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Directed Differentiation of Oligodendrocyte Precursor Cells Using Rationally Designed Solid State Peptide Materials

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Directed Differentiation of Oligodendrocyte Precursor Cells Using Rationally Designed Solid State Peptide Materials Carolyn Scott¹, Sabrina S. Jedlicka^{1,2}

¹Bioengineering Program, ² Materials Science & Engineering

ABSTRACT

Oligodendrocytes are neuroglial cells whose function is to support and myelinate axons in the CNS. Oligodendrocytes have been found to arise from oligodendrocyte precursor cells (OPCs) during late embryogenesis and early post natal development. A single oligodendrocyte can myelinate as many as 40 or more different axons, wrapping the axon with between 20 and 200 layers of highly modified membrane processes¹. The differentiation of OPCs into myelin-synthesizing oligodendrocytes is not well understood, and research suggests that cues for differentiation involve mechanical and chemical signaling from astrocytes and neurons. Many proteins are known to be involved in the migration, proliferation, survival, and differentiation of oligodendrocyte precursors, but their specific roles are not well defined or understood. A better understanding of the mechanism through which these proteins affect the differentiation of OPCs will allow us to more effectively differentiate OPCs to oligodendrocytes, allowing us to better assess the potential for using OPCs as a neurological therapy.

OBJECTIVES

The objective of this study is to determine the effects of 2D silica-based materials containing bioactive extracellular matrix peptides on the differentiation of CG4 oligodendrocyte precursor cells, and to determine the ability of these synthetic peptides to enhance the expression of genes relevant to myelin production.

RESULTS

Proof of Concept

Staining cells on PLO (CNPase and Myelin Basic Protein) show that cells cultured in the absence of growth factors express oligodendrocytic markers

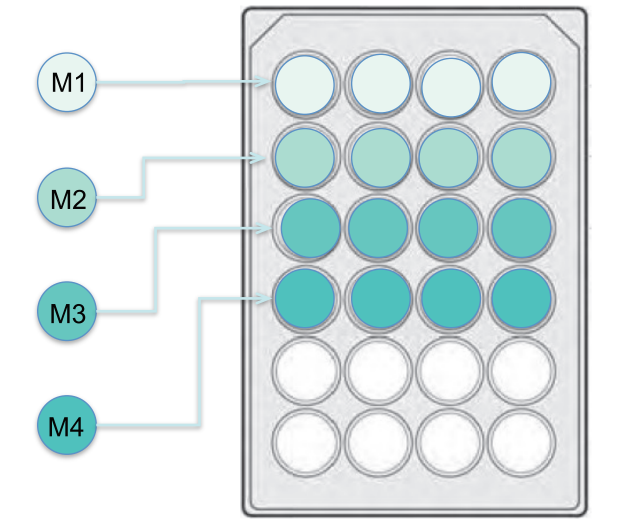
INTRODUCTION

The cells used in this study are CG4s, a bipotential glial cell line capable of differentiating into oligodendrocytes². Various peptide materials are being used to enhance differentiation of CG4 OPCs into mature oligodendrocytes with myelinating capabilities as well as to support mature oligodendrocytes in culture for further study.

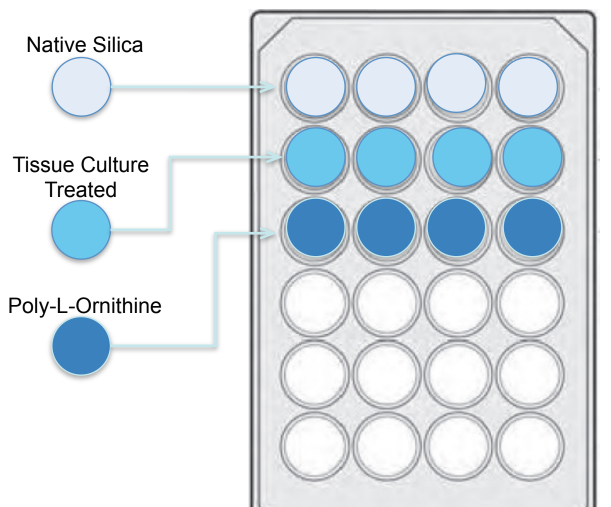
MATERIALS AND METHODS

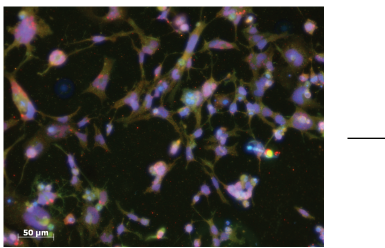
Material Preparation: Dip-coated coverslips in silica-based, peptide solgel materials

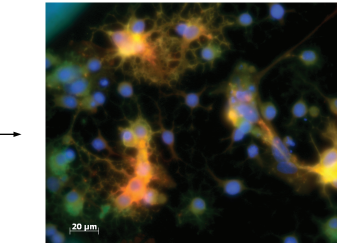
Experimental Material Dip-coated Coverslips



Control Material Coverslips







Cultured with growth factors bFGF and PDGF

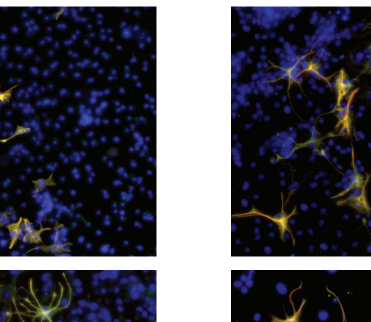
Cultured without growth factors

The images below are CG4 cells cultured on materials for 14 days. Stains for CNPase and Myelin Basic Protein (MBP) have been applied.

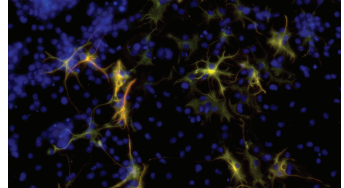
MBP = Red

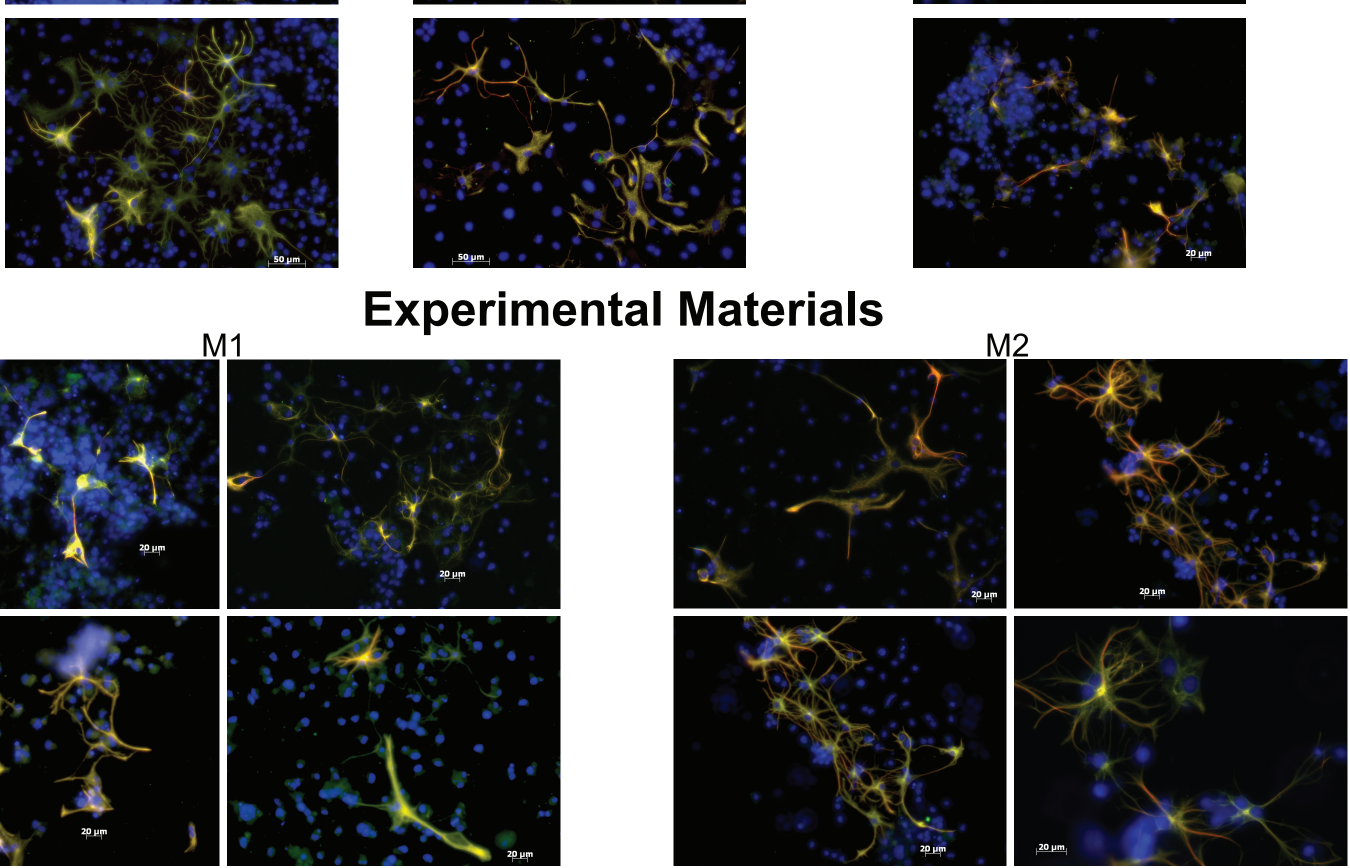
CNPase = Green Negative Control

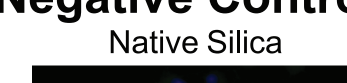
Poly-L-Ornithine Tissue Culture Treated Glass



Positive Controls







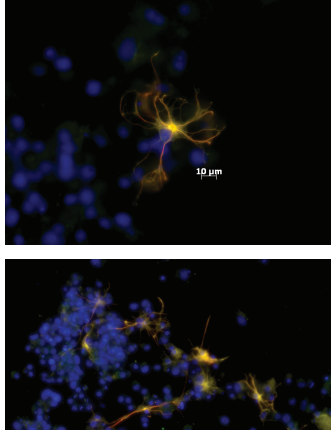


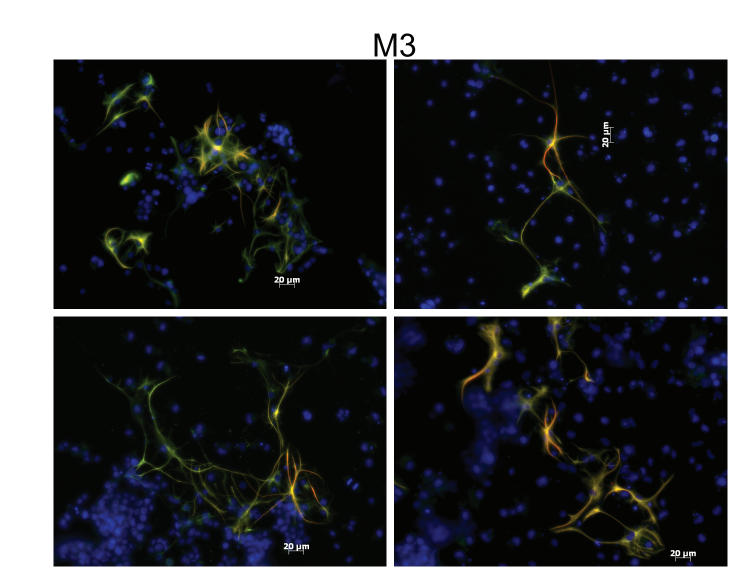
Table 1: Experimental Materials

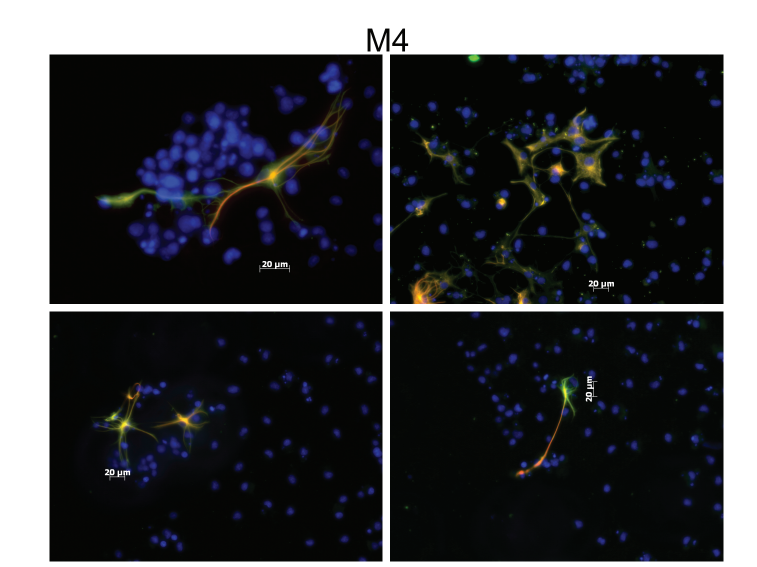
Control Materials Negative Control	Materials	Peptides in Materials
Native Silica	M1	YIG
Positive Control	M2	YIG, AGP
Poly-L-Ornithine	М3	YIG, AGP, SNR
Tissue Culture Treated glass	M4	YIG, AGP, SNR, NID

Table 2: Peptide Amino Acid Sequences		
Peptide	Sequence	
YIG	ACDPGYIGSRGA	
AGP	AGPHSRNAGA	
SNR	ASLVRNRRVITIQG	
NID	ANDNIDPNAVA	
RGD	AYAVTGRGDSPAS	

Cell Culture: CG4 Cells

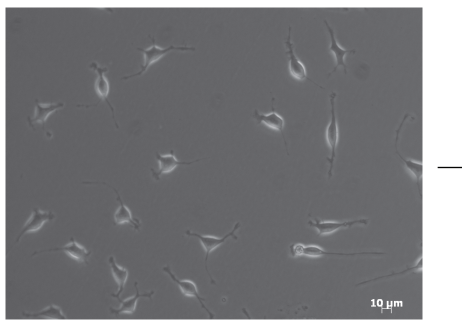
- Cells are cultured on PLO-coated plates supplemented with 10ng/mL bFGF and PDGF
- Cells are seeded on material coverslips, 10, 000 cells/cm³, without growth factor supplements
- Cells fed every 2 days, removing and replacing only half of the media each time so that cells are exposed to the growth factors and signaling

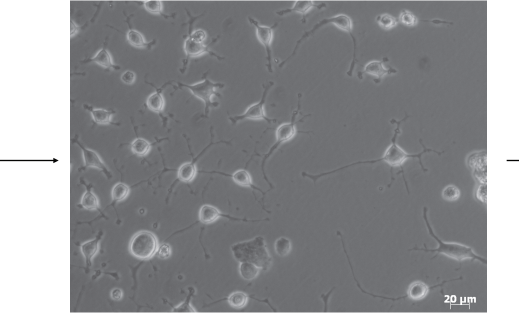




molecules they have secreted

• Cells cultured on materials 10-14 days before fixation





Undifferentiated CG4 cells
Cell Analysis:

Day 3, CG4 differentiation on PLO, induced by growth factor removal

Day 7, Differentiated CG4 cells

Immunocytochemistry used to analyze protein expression of CNPase, Myelin Basic Protein, Proteolipid Protein, Glial Fibrillary Acidic Protein, Actin, and Vinculin (focal adhesion protein)

References

¹ Widmaier, Eric P., Hershel Raff, Kevin T. Strang. *Vander's Human Physiology The Mechanisms of Body Function*. New York: McGraw-Hill Science/Engineering/Math, 2007.

² Louis, J. C., E. Magal, M. Manthorpe, and S. Varon. "CG-4, A New Bipotential Glial Cell Line From Rat Brain, Is Capable of Differentiating In Vitro Into Either Mature Oligodendrocytes or Type-2 Astrocytes." Journal of Neuroscience Research 31 (1992): 193-204.

CONCLUSIONS

- More cells positive for MBP on material 2, containing YIG and AGP, and on material 4, containing YIG, AGP, ANR, NID
- Indicates that these combinations of peptides may enhance oligodendrocyte differentiation

FUTURE WORK

qt-PCR

• Repeat experiment using P19 cell line, another embryonic mouse cell line with a neural-oligodendrocytic potential

