LAB MODULE 4: Controlled Release

In this lab students will investigate how polymeric cross-linked networks can be used to controllably deliver target molecules through diffusion. The relation between cross-linking density and model drug release (bovine serum albumin: BSA) will be highlighted. In addition, students will draw comparisons between diffusion and tether mediated release strategies.

REFERENCES:

1. Lin, C.C. Pharmaceutical Research. 26(3): 631-643. 2009

PRE-LAB REQUIREMENTS:

- 1. Read Lin et al. 2009 in preparation for a brief quiz
- 2. Do prelab calculations (see below)

MATERIALS:

1. Methacrylated poly(ethylene glycol) (PEGDM) (2, 10, and 20kD)

	MW	
PEGDM 2kD	"linear" 2kD PEG with terminal MA functionality	2,172 g/mol
PEGDM	"linear" 10kD PEG with terminal MA functionality	10,172 g/mol
10kD		
PEGDM	"linear" 20kD PEG with terminal MA functionality	20,172 g/mol
20kD		

Figure 1. Methacrylated poly(ethylene glycol)

The precursor molecules that will be used in this lab are all linear PEG molecules of varying molecular weight (i.e. length). We will be using a "linear" 2kD PEG, a "linear" 10kD PEG, and a "linear" 20kD PEG. Each end of these PEG molecules has been terminally functionalized with methyl methacrylate, through which radical initiated cross-linking will occur. As previously performed in lab #1, the synthesis of the mesh-networks will take place by exposing the solution phase precursor mixture to UV light in the presence of a radical generating photo-initiator.

PROTOCOL:

This lab is broken into four sections: (1) in lab synthesis of PEGDM networks, (2) time course collection and replacement of protein containing PBS, (3) end-stage assay for extent of protein release, (4) visual

	Day 1	Day 2	Day 3	Day 4	Day 9
	[protein]	[protein]	[protein]	[protein]	[protein]
PEGDM 2kD					
PEGDM 10kD					
PEGDM 20kD					

Prelab Calculations:

For each PEG MW being investigated (2, 10, 20 kDa), perform the necessary calculations to make a solution that is: 10wt% PEG in PBS + BSA with 10% photo-initiator by volume and a total volume of 250μ L. Assume the PEG does not add significantly to the volume of the solution; assume the PBS+BSA solution and photo-initiator has the same density as plain PBS. Determine the density of PBS through individual research (cite your source). Neatly write up your prelab calculation; it will be collected at the beginning of the lab and will contribute to your grade.

(1) PEGDM Network Synthesis:

- 1. In labs teams calculate and check 10wt% calculations with TA prior to beginning lab. Each lab group must have checked with a TA before receiving the ok to proceed with the lab. Each group will be assigned a specific molecular weight PEGDM to determine protein release. All the group data will be collected at the end of lab and placed on blackboard for the entire class to analyze in their reports.
- Prepare stock solutions: 10wt% PEGDM in <u>PBS+10 mg/mL BSA</u> with 10% photo-initiator by volume; total volume is to equal 250μL. NOTE: the photo-initiator should be added after PEG is dissolved, adding it too early can cause the gel to polymerize in the tube.
- 3. Mix the solution thoroughly to solubilize the PEG (solutions may need to be aggressively vortexed to fully solubilize the PEG).
- 4. Add 40μL of solution to a cylindrical mold (1mL syringe with tip cut off). Run each gel in triplicate at least (n=3). Place molds under UV-lamp for 10min to induce polymerization.
- 5. Once polymerization is complete transfer gels into <u>1mL PBS</u> (exact) in a 24-well plate; label samples with team name as well as PEG molecular weight

(2) Collection of Released Protein: (time course)

- 1. Once the BSA containing hydrogels have been synthesized in lab the PBS "bathing solution" will be periodically collected and replaced over a period of 9 days. This is done to ensure that the surrounding PBS does not become saturated with BSA, thus preventing diffusive release from the hydrogel.
- 2. The BSA containing PBS will be labeled and stored in eppendorf tubes in a -20℃ freezer
- 3. The process of PBS collection and storage will be carried out by the lab TAs

(3) Assay for Protein Release:

- 1. To determine the amount of protein released from each PEG network a Coomassie Protein assay will be performed against a BSA standard curve
- 2. 10µL of standard or unknown sample will be pipetted into a clear 96-well plate

- 3. 300µL of Coomassie Plus reagent will be added to each well in the 96-well plate
- 4. The plate will be covered and allowed to incubate at room temperature for 10min
- 5. Using a plate reader absorbance of the reagent will be measured at a wavelength of 590nm

POST-LAB:

Once the in lab requirements are complete students are required to write a formal lab report summarizing their experimental process and presenting their findings. In addition students must answer the following questions:

- 1. Compare and contrast the key differences between "diffusive release" and "tether mediated" release with respect to controlled drug delivery. Be sure to highlight and discuss rate-limiting steps in each strategy.
- 2. Make a table and graph of the calibration data used to convert absorbance to concentration of BSA in solution. Show the calibration equation used, being sure to discuss its accuracy.
- 3. Make a table and graph the relation between total protein release and time for each of the experimental PEGDM groups. Discuss any trends observed.
- 4. Discuss the mechanism by which BSA was released from the PEGDM hydrogels. Fickian? Kinetic? What are the governing equations that can be used to describe the release, and what are the key parameters?
- Using your experimental data (for each PEG molecular weight) and the governing equation from question 4, determine the diffusion coefficient (D_{BSA,PEG}) for the BSA in each PEG MW hydrogel.
- 6. How is the BSA release profile for each PEG molecular weight affected by the hypothetical meshsize/cross-linking density of the polymeric network?
- 7. Using the Peppas scaling equation

$$\frac{D_{Solute,Gel}}{D_{Solute,PBS}} = 1 - \frac{r}{\xi}$$

where r= the hydrodynamic radius of the solute (r=3.3 nm for BSA), $D_{Solute,PBS}$ is the diffusion coefficient for the solute in solution ($D_{BSA,PBS}$ =6.4*10⁻⁷ cm²/s) and ξ is the mesh size of the gel. Using your D_{BSA,PEG} estimates from 5, calculate the mesh size of each hydrogel [1-2].

- 8. If a larger protein (bigger than BSA) were used to model release from the PEGDM gels how would the release profiles change?
- 9. Discuss at least three parameters that can be altered to obtain different release profiles under diffusive conditions.
- 10. Discuss the advantages and disadvantages of using photoinitiated polymerization to produce polymers/hydrogels for protein delivery and tissue engineering applications.

11. Find one other example (from the literature) where molecular delivery (protein, growth factor, drug, etc.) is accomplished through the application of biomaterials. Briefly explain the delivery strategy used and compare/contrast it to the PEGDM gels.

References

- [1] Peppas, Nicholas A. Hydrogels in Medicine and Pharmacy. Boca Raton, FL: CRC, 1986.
- [2] Pluen, Alain, Paolo Netti, Rakesh Jain, and David Berk. "Diffusion of Macromolecules in Agarose Gels: Comparison of Linear and Globular Configurations." *Biophysical Journal* 77 (1999): 542-52.