A Request for Elevating Horae Gene Therapy Center to Department of Genetic & Cellular Medicine

Guangping Gao, PhD, Executive council, 11/21/2024





HORAE GTC HISTORY & MILESTONES







HORAE GTC CURRENT FACULTY, TRAINEE AND STAFF

Faculty members	20
Interns, graduate and postdoc train	nees 48
Research staff	54
Administrative staff	6
	Total: 128





HORAE GTC CURRENT FACULTY & RESEARCH PROGRAMS





Tenured faculty (5)





Terry Flotte, MD Prof

Tenure Track (7)



Heather GradEdwards VMD.PhD, Asst Prof



Allison Keeler PhD, Asst Prof



Prof & Director Prof & Assoc Director

Dan Wang PhD, Asst Prof

Jen Adair, PhD



Phil Tai PhD, Asst Prof



Miguel Esteves, PhD

Assoc Prof

Guocai Zhong PhD, Asst Prof



Jae Shim, PhD

Assoc Prof

Qinglan Ling PhD, AsstProf



Dominic Gessler MD.PhD, Asst Prof

Research Track (8)



Rob Kotin PhD, Aff Prof



Jun Xie

Sylvain Cecchini PhD, AssocProf





Alisha Gruntman PhD, Assoc Prof VMD.PhD, Asst Prof



Rita Batista PhD, Instr



PhD, Instr





Mohan Parsi Hector Benatti PhD, Instr VMD. PhD, Instr



INNOVATIVE INTERDISCIPLINARY SCIENCE IS THE MAIN DRIVE FOR HGTC RAPID GROWTH

- Human genetic diseases and pathomechanisms
- Animal modeling and translational science in large animals
- > DNA and RNA virus biology and vectorology (Ad, AAV & Lenti)
- Gene and genetically modified Cell Therapy platform Technologies
- Gene Therapy immunology and host interactions
- Vector manufacturing & characterization platform technologies
- Pre-clinical proof-of-concept and clinical gene therapies





PIS & RESEARCH PROGRAMS

PI(s)	Primary Research Area	Gene Tx Products in pipeline	Academic Department
Flotte	AAV biology, preclinical & clinical gene therapies, gene therapy immunolo	gy A1ATD, FAOD, Cockayne Syndrome, DMD	Pediatrics
Adair	Lentivirus vectorrology, genetic modified cell therapy, Car-T, gene editing	Cancer, Sickle cell anemia	Medicine
Esteves	CNS & neuromuscular gene therapy, capsid engineering	Dravet, Rett, SPAST, GM1, GM2, LGMD2g, GNAO1, LMNA, TNNT1, NF1	Neurology
Shim	Gene therapy for skeletal disorders Gen	ne therapies for arthritis, FOP-ACVR1, Bone fracture & graft defects	Medicine
Guangping Gao	Gene Therapy in general, vector engineering and platform technologies	Canavan, GM3S, AxD, H-ABC, 4HLCN, MSUD, SMA, ALS	MAPs
GrayEdwards	CNS gene therapy, large animal modeling	Sialidosis-NEU1, GM1, GM2, MSUD	Radiology
Keeler	Gene Therapy and gene therapy immunology	FAOP, AAVCar T cells, gene therapy immunology	Pediatrics
Wang	Gene editing & replacement Tx, stRNA Tx, animal modeling, AAV production platform technology	ogies Hurler, Pitt-Hopskins, NGly1, FoxG1, MSUD, Dysferlinopathy, AARS2D	RNA Therapeutics Inst
Tai	Vector innovation, development, & engineering, Bioinformatics	Novel vectors &QC pipeline, AMD, Glaucoma	MAPs
Zhong	RNA switches for temporally regulatable gene delivery	Ribozyme on- & Aptazyme off- switches for regulated gene therap	by RNA Therapeutics Inst
Ling	Gene Therapy for neurological mitochondrial disorders	MTATP6-related mitochondrial disease, SURF1-&TACO1-related Leigh Syndrome	MAPs
Gessler	Neuro-metabolome, neuro-energetics, CNS Gene therapy	Canavan, AxD, CoX20D	Neurosurgery
Xie	Vector biology, capsid engineering, vector development, gene therapy	H-ABC, 4HLCN, AxD, SMA, ALS	MAPs
Cecchini	Gene Therapy vector process development and large-scale manufacturing	ng Production of toxicology lots for GM2 gene therapy	MAPs
Gruntman	Respiratory biology and Gene therapy	A1ATD	Pediatrics
Kotin	AAV evolution & vector integration safe harbor	HIV vaccine	MAPs
Batista	CNS gene therapy	Stroke, TTR Alzheimer's disease	Neurology
Taghian	CNS gene therapy, neuroimaging, animal modeling	UBA5 disorder	Radiology
Parsi	AAV evolution & vector integration safe harbor	HIV vaccine	MAPs
Benatti	Gene delivery to the CNS and animal modeling	CNS disorders	MAPs.

UMASS CHAN GENE TX PIPELINE

- 14 DISEASE CATEGORIES AND 42 DRUGS IN DEVELOPMENT

													Syndrome	ERCCO	replacement	IVIICE	Esteves				
										Preclinical			Dravet Syndrome	SCN1A	AAV gene replacement	Mice	Sena- Esteves and Wolfe				
Type of Disorder	Disease	Target Gene(s)	Therapeutic Strategy	Preclinical Proof-of- Concept	UMMS Pis	Type of Disorder	Disease Cytochrome C	Target Gene(s)	Therapeutic Strategy	Proof-of- Concept	UMMS Pls		GNAO1- related neurodevelop mental	GNAO1	AAV gene replacement	Mice	Sena- Esteves				
Cardiovascular	Atrial Fibrillation	KCNH2 and CX43	AAV gene addition	Pigs	Donahue and Tai		Oxidase Deficiency	COX20	AAV gene replacement	Mice	Gao	Neurological/	disorder Hypomyelinati								
Connective Tissue Disorder	Dermatosp araxis Ehlers Danlos	ADAMTS2	AAV gene replacement	Mice	Gray- Edwards	Metabolic Diseases	Leigh Syndrome	SURF1	AAV gene replacement	Mice, patient- derived iPSCs	Ling	Epileptic	on with Atrophy of the Basal Ganglia and Cerebellum	TUBB4a	AAV gene silencing AAV gene replacement AAV gene editing	Mice	Xie, Gao				
Infectious Diseases	HIV	N/A	Neutralizing antibodies	NHPs	Xie and Gao		Maple Syrup BCKDHA an Urine Disease BCKDHB	BCKDHA and	AAV gene replacement	Mice, cows, patient-	J. Wang, D. Wang, Strauss,		ang, D. ang, auss,	(H-ABC) Neurofibromat osis	NF1	AAV gene replacement	Mice, pigs	Sena- Esteves			
	Aicardi-		AAV gene addition	Mice	Xie, Gao			Urine Disease	BCKDHB	, tru gene replacement	derived iPSCs	Gray- Edwards,	Gray- Edwards,		MECP2	AAV gene	Mice	Sena- Esteves			
	Goutières Syndrome	ADAR1 and others	AAV gene replacement								Gao	Gao	UBA5 Deficiency	UBA5	AAV gene	Mice	Taghian				
Leukodystrophie s	(AGS) Alexander Disease (AxD)	GFAP	AAV gene silencing	Mice, rats	Xie, Gao		Mucolipidosis IV	COLN1	AAV gene replacement	Mice	Sena- Esteves/ Gray- Edwards	Neuropathy	Hereditary and Sensory Autonomic Neuropathy	SPTLC1	ASO knockdown	Mice	Brown, Watts				
	Canavan Disease	ASPA	AAV gene replacement	Mice	Gessler, Tai, Gao		Sialidosis	NEU1	AAV gene replacement	Mice, sheep (planned)	Gray- Edwards		Type 1				Keeler				
	Acetamino phen-						VLCAD	VLCAD	AAV gene replacement	Mice	Keeler/Flotte		Uveitis	N/A	CAR Treg therapy	Mice	Darren Lee				
Liver Diseases	Induced Acute Liver Failure	miR-375 and others	siRNA	Mice	Xie, Gao	Myopathy	Nemaline myopathy	TNNT1	AAV gene replacement	Mice and sheep ongoing	Edwards/ Sena- Esteves	Ocular	Glaucoma	RhoA and sFasL	AAV gene knockdown	Mice	Gregory- Ksander, Lin,				
	Syndrome	SOX4	AAV gene silencing	Mice	Xie, Gao		TK2 deficiency	TK2	AAV gene replacement	Mice	Xie and Gao	Diseases			AAV gene addition		Rothstein, Tian, Xie,				
	(ALGS) Galactosial idosis	CTSA	AAV gene replacement	Mice	Gray- Edwards		Hereditary Amyotrophic Lateral	C9orf72	ASO knockdown	Mice, sheep, NHPs, 1 human	Brown		Wet AMD	VEGF	AAV gene addition	Mice	Tai, Gao Lin, Punzo, Tai, Xie, Gao				
	GM1 Gangliosid osis	GLB1	AAV gene replacement	Mice, cats, NHPs, human Phase I/II complete	Sena- Esteves and Gray- Edwards		SOD1 ALS	SOD1	AAV artificial miRNA silencing	Mice, NHPs, 2 humans	Brown, Flotte, Gao, Sena- Esteves	Pulmonary Diseases	Alpha-1 Antitrypsin Deficiency	A1AT	AAV gene augmentation AAV gene replacement/knockd	Mice and ferrets	Flotte, Gruntman				
	GM2 Gangliosid oses		A A) (gopo	Mice, cats, sheep,	Sena-	Nourodogoporativ	00011120		Bivalent siRNA	Mice	Brown, Khvorova		Surfactant B Deficiency	proSFTPB	AAV gene addition	Mice	Gruntman, Flotte				
Lysosomal	(Tay- Sachs and	HEXA and HEXB	replacement	human	Gray-	e Diseases			AAV gene silencing	Mice	Xie		Fibrodysplasia		AAV or siRNA gene	Humanized mice,					
Storage Disorders	Sandhoff Disease)				Phase comp			Phase I/II complete	Edwards	Para		Spastic Paraplegia Type SPG4	AAV silencing with gene replacement	Mice, cow (planned)	Sena- Esteves/ Gray-		Ossificans Progressiva	Ossificans ACVR1 Progressiva	silencing	patient- derived iPSCs	Shim
	GM3 Synthase Deficiency	ST3GAL5	AAV gene replacement	patient- derived iPSCs	Gao, Strauss, Wang, Tai,		Spinal Muscular Atrophy (SMA)	SMN1	AAV gene replacement	Mice	Edwards Xie, Gao	Skeletal Diseases	Fibrodysplasia Ossificans	ACVR1	AAV or siRNA gene	Humanized mice, patient-	Shim				
	Hypomyeli nating							STMN2	STMN2 ASO knockdown Rats, 1	Rats, 1	Brown		Progressiva		silencing	derived iPSCs					
	Leukodystr ophy (POLR3-	POLR3b	AAV gene replacement	Mice	Xie, Gao		Sporadic ALS	ATXN2 and others	AAV gene silencing AAV gene replacement	Mice	Xie		Osteogenesis Imperfecta	COL1A1 and COL1A2	AAV gene replacement or editing	Mice	Shim				

Type of Disorder

Disease

Cockay

Target Gene(s)

Therapeutic Strategy

AAV den

Proof-of-

UMMS PIs Batista and

FIRST-IN-HUMAN GENE THERAPY CLINICAL TRIALS AT UMASS CHAN PRIMARILY LED BY TERRY FLOTTE (7 TOTAL)

Vector	Years	Description
AAV1-AAT	2006–2017	Nine Alpha-1 Antitrypsin Deficiency patients were treated in a Phase I trial and a Phase IIa 5-year follow up.
AAV2-RPE65	2007–present	Eight patients were treated in a Phase I trial and two patients were treated in a Phase I/II trial for Leber Congenital Amaurosis.
AAV2-sFlt1	2009–2014	Two patients were treated for AMD.
AAVrh10-antiSOD1miR	2019–2020	Patients with familial ALS caused by <i>SOD1</i> mutations.
AAVrh8-HexA/HexB	2018–2024	Nine patients with Tay-Sachs/Sandhoff disease were treated in a Phase I/II trial and two patients were treated in an expanded access trial.
AAV9-dCas9/VP64-DMD	2022	N of 1 trail for a DMD patient treated with a custom-designed CRISPR therapy.
AAV9-Bi_HexA-HexB	2024 (planned)	N of 1 trial, then 12 patients with Tay-Sachs/Sandhoff Disease.

Treated during this 5-year review period





AAV GENE THERAPY PIPELINE AT TRANSLATIONAL INSTITUTE OF MOLECULAR THERAPEUTICS LED BY MIGUEL ESTEVES







HORAE GTC PRODUCTIVITIES IN TRAINING, RESEARCH & INNOVATION





HGTC TRAINEE AWARDS FROM ASGTC IN THE PAST 5 YEARS

- CURRENT TRAINEES 48

	2020	2021	2022	2023	2024	Totals
Career Development Award	0	1	0	2	0	3
Travel Award	4	*	5	10	9	28
Excellence in Research Award	1	2	2	1	4	10
Outstanding Poster Award	0	0	3	1	**	4

*No Travel Awards were given in 2021 because the meeting was held virtually due to the COVID-19 pandemic. **Not yet selected.





HGTC GRANT AWARDS/REVENUES

- A 30% INCREASE IN THE PAST 5 YEARS WITH \$135 MILLIONS TOTAL PLUS \$43 MILLION PATENT LICENSURE REVENUE IN THE PAST 15 YEARS







HGTC PUBLICATIONS IN THE PAST 5 YEARS

Year	Total Manuscripts Published	Peer Reviewed	Non-Peer Reviewed	Books/ Chapters
2019 (May–Dec)	24	21	3	2
2020	61	52	9	0
2021	53	40	13	0
2022	54	46	8	1
2023	46	36	10	0
2024 (Jan–Apr)	15	14	1	3
Totals	253	209	44	6





HGTC PATENTS IN THE PAST 5 YEARS

Name	Total Patents	Patents Issued	Patents Pending
Dominic Gessler	1	0	1
Guangping Gao	294	97	197
Allison Keeler-Klunk	2	0	2
Jae-Hyuck Shim	12	12	0
Toloo Taghian	14	2	12
Phil Tai	7	7	0
Dan Wang	10	0	10
Jiaming Wang	4	3	1
Jun Xie	15	0	15
Guocai Zhong	2	0	2
Totals (some duplicate)	361	114	232





RESEARCH INSTITUTES AND RESEARCH SERVICE CORES RUN BY HGTC PIS

- Translation Institute of Molecular Therapeutics directed by Miguel Esteves
- Li Weibo Research Institute of Rare Diseases directed by Guangping Gao
- Research Service Cores
 - Respiratory Biology Core to be set up by Alisha Gruntman
 - Transgenic Animal Core directed by Heather Grayedwards
 - Viral Vector Cores
 - Research Vector Core directed by Jun Xie & Guangping Gao
 - Large Scale Vector Manufacturing Core directed by Sylvain Cecchini & Guangping Gao





HGTC VIRAL VECTOR CORE (NON-SUBSIDIZED CORE) REVENUE

- > Doubled in past five years with \$45 million total revenue since funded 15 years ago
- Impacted by recent Biotech/Biopharma contractions
- > Started advertisement for the first time to expand the client base







PERFORMANCE OF HOARE VECTOR CORE IN PAST 5 YEARS







DEPARTMENT OF GENETIC AND CELLULAR MEDICINE WOULD ALLOW US TO

- Grow and better support our faculty
- Expand and strengthen our Interdisciplinary Science
- Promote science and technology innovations
- Increase & diversify grant awards, licensing and core revenues
- Develop gene therapy-focused training programs & funding resources to train scientist and clinician-scientists as future leaders for gene and cell therapy innovation and clinical translation
- Increase opportunities for research collaborations and engagement among faculty members and trainees





Thank You!

Questions?



UMass Chan MEDICAL SCHOOL