MTR/REG 621 Cell & Gene Therapy

Spring 2022

Time: Wednesdays 10:15 AM-12:15PM

Dates: Jan 19 - Apr 27

Location: SPE 8-100
8th Floor South Pavilion Extension, 3400 Civic Center Blvd

Instructor Information

Course Director
Michael Milone, MD, PhD
Associate Professor of Pathology and Laboratory Medicine
University of Pennsylvania
milone@pennmedicine.upenn.edu

Assistant Course Director
Elizabeth Hexner, MD, MSTR
Associate Professor of Medicine
University of Pennsylvania
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Course Coordinator
Megan Maxwell
Associate Director, ITMAT Education
215-662-4581, mmaxwell@upenn.edu

General Information

Description
This course will provide students with a general overview of translational research in the area of gene and cell therapy. This includes technical considerations, translating preclinical investigation into therapeutics, the execution of gene and cell therapies clinical trials, and key regulatory issues. Entrepreneurial considerations will be discussed as well. By the end of this course, students will understand the basic technologies employed for gene and cell therapy along with approaches and pitfalls to translating these therapies into clinical applications including regulatory and commercial aspects of this emerging area.

Evaluation Methods:
Students will be graded based on class attendance, participation, a paper presentation and two group project presentations.
30% - Attendance and Class Participation
30% - Paper Presentation
20% - Project Presentation #1
20% - Project Presentation #2

Attendance & Participation (30%)

Attendance:
Students are expected to attend and participate in all classes. If for any reason a student will not be in class, they should contact Dr. Milone (milone@pennmedicine.upenn.edu) or Dr. Hexner (hexnere@pennmedicine.upenn.edu) prior to class to alert them of the absence and make arrangements to make up course content. Two excused absences are allowed
during the course which will not affect the attendance grade. Absences in excess of two will result in 1 point deducted from the attendance portion of the grade.

**Participation:**

Prior to each class, materials will be posted online to the Canvas course site. Students are expected to watch and read the materials and **come to class prepared with to discussion the material**. Each class will involve discussion in which you are expected to participate. Your engagement and participation are important not only for your own learning but also for the learning of others.

**Paper discussion (30%):**

Students will be responsible for reading the assigned papers prior to class. The format of the paper discussion will be a brief introduction of the key background and underlying question(s) followed by an in-depth analysis of the paper by all members of the class. Each student should be prepared to present the brief background if called upon. This background should include:

- What question did the study set out to address?
- What is the significance of the question to the field?
- What was the overall approach used to address the question?

In addition, the students should be able to explain each figure of the assigned paper if called upon during the discussion. Students will be graded based on their introductory presentation and active participation in the figure discussions.

**Project (40% divided equally over two presentations)**

The students will be assigned to a group of 3-4 students. As a group, the students will develop a plan to translate a product based upon a gene and/or cell therapy technology from the basic preclinical phase into a phase I clinical trial. The group will be required to prepare two (2) presentations. The first presentation will focus on assessing the basic science around the technology, the adequacy of the pre-clinical testing as it relates to safety and efficacy prior to entering a human clinical trial and critically evaluating approaches to manufacturing the product appropriate for use in humans. The second presentation will focus on the regulatory and ethical aspects to translating the technology into the clinic. This will include proposing a clinical trial strategy for a first-in-human study with a critical evaluation of the pitfalls and challenges associated with the proposed strategy.

**Course Policies:**

**Academic Integrity:**
As a student at The University of Pennsylvania, you are required to uphold the Code of Academic Integrity. Specifically, this means that materials that you submit either online or in person should be independent works created by you that uphold all tenets of academic integrity (i.e. do not cheat, fabricate, or plagiarize, amongst others). We encourage you to reach out to the course director or coordinator if you are not clear on what potential violations are.

**Canvas:**
All course materials (ppts, announcements, lecture recordings) and assignments will be posted on Canvas. We recommend that you choose the “Notify me right away” option for your most frequently checked email address in the “Announcements” area of the “Notification Preferences” page: [https://canvas.upenn.edu/profile/communication](https://canvas.upenn.edu/profile/communication).

**Course Evaluations:**
Course evaluations are completed via Blue at the end of the semester. These are a required part of course participation. Students can access evaluation forms with their PennKey and password and will also receive emails when forms are available.

**Student Disabilities Services:**
The University of Pennsylvania provides reasonable accommodations to students with disabilities who have self-identified and been approved by the office of Student Disabilities Services (SDS). Please make an appointment to meet with me as soon as possible in order to discuss your accommodations and your needs. If you have not yet contacted SDS, and would like to request accommodations or have questions, you can make an appointment by calling SDS at 215-573-9235. The office is located in the Weingarten Learning Resources Center at Stouffer Commons 3702 Spruce Street, Suite 300. All services are confidential.
## Course Schedule

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<thead>
<tr>
<th>Week</th>
<th>Topic</th>
<th>Lecturer</th>
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<tbody>
<tr>
<td>Jan 19</td>
<td>Overview of Gene and Cell Therapy Field &amp; Orientation to Course &lt;br&gt;Paper discussions (Instructor Lead) Horowitz 1990 HSCT and T cells</td>
<td>Michael Milone, Elizabeth Hexner</td>
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<td>Jan 26</td>
<td>Gene Therapy Vectors &lt;br&gt;Paper Discussions (Student Lead)</td>
<td>Michael Milone</td>
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<td>Feb 02</td>
<td>Gene Editing &lt;br&gt;Paper Discussions (Student Lead) NYCE T cell paper</td>
<td>Jim Riley</td>
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<td>Feb 09</td>
<td>Stem Cells &amp; Tissue Engineering &lt;br&gt;Paper Discussions (Student Lead)</td>
<td>Elizabeth Hexner, Roddy O’Connor</td>
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<td>Feb 16</td>
<td>MAGE-A3 TCR toxicity &lt;br&gt;Correlative Sciences and its Importance to Successful Clinical Trials - Tet2 story &lt;br&gt;Paper Discussions (Student Lead)</td>
<td>Michael Milone</td>
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<td>Feb 23</td>
<td>Pre-clinical Safety Testing of Vector-based Gene Therapies &lt;br&gt;Paper Discussions (Student Lead)</td>
<td>Valder Arruda</td>
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<td>Mar 02</td>
<td>Manufacturing of Complex Cell-based Therapeutics &lt;br&gt;Apheresis/Tour of South Tower cGMP Facility</td>
<td>Bruce Levine, Andrew Fesnak</td>
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<td>Mar 09</td>
<td>No class Spring Break</td>
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<td>Mar 16</td>
<td>Project Presentations - Part 1</td>
<td>Jim Bowen, Kathryn Doyle, PCI</td>
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<td>Mar 23</td>
<td>Patenting Gene &amp; Cell Therapies</td>
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<td>Mar 30</td>
<td>Ethical Considerations in Gene and Cell Therapy Clinical Trials</td>
<td>Megan Kasimatis Singleton (JHMS)</td>
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<td>Apr 06</td>
<td>Overview of Regulatory Review for Gene and Cell Therapies &lt;br&gt;FDA Guidance on Cell &amp; Gene Therapies &lt;br&gt;How-to Workshop: Writing an IND for Gene Therapy Products</td>
<td>Julie Jadlowsky</td>
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<td>Apr 13</td>
<td>Translating Gene &amp; Cell Therapies into Viable Commercial Products</td>
<td>Arun Das</td>
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<td>Apr 20</td>
<td>Final Project Presentations</td>
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<td>Apr 27</td>
<td>Last Day of Class</td>
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