—one embedding an active device level—provides the most explicit means for constructing efficient phase change based actuators.







Figure 2: Front view and top view of the side by side Ru-PAA/PHEMA gel: (a) original particle; (b) the particle went through water-splitting reaction; (c) the recovered particle. (d) The distance in between the ends of the particle and the edge of the petri-dish.



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3. NORTHWESTERN UNIVERSITY (GRZYBOWSKI GROUP)

1. Programmable Self-Assembling Systems: Directed assembly of colloids into crystals.

Assembly of colloidal building blocks into complicated structures is of fundamental and technological importance for their applications in photonics¹ and devices². External fields are also frequently used for directing colloids not only for studying the equilibrium behavior³ but also for drug delivery⁴ and cancer therapy⁵. Here we employ precisely controlled, to a micrometer scale, external magnetic fields to direct assembly of nonmagnetic and magnetic colloidal systems into colloidal crystals or core-shell assemblies. It is known that non-magnetic particles dispersed in a ferrofluid can be directed by magnetic field gradients since the pioneering work of Skjeltrop⁶. There is few reports since the work of Skjeltorp, we use nonmagnetic and magnetic particles or a mixture of both in a ferrofluid made from an aqueous solution of holmiumnitrate (which has no durability issues) to control the assembly of these colloids on an alternating magnetic field surface.

We used colloidal particles with different sizes, dispersed in refractive index matching DMSO-water or glycerol-water mixtures. Refractive index matching of the particles was necessary for the imaging purposes but more for the stability of charge-stabilized particles due to the added holmium salt, which decreases the screening length and causes particle aggregation. Refractive index matching enabled us to increase the salt concentration up to 1M without any aggregation of particles. Holmium nitrate concentration was kept between 0.2M-0.4M. This causes an optimum magnetophoretic force for micron-sized spheres.

Colloids dispersed in holmium nitrate solution were let to sediment on top of a nickel grid or pattern, which was placed on top of a magnet as depicted in Figure 1a. Nickel is a ferromagnetic metal and is polarized upon exposure to external magnetic field. When a nickel grid is placed on top of a magnet the metal bars will be polarized and it will form a magnetic grid above the nickel grid. The magnetic field flux over a nickel grid is shown in Fig 1h. As shown also in Fig 1i, numerical calculations of the magnetic field on a nickel grid, the magnetic field fluctuates along the grid holes and bars of the grid. When non-magnetic colloids, dispersed in a paramagnetic fluid, are exposed to such magnetic field pattern, particles will prefer to go towards the low field, as the paramagnetic fluid would be attracted towards the higher fields. The magnetophoretic drive of the colloids to the lower field was observed for 1.2 μ m particles on a grid with 7 μ m holes, seen in Fig 1b. We also managed to do this assembly on a single particle level by sedimenting 50 μ m particles on square and hexagonal grids with 43 μ m holes, seen in Fig. 1e and Fig. 1g, respectively. Fig 1c, 1d and 1f show other types of assembly where 1.2 μ m particles assemble on 1.4 μ m wells in 1c, on 3 μ m wells in 1d and on 5-fold quasicrystalline 2 μ m wells in 1f.



Fig 1. (a) Side view of the sample cell construction. Single type of colloids directed towards the holes of different types of TEM grids, **(b)** 1.2 μ m colloids assembled on the holes of a TEM grid with a 7 μ m hole size, **(c)** 1.2 μ m colloids assembled on the holes of a lithographically made nickel grid with holes of 1.4 μ m, **(c)** 1.2 μ m colloids assembled on the holes of a TEM grid with a 7 μ m hole size, **(d)** 800 nm silica particles assembled in 3.0 μ m sized magnetic wells, where only 7 particles fit into the well, **(e)** 50 μ m colloids assembled on the holes of a square TEM grid with 43 μ m hole size, **(f)** 1.2 μ m colloids assembled on 5-fold quasicrystalline ordered wells, **(g)** 50 μ m colloids assembled on the holes of a hexagonal TEM grid with 43 μ m hole size, **(h)** Numerical calculation of the magnetic flux density over a Nickel grid, placed on top of a magnet, **(i)** The magnetic flux density on the lines (red and green) depicted at (h) along the holes of the grid (green) and along the intersections of the grid (red).

When a mixture of non-magnetic colloids with different sizes was sedimented on nickel grids, we observed segregation of these particles in a core-shell manner. Magnetophoretic forces depend on the volume of the spheres and for a larger sphere the force is larger to the r^3 , where r is the radius of the colloid. This difference in the applying forces plays the role to push the larger 1.2 µm particles to the holes first and the smaller 800 nm ones afterwards, which causes the formation of the core-shell construction seen in Fig 2a-c. Fig 2a shows the confocal microscopy image of the 800 nm and Fig 2a shows the image of the 1.2 µm particles that form the core of the assembly. Fig2c is an overlay of the images given in Fig 2a and Fig 2b that clearly shows the core shell structure. Fig 2d is a constructed image from the same z-stack of the images shown in Fig2a-c and it shows the side view of the particles in z-axis. We made the same experiment with 5 µm and 1.2 µm particles` mixture on a grid with 7 µm holes and the result was formation of raspberry like structures due to the size segregation, as shown in Fig. 2e.



Fig 2. Binary mixtures of colloidal particles driven towards the low magnetic field regions with a phase separation due to size difference, (**a**) fluorescence image of RITC labeled 800 nm silica particles, which form a shell around the 1.2 μ m FITC labeled silica colloids shown in (b). (**b**) Fluorescence image of FITC labeled particles with a size of 1.2 μ m, these particles are driven to the core of the assembly due to their larger size. (**c**) Overlay of the images shown in (a) and (b), which shows clearly the core-shell structure, (**d**) side view of the z-stack along a line shown in (c) (Scale bar 40 μ m). (**e**) Raspberry-like assembly of 5 μ m silica particles labeled with RITC, which are surrounded by 1.2 μ m FITC labeled particles (Scale bar 10 μ m).

With the fact of an underlying magnetic grid it is expected that a mixture of magnetic and non-magnetic particles will phase separate. When we sedimented a mixture of magnetic and non-magnetic particles of about 1μ m size on a grid with 43 μ m holes, particles phase separated on the grid bars and the holes. Magnetic particles were driven towards the grid bars with a higher concentration on the crosses of the bars, whereas the non-magnetic particles filled the holes of the grids, as seen in Fig. 3a. It should be noted that the magnetic field is stronger on the crossings of the bars as can also be seen in Fig. 1f and this causes the magnetic particles to concentrate more on these spots.



Fig 3. Confocal microscopy images of binary mixtures of colloids with different magnetization (magnetic susceptibility), **(a)** non-magnetic 900nm silica particles labeled with RITC (red) and 800 nm magnetic silica particles labeled with FITC (green), that are phase separated on a nickel grid with 43 μ m holes, **(b)** non-magnetic particles of 5 μ m size labeled with FITC(green), and 5.8 μ m magnetic particles labeled with Alexa Fluor 660 (red), **(c)** and **(d)** 1 μ m sized non-magnetic (green), and magnetic (red) particles on 1.4 μ m pitch sized grids.

Magnetic and non-magnetic particles with 5 μ m and 5.8 μ m sizes, respectively, formed a 2D binary crystal when sedimented on a grid with 7 μ m holes and 12 μ m pitch size. Here again the magnetic particles mostly preferred the intersection of the bars which enabled the formation of the binary square lattice shown in Fig. 3b. Fig. 3c and 3d shows 1 μ m sized non-magnetic (green), and magnetic (red) particles on 1.4 μ m pitch sized grids, these particles assemble into AB crystals in Fig 3c whereas they form an AB₂ type crystal in Fig 3d. Formation of these ordered assemblies are highly coordinated with the underlying nickel grid which gives the opportunity to design diverse structures.

We also investigated the behavior of magnetic and non-magnetic particle mixtures with different sizes. We used 1.2 μ m non-magnetic silica particles and 5.8 μ m magnetic particles. In a uniform magnetic field these particles were oppositely polarized due to the magnetic susceptibility difference between the

particles and the solvent⁷. For nonmagnetic particles and magnetic particles dispersed in a paramagnetic fluid the induced dipoles are in opposite directions and this causes an attraction between the particles. Here due to the size ratio smaller particles cover the big one and form raspberry like structures, shown in Fig 4a. These raspberries were similar to the ones shown in Fig 2e, but the underlying physics was totally different. However, this inverse-dipolar attraction between the magnetic and non-magnetic particles was overcome when particles were on a gradient of magnetic field as on a grid shown in Fig 4b. The grid with the 7 μ m hole sizes broke the attraction between the particles and directed the non- magnetic particles to the holes and the magnetic particles to the grid bars. Interplay of the two different phenomena could be used in a ternary system to make raspberry structures both in a gradient and in homogenous field. Ternary system composed of magnetic 5.8 μ m particles and non-magnetic 5 μ m and 1.2 μ m particles. In a homogenous field magnetic particles were mostly left out of the attraction shell. When the ternary particles system was brought on the grid the magnetic particles were driven on the grid bars and the nonmagnetic large and small particles formed raspberry structures in the grid holes, both raspberry structures in homogeneous and alternating field strengths were shown in Fig 4c and d, respectively.

In summary, we have shown different assemblies of nonmagnetic and magnetic particles at different length scales between 1-50 μ m by using magnetic field gradients for directing both magnetic and non-magnetic particles. We would like to point out that our results could be of high interest for potential applications in optical devices or in delivery systems.



Fig 4. Confocal microscopy images of binary mixtures of binary (a,b) and ternary (c,d) colloids with different magnetization (magnetic susceptibility) and different size, **(a)** non-magnetic 1.2 μ m silica particles labeled with FITC (green) and 5.8 μ m magnetic particles labeled with Alexa Fluor 660 (red), that are in an homogenous magnetic field, **(b)** same particle system separated on a nickel grid with 7 μ m hole sizes, **(c)** non-magnetic particles of 5 μ m size labeled with RITC (purple), 5.8 μ m magnetic particles labeled with Alexa Fluor 660 (red) and non-magnetic 1.2 μ m FITC labeled particles (green) in a homogenous magnetic field, **(d)** same particles system as in (c) separated on a nickel grid with 7 μ m hole sizes.

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