“The program has no set time limits. Research is a lifelong experience and we hope to remain a resource to our students long after ‘graduation’”

The Garcia Center for Polymers at Engineered Interfaces was founded in 1996 and is named after the late Queens College professor Narciso Garcia, who was a pioneer in the integration of education and research. The Center focuses on the integration of materials research with tissue engineering, biomaterials, drug delivery systems, sustainable energy, nanocomposites, and recently, additive manufacturing. The Center also supports innovation through entrepreneurship and has multiple collaborations with industry and national laboratories, both in the US and abroad. For information on the numerous programs that are available please see our website at: http://polymer.matscieng.stonybrook.edu

The research scholar program offers the opportunity for high school teachers, undergraduate, and high school students to perform research on the forefront of polymer science and technology together with the Garcia faculty and staff. Students work as part of focus research teams and are taught to make original contributions of interest to the greater scientific community. In addition to entering national high school STEM competitions, the students are encouraged to publish in refereed scientific journals, present their results at national and international conferences, and file for disclosures and patents to protect their intellectual property.

Our goal is to convey to the students the excitement we enjoy daily in research and provide for them a supportive network within the scientific community. Research is a lifelong experience and we hope to remain a resource to our students long after ”graduation”.

Miriam Rafailovich & Jonathan Sokolov August 8, 2018

Credits: Sahith Vadada and Kuan- Che Feng, AFM topographical image of PLA crystalline film spun cast on single crystal silicon.
High School Students

Auerbach, Yehoshua

Blatt, Ashley

Brahmbhatt, Isha

Chang, Benjamin

Chao, Daniel V

Chao, Elaine Y

Chen, Guan Hao (Michael)

Chen, John

Chen, Kevin

Chen, Thomas

Choi, Inyoung (Erica)

Chowdhury, Saajid

Deluxe, Antony J

Fontaine, Alice S

Gan, Alan Q

Goldschlag, Jonathan A

Hsu, Allison L

Hulyalkar, Miguel F

Iyer, Shreyas A

Jacobsen, Nicole B
High School Students

Jiang, Lan
Jindal, Varun
Kelly, Danielle M
Khan, Zaiff
Lam, Pik Hoi
Lee, Joon Young
Li, Mindy Z
Limjoco, Ian J
Lin, Lucia
Liu, Grace
Liu, Zipei
Lu, Evan C
Millstone, Eliana
Murthy, Bhuvna R
Nitta, Frederick
Oh, Brandon
Patel, Rushikesh
Phan, Byron M
Pisipati, Anish V
Popova, Anastasia N
High School Students

Prasad, Anisa V
Prasad, Sirina V
Puri, Isha
Qu, Matthew T
Radinsky, Avi
Rai-Gersappe, Diya
Rao, Kavya G
Sandhu, Bhawan K
Shine, Audrey
Schein, Tzipora C
Schneider, Gemma J
Shi, Zhining (Clara)
Shin-An, (Ethan) Ho
Singh, Anjali
Singh, Vedant
Sun, Aaron
Tian, Albert
Todt, Andrew D
Tong, Christina Y
Tran, Kimberley P
High School Students

Upadhyay, Aum D
Urquiola, Elena
Vadada, Sahith
Vilkas, Joshua S

Wang Xiaoxin
Winkler, Ethan S
Wu, Kenton
Wu, Steven

Xu, Kevin J
Xu, Vicki
Yin, Ruilin
Ying, Samantha

Zeng, Caroline
Zhang, Michael S
Zhang, Vincent
Zhao, Wenqi (Jennifer)

Zheng, Derek, N
Zhou, Anson Y
Zhou, Justin
Zhu, Albert Y
Research Experience for Undergraduates

Albert, Deena B
Chan, Kenneth K
Gao, Ryan
Hofflich, Jessica

Jenkins, Delivrance
Labiak, Kaitlin C
Lo, Liana
Mora, Anyerlin M

Moulton, Matthew
Orduna, Cecilia
Passariello, Diana
Shata, Ahmed

Sutaria, Jainil
Waqr, Kasim M
York, Matthew J
Zaidi, Mustafa

Zumba, Nicholas
Ossip, Jonathan
Graduate Students

Budassi, Julia
Chuang, Ya-chen
Clayton, Nicholas
Feng, Kuan-che
Li, Juyi
Li, Kao
Ricotta, Vincent
Shmueli, Yuval
Wang, Likun
Xue, Yuan
Yang, Fan
Yin, Yifan
Zhang, Linxi
Zhou, Yuchen
Zuo, Xianghao
Faculty Mentors

Davis, Raphael
Fiore, Susan
Galanakis, Dennis
Gersappe, Dilip
Jerome, John Luckner
Tuminello, Donna
Pinkas-Sarfova, Adriana
Phillips, Reed
Simon, Marcia
Walker, Stephen
Bliznakov, Stoyan
Tae Jin Kim
Cuiffo, Mike
Grella, Rebecca
Isseroff, Rebecca
Weiss, Herb
It’s Lit

Yearbook Picture Credits: Sahith Vadada
Lectures and Presentations

Yearbook Picture Credits: Sahith Vadada
In the Lab!

Yearbook Picture Credits: Sahith Vadada
ADRIANA STUCK IN ELEVATOR!

Yearbook Picture Credits: Sahith Vadada
## Garcia: Polymers at Engineered Interfaces-2018
### Summer Scholar Program Schedule of Activities

_EVERY DAY STARTS WITH A GROUP MEETING_

**CHECK SCHEDULE DAILY!**

<table>
<thead>
<tr>
<th>Week of 6/25</th>
<th>MONDAY</th>
<th>TUESDAY</th>
<th>WEDNESDAY</th>
<th>THURSDAY</th>
<th>FRIDAY</th>
</tr>
</thead>
<tbody>
<tr>
<td>6/25</td>
<td>10:00am - 10:15 Judith Berhannan Dean of Admissions <strong>Welcome to Stony Brook</strong></td>
<td>10:00am</td>
<td>10:00am - 10:15 General meeting</td>
<td>10:00am - 10:15 General meeting</td>
<td>10:00am General meeting</td>
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<tr>
<td></td>
<td>10:15-10:50 Intro to Garcia - Miriam Rafailovich Grad and REU introduction</td>
<td>10:15 am - 11:00 am Michael Hadjijargyrou - NYIT</td>
<td>10:15-12:45 Mandatory Lab Safety Training w/ EH&amp;S</td>
<td>10:30 am-12:30 pm Library Resources and Intro to Excel Courses (2 Groups) (Please have your laptop with you)</td>
<td>10:00 am General meeting</td>
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<td>11:00 – 11:30 pm Robert Harrison Theoretical Modelling</td>
<td>11:30 - 12:00 Vincent Ricotta, PhD Engineering Biosensors</td>
<td>12:45pm-1:45pm Lunch</td>
<td>12:30 pm-1:15pm Lunch</td>
<td>10:05 am-11:05 am Lab Safety and Facilities Quiz</td>
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<td></td>
<td>12:15pm-1:15pm Lunch</td>
<td>1:15pm-4:00pm Facilities Tour: AFM - Ya-Cen Chung FTIR - Harry He LB trough - Yuchen Zhou 3D printer - Yuval Shmueli Cell Lab - Kuan-Che Feng Rheology - Jui Li Electrospraying - Kao Li, Fuel cells - Likun Wang Hands on safety training specific to Garcia labs - Linxi Tensile - Xianghao Zuo Brands ender - Yuan Xue Spinner/ Oven Ellipsometer - Jonathan Sokolov Confocal microscope - Julia UV vis and DLS - Fan Yang</td>
<td>1:45 pm-2:30 pm Richard Clark Dermatology, SBU</td>
<td>1:15 pm-2:30 pm Steffen Mueller, Ph.D. President &amp; CSO Codagenix Inc.</td>
<td>11:05-11:50 am Gary Halada Nanoscale characterization SBU</td>
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<td>1:30pm-2:15pm Madeline Augustine, BOEING</td>
<td>2:30pm–3:15 pm Steffen Mueller, Ph.D.</td>
<td>2:30-2:50 PM Dr. Linxi Zhang Engineering scaffolds for mineralization</td>
<td>2:00–2:30 pm Dr. Reed Philips Energy Systems</td>
<td>PIZZA (Hunkis)</td>
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<td>2:15pm-4:00pm University Tour</td>
<td>2:30-4:00pm Ms. Rebecca Isseroff, Lawrence HS Science Research Coordinator (i) Magic of Graphene (ii) Keeping a Laboratory Notebook (iii)box presentation</td>
<td>2:30-2:50 PM Dr. Reed Philips Energy Systems</td>
<td>1:00–2:00 pm Jonathan Sokolov DNA Research at SBU</td>
<td>12:00- 12:30 Herb Weiss ISEF Ethics regulations</td>
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<td>11:00am-1:30pm ID cards at SAC and lunch at East Side Dining</td>
<td>3:15-4:00 pm Sherif Abdelaziz Civil Engineering, SBU</td>
<td>2:00-2:30 pm Dr. Linxi Zhang Engineering scaffolds for mineralization</td>
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<td>12:30- 1:00 pm Jainil Sutaria Hybrid hydrogels</td>
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<td></td>
<td>1:30pm-2:15pm Madeline Augustine, BOEING</td>
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<td></td>
<td></td>
<td>1:00-2:00 pm Jonathan Sokolov DNA Research at SBU</td>
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<td></td>
<td>2:15pm-4:00pm University Tour</td>
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<td></td>
<td></td>
<td>1:00-2:00 pm Jonathan Sokolov DNA Research at SBU</td>
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<td>2 groups - bus tour to Dental school, Marine science bldg, and Energy center</td>
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<td></td>
<td></td>
<td>1:00-2:00 pm Jonathan Sokolov DNA Research at SBU</td>
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*Week of 6/25*
<table>
<thead>
<tr>
<th>Date</th>
<th>Time</th>
<th>Event</th>
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<tbody>
<tr>
<td>7/2</td>
<td>10:00-10:15 am</td>
<td>General Meeting</td>
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<td>10:15 -11:00 am</td>
<td>Frederick E. Grine SBU</td>
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<td>11:00-11:45 am</td>
<td>Dilip Gersappe SBU Theory and Modeling</td>
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<td>11:45 am – 12:00 pm</td>
<td>Yuval Shmueli: FDM printing</td>
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<td>12:00-12:15</td>
<td>Yuchen Zhou: Perovskite Solar Cells</td>
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<tr>
<td>7/3</td>
<td>10:00 am</td>
<td>General Meeting</td>
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<td></td>
<td>10:30 am-1:00 pm</td>
<td>Divide into groups for Journal Club Student Presentations</td>
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<tr>
<td>7/4</td>
<td>10:00 am</td>
<td>Happy 4th of July</td>
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<td>10:00 am</td>
<td>Proper Laboratory Attire</td>
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<tr>
<td>7/5</td>
<td>10:00 am</td>
<td>Data Analysis and Statistics Part II Dr. Miriam Rafailovich</td>
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<td>Distribution of boxes</td>
<td>10:30 -4:00</td>
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<td></td>
<td>Group Experiment: Polymer Thin Film Processing:</td>
<td>Recycling, Rheology, Determination of Mw Si Single Xtals Lab Data Analysis</td>
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<td>Dr. Jonathan Sokolov The ellipsometer: coherent and incoherent light--a hands on experience</td>
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<tr>
<td>7/6</td>
<td>10:00 am</td>
<td>General Meeting</td>
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<td>Proper Laboratory Attire</td>
<td>10:00am</td>
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<tr>
<td></td>
<td>Data Analysis and Statistics Part II</td>
<td>Dr. Miriam Rafailovich</td>
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<tr>
<td></td>
<td>Group Experiment Cont:</td>
<td>Polymer Thin Film Processing:</td>
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<tr>
<td></td>
<td>Dr. Jonathan Sokolov The ellipsometer: coherent and incoherent light--a hands on experience</td>
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<td></td>
<td>Preparation of group ppt and lab reports. General meeting</td>
<td>3:00pm-4:00pm</td>
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<td>11:30am</td>
<td>Pizza Lunch</td>
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</tbody>
</table>

### Week of 7/2

**HOMEWORK for 7/2**

- Prepare ppt for journal club.
  - a. Choose a research paper of interest to you.
  - b. Prepare a 5 minute Power Point presentation summarizing the paper.
    - Slides:
      1. your name, school, title of paper, reference
      2. Hypothesis
      3. Materials/Method
      4. Results and Discussion
      5. Conclusion and evaluation

---

**Monday, 7/2**

- 10:00-10:15 am General Meeting
- 10:15 -11:00 am Frederick E. Grine SBU
- 11:00-11:45 am Dilip Gersappe SBU Theory and Modeling
- 11:45 am – 12:00 pm Yuval Shmueli: FDM printing
- 12:00-12:15 Yuchen Zhou: Perovskite Solar Cells

**Tuesday, 7/3**

- 10:00 am General Meeting
- 10:30 am-1:00 pm Divide into groups for Journal Club Student Presentations
- 1:00-2:00 Lunch (Lab safety Quiz make up exam in room 173 Light Engineering)
- 2:00-2:30: Sherif Abdelaziz Biodegradable Polymers in civil engineering

**Thursday, 7/5**

- 10:00 am General Meeting Dr. Miriam Rafailovich
- 10:30 am-2:00pm Group Experiment Cont: Polymer Thin Film Processing:
  - Recycling
  - Rheology
  - Determination of Mw Si Single Xtals Lab Data Analysis
- 3:00pm-4:00pm Preparation of group ppt and lab reports. General meeting
- 11:30am Pizza Lunch

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**Friday, 7/6**

- 10:00am General Meeting Dr. Miriam Rafailovich
- 10:30 am-4:00 Polymer Thin Film Processing:
  - Recycling
  - Rheology
  - Determination of Mw Si Single Xts Lab Data Analysis
- Dr. Jonathan Sokolov The ellipsometer: coherent and incoherent light--a hands on experience

---

**Remember**

- Proper laboratory dress on Thursday.
<table>
<thead>
<tr>
<th>Week of `7/9</th>
<th>7/9</th>
<th>7/10</th>
<th>7/11</th>
<th>7/12</th>
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<tr>
<td>10:00am</td>
<td>10:00am</td>
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<td>10:00-10:10 am: General meeting</td>
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<td>General meeting</td>
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<td>General meeting</td>
<td>10:10-11:00 Student talk I: Cellulose hydrogen fuel cell group</td>
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<td>10:15 am-11:00 am</td>
<td>10:15-11:00 am</td>
<td>10:15 am</td>
<td>10:30am</td>
<td>10:00-10:10 am: General meeting</td>
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<tr>
<td>Gordon Taylor: RAMAN Spectroscopy</td>
<td>SOP Writing Ying Liu (EHS)</td>
<td>WORK!</td>
<td>WORK!</td>
<td>WORK!</td>
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<tr>
<td>11:00 am-11:45 am</td>
<td>11:00-11:15 pm</td>
<td>11:15-11:30 pm</td>
<td>11:30-12:30 pm</td>
<td>12:30pm: Lunch 1:30pm WORK!</td>
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<tr>
<td>Steve Walker: Microbiology</td>
<td>Harry Shan He: Electrosyn fibers for trapping chemical free trapping of insects</td>
<td>Lev Neymotin Laser Lithography</td>
<td>Selection of Projects II</td>
<td>Selection of Projects III</td>
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<tr>
<td>11:45-12:30 pm Lunch</td>
<td>Lunch</td>
<td>Lunch</td>
<td>1:30-2:30 pm Lunch</td>
<td>2:30-4:00 pm Writing and submission of SOPs with student signatures</td>
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<tr>
<td>12:30-2:00 pm Student presentation on Spin Casting Lab</td>
<td>12:30-2:00 pm Graduate student presentations-- Yuan Xu/ Xianghao composites Juyi--hydrogels Ya-Chen--DPSC nanoparticles</td>
<td>Selection of Projects I</td>
<td>Selection of Projects II</td>
<td>Official date for ISEF/Regeneron project paper work</td>
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<tr>
<td>2:15 pm-3:00 pm</td>
<td>11:15-11:30 Lev Neymotin Laser Lithography</td>
<td>11:30-12:30 Selection of projects II</td>
<td>12:30pm-1:30 pm Lunch</td>
<td>2:00pm End of Day</td>
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<tr>
<td>Selection of Projects I</td>
<td>Selection of Projects II</td>
<td>Selection of Projects III</td>
<td>Selection of Projects III</td>
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</tbody>
</table>

**Homework**

Each student must submit a lab report by **Midnight on Monday Morning.**

Remember to give proper acknowledgements to your group partners, REUs, Grads.

**Proper Laboratory Attire**

- Week of 7/6
- Week of 7/7
- Week of 7/8
- Week of 7/9
- Week of 7/10
- Week of 7/11
- Week of 7/12
- Week of 7/13
<table>
<thead>
<tr>
<th>Date</th>
<th>Event Details</th>
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<tbody>
<tr>
<td>7/16</td>
<td>General meeting:</td>
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<tr>
<td></td>
<td>Science moment:</td>
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<tr>
<td></td>
<td>10:30am WORK!</td>
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<td></td>
<td>12:30pm: Lunch</td>
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<td></td>
<td>1:30pm WORK!</td>
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<td>Proper Laboratory Attire</td>
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<tr>
<td>7/23</td>
<td>10:00am Science moment: Dr. Rebecca Grella -- Plastics and the environment</td>
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<td>WORK!</td>
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<td>12:30pm: Lunch</td>
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<td>Proper Laboratory Attire</td>
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<tr>
<td>7/24</td>
<td>10:00am General meeting</td>
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<td>Trip to VJ Technology for 3-D PLA printing group</td>
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<td>12:30pm: Lunch</td>
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<td>Proper Laboratory Attire</td>
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<tr>
<td>7/25</td>
<td>10:00am General Meeting</td>
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<td>WORK!</td>
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<td></td>
<td>12:30pm: Lunch</td>
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<td>Proper Laboratory Attire</td>
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<tr>
<td>7/26</td>
<td>10:00am General Meeting</td>
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<td>Student talks Fibrinogen Nanocomposites (2 groups)</td>
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<td>Stem Cell Education Nanoparticles and stem cell differentiation</td>
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<td>Pizza Lunch</td>
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<td>2:00pm End of Day</td>
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<tr>
<td>7/27</td>
<td>10:00am General meeting</td>
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<td>Student talks Fibrinogen Nanocomposites (2 groups)</td>
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<td>Stem Cell Education Nanoparticles and stem cell differentiation</td>
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**Abstracts**

1. **One page**, (one inch margins, min. 11 pt font.)
2. One or more Figures with fig. captions
3. Two or more references from peer reviewed sources.
4. Due Sunday at **8:30 PM**

**Symposium:**

Please come at 9:30 for coffee and great music.

Program starts promptly at 10:00

Business formal attire please.
Annual Summer Scholar
Research Symposium & Musicale
August 8th, 2018
10:00am — 2:00pm
Student Activity Center
Ballroom A
Buffet Luncheon: Wing Wan of West Hempstead
Opening Speaker
Prof. Peter Brink
Swimming with sharks and
Learning about cancer
RSVP Adriana.Pinkas-Sarafova@stonybrookmedicine.edu
### SAC Ballroom A

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| 9:30 AM – 10:00 AM | Breakfast  
Musical Presentation  
Arranged by Professor John Luckner Jerome and Garcia Students |
| 10:00 AM        | Professor Peter Brink – *Swimming with Sharks and Learning About Cancer* |
| 10:10 AM – 10:20 AM | **Biosensors**  
Chairs: Deena Albert, Queens College of the City University of New York  
Kaitlin Labiak, University of Michigan |
|                 | *Microfluidic Delivery of Enzymes for Ordered DNA Fragmentation*  
Alan Gan, East Brunswick High School, East Brunswick, NJ |
|                 | *Developing Potentiometric Biosensors Based on Surface Molecular Imprinting Utilizing Surface Roughness for The Detection of Bio-Macromolecules*  
Yehoshua Auerbach, Lawrence High School, Cedarhurst, NY  
Miguel Hulyalkar, Southside High School, Rockville Center, NY  
Andrew Todt, Lake Oswego High School, Lake Oswego, OR |
|                 | *A Pilot Study of the Analysis of Facial Motion using Digital Image Spectroscopy Following Acoustic Neuroma Surgery*  
Ian Limjoco, Sachem East High School, Farmingville NY |
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<td>Avi Radinsky, Davis Renov Stahler Yeshiva High School For Boys, Woodmere, NY</td>
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<td><strong>Optimizing Gelatin-Based Hydrogels for Use in 3D Bioprinting of Human Dermal Fibroblasts</strong></td>
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<td>Elaine Chao, Northwood High School, Irvine, CA Eliana Millstone, Stella K Abraham High School for Girls, Hewlett, NY Kenton Wu, The Wheatlew School, Old Westbury, NY</td>
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<td><strong>Synthesizing and Characterizing Novel Gelatin and Poloxamer PF127, PF108, PF98 Hybrid Hydrogels</strong></td>
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<td>Erica Choi, St. Paul’s School, Concord, NH Christina Tong, Fairview High School, Boulder, CO</td>
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<td><strong>Analyzing the In Vitro Viability of Novel Gelatin-Pluronic ® F127 Hybrid Hydrogels as Cell Barrier Membranes</strong></td>
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<td>Kevin Chen, Mira Costa High School, Manhattan Beach, CA Joon Young Lee, Seoul International School, Seongnam City, South Korea Aaron Sun, Ed W Clark High School, Las Vegas, NV</td>
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<td><strong>Modeling the Diffusion of Drugs Through Hydrogels For Applications in Drug Delivery</strong></td>
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<td>Isha Puri, Horace Greeley High School, Chappaqua, NY Zhining Shi, The Hockaday School, Dallas, TX</td>
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<td><strong>Lattice Boltzmann Modeling of Fluid Flow Dynamics in Metal-Organic Frameworks</strong></td>
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<td>Shreyas Iyer, Munster High School, Munster, IN</td>
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<td>10:35 AM –</td>
<td>Cellular/Bacterial Interactions</td>
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<td><strong>Characterization of Antibiofilm Activity of Silver Nanoparticles Deposited in Situ on 3D Printed Polylactic Acid Implant Models</strong></td>
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<td>Anastasia Popova, Hackley School, Tarrytown, NY Isha Brahmbhatt, Ardsley High School, Ardsley, NY</td>
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<td><strong>Bactericidal Efficacy and Cytotoxic Effects of Newly Developed Calcium Hydroxide Antibacterial Compound (CASA)</strong></td>
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<td>Naturally Occuring Fibrinogen Chain Variant γ’ Associated with Decreased Risk for Staphylococcus Aureus Bacteremia</td>
<td>Anish Pisipati, Half Hollow Hills High School East, Dix Hills, NY</td>
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<td>Isaias Lipa, Brentwood High School, Brentwood, NY</td>
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<td>Anson Zhou, Patchogue-Medford High School, Medford, NY</td>
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<td>Investigating the Nanotoxicity of Titanium Dioxide via Fibroblast Growth and Infection with Staphylococcus Aureus</td>
<td>Justin Zhou, Patchogue-Medford High School, Medford, NY</td>
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<td>Vincent Zhang, Sachem High School East, Farmingville, NY</td>
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<td>Ethan Winkler, Hebrew Academy of the Five Towns and Rockaway, Cedarhurst, NY</td>
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<td>Jonathan Goldschlag, Hebrew Academy of the Five Towns and Rockaway, Cedarhurst, NY</td>
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10:50 AM – 11:10 AM  **Fuel Cells**  
Chairs: Matthew Moulton, Cooper Union, New York, NY  
Kenneth Chan, Boston University, Boston, MA

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<td>Coating Nafion 117 Membranes with Graphene Oxide using Various Surface Pressures to Increase Performance of Proton Exchange Membrane Fuel Cells</td>
<td>Lucia Lin, Townsend Harris High School, Flushing, NY</td>
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<td>Kimberley Tran, Clayton High School, St. Louis, MO</td>
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<td>Alteration of the Surface Structure of the Nafion Membrane to Increase PEMFC Efficiency</td>
<td>Evan Lu, Clear Lake High School, Houston, TX</td>
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<td>Brandon Oh, Seoul International School, Seoul, South Korea</td>
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<td>Matthew Qu, Wayzata High School, Plymouth, MN</td>
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<td>Michael Zhang, Saratoga High School, Saratoga, CA</td>
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<td>Varying Graphene Oxide Deposition Mass in Polymer Electrolyte Membrane Fuel Cells to Optimize Fuel Cell Performance and Durability</td>
<td>Saajid Chowdhury, MacArthur High School, Levittown, NY</td>
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<td>Kevin Xu, Ward Melville High School, East Setauket, NY</td>
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<td>Application of Electrospinning Deposition to Enhance Single and Triple MEA Performance in Polymer Electrolyte Membrane Fuel Cells</td>
<td>Zipei Liu, Shenzhen Middle School, Shenzhen, China</td>
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<td>Guan Hao Chen, St. George’s School, Vancouver, Canada</td>
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<td>Audrey Shine, Plainview Old-Bethpage JFK High School, Plainview, NY</td>
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<td>Danielle Kelly, Friends Academy, Locust Valley, NY</td>
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| 11:30 AM     | *Investigating the Effects of ALD-Coated Titanium Dioxide on Dental Pulp      | Nicholas Zumba, Columbia University, New York, NY  
|              | Stem Cell Response to 3D-Printed PLA Scaffolds**                             | Delivrance Jenkins, Suffolk Community College, Brentwood, NY  
|              | **Evaluating the Effects of Graphene-Loaded Polylactic Acid Electrospun      | Benjamin Chang, Woodbridge High School, Irvine, CA  
|              | Fibers and Spun-cast Thin Films on the Proliferation and Differentiation of  | Antony Deluxe, Wheatley High School, Old Westbury, NY  
|              | Dental Pulp Stem Cells**                                                    | Ethan Ho, Northfield Mount Hermon School, Gill, MA  
|              | **Investigating the Effects of Substrate Mechanical Patterns on Proliferation  | Bhuvna Murthy, Huron High School, Ann Arbor, MI  
|              | and Differentiation of Dental Pulp Stem Cells**                            | Rushi Patel, Herricks High School, New Hyde Park, NY  
|              | **The Effects of TiO\(_2\) Nanoparticles on the Proliferation and            | Wenqi Zhao, Milton Academy, Milton, MA  
|              | Differentiation of Dental Pulp Stem Cells**                                |                                                                                                   |
|              | **Utilization of P4VP, Graphene, and Polyornithine to Potentially             | Mindy Li, Princeton High School, Princeton, NJ  
|              | Differentiate Dental Pulp Stem Cells into Functional Neurons**              | Samantha Ying, South Side High School, Rockville Centre, NY  
|              | **Utilizing ALD-Coated Titanium Dioxide as a Gold Nanoparticle Catalyst      | Alice Fontaine, Thomas Jefferson High School for Science and Technology, Alexandria, VA  
|              | Support in Electrodes to Increase Efficiency Proton Exchange Membrane Fuel   | Allison Hsu, Syosset High School, Syosset, NY  
|              | Cells**                                                                      | Anjali Singh, Novi High School, Novi, MI  
|              | **Investigating the Effects of Substrate Mechanical Patterns on Proliferation  | Albert Zhu, Our Lady of Lourdes High School, Poughkeepsie, NY  
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<p>|              | <strong>The Effects of TiO(_2) Nanoparticles on the Proliferation and            |                                                                                                   |
|              | Differentiation of Dental Pulp Stem Cells</strong>                                |                                                                                                   |
|              | **Utilization of P4VP, Graphene, and Polyornithine to Potentially Differentiate|                                                                                                   |</p>
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<td>Ryan Gao</td>
<td>Pik Hoi Lam, Wilson Area High School, Easton, PA</td>
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<td>Nicole Jacobsen, Plainedge High School, Massapequa, NY</td>
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<td>Joshua Vilkas, Hebrew Academy of Nassau County High School, Uniondale, NY</td>
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<td>Nanocomposites</td>
<td>Mustafa Zaidi, Kasim Waqar</td>
<td>Ruilin Yin, St. Anthony's High School, South Huntington, NY</td>
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<td>Elena Urquiola, Hunter College High School, New York City, NY</td>
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<td>Frederick Nitta, Henry M. Gunn High School, Palo Alto, CA</td>
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<td>Vicki Xu, Mission San Jose High School, Fremont, CA</td>
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<td>11:50 AM - 12:00 PM</td>
<td>Perovskite Solar Cells</td>
<td>Cecilia Orduña, Liana Lo</td>
<td>Albert Tian, Ward Melville High School, East Setauket, NY</td>
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<td>Varun Jindal, Ward Melville High School, East Setauket, NY</td>
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<td>12:00PM-12:10PM</td>
<td><strong>3-D Printing</strong></td>
<td>Chairs: Matthew York, Case Western University, Cleveland, OH Ahmed Shata, Stony Brook University, Stony Brook, NY</td>
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<td><strong>Characterization of Electrical and Mechanical Properties of Thermoplastic Poly(Lactic Acid) Filaments Infused with Graphene Nanoplatelets through Fused Deposition Modeling Techniques</strong></td>
<td>Xiaoxin Wang, Padua Franciscan High School, Parma, OH Steven Wu, Clear Lake High School, Houston, TX Derek Zheng, Monta Vista High School, Cupertino, CA</td>
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<td><strong>Evaluating the Effects of Material Composition on Molded and 3D Printed Samples</strong></td>
<td>Vedant Singh, The Wheatley School, Old Westbury, NY Daniel Chao, Hunter College High School, New York, NY</td>
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<td><strong>Optimizing 3D Printing of PLA Filaments Through Lattice-Boltzmann Modeling and Graphene Nanocomposite Integration</strong></td>
<td>Lan Jiang, University High School, Irvine, CA Caroline Zeng, Wayzata High School, Plymouth, MN</td>
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**Acknowledgement:** Sponsored in part by the National Science Foundation and the Morin Foundation Trust.
Session 1: Biosensors

Chairs: Fan Yang & Julia Budassi
Next generation sequencing, with all its improvements over older and less efficient and cost-effective methods of genome sequencing, remains unoptimized due to the neglect of preserving the order of the fragmented DNA. This greatly complicates the reassembly process. A possible method of preserving order involves the cutting of linearized parallel strands of DNA adhered to and immobilized on a polymethyl methacrylate (PMMA) coated surface. The PMMA coated surface is lowered into a DNA solution and then raised slowly. The DNA is pulled upwards and stretched into a line as the meniscus follows the surface. A polydimethylsiloxane (PDMS) stamp with microfluidic channels running in a perpendicular direction to the strands of DNA is placed on the surface. Flowing a solution of an enzyme that cuts DNA (DNase I) through the channels will cleave the DNA wherever it is exposed to the channels while preserving the sequence order and therefore simplifying the sequencing process.

A problem with this method is that the DNase I enzyme is strongly attracted to the PDMS stamp used in the filling process, hindering its ability to travel the full length of the stamp. A coating to which DNase does not absorb must be applied to the inside of the channels first. Bovine serum albumin (BSA) in a solution can be used for this purpose. An experiment was conducted to test how far BSA would travel through the microfluidic channels when allowed to diffuse for variable amounts of time. The stamps that were created were cured from a mixture of PDMS and a curing solution at 60-70°C. They were poured onto a “master” pattern which is a silicon wafer etched with the negative replica of the desired microfluidic channels. After the PDMS is cured, it is cut into a stamp shape and a reservoir which holds the buffer, BSA solution, and DNase I solution is glued on with more liquid PDMS and left to cure. 150 µL of 1X buffer is pipetted into the reservoir and the entire stamp is left inside a lower pressure vacuum of approximately 115 torr for 15 minutes to promote filling of the channels. After returning to atmospheric pressure, the stamps are left on a 40°C hot plate with an additional 7.5 µL of BSA, which is added and mixed with a pipette.

For this experiment, PMMA was coated onto a cleaned #2 glass coverslip (thickness from 0.17 mm to 0.25 mm) so that microscopy could be used to examine the samples without removal of the stamp. A fluorescent BSA was used so that the distance the BSA traveled could be determined under a microscope. The samples were left on the heat for 6, 12, 24, 30, and 60 minutes. Each samples’ reservoir is drained of the buffer and BSA, and examined. The distance the BSA traveled was measured by finding the coordinates of the beginning of the channels (Fig. 1) and subtracting it from the coordinates of where most of the channels end (Fig. 2). Additional testing includes adding more data points to increase accuracy as well as varying channel size to determine its effect on the diffusion rate of BSA.

Developing Potentiometric Biosensors Based on Surface Molecular Imprinting
Utilizing Surface Roughness for The Detection of Bio-macromolecules

Yehoshua Auerbach¹, Nick Clayton², Miguel Hulyalkar³, Dr. Miriam Rafailovich²,
Dr. Vincent Ricotta², Andrew Todt⁴
¹Lawrence High School, Cedarhurst, NY; ²Stony Brook University, Stony Brook, NY;
³South Side High School, Rockville Center, NY; ⁴Lake Oswego High School, Lake Oswego, OR

The development of fast, inexpensive biosensors holds significance in both medical and biochemical fields of study and industry as they are non-invasive and accurate tools to detect bio-macromolecules. While it is possible to detect flaviviruses like Zika or Dengue virus via antibody- or RNA-based methods, biosensors utilize surface molecular imprinting to differentiate bio-macromolecules of similar size and shape with high selectivity and sensitivity. These biosensors are extremely versatile and can be tailored to identify a large range of molecules, which can lend itself for use in analyses of processes including stem cell differentiation.

A scanning electron microscope (SEM) was used to analyze the surface roughness of both polished (smooth) and unpolished (rough) Ti/Au-coated Si chips (Fig.1). Biosensor chips were fabricated using both smooth and rough Si wafers coated with 3 nm Ti and 50 nm Au, then diced into 8 × 20 mm chips. The chips were masked with adhesive PTFE tape hole punched with 6 mm holes and were imprinted in 19/1 (v/v) of known concentrations of analytes in 1X dPBS/100 uM 11-mercapto-1-undecanol in dimethyl sulfoxide. Analytes were imprinting at the following concentrations: EhV-163 with a final imprint concentration of 10² P/mL; S17 fibrin-depleted fibrinogen at 3 µg/mL and 30 µg/mL; 50P fibrin-rich fibrinogen at 2.3 µg/mL and 23 µg/mL, then washed with 3M NaCl overnight. Re-adsorption of the target analytes to the washed chips were measured in real-time potentiometrically; after 20 minutes of stabilization in 1X dPBS, serial dilutions of target analyte was added every 5 minutes in 100 µL aliquots, successively increasing in concentration 10-fold. Various tests of EhV-163 were performed, including testing washing methods using ethanol, DI H₂O, and Tween-20. The smooth imprints yielded no voltage response, as expected as its vertical roughness of about 3.5 nm (1), and cavity length in the range of 6 nm (1), is not in the size range of the ~150 nm EhV-163 (2). The rough EhV-163 imprints gave highly erratic responses, leading us to test instead with fibrinogen. Because of the low imprint concentration, these sensors were not very sensitive and only had a notable drop in open circuit potential (OCP) after additions of 230 µg/mL for 50P (Fig. 2a) and 300 µg/mL for S17 (Fig. 2b). However, the S17 imprints gave a larger average final addition response of -25.7 mV, compared to the 50P’s -11.8 mV average. Attempting to increase the sensitivity of the fibrinogen biosensors, they were imprinted again at 10-fold lower concentration. This did not noticeably affect the sensitivity of the 50P sensors, only yielding an average of -4.1 mV final response (Fig. 2a), but we hypothesize that the low-concentration S17 sensors will yield better results, as they lack the disordered clusters of soluble fibrin which are present in 50P solutions. Our future work will focus on using imprinted biosensors to detect stem cell differentiation through detecting the presence of alkaline phosphatase and osteocalcin in growth medium, as well as on completing low-concentration fibrinogen testing.

A Pilot Study of the Analysis of Facial Motion using Digital Image Spectroscopy following Acoustic Neuroma Surgery

Deena Albert1  Ian Limjoco2  Diya Rai-Gersappe3  Avi Radinsky4  Fan Yang5  Miriam Rafailovich6
1 CUNY Queens College Flushing NY; 2 Sachem East High School, Farmingville NY; 3 Huntington High School, Huntington, NY; 4 Davis Renov Stahler Yeshiva High School For Boys, Woodmere, NY; 5 Department of Material Science & Engineering, Stony Brook University, Stony Brook, NY; 6 Department of Materials Science & Engineering, Stony Brook University, Stony Brook, NY

Vestibular schwannoma (VS), also known as acoustic neuroma, is a benign brain tumor that develops along the cranial nerves running from the inner ear to the brain, affecting the facial nerve (cranial nerve VII) and the vestibulocochlear nerve (cranial nerve VIII). While the most common diagnostic methods of VS include MRI or CT scan, these methods prove to be expensive and time-consuming. The purpose of this study is to determine the extent to which VS influences facial nerve function by using digital photography. Digital Speckle Correlation (DISC) is an imaging technique that calculates facial motion from two pictures taken in rapid succession. In contrast to MRI and CT scans that analyze the internal nerve network of the head through electromagnetic frequencies, DISC inspects only the external skin structure, providing insight to the progression of the VS tumor via optical examination of facial morphology. DISC can furthermore be applied to patients that undertook a surgical resection, a gamma knife procedure, or no operation to compare the effects of each treatment method on VS progression and facial nerve functionality. In order to compare the facial structure of the participants, a brief medical history, list of medications currently being taken, and an abbreviated cranial nerve exam were performed and recorded from the patients who had VS (n=35) and the control group, who had no cranial nerve deficits (n=36).

For each participant, two pictures were taken: one expressionless and one slightly smiling. When comparing DISC-analyzed photographs of control participants to those of patients with VS, the results indicated that patients with VS had asymmetrical facial musculature, whereas control patients featured more symmetrical facial muscular structures. As seen in Figure 1.a, the patient with VS displays much higher muscle movement on the left side of their face whereas the control subject in Figure 1.b portrays a more equal amount of facial movement on both hemispheres of the face. As shown in Figure 2, we can conclude that DISC can successfully correlate severe VS progression to a detectable level of asymmetry that may not necessarily be observable to the human eye. Future applications of this study may be tested on the recovery speed of brain surgery patients via analysis of facial nerve repair changes and stroke victim aid in damage assessment.

Session 2: Hydrogels

Chairs: Juyi Li & Hongyu Li
Hydrogels are networks of polymers that are swollen with water and can be composed of natural or synthetic polymer chains. These biomaterials have several applications in tissue engineering and regenerative medicine as they are very biocompatible. Additionally, hydrogels can be crosslinked to create an entangled polymeric network that can be used to encapsulate cells and other compounds. Hydrogels can be crosslinked ionically through interactions between ions and polymer chains, enzymatically through the creation of covalent bonds, or through photopolymerization from exposure to UV radiation. The ease of modification and biocompatibility of hydrogels makes it an especially useful material in bioprinting.

Bioprinting is a revolutionary new 3D biofabrication technique used to precisely dispense biomaterials, including hydrogels and cell-laden mixtures. This study utilized a modified version of the Inkredible+ extrusion-based bioprinter, which prints using air pressure and has both heating and cooling capabilities for the cartridges, allowing for the printing of a wide array of materials. Specific advantages of extrusion-based bioprinters include high deposition and printing speed and its ability to print porous structures. In this study, gelatin, Pluronic® F127, polymerized F127, alginate, crosslinked gelatin, and gelatin-F127 hydrogels were synthesized to optimize mechanical properties for bioprinting.

For the crosslinked gelatin and hybrid gelatin-F127 hydrogels, several different ratios of gelatin to microbial transglutaminase (mTG) and gelatin to F127 were synthesized. The printing process for each gel was optimized by finding the extrusion pressure, temperature, and speed that allowed for the extrusion of a single, continuous line. The crosslinked gelatin and hybrid gelatin-F127 hydrogels were also optimized to find the ratio that would most consistently and effectively print a single, continuous line. The optimized printing conditions were then used to attempt to print a 3D cylindrical structure. The 3D bioprinted designs were created in AutoCAD Fusion 360 and processed in Cellink Heartware to create gcode files for printing.

Rheological analysis was performed on the crosslinked gelatin hydrogels to determine the change in elastic modulus over time. This data was used to determine the best possible crosslinked gelatin hydrogel combination for use in bioprinting and determine ideal times for gelatin-mTG crosslinking before printing.

To analyze the effects of bioprinting on cell viability, sterile gels were mixed with human dermal fibroblast cells and printed at each gel’s optimal printing condition [see figure 1]. The printed gels were then analyzed with fluorescence microscopy to determine cell viability during the printing process, which may be affected by sheer pressure, print speed, and nozzle gauge.

Future research will involve further analysis of embedded cells and culturing of cells from bioprinted gels. Additional tests need to be done to print mechanically stable 3D structure.

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**Figure 1.** Fluorescent microscopy was performed after 20% Pluronic F127 was mixed with human dermal fibroblasts and shows significant cell viability after being printed at optimal pressure, temperature, and speed.

**Figure 2.** Successful blood vessel print test run was performed on 30% Pluronic F127 at optimal speed, temperature, and pressure.

**Figure 3.** Inkredible+ extrusion based bioprinter with heating and cooling capabilities.

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Hydrogels are three-dimensional cross-linked polymeric networks. They have been used largely for regenerative medicine and still continue to show very promising applications in the biomedical field due to its biocompatibility and unique properties. Gelatin, a protein derived from collagen in animal tissue, is widely used in the creation of hydrogels; it has excellent biocompatibility, non-immunogenicity, biodegradability, and is relatively low cost. However, gelatin possesses poor mechanical strength and instability, especially under physiological conditions. Cross-linking gelatin with polymers has been shown to increase these properties. Poloxamers, tri-block copolymers composed of PEO$_x$-PPO$_y$-PEO$_z$ chains, are being explored for their biomedical applications due to their unique thermo-gelling properties, biocompatibility, and self-assembling micelle structures. This study had two major goals: synthesize and characterize hybrid gels using poloxamers PF127, PF108, and PF98 in order to evaluate their potency and characterize these new poloxamers.

Hybrid hydrogels were synthesized by mixing 15% w/v gelatin, 20% w/v poloxamer solution. 10% of Microbial transglutaminase (mTG) was added to crosslink the gelatin. Gel solutions were plated, incubated at 37°C, and swelled with 1.5 mL DI water in 30 minute intervals over 3 hours. Rheological analysis was done to evaluate the mechanical strength of these gels, revealing that PF108 had a significantly higher elastic modulus compared to the other gels [Fig 1]. The laser microscope imaging supported this result as the structure of the PF108 hybrid gel displayed branching fibers, while the structures of the other hybrid gels displayed gelatin mesh networks [Fig 1]. The significantly lower surface roughness of the PF108 hybrid gel is associated with decreasing cell attachment, thus, gelatin-PF108 hybrid gel may possess promising characteristics to serve as a cell barrier layer.

To further characterize the poloxamers, a variety of tests were used on the solutions using the rheometer. The amplitude sweeps revealed important properties about each poloxamer regarding elastic and viscous modulus. Frequency sweeps were performed to determine the critical frequency at which $G'$ (elastic modulus) = $G''$ (viscous modulus) [Fig 2]. At amplitude sweeps above such frequency, $G'>G''$ held true for each poloxamer, while below such frequency, PF108 and PF98 had $G''>G'$. This suggests that the top layers of collapse at low frequencies while all layers stay elastic at high frequencies [Fig 2]. Temperature sweeps utilized the critical frequency to determine the temperature at which each poloxamer solution transitions from liquid to gel [Fig 3].

Future research for hybrid hydrogels will include assessing the cell attachment and testing the clinical viability of the gels through in vivo studies and under physiological conditions. For the poloxamer gels, future research will include further determination of its structures.
Analyzing the In Vitro Viability of Novel Gelatin-Pluronic® F127 Hybrid Hydrogels as Cell Barrier Membranes

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Hydrogels are hydrophilic macromolecular natural or synthetic polymer networks that are crosslinked to produce an elastic, gel-like structure. They are biocompatible, can mimic the characteristics of naturally occurring gels, and can be engineered with desirable properties, creating potential for a variety of promising biomedical applications. Gelatin is a natural polymer with high biocompatibility frequently used in hydrogel synthesis; however, it has poor mechanical strength. Pluronic® F127, a hydrophilic triblock copolymer, can be mixed with gelatin for increased mechanical strength and tunable biocompatibility; additionally, unlike gelatin hydrogels, F127 hydrogels have demonstrated decreased cell attachment. The resulting potential for a novel Gelatin-Pluronic F127 hybrid hydrogel to possess tunable biocompatibility, high mechanical strength, and impermeability to cells makes it ideal for biomedical use as a cell barrier membrane; thus, this study focused on analysis of their in vitro viability.

Based on previous research at Stony Brook University that determined an optimal Gelatin-F127 hybrid mix, 15% w/v Gelatin-20% w/v F127 hybrid hydrogels were synthesized with 1%, 5%, 10%, and 15% w/v microbial transglutaminase (mTG), a cross-linking enzyme that synthesizes mesh networks from polymer strands. Characterizing hybrid hydrogels with rheological analysis demonstrated drastically increasing average elastic modulus with increasing mTG concentration (Figure 1). Further characterization using surface analysis with laser microscopy demonstrated decreasing surface roughness as mTG concentration increased.

Cell attachment was analyzed to determine the impermeability of the hybrid hydrogels to cells. Specifically, green fluorescent human dermal fibroblasts were plated on pure gelatin and hybrid hydrogel samples. Fluorescence imaging indicated a clear decrease in cell attachment with increased mTG concentration; for instance, while cells remained attached on highly crosslinked 15% gelatin, none remained attached on 15% hybrid hydrogels (Figure 2). Further analysis with confocal microscopy indicated similar results. To determine whether reduced cell attachment was due to barrier membrane capabilities or possible toxicity, cytotoxicity tests were conducted with fibroblasts plated in both hybrid gel cell mediums and in direct contact with hybrid gels. Similar fluorescence cell plate readings across the data for both tests indicated hybrid hydrogels are not toxic to cells and are biocompatible. In vitro membrane barrier testing was done with three-part gels plated in wells, with a control setup consisting of gelatin/gelatin/gelatin and an experimental setup of gelatin/hybrid/gelatin, with 5%, 10%, and 15% mTG concentrations for gels plated in the middle. Gelatin on the sides of the gels had a 10% mTG crosslinking density. Red and green fluorescent human dermal fibroblasts were plated on the side gelatin gels. Fluorescence readings revealed mixing of the fibroblasts in gelatin controls while the hybrid gels, particularly 15% mTG, prevented mixing of the fluorescent fibroblasts, demonstrating the in vitro effectiveness of the hybrid hydrogel as a cell membrane barrier.

Due to their non-cytotoxicity, cell impermeability, and high mechanical strength shown in vitro, hybrid hydrogels, particularly 15% mTG crosslinking density samples, have high potential for use as cell barrier membranes in guided bone regeneration following periodontitis, a disease that is the leading cause for tooth loss and affects nearly half of adults in the United States. They have further potential as burn wound dressings, as cell impermeability allows for both blocking of infections and easy changing of gel dressings with minimal patient pain. For further analysis of biomedical applicability, the ability of the synthesized gelatin-Pluronic® F127 hybrid hydrogels to entrap and release small hydrophobic drugs may be tested, as well as the clinical in vivo viability of the gels on animals with periodontitis or burn wounds.

Modeling the Diffusion of Drugs Through Hydrogels For Applications in Drug Delivery
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In recent years, materials known as hydrogels, or hydrophilic networks of polymer chains, have been increasingly used to release medication slowly over time to a specific part of the body. Hydrogels have many physical properties that contribute to their utility for drug delivery; besides their highly porous structures which allow for straightforward loading and diffusion of drugs, hydrogels are easily controllable, highly biocompatible, and relatively flexible.¹ The goal of this research was to model how changes in the structure and behavior of hydrogels and various drugs would affect drug delivery via diffusion. The modeling throughout this study was done with LAMMPS (Large Scale Atomic/Molecular Massively Parallel Simulator), a molecular dynamics software that simulates particle movement.

Two types of physically associating gels were simulated in this study. We started with a simple system of 1000 telechelic polymers chains of length 20 segments (12.2% volume fraction)² and formed an organic gel via end group attractions. Then spherical nanofillers that had strong attraction to the end groups of the polymer chains were introduced to simulate an organic–inorganic network. To analyze the structures of the two gels, we examined the radial distribution functions for both and discovered that the gel with nanofillers had a significantly more uniform intercluster distance, suggesting that nanofillers facilitated the formation of ordered local structure.

Afterwards, we studied how drug particles of different sizes and shapes diffused through both gels. The drugs were first modeled as rigid solid spheres. To ensure that the diffusion coefficient only depended on the structure of the drug and the hydrogel, we minimized the concentration of drug particle. Mean squared displacement was measured over 500000 timesteps, and linear regression was performed to obtain the diffusion coefficient. The results (Figure 1) showed that the diffusion coefficient decreased as the size of the drug particles increased, and drug particles of the same size diffused at a slower rate in the nanocomposite gel due to higher crosslinking density. Then we used the same method to model drugs as 30 oligomers of length 5 segments, which also diffused at a slower rate in the nanocomposite gel.

The simulations provided information on how different parameters changed the diffusivity of solutes through hydrogels, which can lead to more effective uses of hydrogels in drug delivery therapies. For further study, we plan to study more parameters that can change the diffusivity, run the simulations on a bigger scale, and increase the complexity of the shapes of the drugs to better mirror drugs in the market.

![Figure 1](image)

Figure 1: graphs of mean squared displacement vs time for diffusion of drug particles through hydrogel without (left) and with nanofillers (right)

Lattice Boltzmann Modeling of Fluid Flow Dynamics in Metal-Organic Frameworks

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Metal-organic frameworks (MOFs) are large crystalline networks consisting of inorganic metallic ions bonded together by organic linkers. These structures have nanoscale pores, extremely high surface areas, and can be reused many more times than traditional materials. Thus, MOFs are emerging in various applications such as catalysis, sensing, gas storage, and gas separation \(^1\). Indeed, over 50% of industrial catalysts used today are MOFs \(^2\). Unfortunately, there are two main problems with the usage of MOFs. First is that the permeability of a MOF is hard to control or predict without experimentation or expensive laboratory testing. Second is that it is nearly impossible to create a perfect MOF without any defects \(^3\).

To address these issues, we used a Lattice Boltzmann model of fluid flow through a MOF-808 crystalline structure. Analyzing fluid flow is a necessary step toward understanding the mechanisms of their activities as catalysts, sensors, and filtration materials. The flow was initialized by creating a higher density at one side of the lattice and a lower density at the other, forcing the liquid to move through the pores of the MOF. Defects were introduced in the structure by creating a random number generator and constraining the outputs to certain probability values. Most MOFs have 2%-14% defects, so preliminary results were calculated for 0% defects, 2% defects, and 5% defects. The pressure gradient was tested at values of 0.02, 0.04, 0.06, 0.08, and 0.10. A surface integral was introduced into the code to calculate the flux of the fluid exiting the lattice.

The most relevant data existed at the points when the fluid flow had reached steady-state equilibrium. Flux versus negative pressure gradient was graphed and analyzed (Figure 1). Because effective permeability is directly proportional to the slope of the curve, this data was also yielded permeability values (Figure 2). To further contextualize these results, a simulation was done to determine the approximate amount of time that each lattice took to reach equilibrium (Figure 3). Velocity distributions were then characterized along a single, straight line using Paraview simulation software (Figure 4).

The data indicate that the higher defect value lattices have increased permeability but take longer to reach equilibrium and have lower fluid velocities overall in this state, which is a novel result. This is likely due to the abnormal channels that are created when defects are introduced. In addition, these results are applicable to other anisotropic MOFs and suggest that MOFs can be designed with desired fluid permeabilities. Further development includes testing greater defect and pressure values, changing the defect locations on the lattices, and analyzing the channels created when defects are introduced.

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Session 3: Cellular/Bacterial Interactions

Chairs: Kao Li & Fan Yang
The formation of bacterial biofilms on implants is a great problem, because once formed they cannot be removed. Thus, there is a need of an approach to modify implant materials in order to prevent biofilm formation. There has been a lot of research done on the antimicrobial properties of silver nanoparticles [1]. However, when silver nanoparticles have been mixed with a polymer before implant preparation, it has been observed to be unable to release silver ions which carry the antimicrobial effect. Polyactic acid (PLA) is biodegradable and FDA-approved for use in the human body. The Brunst method has recently been patented for in situ synthesis of Ag nanoparticles coating PLA [2]. Thus silver nanoparticles, when incorporated on the PLA surface, will allow release of silver ions for an extended period of time, inhibiting biofilm formation. The efficiency of coating can differ based on AgNO₃ concentration used, and PLA filament content related to different 3D printers.

We hypothesized that an increased concentration of silver nanoparticles would result in an increased ability to inhibit biofilm formation and will not be affected by the difference between the filaments.

PLA scaffolds were printed using two 3D printers, a Makerbot Replicator 2X and Ultimaker 2 Extended+, with corresponding filaments, and characterized with a laser microscope and FTIR. Two concentrations of AgNO₃ (0.1M and 0.01M) and 0.001M NaBH₄ were used with the Brunst method simultaneously, for both scaffolds. The in situ deposition of silver nanoparticles was observed with SEM. The PLA-AgNO₃ scaffolds were tested for ability to prevent biofilm formation against two strains of bacteria, Staphylococcus aureus (gram-positive) and Escherichia coli (gram-negative) in TSB media. Biofilm formation was identified by crystal violet staining, life/dead fluorescent stain, and confocal microscopy at 6, 24 and 48 hours. The PLA-AgNO₃ scaffolds were also tested for oral pulp stem cell (DPSC) adhesion. After plating, DPSCs were cultured in media (αMEM, 10% FBS, 0.002M Ascorbic acid, 0.01M β-Glycerol phosphate). Alamar blue staining and EVOS FL microscope were used to monitor cell attachment and proliferation.

Analyses of both types of scaffolds show significant differences in surface roughness (fig.1), and chemistry. These reflected the adhesion and biofilm formation of both bacterial strains, surprisingly more attracted to smoother scaffolds printed on Ultimaker 2 Extended+. PLA-AgNO₃ nanoparticle coated scaffolds at 22 and 48 hours after bacterial inoculation showed significantly smaller size of the biofilms on average for both E. coli (fig.2) and S. aureus (fig.3) with 0.1M than the one with 0.01M nanoparticles (fig.2, fig.3) as proposed. The DPSCs had the same correlation and attached significantly better to the Ultimaker samples in comparison to the Makerbot samples (fig.4). Future work will include observing DPSC attachment on PLA scaffolds coated with AgNO₃, as well as analyzing the live-dead ratio of bacteria in the biofilms.

Characterization of Antibiofilm Activity of Silver Nanoparticles Deposited in situ on 3D Printed Polylactic Acid Implant Models

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The formation of bacterial biofilms on implants is a great problem, because once formed they cannot be removed. Thus, there is a need of an approach to modify implant materials in order to prevent biofilm formation. There has been a lot of research done on the antimicrobial properties of silver nanoparticles [1]. However, when silver nanoparticles have been mixed with a polymer before implant preparation, it has been observed to be unable to release silver ions which carry the antimicrobial effect. Polylactic acid (PLA) is biodegradable and FDA-approved for use in the human body. The Brunst method has recently been patented for in situ synthesis of Ag nanoparticles coating PLA [2]. Thus silver nanoparticles, when incorporated on the PLA surface, will allow release of silver ions for an extended period of time, inhibiting biofilm formation. The efficiency of coating can differ based on AgNO₃ concentration used, and PLA filament content related to different 3D printers.

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Naturally Occurring Fibrinogen Chain Variant $\gamma'$ Associated with Decreased Risk for *Staphylococcus aureus* Bacteremia

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*Staphylococcus aureus* bacteremia has become one of the most prevalent and difficult medical conditions to treat. Over time, the number of antibiotic-resistant *Staphylococcus aureus* strains has increased, posing a serious clinical challenge and public safety concern.¹ Bloodstream infections (BSI) is ranked as the 11th leading cause of death in the United States, accounting for almost 36,000 deaths in 2008 alone. It has been suggested however, that the number of BSI-related deaths in the United States is severely under-reported and actually numbers upwards of 70,000 given that in many cases bloodstream infections are overlooked as a cause of death amongst terminal conditions (eg. end-stage kidney disease, advanced malignancy) and underlying sources of infection.² With the viability of conventional treatments diminishing, naturally occurring therapies have become a target. Fibrinogen is the soluble blood precursor of the fibrin clot and plays crucial roles in mediating clot formation and wound healing.³ Each molecule is symmetrical and consists of a set of three different chains termed $\alpha$, $\beta$, and $\gamma$. In approximately 10% of all fibrinogen, a natural variant called $\gamma'$ arises due to an alternative processing of the $\gamma$-mRNA.⁴ *Staphylococcus aureus* has the ability to bind to fibrinogen via membrane-bound bacterial virulence proteins, an important mechanism that contributes to bacteremia. However, studies in mice have shown that a higher incidence of mutant fibrinogen $\gamma^{\Delta5}$ - which lacks certain bacterial binding motifs - actually significantly increases host survival rates from BSIs.⁵ In this study, it was hypothesized that the human variant $\gamma'$ could possibly demonstrate a similar capacity to combat bacterial infections. Solutions of polystyrene ($M_w=200k$, monodispersed; $M_w=400k$, monodispersed; $M_w=35k$; $M_w=280k$) dissolved in toluene was created and silicon wafers were treated via piranha etching. The wafers were then treated with HF to turn the surface more hydrophobic and the polystyrene was spun cast on to the wafers for 30 seconds at 2500 RPMs and annealed overnight at 120°C. Two types of fibrinogen-fibrin solutions (rich in soluble fibrin and depleted in soluble fibrin) were deposited on the various polystyrene thin film samples. Fiber growth occurred overnight and was quantified using optical microscopy (Fig. 1), identifying polystyrene ($M_w=200k$, monodispersed) to be the optimal type for fiber development (Fig. 2). *Staphylococcus aureus* was deposited on fibers grown in polystyrene ($M_w=200k$, monodispersed) and incubated at 37°C for 1 hour. A fluorescent live/dead stain was completed on the samples and observed with fluorescent microscopy. A subsequent trial was conducted; prior to bacterial deposition, the fibers were treated with anti-$\gamma$(80-411) to block a large portion of the gamma chain. A secondary anti-sheep antibody treatment was done after bacterial deposition to fluoresce the primary anti-$\gamma$(80-411) antibody. In future studies, anti-$\gamma'$ antibodies will be utilized and their impact on bacterial adhesion will be observed through AFM and SEM.

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Investigating the Nanotoxicity of Titanium Dioxide via Fibroblast Growth and Infection with *Staphylococcus aureus*

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Titanium dioxide (TiO₂) is commonly used in paints, toothpaste, sunscreen, cosmetics, pharmaceutics, cosmetics, and food additives. [1] While there has been an abundance of research on its anti-cancer and anti-bacterial applications with irradiation with UV light, there is little investigation on how TiO₂ may actually be harmful to cells. In a previous study, HeLa cells were exposed to TiO₂ nanoparticles, and it was concluded that exposure might significantly increase the risk of bacterial invasion. [2] Because of its wide array of applications, constant contact with these nanoparticles could be extremely harmful. In our study, we sought to corroborate the conclusions made by researching the effects of exposure of TiO₂ rutile nanoparticles on cell growth as well as bacterial infection behavior.

Human dermal fibroblasts and a gelatin hydrogel substrate were utilized. We sought to analyze how these cells would grow in four different conditions—normal tissue culture plate (TCP), TCP with TiO₂, Gel, and Gel with TiO₂. The titanium dioxide was characterized with X-ray diffraction, TEM, and UV-Vis spectrophotometry. We measured cell proliferation with the hemocytometer and correlated the doubling time calculated with EVOS and confocal microscopy.

In both our cell counting data and fluorescent microscopy images, it was apparent that there was little contrast between the TCP and TCP+TiO₂ samples, which may be due to the relatively low concentration of 0.1 TiO₂/mL. On the contrary, the Gel+TiO₂ samples showed a much lower cell count than the Gel samples (Figure 1). Moreover, the narrowness and clumping of fibroblast ends exposed to TiO₂ indicate cell deficiency (Figure 2). While we have not received bacterial colony formation units (CFU) data yet, it seemed that the bacteria are more clustered together with TiO₂.

From the data of the hemocytometer and the fluorescent microscopy, we can conclude that exposure to TiO₂ nanoparticles may potentially be toxic. Because a hydrogel substrate is meant to simulate physiological conditions, this provides compelling evidence that TiO₂ does decrease cell proliferation rate, while inhibiting some cellular mechanism to affect fibroblast morphology. Future research involves investigating why the fibroblast samples on a hydrogel substrate are more heavily affected than TCP samples, as well as the mechanisms behind different bacterial invasion behaviors.

Session 4: Fuel Cells

Chair: Likun Wang
Coating Nafion 117 Membranes with Graphene Oxide using Various Surface Pressures to Increase Performance of Proton Exchange Membrane Fuel Cells

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Proton exchange membrane fuel cells (PEMFCs) present a clean energy alternative to fossil fuels for generating electricity. In PEMFCs, hydrogen protons pass through the proton exchange membrane (PEM). To maximize performance, PEMs must have high proton conductivity and low gas permeability (Xue et al., 2014). Currently, Nafion, a perfluorosulfonic acid membrane, serves as the PEM in fuel cells. While Nafion has high proton conductivity, it is susceptible to hydrogen gas permeability which interferes with fuel cell performance (Francia et al., 2010). When coated on the PEM, graphene oxide (GO) can increase PEMFC performance due to oxygen-containing acidic functional groups which provide proton transport channels and smaller nanopores which decrease gas permeability (Farooqui et al., 2018; He et al., 2016).

The purpose of this research was to determine the effects of different surface pressures of a KSV Instruments Langmuir-Blodgett (LB) trough when coating Nafion 117 membranes with GO. 40 mL of GO solution was dropped onto the air-water interface using a syringe. The dipping mechanism of the LB trough was used to coat Nafion 117 membranes at 1 mN/m, 3 mN/m and 5 mN/m surface pressures.

The Nafion membranes were tested in a single-stack demonstration fuel cell kit from H-TEC. Four membranes were tested for fuel cell performance: an unmodified Nafion membrane as the control and Nafion membranes coated in GO at surface pressures of 1 mN/m, 3 mN/m, and 5 mN/m, respectively. The BK Precision 7601 DC Electronic Load controlled the resistance of the demonstration kit, and the BK 8600 Series Software recorded the current, voltage, and power. Current density and power density were later calculated.

The membranes coated with GO at surface pressures of 1 mN/m and 3 mN/m displayed similar decrease in voltage (Figure 1) and remained stable at the highest current densities (Figure 1 and 2). However, 1 mN/m showed the highest peak power density (Figure 2). When coating using the LB trough, the GO monolayer must be stable and homogeneous in order to best enhance the PEM’s proton conductivity. From these results, it appears that within our experimental groups, 1 mN/m is an optimal surface pressure for coating Nafion with GO. Surface pressures higher than 3 mN/m can possibly cause the GO monolayer to collapse, which does not allow the LB trough to lay a fully homogeneous layer of GO onto Nafion. More tests will need to be conducted for surface pressures between 1 mN/m and 3 mN/m to determine the exact value for optimal surface pressure.

Further experimentation will be done to analyze the surface morphology and thickness of the GO layers when coated on Nafion 117 membranes with different surface pressures. Additionally, the effect of the optimal number of monolayers of GO on Nafion 117 can be determined to effectively enhance the performance of proton exchange membrane fuel cells.

Proton-exchange membrane fuel cells (PEMFCs) have recently generated renewed scientific interest as a source of alternative energy due to their ability to generate electrical energy at a relatively low temperature and pressure without harmful by-products. Specifically, the polymer electrolyte membrane of PEMFCs remains a focus of current research, with Nafion being the most common polymer used in such membranes. However, the current efficiency of PEMFCs is well under the desired threshold to be widely used in a variety of applications. With increased utilization efficiency of platinum content in PEM fuel cell, this research investigated the effects of the deposition of platinum nanowires onto the Nafion membrane at various surface pressures.

Platinum nanowires were synthesized by the Stony Brook Department of Chemistry and were sonicated in ethanol to create 1.0 mg/mL solutions. The resultant solution was used with a Langmuir-Blodgett (LB) trough (KSV 3000) to deposit a platinum nanowire monolayer onto the Nafion membranes at surface pressures of 1 mN m\(^{-1}\), 3 mN m\(^{-1}\), and 5 mN m\(^{-1}\) with respect to the baseline (Figure 1). All Nafion membranes were tested at the same fuel cell test station connected to a programmable DC electronic load (BK Precision 8601) with 4 cm x 4 cm electrodes of 0.1 g/cm\(^2\) Pt/C. The hydrogen release rate was set to 80 cm\(^3\) min\(^{-1}\), and voltage and power outputs were measured starting from 0.000 A with a 0.050 A increase every 30 seconds.

The isothermal curve has a baseline at around 3 mN m\(^{-1}\); the irregularities of the curve past 350 cm\(^2\) on the horizontal axis are due to the addition of platinum-ethanol solution disrupting the surface of the water bath. The baseline remains at approximately 3 mN m\(^{-1}\) because the ethanol has not yet evaporated and thus retains surface pressure on the water. Nevertheless, the abrupt spike in surface pressure confirms the presence of a platinum nanowire monolayer, and deposition occurred at the labelled points. Figure 2 displays the polarization curve of PEMFCs using an uncoated control in addition to the three membranes coated with platinum nanowires at the varying surface pressures. The control membrane performed the worst, as expected. Membranes coated with surface pressures of 3 mN m\(^{-1}\) and 5 mN m\(^{-1}\) returned similar results, so it is hypothesized that the optimal coating surface pressure lies between these two values.

Future work includes deposition of platinum nanowires at surface pressures between 3 mN m\(^{-1}\) and 5 mN m\(^{-1}\) to optimize the power output of the PEMFC. In addition, scanning electron microscopy can be used to detect differences between the various platinum nanowire-coated membranes and elucidate the properties of the coated Nafion membranes.

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Varying Graphene Oxide Deposition Mass in Polymer Electrolyte Membrane Fuel Cells to Optimize Fuel Cell Performance and Durability

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While, in theory, Polymer Electrolyte Membrane Fuel Cells (PEMFCs) offer a clean and highly efficient form of alternative energy, there still remains several obstacles in their development that must be overcome to compete with traditional energy sources, including cost, power output and durability.¹ Studies have shown that GO has positive effects on both power density and durability of PEMFCs.² However, previous research involving GO deposition does not compare the effects of various mass loadings of GO on the electrodes or membrane to find the optimal amount needed to maximize efficiency. In this study, a calibration curve was created to correspond mass loading of GO with area below Raman spectroscopy peaks of different GO samples. Using this curve, the performance of various tested mass loadings of GO were compared to a control to observe differences in power output.

A GO solution of 5mL Isopropyl alcohol, 5mL DI water and 10mg of GO flakes was sonicated for two hours before being sprayed onto 1x1cm squares of aluminum sheets at masses of 2, 12, 27, 60, 98 µg per square centimeter. Raman spectroscopy was conducted on the individual samples with three scans of equal area size being taken of each sample. The areas below the spectroscopy peaks for each sample were recorded and a calibration curve was constructed, which correlates area below peaks with the mass loading of GO (Fig. 1).

Four fuel cells were constructed with a platinum loading on the 5cm² electrodes of 2.6mg and 5.1mg for the anode and cathode, respectively. Two separate membrane electrode assemblies (MEAs) with 10µg and 20µg of GO solution sprayed on their electrodes were produced. Another MEA was also constructed with 1µg of GO solution spray-coated on the Nafion membrane. Each MEA was hot-pressed at 266°F and compared to a control cell that contained no GO.

Initial polarization curves and CV curves for each MEA were plotted at 60°C and 80°C, as well as after 5000 and 10000 cycles. As shown in Figure 2, both MEAs with the modified electrodes produced significantly lower power outputs than the control MEA. It is believed that the mass-loadings of GO may have been too high and were inhibiting electron movement within the fuel cell as evidenced by a high impedance spectrum. The MEA with 1µg of GO on its membrane, however, performed significantly better than the control fuel cell, validating that a lower mass loading than the other GO fuel cells would better optimize performance.

Further work will be done to compare the reduction in active platinum catalytic area within the cells after running many cycles. More MEAs will also be produced to find the mass loading of GO on both the electrodes and the membrane that will have the greatest power output. Tests could also be run with gas feeds containing carbon monoxide to test GO’s ability to prevent CO poisoning of the MEAs.

From the dawn of the industrial revolution, fossil fuels have remained the primary source of energy, threatening the fragile balance on our planet through ever-increasing carbon emissions. Standing at the cutting edge of material science, Polymer Electrolyte Membrane Fuel Cells (PEMFCs) provide an alternative source of clean energy through the reduction-oxidation reaction of hydrogen and oxygen gas. Although Langmuir-Blodgett through and air spraying techniques have served as the primary methods of catalyst ink deposition, this study applies electrospinning techniques to enhance electrical conduction across electrodes. By producing 3D nanofiber structures, we aim to increase the surface area to volume ratio for catalysis, promote proton transport, and ultimately, optimize hydrogen fuel cell performance.

To accomplish this, different polymers were tested until (Poly(acrylic acid)) was chosen for its optimal viscosity and conductivity. When mixed in a 4:1 weight ratio with Nafion to create a base solution, fibers of consistent diameter (1 µm) were produced with Pt/C nanoparticles effectively attached, as demonstrated by SEM imaging (Fig. 1). The final solution consists of 4.10 wt % Pt/C, 8.73 wt % Nafion and 2.18 wt % PAA and was electrospun at a potential difference of 12.5 kV and a syringe pumping rate of 0.5 mL/hr onto microporous carbon electrodes, achieving a Pt loading of around 0.100 mg/cm².

The electrospun electrodes, commercial electrodes, and airsprayed electrodes were then tested in an H-tech PEMFC kit in an open-air environment at the cathode with pure hydrogen being fed into the anode. The PEMFC tests indicate overall that electrospinning enhanced fuel cell efficiency. 0.1 mg/cm² Pt loading from electrospinning exhibited a 108% increase in max power output compared to 0.1 mg/cm² Pt loading from airspraying. Going forward, electrospinning solutions will be continually prepared to perfect the ratios of PAA, Nafion and Pt/C and optimize fiber formation. Polarization measurements will be recorded in pure air, 0.1% carbon monoxide, and triple MEA settings to determine if the fiber structure consistently enhances performance. Finally, Graphene Oxide and Au-Ag nanoparticles, which has previously been shown to mitigate CO poisoning, will be incorporated into the electrospinning solutions. We will proceed to compare the surface morphologies and performance of the fiber composites with the ultimate goal of identifying the most promising combination.

Utilizing ALD-Coated Titanium Dioxide as a Gold Nanoparticle Catalyst Support in Electrodes to Increase Efficiency Proton Exchange Membrane Fuel Cells

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Proton exchange membrane fuel cells (PEMFCs) are increasingly being used for their lack of pollution and clean reactants and by products¹. However, in real-life applications, the PEMFCs face carbon monoxide (CO) poisoning from hydrogen gas and atmospheric air impurities. The CO binds to the platinum at active catalyst sites of the electrodes, greatly lowering their electrochemical ability. Previous studies have found that gold nanoparticles (AuNPs) effectively catalyze the CO oxidation at low temperatures. Additionally, thin layers of titanium dioxide (TiO₂) are effective catalytic support². In this study, our team investigated the effect of coating the electrodes with ALD-coated TiO₂ and AuNPs on the PEMFCs and the optimal number of layers of TiO₂ to allow the highest power output in the fuel cell.

Atomic layer deposition (ALD) of TiO₂ was done on two 4cm x 4cm electrodes of 0.1 g/cm² Pt/C at 80°C using precursors of titanium tetraisopropoxide (TTIP) and water from 1 to 5 cycles. An AuNPs solution of 1mg/mL in toluene was created and synthesized through the Brust method. A Langmuir-Blodgett (LB) trough was used to coat a gold nanoparticle monolayer onto the TiO₂-coated electrodes. The electrodes were tested in a fuel cell test station (BK Precision 8601 DC Electronic Load single stack) with Nafion membrane. Tests were run with currents from 0.000A to 0.700A at 0.050A intervals timed at 30 seconds per interval to determine voltage and power output.

The power output of electrodes with a single layer of TiO₂ and AuNP coating had the highest power output of those tested, as shown in figure 1, with a peak of 0.1387 watts. This wattage is a 27% increase in power output in comparison to the power output of the control, which was found using uncoated electrodes. In general, the AuNP coat increased the power output for each amount of TiO₂. The trial with the worst power output, having a peak power output of 0.032 watts, 5 cycles of TiO₂ and no AuNPs confirms the previous research that TiO₂ is an effective catalyst support for AuNPs in the oxidation of CO, as TiO₂ alone blocks the flow of protons through the PEM, but helps AuNPs catalyze CO. To confirm an even coat of AuNPs, photos were taken under a transmission electron microscope, as shown in figure 2, and analyzed using ImageJ. The average particle size was found to be 2.46 nanometers.

Further characterization tests such as ellipsometry to determine the thickness of TiO₂ and TEM imaging of the TiO₂ to find the distribution of particles will be done. Other future work includes testing the durability of the electrodes by running it through 33,000 cycles in the fuel cell test station, and CO will be cycled through the hydrogen fuel cell to test the effectiveness of the AuNPs.

Session 5: Dental Pulp Stem Cells

Chairs:
Kuan-Che Feng, Linxi Zhang,
Ya-Chen Chuang, Xianghao Zuo,
Juyi Li
Despite myriad advances in materials, techniques, and implant design, implant failure remains a significant concern in the clinical world. In an effort to reduce implant failure rates, researchers have been investigating new ways to modify the surface of implants in order to promote implant-tissue integration, or osseointegration [1]. Many of these methods, such as an already FDA-approved laser rostering method, are expensive and time-consuming. We are thus looking at the potential of a novel, cost and time effective method to promote osseointegration: chemical etching using Vilella’s Reagent. Commercially available 316L stainless steel (316L SS) is used for orthopedic implants due to its desirable structural properties, biocompatibility, and proven success in load bearing and fixation [2]. Because of its known benefits, we chose to utilize 316L SS in our study in which we investigated four prototypes: chemically etched wrought 316L SS, polished wrought 316L SS, chemically etched 3d-printed 316L SS, and polished 3d-printed 316L SS.

Each 316L SS sample was printed under identical conditions using a Renishaw AM250. Scanning electron microscopy (SEM) was used to assess the surface roughness (figure 1). All four prototypes were compared for ability to promote dental pulp stem cell (DPSC) attachment, proliferation, and differentiation. DPSCs were plated onto 3d printed and wrought 316L etched and non-etched samples and were cultured in the same medium (αMEM, 10% FBS, 0.002M Asc. acid, 0.01M β Glycerol phosphate). Alamar blue staining and EVOS FL imaging for fluorescence and transmitted light applications were used to monitor cell proliferation and cell-substrate interaction, respectively.

We performed x-ray photoelectron spectroscopy (XPS) on the 3d printed steel before and after etching in order to confirm that the etching process did not yield an unfavorable surface chemistry. The contact angle of each surface was then measured, which allowed us to confirm that the etching process increased the hydrophilicity of the surface. Thus far, our cellular study has suggested that the etched 3d printed 316L SS is the most favorable substrate in terms of cell growth (figure 2). Future work includes RT-PCR analysis of mRNA expression for DPSC differentiation, confocal imaging of actin-stained cells 24 hours after plating in order to assess cell adhesion, and a scratch test for evaluation of osseointegration.
Poly-lactic acid (PLA), a synthetic biodegradable poly-lactone, is commonly used in the tissue engineering field for its suitable mechanical properties and biocompatibility. In particular, its thermoplastic characteristics can be harvested by 3D-printing technology to create scaffolds for cell growth. However, the polyester surface is characterized by a suboptimal hydrophobicity for cell adhesion, which may limit its use. In this study, 3D-printed PLA scaffolds are coated with a thin, uniform layer of titanium dioxide (TiO₂) through Atomic Layer Deposition (ALD) to create an environment for dental pulp stem cells (DPSCs) attachment, proliferation, and differentiation.

Circular disks of 12 mm in diameter and 2 mm in height were prepared using PLA filaments through 3D-printing at 50 mm/s and 230°C as well as molding at 180°C and 7 psi to compare the cell response on rough and smooth surfaces. ALD was performed at 85°C for 50 cycles to produce a 5 nm layer of TiO₂ on the surface of the scaffolds. Scanning electron microscopy (SEM) was conducted on the scaffolds, and the results confirmed that ALD of TiO₂ was uniform (Fig. 1). The Atomic Force Microscopy (AFM) analysis revealed that the TiO₂-coated 3D-printed scaffolds have uniform, nanoscale roughness compared to the uncoated counterparts. DPSCs were plated on the scaffolds in Alpha MEM medium plus 10% Fetal Bovine Serum (FBS) at an initial 9*10^4 cell/mL. After day 1, the medium was changed every other day with Alpha MEM, 10% FBS, 200mM ascorbic acid and 10mM β-glycerophosphate until the end of the experiment. Cells were counted and visualized on day 1, 4, and 7 through fluorescence staining and microscopy (Fig. 2). The calculated doubling time between day 1 and day 7 displayed no significant difference from each other regardless of scaffold type, signifying overall cell health (Fig. 4). Future research will be conducted on the differentiation through RT-PCR analysis and biomineralization of DPSCs through SEM analysis on Day 28 and 42.
Evaluating the Effects of Graphene-Loaded Polylactic Acid Electrospun Fibers and Spun-cast Thin Films on the Proliferation and Differentiation of Dental Pulp Stem Cells

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As research in biomedical engineering continues to peak the interest of many scientists, studies utilizing dental pulp stem cells (DPSCs) have played a major role in the development of scaffolds. DPSCs are easily accessible and have a multilineage differentiation potential. The polymer scaffold we used is polylactic acid, rather than the more commonly used polystyrene(PS)-based tissue culture plastic. PLA is a biodegradable, bioabsorbable, and biocompatible polymer that has several applications in stem-cell based tissue engineering and drug delivery.2 Adding a material like graphene, which holds many unique mechanical, chemical, and electrical properties, can have a powerful effect on how cells interact on biocompatible scaffolds.

The primary goal of our research was to test the effect of graphene in PLA electrospun scaffolds on the differentiation and proliferation of DPSCs. To initiate the experiment, we created four different solutions of varying concentrations of PLA and graphene. Two samples of 12.5% PLA/chloroform and 5 mg/ml PLA/chloroform solutions were created for electrospinning and spincasting. These solutions were reproduced with the addition of 3% graphene. PLA thin film substrates were spun-cast with and without graphene at 2500 rpm for 30 seconds each. Fibers were electrospun on silicon wafers and glass slides at 1 mL/hour and 20 kV. After electrospinning and spincasting were completed, DPSC cultures were prepared for a 28-day analysis to study cell proliferation and differentiation.

Scaffolds were characterized by SEM and AFM images. Graphene aggregations formed on thin films. After determining the diameter of the PLA fibers from Figure 1 SEM images, we discovered PLA/graphene fibers had a high dispersion rate. Cell counts showed that growth rates were similar for both fiber samples. Confocal microscopy was performed to view cell morphology, and 40x magnification images in Figure 2 displayed that cells grew linearly along electrospun fibers. PLA/graphene thin films inhibited the extension of some actin filaments. PLA and PLA/graphene fibers had low proliferation rates, as shown in Graph 1 , indicating a possible cell confluence on day 7. On days 7, 14, and 28, RT-PCR, SEM/EDX, and confocal microscopy will be conducted to measure differentiation and biomineralization of DPSCs. While microscopy will determine if the deposits and proteins display differentiation, RT-PCR will measure several differentiation gene markers, including osteocalcin and dentin sialophosphoprotein. These genes will indicate relative levels of osteogenic and odontogenic differentiation.

Figure 1: AFM characterization
Figure 2: DPSCs are elongated on fibers
Graph 1: DPSC growth curve

Investigating the Effects of Substrate Mechanical Patterns on Proliferation and Differentiation of Dental Pulp Stem Cells

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Previously, stem cell differentiation control has relied heavily on chemical signals, but new research has demonstrated that it can be controlled by the mechanical properties of a cell’s surroundings. Multipotent dental pulp stem cells (DPSCs) can differentiate into odontoblasts, osteoblasts, and adipocytes and are sensitive to the hardness of their substrate. Odontogenesis is supported by hard thin spin-cast polybutadiene films and inhibited by soft thick films. It is hypothesized that, during odontogenic differentiation, stem cells transmit signals that promote or inhibit differentiation. Hence, cells were plated in touching and non-touching conformations to determine whether these signals are transmitted physically or through the medium.

3 mg/mL and 20 mg/mL concentrations of polybutadiene/toluene were prepared and spun cast onto Si wafers to create 20nm and 200nm thin films. 2x2 cm wafers with thick or thin films were used as controls in isolated wells, and 4 1x1 cm wafers were placed with alternating thick and thin films so that they were all touching or all non-touching. Strain 13 DPSCs were plated at 5x10³ cells/cm², fed every 2 days with osteogenic media of α-MEM, 10% FBS, Pen Strep, L-ascorbic acid, and β-glycerol phosphate and incubated at 37°C and 5% CO₂. On day 7, the relative moduli of the cells were measured using shear modulation force microscopy. On days 7, 14, and 28, samples were stained for confocal microscopy with AF488 and DAPI for F-actin and nuclei respectively and prepared for reverse transcription polymerase chain reaction (RT-PCR) using QIAzol Lysis Reagent.

On day 7, the relative modulus was low for cells on thick films and high for cells on the thin film control. However, all cells on touching thin films and some on non-touching thin films had low relative modulus (Fig. 1). In confocal images, all cells had abundant amounts of actin and were therefore healthy. However, actin on thin films tended to be randomly oriented while actin on thick films were oriented in one direction, which may explain the differences in moduli (Fig. 2). These results suggest that cells on thick films influenced differentiation of cells on thin films in co-cultures through media.

On day 28, biomineralized crystals will be characterized using Raman and scanning electron microscopy, and mRNA will be analyzed using RT-PCR to determine if odontogenic and osteogenic differentiation have occurred. This is identified by gene markers, specifically ALP, OCN, and DSPP. To produce more suitable in vivo conditions, microscale patterns will be created through laser etching, and nanoscale patterns by spin casting polybutadiene-polystyrene mixtures, which will phase-separate and create bumpy surfaces.

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Utilization of P4VP and Graphene to control Dental Pulp Stem Cell Neurogenesis

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Currently, the central nervous system is unable to heal effectively by itself, leading to a search for ways to regenerate or grow neurons for replacement. Dental Pulp Stem Cells (DPSCs) are multipotent stem cells that have exhibited the ability to differentiate into osteoblasts, cardiac cells, and neurons. Since DPSCs are easily acquired from extracted wisdom teeth and can provide cells in various areas resulting in an autologous implant, preventing rejection by the patient, DPSCs show a promising hope in biomedical research and engineering.¹ P4VP, a polymer shown to retain cell morphology, can help cells adhere to a substrate, potentially eliminating the need for polyornithine.² Graphene, known for its electrical conductivity, may prove useful in the differentiation of DPSCs into neurons, since neurons communicate through electrical impulses called synapses.³ This experiment compares the effects of P4VP, graphene, polyornithine, and the presence of fibers, on DPSCs and their differentiation into neural cells. Cell functions will be monitored using microscopy and differentiation verified by Real-Time Polymerase Chain Reaction (RT-PCR) using primers for neurogenic markers (e.g., nestin, tubulin III, NF-M).

For spincoating, P4VP was dissolved in DMF at 7 mg/ml with or without 3% graphene by mass to P4VP. For electrospinning, P4VP at 25% by mass was dissolved in a 5:1 ratio of ethanol and DMF with or without 3% graphene by mass to P4VP. For evaluations of mRNA expression, 2x2 cm silicon (Si) wafers were used. These wafers were first cleaned using a 3:1:1 ratio of piranha solutions. Si wafers were spuncast at 2000 rpm for 30 sec along with glass coverslips, for optical microscopy, for 45 seconds. Half of the samples were also electrospun at 15k V and 3 mL/hr to create fibers (Figure 1). This experimental setup was replicated, one setup coated with polyornithine, the other without. All samples were treated with laminin (5 µg/mL) overnight. Glass coverslips were placed in 24 well plates and plated with 33,000 cells in 1 mL Growth Media (alpha MEM supplemented with 10% fetal bovine serum, 0.1 mM L-ascorbic acid 2-phosphate, 2 mM L-glutamine), and Si wafers were placed in 6 well plates, plated with 150,000 cells in 3 mL growth media. A positive control of tissue culture plastic was also plated. Cells were cultured in Growth Medium for 2 days, after which the media was replaced with neuronal growth media (Neurobasal A Medium, B27, 20 ug/ml EGF, 40 ug/mL FGF2), changing every 2-3 days for 21 days. During the first week there was no significant difference between the control and thin films. The cells growing in fiber samples tended to elongate and grow parallel to the fiber structures surrounding them with a higher confluency on P4VP fibers alone, compared to P4VP with graphene fibers (Figure 1). Future Work includes conducting confocal microscopy with F-actin staining and RT-PCR with early, middle, and late neurogenetic markers after 21 days of growth.

Figure 1: Control Tissue Culture Plastic (Top-Left), P4VP (Top-Right), P4VP + Graphene (Bottom-Left), P4VP Fiber (Bottom-Right), Electrospun Samples (Far-Right)

The development of stem cell research has had a tremendous impact on the advancement of regenerative medicine, drug testing, and our understanding of the human body. Although stem cells were previously retrieved from embryos, several studies show that mesenchymal stem cells (MSC) have multipotency. Scientists have been investigating the abilities of dental pulp stem cells (DPSC), a type of MSC that can be found in the third molar\(^1\). DPSCs can be cryopreserved for future use and, when given the right environment, can develop into a myriad of different cells\(^2\). Titanium dioxide is most commonly used as a whitening agent in toothpastes and food products, with a small portion present in the form of nanoparticles (NPs). Recently, a study showed that the cellular uptake of TiO\(_2\) NPs had adverse effects on the cell viability and growth of MSCs\(^3\).

This study was conducted to further examine the effects of TiO\(_2\) NPs on the proliferation as well as the differentiation of stem cells, specifically for osteogenesis and odontogenesis. To create a suitable substrate for the cells to attach to, thin (TN) and thick (TK) polybutadiene (PB) films were prepared by spin-casting 3 mg/mL and 20 mg/mL solutions on 1-by-1 and 2-by-2 cm\(^2\) silicon wafers of Miller index [100]. After samples were annealed for five hours at 150°C at 2.0 \(\times\) 10\(^{-7}\) torr, the PB films were determined to be \(\sim\)20 nm thick and \(\sim\)200 nm thick for TN and TK, respectively, through ellipsometry. The AV3 strain DPSCs were plated with a density of 5000 cells per cm\(^2\), and rutile TiO\(_2\) NPs were added at a concentration of 0.1 mg/mL the following day. Cell count was conducted on days 1, 3, and 6 with a hemocytometer. Atomic force microscopy (AFM) was used on day 5 to measure cell modulus (see Figure 1). A strange phenomenon was found where the addition of NPs made cells on the thick film harder and on the thin film softer. When the stained DPSCs were further examined with confocal microscopy (see Figure 2), the actin fiber growth supported the moduli data. Samples were collected for confocal microscopy and for real-time polymerase chain reaction (RT-PCR) on days 7, 14, and 28 and for scanning electron microscopy (SEM) on day 14 and 28. Confocal microscopy was used to examine actin fiber growth on day 14 (see Figure 2) as well as the nuclei of the DPSCs. SEM will allow us to examine the biominalization on the films, and Raman microscopy to further confirm its crystal structure. Using RT-PCR, we can observe the mRNA expression for up-regulation of alkaline phosphatase, osteocalcin, and dentin sialophosphoprotein gene marker to monitor the differentiation pathways.

### Figure 1
Relative moduli of cells measured on day 5 with AFM. TK/NPs saw a higher modulus than its control while TN/NPs had a lower modulus.

### Figure 2
Confocal microscopy of actin stained with AF488 on day 14 samples for (A) TN (B) TN/NPs (C) TK (D) TK/NPs. Thick actin fibers are visible, indicating healthy cell growth.

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Session 6: Environmental Protection

Chairs:
Yuan Xue & Xianghao Zuo
A Novel Fire Retardant Solution to Mitigate the Onset of Wildfires
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The dangers of wildfires have been brought to the public’s attention recently as a Greek forest fire took the lives of nearly 91 innocent civilians[1]. The demand for a proper fire retardant to stop the spread of a blaze before it reaches residential areas is growing, and it is important to therefore ensure that the fire retardant is safe (non-toxic, etc.) for the environment and the organisms that inhabit it. The experiment’s goal is to engineer an aqueous solution that can coat leaves and actively prevent the spread of wildfires. The leaf species used is Cornus kousa, a subspecies of the Cornus, or Dogwood family, and is freshly-picked from a local tree.

The leaves are divided into three categories: fresh leaves, oven-dried leaves, and naturally fallen leaves. Fresh leaves are manually selected and used as a sample less than 24 hours after selection. Dried leaves are curated in a CIT Alcatel 2004A Vacuum Oven for 25 min at 90 Celcius and 400 mTorr. It is proven through burning tests that fallen leaves and oven dried leaves burn faster and easier than fresh leaves, due to the fact that fresh leaves contain 66.28% of water on average.

The research investigates the effects of solutions that consist of combinations of different concentrations of component m, component n, component o, and deionized water. The ingredients were hypothesized to form a fire retardant solution and in our current study, it is made into an aqueous solution that can be distributed onto leaves for application on live trees.

The effects of pyrolyzing the leaves are measured through the time required for self-extinguishing (Table 1), charred area, and mass of leaves before and after burning (Figure 1). The ability of the solution to help yield char residue is analyzed through thermogravimetric analysis (TGA), shown in Figure 2. At 800°C, we could see that leaves coated with component m/DI water solution or component m/n/DI water solution yields more char than the ones without coating. Contact angle tests are performed to determine the hydrophobicity of the solutions on the leaf surfaces, seen in Fig. 3.

In addition, a toxicity test on Brassica rappa (Wisconsin Fast Plants) in relation to seed germination and plant growth is performed to study the ramifications of the solutions on plants. It has been noted that varying concentrations of component m and n do not completely stop germination, but rather retard the progress of germination. Component n has been observed to retard germination the most. Further studies are planned to test the toxicity of the solutions on animals living in the forested areas.

Table 1 Burning test of leaves

<table>
<thead>
<tr>
<th>Solution</th>
<th>Time to self-extinguish fresh leaf</th>
<th>Time to self-extinguish dry leaf</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>56.14 sec</td>
<td>6.94 sec</td>
</tr>
<tr>
<td>Solution A</td>
<td>&lt;1 sec</td>
<td>&lt;1 sec</td>
</tr>
<tr>
<td>Solution B</td>
<td>&lt;1 sec</td>
<td>&lt;1 sec</td>
</tr>
<tr>
<td>Solution C</td>
<td>&lt;1 sec</td>
<td>&lt;1 sec</td>
</tr>
</tbody>
</table>

Figure 1 Pyrolyzing Fresh Control

Figure 2 TGA results

Figure 3 contact angle measurements

The Effect of Graphene Oxide on Pheromone Evaporation in *Cimex lectularius* Traps

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The infestation of *Cimex lectularius*, commonly known as bed bugs, has been plaguing homes for centuries. Rates of bed bug infestation have recently been on the rise due to resistance to the chemicals and toxins used for their extermination. A new tool against *C. lectularius* is the Fibertrap, which is considered to be more effective and less expensive than the chemicals currently used to eradicate the insects.

Fibertrap incorporates fibers created by electrospinning to ensnare the fine leg hairs of *C. lectularius*. The insects are attracted to the traps because of the pheromones included, 1-Octen-3-ol and histamine. The problem with using the cost-effective pheromone 1-Octen-3-ol is that its evaporation rate is very fast, restricting the traps effectiveness to short periods of time. This research project examines using graphene oxide (GO) to prolong the effect of pheromones in *C. lectularius* traps.

Graphene oxide was first synthesised using a modified version of the Hummer’s Method. Graphite was dissolved using sulfuric acid (H₂SO₄) and was oxidized with potassium permanganate (KMnO₄) and sodium nitrate (NaNO₃). After oxidation, the GO was washed with distilled H₂O until the filtrate pH reached between 4 and 5. The GO was then dried in an oven at 50 degrees Celsius. Control concentrations of GO (10 mg/ml, 14 mg/ml, 20 mg/ml) were combined with H₂O. The same concentrations of GO were made with 1-Octen-3-ol. Another set was made with an additional pheromone of *C. lectularius*, histamine, along with 1-Octen-3-ol, the pheromone known to evaporate. A second set of samples was prepared using a solvent of 25% ethanol and 75% H₂O.

Fourier-transform infrared spectroscopy (FTIR) was used to identify the chemical bonds of the 1-Octen-3-ol in the 10 mg/ml solutions of GO. However, the solvents overpowered the FTIR data and no other bonds were visible. A new test was introduced to measure the decrease in mass over time of the 14 mg/ml solutions of the GO on a microgram scale, plotting the rate of evaporation for each sample. The results are shown in Figure 1. Future work will involve creating samples of pheromones without GO or ethanol solvent to compare the results of the mass lost and to test the bedbug’s attraction to each sample.

![Figure 1: The mass of the 14 mg/ml samples of GO and pheromones shown decreasing over time.](image-url)

Session 7: Nanocomposites

Chairs:
Xianghao Zuo & Yuan Xue
PLA is a 100% recyclable and biodegradable but weak polymer that has been used as an alternative to polystyrene in disposable cups, lids, and containers, and eating utensils. Commonly, binary blends including PLA and another polymer additive are created to enhance the mechanical properties of PLA, but this has found to be ineffective due to the high interface energy between the polymers. We hypothesize that adding small concentrations of PMMA or SMMA as compatibilizers to the binary mixes will lower the interface energy between the polymers.

To create the ternary blends, the polymers were added to a C.W. Brabender and then molded into the tensile and impact test shapes using a Carver Hot Press. The blends were also run through an extruder to create filaments for 3D printing. The mechanical properties of these molded and printed ternary blends were tested using impact and tensile strength tests. We hypothesize that due to the acrylic and styrene group present in SMMA, it will act as a compatibilizer that is miscible with PLA and the styrene based polymers, lowering the interface energy and improving mechanical properties. Further analysis about the interfacial properties is done by SEM imaging.

Each ternary blend contains 70% PLA and 30% styrenic polymer. The overall results of the impact test are shown in Figure 1. Among the materials we used, HIPS and ABS show a significant mechanical increase. The PLA/HIPS blend had maximum mechanical strength with 2wt% SMMA added, but as more SMMA was added, its mechanical strength weakened. A possible explanation for this is that the SMMA acts as compatibilizer and first gathers in the interface of the binary blend. As the amount of SMMA increases, the SMMA occupies the whole interface and forms a third polymer layer, because at high concentrations it is not totally miscible with either polymers.

![Figure 1](image1.png)

**Figure 1**: Impact test results for the 4 polymer blend vs compatibilizer weight

The SEM image of the PLA/HIPS blend with different SMMA concentrations confirmed these results, as shown in Figure 2. In Fig. 2a the blend with no SMMA shows the domain size of the HIPS matrix is ~11µm, in Fig. 2b the added 2wt% SMMA displays a significant decrease in size of the domain to ~5µm and the wrapping of HIPS by SMMA can be observed, in Fig 2c the domain size increased to ~9µm and the high concentration of SMMA formed an apparent layer around the HIPS, both contributing to the decrease of mechanical properties of the polymer blend at a high SMMA concentration. A 286% increase for the molded sample in the impact test was observed when 4wt% PMMA was used with the PLA/ABS blend. Figure 3 shows the significant improvement of impact strength for both the molded and 3D printed samples over the pure PLA and the binary PLA/ABS mix. It also shows that this ternary polymer material can maintain high mechanical properties when 3D printed.

For future study, interfacial properties in the ternary blends could be investigated to reveal interactions that contribute to mechanical strength. This could be conducted using TEM and SIMS analysis.
Enhancing Thermal Conductivity of Polylactic Acid and Polybutylene Adipate Terephthalate Blends with Boron Nitride and Graphene

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Thermal management is critical to the continually growing electronics industry to prevent devices from overheating and losing their functionality. As low-cost, lightweight, and highly versatile materials, polymers exhibit great potential for heat exchange applications, but their low thermal conductivity is a barrier to their effectiveness.1 To overcome this obstacle, we can supplement polymers with thermally conductive fillers such as graphene and boron nitride.

In this project, we investigated the thermal conductivity of loading blends of PLA (polylactic acid) and PBAT (polybutylene adipate terephthalate) with graphene and hexagonal boron nitride (hBN). Graphene and hBN have thermal conductivities of approximately 5300 W·m⁻¹·K⁻¹ and 360 W·m⁻¹·K⁻¹, respectively.2,3 We hypothesized that the thermal conductivity of the PLA/PBAT nanocomposite would be elevated compared to the thermal conductivity of the PLA/PBAT blend without the fillers. We also hypothesized that, since graphene had a greater thermal conductivity than hBN, the thermal conductivity would increase directly with the graphene-boron nitride ratio within the PLA/PBAT nanocomposite.

The CAM 200 Optical Contact Meter was first used to measure the contact angle between polymer droplets on hBN and graphene layers deposited on a silicon wafers to determine the polymers’ relative affinities for the fillers. A twin-screw extruder (C.W. Brabender Intelli-Torque Plasti-Corder®) was used to melt-mix PLA (NatureWorks Ingeo™ Biopolymer 4043D) and PBAT (ecoflex® F Blend C1200) with varying concentrations of hBN powder (Sigma-Aldrich®, ~1 µm) and graphene nanoplatelets (xGnP®, Grade H, 5 µm). We first tested PLA/PBAT and PBAT blends with hBN alone. Seeing a slow increase in thermal conductivity with increasing weight fraction of hBN, we added the graphene to see if graphene-hBN interactions would lead to greater improvements in thermal conductivity. Then, the samples were molded with a hot press (Carver Model C) set at 180°C and 7 psi. The thermal conductivities of the specimen were tested using the Anter Unitherm Model 2022 (ASTM E1530).

We discovered that adding the fillers did increase thermal conductivity. In addition, as the graphene content of the nanocomposite increased relative to the hBN content, the thermal conductivity increased as well.

For future research, we plan to conduct SEM and TEM imaging on our current samples to investigate how graphene and boron nitride interact within the PLA/PBAT blend. We also plan to investigate the mechanical properties of PLA/PBAT blends with graphene and boron nitride using impact and tensile strength tests. Ultimately, we hope to engineer a low-cost, biodegradable polymer nanocomposite with applications in the electronics industry.

Creating Flame Retardant and Biodegradable Polymer Blends for Industrial Use

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White pollution remains an ever-present issue and points to the need for biodegradable polymers that can fit a variety of applications. PLA (Poly(lactic acid)) presents a solution: derived from renewable resources such as corn starch, cassava roots, and sugarcane, PLA has low production costs, exhibits biocompatibility and biodegradability, and still maintains comparable mechanical properties to other conventional polymers. However, due to its poor impact strength, PLA is not used for most industrial applications, but rather for packaging, bio-implants, and 3D printing. To increase impact strength, PLA is often mixed with flexible polymers such as PBAT¹ (polybutylene adipate terephthalate). To broaden its applications to electronics and automotive industries, PLA’s flammability must be addressed.

We achieved V0 grade flame retardancy while maintaining and sometimes enhancing the mechanical properties of a PLA/PBAT blend using low RDP, APP and C-30B loading. Afterwards, we found the optimal settings for a 3D printer to maximize the mechanical properties.

As expected, the addition of PBAT to PLA enhanced the impact strength of the polymer blend (see Figure 1). However, addition of the flame retardants (APP, RDP) and compatibilizer (C-30B) reverted any mechanical advantage provided by the aforementioned PBAT addition. Alternatively, if the flame retardant blend is 3D printed with a vertical infill alignment, the impact strength increases by 15 J/m. We speculate the printing, as opposed to molding, creates a reinforced internal structure due to the directionality of the filament. Since the filament is aligned completely perpendicular to the direction of impact, there are no natural cleave lines in the sample.

Additionally, we notice while testing the tensile properties of our flame retardant blend, the printed samples tend to become fibrous under tension. However, the molded samples simply gain stretch marks and break cleanly. This can be easily seen through the tensile curves (see Figure 2 and 3). The printed sample has many jagged drops after the initial drop indicating how each large fiber breaks and passes on the stress to the next connecting fiber. In contrast, the molded sample has a much cleaner plastic deformation phase and breaks all at once.

Through the addition of PBAT, flame retardants (APP, RDP) and a compatibilizer (C-30B), the applications of PLA were successfully broadened. Not only is this blend flame retardant and thermoplastic, it is also biodegradable; it can be 3D printed and used in electronics all while representing a eco-friendly alternative to commercial plastics.

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Session 8: Perovskite Solar Cells

Chairs: Yuchen Zhou & Yifan Yin
Optimization of Organic-Inorganic Halide Perovskite Solar Cells via a Novel Polycaprolactone Additive Pathway

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The recent global energy crisis has resulted in a push for the development and commercialization of renewable energy sources, most notably solar cells. In 2009, Kojima et. al. revolutionized the field of photovoltaics by synthesizing the first perovskite-based solar cell.¹ Perovskites are semiconducting materials with the crystal structure ABX₃ whose unique optoelectronic properties make them very suitable as the light-absorbing layer in photovoltaic devices. Previous research has shown that additives can optimize the perovskite crystallization rate, thus producing larger crystal grains and full coverage.² This results in a perovskite layer with fewer defects and a solar cell with a higher power conversion efficiency (PCE). This study investigates the effect of a biodegradable, environmentally-friendly polymer additive, polycaprolactone (PCL), on the performance of planar, thin-film methylammonium lead iodide (MAPbI₃) perovskite solar cells (PSCs).

Planar, thin-film PSCs were fabricated with a FTO glass/TiO₂/PCL-doped MAPbI₃/spiro-OMeTAD/Au structure (Figure 1). The TiO₂, PCL-doped MAPbI₃, and spiro-OMeTAD layers were spin-coated, followed by PVD for the gold electrodes. To observe the effect of the additive on device performance, PCL/MAI/PbI₂ in dimethyl sulfoxide (DMSO)/γ-butyrolactone (GBL) precursor solutions were prepared with 5 different concentrations of PCL: 0.0, 0.1, 0.3, 0.6, and 1.0 mg/mL. UV-Visible Light (UV-Vis) spectrometry revealed comparable spectral shapes and absorption edges, indicating that doping PCL did not compromise the light-absorbing ability of the perovskite layer. Moreover, the PCL additive did not deform the crystal structure of the perovskite, as major peaks were seen in X-ray diffraction (XRD) graphs for all the samples.

The root mean square (RMS) values obtained from atomic force microscopy (AFM) data were much lower for higher concentrations of PCL, suggesting that the additive decreases surface roughness. To further characterize surface morphology, perovskite crystal sizes were compared using scanning electron microscopy (SEM) images. Distributions of grain size show that all films with the PCL additive exhibit much larger grain sizes, and thus fewer grain boundaries (GBs), than pure perovskite film (Figure 2). A smoother perovskite layer with fewer GBs facilitates the diffusion of excitons to their respective transport layers; thus, based on improved surface morphology, the PCL additive is expected to enhance PSC performance.

J-V curves and data from efficiency tests are shown in Figure 3 and Table 1, respectively. PSCs fabricated from perovskite precursor solutions with 0.1 and 0.3 mg/mL PCL exhibited the highest PCE of 10.7%. The peak in efficiency at lower concentrations of PCL and subsequent drop at higher concentrations suggest that while small amounts of additive enhance PCE, too much can decrease cell performance. The increase in PCE of cells with the novel PCL additive suggests its viability as a promising component of future, high-efficiency PSCs.

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Improvement of Perovskite Solar Cell Efficiency through Polylactic Acid Additive
Induced Boundary Passivation

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In recent years, there has been an unprecedented large improvement in the power conversion efficiency (PCE) of perovskite solar cells, which rivals that of established silicon and thin-film cells[1]. Perovskite solar cells have many advantages over established photovoltaic (PV) technologies, such as its low-cost, rapid manufacturing process and its exceptional optoelectronic properties: absorption, photoluminescence, & low charge-recombination rates[1, 2]. Perovskites are hybrid organic-inorganic metal halides with structural formula ABX₃, where A is an organic cation, B is an inorganic cation, and X is a halide anion[2].

In this research, polylactic acid (PLA) was added to the perovskite layer in hopes of improving PCE and perovskite film morphology. We hypothesize that its carbonyl functional groups will exhibit a weak interaction with ionic PbI₂ in solution. This would form a PLA-PbI₂ suspension which could lower the interfacial energy and passivate the crystal grain boundaries. By passivating the crystal grain boundaries the crystal areas could be increased, which reduces the defects in the perovskite layer and in turn increases the PCE.

The fluorine doped tin oxide glass was spin-coated with the TiO₂ precursor solution and then annealed. The perovskite solution was spin-coated in a 2 step process at 1000 and 5000 rpm. Chlorobenzene was added 3 to 4 seconds before the end of the perovskite spinning as an antisolvent to induce crystallization. The Sprio - OMeTAD was spin coated at a constant speed of 4000 rpm, and physical vapor deposition was used to add the gold electrodes. The final device is depicted in Figure 2. Scanning Electron Microscopy (SEM) was conducted to analyze the crystal grain sizes. The crystal sizes increased by up to 226% after the addition of PLA. Using a 2-sample T-test, both the 0.6 mg/mL 16K and 30K molecular weight PLA show a statistically significant increase in the crystal sizes at an alpha level 0.01 (p < 0.00001 for 16k, p < 0.0001 for 30k).

Since both molecular weights had a positive effect on the crystal size, the increase is likely due to an intrinsic chemical interaction between PLA and perovskite. Furthermore, bright spots on the SEM images, which indicate the presence of poor-conducting polymer-based aggregates, suggest an interaction at the interface of the crystals as seen in Figure 3. The increase in intensity of the peak at 15 degrees in the XRD data from the control indicates the presence of PbI₂ and PLA complexes. Finally, turbidity resulted in mixing the clear perovskite and PLA solutions, suggesting the formation of a complex. Yet, UV-Vis spectroscopy and XRD peaks at 14 degree and 28 degree indicate that the addition of PLA did not modify the crystal structure of the perovskite. This supports the hypothesis that the PLA-PbI₂ complexes may act as a surfactant between crystal grains. Furthermore, PLA additives significantly increased the PCE of the PV devices up to 17.5% using the 16K 0.3 mg/mL PLA additive. This is consistent with the unanimous increase in crystal sizes, which is correlated with improved efficiency as seen in Figures 4 and 5. The 16K 0.3 mg/mL devices showed the best PCE of 11.10%. This improvement is likely due to the increased charge transport caused by PLA-PbI₂ complexes passivations at the interfaces of crystal grains. The complexes passivate the crystal boundaries, decreasing exciton dissolution and trapping at the boundaries and redirecting exciton flow in the direction of the useful current. Finally, the improvement through adding PLA varies non monotonically with PLA concentration, suggesting that there is an optimal concentration to achieve maximum efficiency.

These results demonstrate that PLA additives are a promising method to improve the performance of the PV devices. They also give evidence for two mechanisms by which PLA achieves these positive effects: boundary passivation and crystal-interface surfactant.

Session 9: 3D Printing

Chairs: Yuval Shmueli & Zhuolin Xia
Characterization of Electrical and Mechanical Properties of Thermoplastic Poly(Lactic Acid) Filaments Infused with Graphene Nanoplatelets through Fused Deposition Modeling Techniques

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Additive manufacturing through fused deposition modeling (FDM), the most common method of 3D printing, has become a significant manufacturing technology utilized in a wide variety of fields, from engineering to biomedicine. Current 3D printing filaments are commonly composed of poly(lactic acid) (PLA), a biodegradable material synthesized from plant matter that suffers from poor thermal diffusion [1], resulting in structurally weak prints. Recent studies on graphene have revealed its high mechanical strength and useful thermal and electrical conductivity properties. Graphene, when infused in 3D prints, can form an improved high tensile strength conductive polymer with increased structural integrity and application to modern electronics [2]. This study focuses on incorporating graphene nanoplatelets (GNP) within PLA filaments across various concentrations in order to characterize their effects on thermal diffusion, porosity, mechanical strength, and electrical conductivity, all of which can influence the structural integrity and characteristics of a 3D printed product.

To determine optimal conditions for thermal diffusion between filaments, rectangular lattice samples with concentrations of 0%-10% GNP were printed with interfilament widths (IFW) of 0.4 mm-0.5 mm with nozzle extrusion temperatures (NET) of 200°C-245°C on the Lulzbot Mini and Ultimaker 2 Extended+ FDM printers. A FLIR A300C thermal camera recorded thermal diffusion and cooling time to Tg (~65°C) during the printing process (Fig. 1), which was corroborated with Scanning Electron Microscope (SEM) imagery of lattice cross sections. Thermal diffusion was generally optimized at 230°C NET, 10% GNP, and 0.4 mm IFW (Fig. 2-4). 245°C NET tended to cool faster than 230°C due to the larger temperature disparity between the print and the surrounding air. SEM imagery revealed that the graphene platelets were oriented in the same direction as the printed filament, allowing heat to transfer quickly lengthwise within the core of the filament and retaining heat for a longer period of time, explaining why higher graphene concentrations improved thermal diffusion. Smaller IFWs minimized voids, surface area for heat diffusion into air, and distances for diffusion between filaments.

Future studies will utilize small angle X-ray scattering to further characterize the relationship between thermal diffusion and crystallinity of GNP lattices. Circuits of 10%-20% GNP will be printed to study electrical conductivity, and 3-point bending tests will be conducted on samples to study the effect of thermal diffusion and filament composition on mechanical strength. Experimentation will also be continued with boron nitride, another promising material similar in structure to graphene, infused as a nanocomposite with PLA to characterize its applications in 3D printing.

Evaluating the Effects of Material Composition on Molded and 3D Printed Samples

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Poly(propylene) is currently a topic of interest in the 3D printing world -- with better mechanical and thermal properties than the leading filament material, Poly(lactic acid), its difficulty to print due to its semi-crystalline nature holds it back from being used more often.1 poly(propylene) is currently used in living hinges, chairs, electrical packaging, batteries, and textiles due to its chemical resistance, elasticity, and toughness, which further demonstrates its potential in 3D printing.

3D printing has shown to create well-defined structures that could be key in improving the thermal properties of a polymer relative to molded polymers, but a point of interest is improving the thermal, mechanical, and electrical properties of said structures to allow for easier and a greater range of uses. One property that might have an effect on these is the orientation of the filaments in 3D printed samples. With each of their proven abilities to conduct heat and electricity, graphene and boron nitride may also amplify the thermal and electrical conductivity or mechanical strength when mixed with poly(propylene), allowing for easier printing and more uses for 3D printed products.

5% and 10% mixtures of graphene and boron nitride with polypropylene, as well as Ultimaker NFC manufactured polypropylene and pure poly(propylene) filaments were used to create samples. For each composition, three differently-oriented samples were made: one with filaments running vertically through the sample, one with filaments running horizontally, and one that was molded. The Lulzbot Mini printer was used to create the 3D printed samples with a 10mm by 5mm base and a 10mm height with a .4mm filament. The hot press was used at 390°F and with 11,000 pounds of pressure to melt pellets of each material into a mold. The prisms were then placed onto a 100°C stage and recorded using a FLIR A300C thermal camera. The temperature was observed from 0 cm, 2 cm, and 4 cm from the bottom of the sample using the Research IR software. The difference was calculated between the 0 cm and 2 cm positions and the 0 cm and 4 cm positions to give a measurement of thermal conductivity.

In Figure 1, the different orientations of the 5% boron nitride affected the thermal conductivity of the samples. The printed versions of the sample had negative effects on the thermal conductivity, as it slowed down the transfer of heat to the top of the sample, leading to larger differences in temperature. Among the printed samples, horizontal samples allowed for quicker heat transfer than the vertical samples.

Similarly, in Figure 2, the 5% graphene samples follow a similar trend. All three samples began with similar temperature differences between the bottom and 20mm above the bottom, but by the end the molded sample had a noticeably smaller temperature difference, while the horizontally and vertically printed samples ended up with similar results.

Finally, Figure 3 shows a comparison of the thermal conductivity of different composites -- it can be seen that 10% graphene has a noticeably stable temperature difference, staying lower than all of the other samples at all times past the start. Following 10% graphene is 5% graphene, ending up with a slightly higher temperature difference, but getting there at a slower rate. 5% and 10% BN are shown to have better thermal conductivity than pure poly(propylene) -- both commercial and manufactured, but not be as conductive as graphene.

Optimizing 3D Printing of PLA Filaments through Lattice-Boltzmann Modeling and Graphene Nanocomposite Integration

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As a rapidly expanding field, 3D-printing technology has several applications ranging from personal prototyping to industrial manufacturing. Optimal filament composition increases filament fusion, which creates stronger structures and decreases printing time¹. Graphene nanocomposites are a promising material to incorporate in 3D printing due to high thermal conductivity and chemically stable configurations. In addition, modeling is more efficient than traditional lab techniques for running and visualizing combinations of various material compositions². The objective of this study was to derive methodologies to determine optimal strength and efficiency in 3D printing fusion through modeling the effects of graphene molecules on heat diffusion and welding in the polylactic acid (PLA) filaments.

Modeling techniques from Lattice Boltzmann, a class of computational fluid dynamics methods for fluid dynamics, were derived in the C++ based software Palabos v2.0 and visualized in Paraview. The general approach of this study was split into two phases: the first focused upon modeling graphene’s effects on fusion through manipulating coded parameters only. In contrast to polypropylene, PLA has poor crystallization and melts/fuses with greater consistency, making modeling through parameter manipulation an accurate method. This approach is simple in that the graphene effects can be directly modeled through changing relaxation coefficients, thus resulting in faster simulation generation because of minimal requirements for computational power. Upon plotting the advection density of the pure PLA and 10% graphene simulations against the length of filament junctions, graphene incorporation proved to increase heat diffusion (Figure 1).

The second phase involved modeling graphene’s effects through creating actual particles in the fluid flow simulation. This simulation is ultimately more accurate than phase one, but required novel coding techniques to create first simulation of multiple particles moving in a streamlined process through a fluid (Figure 2). Currently, the simulation is successful for two graphene particles, but in future directions, techniques for simulating more particles will be explored.

Ultimately, this study determined graphene is a promising material for improving filament fusion and created a novel technique in particle-fluid flow simulation through successful modeling of multiple 3D printing processes involving graphene nanocomposite incorporation.

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