



#### **MGMR** Peptide Delivery

#### MGMR demonstrated to enhance cell penetration of KLA—a cell "impermeable" 1.5kD protein<sup>1,2</sup>

- Proteins are powerful cell-regulators but very challenging for intracellular delivery - Do not efficiently cross the cell's membrane Protein activity is very sensitive and easily
- diminished In this case study: MGMR binds to KLA with high affinity, transports it across the plasma
- membrane, and releases it—triggering apoptosis
- MGMR+KLA peptide produced absolute apoptosis
- MGMR alone has no effect (high tolerability) - KLA alone has no effect (benign externally)

MGMR = Healthy MGMR+KLA = 1Zurita et al. Cancer Research 64, 435-439, 2004 2Yang et al. Methods Mol. Biol. 1266, 29-53, 2015 Apoptotic



**MGMR** Nuclear Staining in Mesenchymal Stem Cells

DAP (30)	l Only 0nm)	MGMR + I (300nn			6
Fluorescence	Bright-field	Fluorescence	Bright-field	Units	50
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		and the state of the second		enc	4(
				Fluoresence	3
				_	2
				Relative	1

DNA-Binding Enzyme (Fluorescent-tagged) Incubated with Stem Cells without MGMR ---NT **Two Views of Same Area:** Bright-field Fluorescence ----MGMR+DAPI (300nM Stem Cells Stem Cells Aggregated Enzymes Remain Outside of Cells Without MGMR for Delivery, DNA-binding **Enzymes Are Not Internalized by Cells** 48 hr 96 hr 24 hr 72 hr **MGNR** Next-generation Transfection Technology How can MGMR<sup>®</sup> improve cell engineering? Eliminates viral-mediated transfection No risk of insertional mutagenesis

Cargo	MGMR™	Viral Vectors	Polymers	Liposomes		
Small Molecules	+	-	+	0		
Peptides	++	+	+	0		
Nucleic Acids	++	++	+	+		
Proteins	++	+	+	0		
ipopolysaccharides	++	-	+	0		Кеу
Log(P) Independent	+	+	-	-	++	Advantage
Properties					+	Compatible
on-immununogenic	+	-	+	+	0	Limited Compatibility
Non-GMO	+	-	+	+	-	Non-Compatible
Simple Optmization	++	-	0	-		
Scalable	++	+	+	-		
Eukaryotes	+	-	0	0		
Prokaryotes	+	-	0	-		

# **MGMR** Versatile Nano-vehicle for Non-viral Cellular Transfection

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### **MGNR**<sup>°</sup> siRNA Delivery



More Dark Space = More Knockdown of Green **Fluorescent Protein** 



 Unlike detergent-based delivery systems, MGMR is non-cytotoxic and compatible across many types of molecular cargo

- No viral toxicity and immunogenicity Non-replicative
- Streamlines manufacturing and quality processes
- Non-biological transfection technology reduces variation - Readily extractable from final cell product
- Eliminates need for new vector development Single transfection vehicle for all cellular gene-editing systems
- Versatile loading profile effective for intracellular delivery of: - Gene-editing complexes
- Oligonucleotides
- Peptides/Proteins
- Improves cell yield
- Reduced toxicity vs. viral or detergent-mediated transfection - Improved cell proliferation







### **MGMR** Supports T-cell Proliferation



#### Endocytotic Trafficking

- MGMR is transported into the cell by active transport
- not damage the cell's external casing MGMR is initially encapsulated into larger
- the interior of the cell molecular cargo
- Once internalized, MGMR is released from vesicles into the intracellular environmen locations inside the cell

#### **MGMR** mRNA Delivery



Unlike other transport systems, MGMR does (plasma membrane) in order to enter inside compartments called vesicles - that traffic into

MGMR is cargo agnostic - it is not prevented from getting inside cells by the size of its

MGMR can deliver cargo to many differen



DNA-Binding Enzyme (Fluorescent-tagged) Loaded on