GRANT SUPPORT

a) Qu as the PI or subcontract to UB PI

Active:

35. W81XWH1910805(Qu) Role: 9/15/2019-9/14/2022 0.12 calendar

sub-PI

DOD DC to Qu Lab: \$526,068

The Network Biology of Pathogen-Host Interactions Driving Exacerbation in Chronic Obstructive Pulmonary Disease The goal of these studies is to develop a novel proteomics pipeline to procure large dataset for surveying COPD clinical samples for modeling purpose.

34. Center for Protein Therapeutics (Qu) Role: PI 9/1/2019-8/31/2020 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$78,000

A 3D-printed micro-scaffold for MS Imaging and Spatially Resolved Determination of Mab and Receptors in Tissues.

The goal of these studies is to develop a novel micro-scaffold for compartmentalized digestion and sample treatment, to enable reliable MS imaging and to create the density map of drug and targets.

33.GSK Research (Qu) Role: PI 8/1/2019 – 7/31/2021 0.2 calendar

Pharma Research grant DC to Qu Lab: \$200,000

Novel LC-MS strategies for comprehensive in vivo investigations of antibody-drug conjugates and toxicity biomarkers. The purpose of this grant is to develop novel proteomics-based strategy to discover novel proteases in the interstitial space and cellular compartments.

32.AbbVie SRA (Qu) Role: PI 7/1/2019 – 12/31/2020 0.2 calendar

Pharma Research grant DC to Qu Lab: \$140,000

Novel cancer-related proteases. The purpose of this grant is to develop novel proteomics-based strategy to discover novel proteases in the interstitial space and cellular compartments.

31.CA234775 (multiple) Role: PI 12/1/2018 – 11/30/2020 0.24 calendar

NIH DC to Qu Lab : \$110.000

LARGE-SCALE PROTEOME-WIDE ANALYSIS WITH HIGH ACCURACY/PRECISION TO GUIDE PANCREATIC CANCER THERAPY DEVELOPMENT. The purpose of this grant is to develop novel proteomics-based method to quantitatively analyze drug-responsive proteins in PDX models.

30.CA224434 (Qu) Role: Subcontract PI 05/15/2018 - 4/30/2023 0.24 calendar

NIH DC to Qu Lab : \$86,500

GMPS-GMPR AXIS MELANOMA PROGRESSION AND THERAPY. Qu's role is to research on the quantitative interactome method to discovery novel interactor of different GMPR isoforms in various biological systems.

29. Center for Protein Therapeutics (Qu) Role: PI 9/1/2017-8/31/2019 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$78,000 Spatially Resolved Determination of Mab and Receptors in Tissues.

The goal of these studies is to develop novel sample treatment methods and cutting-edge LC/MS techniques to enable ultra-sensitive analysis of bi-specific antibodies and receptors in a spatial manner, to create the density map of drug and targets.

28. Amgen Research grant (Qu) Role: PI 4/6/2017 - 4/5/2018 0.5 calendar

DC to Qu Lab: \$54,044

Membrane Receptors as Potential Therapeutic Target

This project proposes to push the analytical sciences for the general analysis of membrane receptors. We will develop and optimize novel LC/MS-based technologies to quantitatively investigate specific cell surface receptors that may serve as potential drug targets. These molecules are of low abundance and hydrophobic, representing a daunting challenge for current analytical techniques.

27. Al129518 (Zand) Role: Subcontract PI 2/1/2017 - 1/31/2022 0.5 calendar

NIH DC to Qu Lab: \$302,935

Modeling Mechanisms of Adjuvanted Influenza Vaccine Induced IgG Repertoire Diversity and Heterosubtypic Immunity

This project proposes to investigate how a new vaccine, which contains the adjuvant (immune system booster) MF59, increases the range of influenza antibodies binding to molecularly different influenza strains. My lab will use a combination of data from mice and human subjects, combined with mathematical modeling, to test hypotheses about how antibodies that bind different influenza strains arise.

26. U24DK11234 (Adkins) Role: Subcontract PI 1/1/2017 - 2/31/2021 0.5 calendar

NIH DC to Qu Lab: \$120,000

Promotr: A Proteomics Center for Motrpac

The proposed research aims to provide a comprehensive map of the protein "molecular transducers" that transmit the health benefits of physical activity by applying high throughput proteomics technologies. This project will be accomplished by a team and facility with an excellent record of accomplishment applying and developing advanced mass spectrometry-based workflows and pipelines for proteomics research for human health applications. My lab will be responsible for the development of high-throughput LC-MS strategy for method validation.

25. UCB scientific research grant (Qu) Role: PI 12/1/2016 - 6/30/2018 0 calendar

UCB of UK DC to Qu Lab: \$100,000

Urine Metabolite Biomarkers for Renal Fibrosis

The goal of this scientific research grant is to establish a series of di-peptide metabolites for staging renal fibrosis caused by kidney diseases and for evaluating of therapeutic efforts. Novel preparation, treatment and analytical methods will be developed advance this important field.

24. R41 GM121174 (Qu, Aletta) Role: co-Pl 9/1/2016-8/31/2018 0.5 calendar

STTR DC to Qu Lab: \$56,000 (phase-I)

Drug Discovery Platform for Protein Arginine Methyltransferase Inhibitors

The long-term objective of this project is the generation of a universal drug discovery platform based on protein arginine methylation mechanisms involved in human disease.

23. BX002659 VA (multiple) Role: co-Pl 10/1/15-9/30/19 0.6 calendar

Department of Veterans Affairs DC to Qu Lab: \$94,600

Dynamic Remodeling from Reversible Ischemia and Sudden Cardiac Arrest

The central hypothesis of this proposal is that ischemia-induced adaptations resulting from the progression of a coronary stenosis leads to dynamic molecular remodeling that transiently increases the vulnerability to VT/VF during sympathetic activation. My lab employs proteomics technique to characterize the dysregulations during brief ischemia and arrhythmia in swine models.

22. La-Roche scientific research grant (Qu) Role: PI 12/1/2015 - 11/30/2018 0 calendar

Roche-Pharmaceuticals EPBA1902731A17 DC to Qu Lab: \$300,000

A High-Throughput LC/MS Method for Quantification of Biotherapeutics

The goal of this scientific research grant is to push the limit of bioanalytical sciences and develop novel high-throughput, ultra-sensitive and robust methods for targeted protein quantification and address the challenges in biotherapeutics investigation.

Completed

21. Center for Protein Therapeutics (Qu)

Peer-reviewed Industry Consortium Funds

Role: PI 9/1/2016-8/31/2017 0 calendar

DC to Qu Lab: \$158,000

Characterization of Plasma PK and Tumor Penetration of Bi-Specific Antibodies Using LC/MS.

The goal of these studies is to develop novel sample treatment methods and cutting-edge LC/MS techniques to enable ultra-sensitive analysis of bi-specific antibodies, and to investigate the tumor penetration, B-cell and T-cell recruitment, activation and depletion.

20. CTSA (Qu) Role: Pl 7/1/2016-6/30/2017 0 calendar

UB-CTSA Award DC to Qu Lab: \$50,000

Novel Circulating Biomarkers for Sudden Cardiac Death

The goal of this grant is to develop targeted LC/MS methods to validate highly promising candidates predictive of the risk of sudden Cardiac arrest.

19. MCR grant (Qu) Role: PI 6/1/2014 - 5/30/2017 0 calendar

Murdoch Children's Research Institute DC to Qu Lab: \$60,000

Vitamin D and Children's Diabetes

The goal is to provide a seed grant to explore the relationship of Vitamin D level and children diabetes.

18. 12SDG9450036 (Qu) Role: Pl 1/1/2012 - 12/31/2016 2.4 calendar

AHA DC to Qu Lab: \$278,766

Biomarker Release after Reversible Ischemia

The goal is to characterize the cTnl release and modification after reversible myocardial injury in large animal models.

17. U54HD071594 (Qu) Role: Core PI 9/30/2011-8/31/2016 0.6 calendar

NIH DC to Qu Lab: \$202,937

Proteomics, Bioanalysis and Bioinformatics Core-E. New York Pediatric Developmental

Pharmacology Research Consortium.

The goal is to perform proteomics biomarker discovery of ROP in clinical investigations.

16. Athenex contract (Qu) Role: PI 1/7/2010-12/31/2016 0 calendar

Athenex Inc. #641624 DC to Qu Lab: \$393,882

Corporal Research Program with Athenex

The goals of this collaborative research grant are i) support the proteomics efforts to elucidate the action mechanisms of new anti-cancer drug candidates and ii) development of a novel Optimized Photo Affinity cleavable linker (OPAL)-LC/MS/ETD technique to find the smoking-gun evidence of drug-protein binding in vivo.

15. Center for Protein Therapeutics (Qu) Role: PI 9/1/2015-8/31/2016 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$158,000

Characterization of ADC in Tissues and Study of MAb Catabolism/Metabolism.

The goal of these studies is to use cutting-edge LC/MS techniques to investigate the tissue distributions of ADC as well as elucidation of the catabolism mechanisms of SC injection.

14. UB CAT (Qu) Role: PI 7/1/2013 - 6/30/2016 0 calendar

UB DC to Qu Lab: \$6,358

Enrichment Toolkit for Proteomic Biomarkers

The goal is to develop a commercial kit for the analysis of arginine methylation in clinical samples.

13. Merrimack Research Contract (Qu) Role: PI 12/1/2014 - 10/30/2015 0 calendar

Merrimack Pharma DC to Qu Lab: \$22,540

Extensive Investigation of Protein Binding in Liposome Preparations

The goal is to use proteomics to find binding partners novel liposome dosage forms.

12. SUNY Cooperate Fund (Qu) Role: PI 6/1/2014 - 5/30/2015 0 calendar

Center for Hearing and Deafness DC to Qu Lab: \$12,500

Brain Network: Membrane Permeable Transcriptional Regulators for Retinal Repair

The goal is to use proteomics to find biomarkers for retinal cell differentiation.

11. Center for Protein Therapeutics (Qu) Role: PI 9/1/2012-8/31/2017 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$235,000

Highly Accurate and Reliable Quantification of Mab Distribution in Various Tissues.

This project addresses some fundamental challenges for investigation of mAb tissue distribution by a LC/MS-based method.

10. CTSA120077 (Qu) Role: PI 8/1/2010-7/31/2011 0 calendar

UB-CTSA Award DC to Qu Lab: \$50,000

Proteomics Investigation of Laser Micro-Dissected Autopsy Samples from Prostate Cancer Patients

The project aims to develop robust, quantitative, accurate, and sensitive proteomic strategies by which to analyze protein biomarker expression patterns in LMD samples obtained by biopsy of CaP patients.

9. W81XWH-10-1-0728 (Qu) Role: co-PI 10/1/2010-9/30/2012 0.6 calendar

DOD DC to Qu Lab: \$76,651

Gene-Environmental Interactions in Progression of Multiple Sclerosis

The goal is to characterize the relationship between Vitamin D metabolites and progression of multiple sclerosis using a LC/MS-based strategy.

8. Center for Protein Therapeutics (Qu) Role: PI 9/1/2011-8/31/2012 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$79,000

Accurate and Sensitive Quantification of Therapeutic MAbs by Trapping-Micro-LC/MS and Stable-Isotope-Labeled, Full-Length Proteins.

This project seeks to understand the prominent problem of poor absolute accuracy associated with LC/MS-based quantification of therapeutic proteins.

7. Center for Protein Therapeutics (Qu) Role: PI 9/1/2011-8/31/2012 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$79,000

Investigation of Levels of FcRn in Various Tissues.

This project seeks to develop a method for accurate and sensitive quantification of FcRn, a molecule that is critical for the PK of mAb, by an efficient precipitation/on-pellet-digestion method, a Trapping-micro-LC/MS and stable-isotope dilution.

6. Center for Protein Therapeutics (Qu) Role: PI 9/1/2010-8/31/2011 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$79,000

Sensitive and Robust MAb Quantification by Nano-LC/MS.

This project develops and evaluates a novel nano-LC/MS-based method for the investigation of mAb in various pharmaceutical matrices, which is highly sensitive and reasonably robust.

5. Center for Protein Therapeutics (Qu) Role: PI 9/1/2010-8/31/2011 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$79,000

Investigation of Anti-CEA MAb in Various Matrices.

This project employs a LC/SRM-MS-based method for the investigation of target-mediated dispositions of an anti-CEA antibody in various pharmaceutical matrices.

4. PSA-contract (Qu) Role: PI 6/1/2010-5/31/2011 0 calendar

Health Research Inc DC to Qu Lab: \$13,900

PSA-Proteomic Analysis of Rb-Associated Proteins

The fund supports the research of a comprehensive and sensitive method to characterize the sub-proteome pulled by Rb protein in rat models.

3. Center for Protein Therapeutics (Qu) Role: PI 9/1/2009-8/31/2010 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$79,000

Quantification of MAb in Tissues.

This project explores the feasibility of quantifying mAb in tissues using a strong-buffer extraction, a gel-free preparation method and a LC/SRM-MS based analytical strategy.

2 .Center for Protein Therapeutics (Qu) Role: PI 9/1/2009-8/31/2010 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$79,000 Quantitative Characterization of in Vivo Immune Complexes of MAb.

This project seeks to develop a novel method to quantitatively analyze immune complexes in circulation by a Blue Native electrophoresis, followed by in-gel-digestion and nano-LC/MS analysis.

1. Center for Protein Therapeutics (Qu) Role: PI 9/1/2008-8/31/2009 0 calendar

Peer-reviewed Industry Consortium Funds DC to Qu Lab: \$79,000

Ultra-Sensitive Quantification of Cytokines.

This project seeks to develop a ultra-sensitive method for the quantification of cytokines in tissue matrices.

b) Qu as the Co-I (direct cost allocated to Qu lab)

Active:

34. DC016869 (Torregrossa) Role: Co-l 12/01/2018-11/30/2023 0.24 calendar

NIH DC to Qu Lab : \$48,000

Salivary Protein Influence on Taste and Feeding The goal is to develop an IonStar-based strategy to provide novel insights into the effect of proteins on the taste, based on global survey of salivary proteomes.

33. DE027073 (Visser) Role: Co-l 09/01/2018-08/31/2023 0.24 calendar

NIH DC to Qu Lab : \$48,000

THE ROLE OF ORAL SPIROCHETE VIRULENCE FACTORS IN THE IMPAIRMENT OF NEUTROPHIL RESPONSES

The goal is to advance our understanding of spirochete pathogenicity by examining common functionality of Msp proteins across oral treponema species, provide novel insight into the contribution of OMVs and the role of Msp in OMV function and interaction with neutrophils.

32.EY028553 (Farkas) Role: Co-l 12/01/2017 – 11/30/2021 0.24 calendar

NIH DC to Qu Lab : \$76,245

Using Functional Homology of RP1 Isoforms to Guide Alternative Therapeutic Strategies.

Qu's role is to research on the quantitative interactome method to discovery novel interactor of different RP1 isoforms in various biological systems.

31. HL103411(Neelamegham) Role: Co-l 08/04/2017 - 05/31/2021 0.24 calendar

NIH DC to Qu Lab : \$114,502

Systems Biology of Glycosylation

Qu's role is to develop a novel nano-LC/CID/HCD/ETD on a ultra-high-field Obitrap analyzer for more efficient fragmentation of glycosylated proteins in complex biological systems, and to participate in the bioinformatics efforts to elucidate the complex sugar structure.

30. CA204192(Balthasar) Role: Co-l 3/1/2017-02/28/2022 0.5 calendar

NIH DC to Qu Lab: \$20,000

Catch and Release Immunotoxins: CAR-Bombs for Cancer

My role is to develop a LC/MS method to characterize peptide toxin in biological systems.

29. Al125746(Read) Role: Co-l 06/17/2016-05/31/2018 0.2 calenda

NIH DC to Qu Lab: \$23,031

Posttranslational Modification of the Regulatory RNA Binding Protein, ZFP3

My role is to develop a *de novo* method to identify the PTM of ZFP3 in a complex biological system.

28. NS096104(Wrabetz) Role: Co-I 04/01/2016-03/31/2018 0.2 calenda

NIH DC to Qu Lab: \$26,541

Pathogenesis of Myelin Protein Zero Neuropathies in Transgenic Mice.

This study will identify some of the pathological mechanisms, and inform potential therapeutic strategies for hereditary neuropathies.

27. EY019949 (Zhang) Role: Co-l 09/01/2015 – 08/31/2019 0.6 calendar

NIH DC to Qu Lab: \$42,650

ER Stress and Diabetic Retinopathy.

The goal of our project is to identify and harness endogenous protective factors to enhance retinal cell survival and improve vascular function in diabetes mellitus.

26. NS094181 (Park) Role: Co-I 09/15/2015 – 06/30/2020 0.3 calendar

NIH DC to Qu Lab: \$93,500

Transcription Mechanism of Myrf for Central Nervous System Myelination.

This proposal aims to unravel the transcription mechanism of Myrf.

25. RSG-14-214-01-TEB (Zhang) Role: co-l 01/01/2015-12/31/2018 0.1 calenda

American Cancer Society (ACS) DC to Qu Lab: \$44,312

PTPN14 and YAP Tyrosine Modification Regulate the YAP Oncogenic Function

Study focuses on the investigation of mechanisms by which PTPN14 and tyrosine phosphorylation regulate the YAP oncogenic function and how these regulatory interactions further affect tumor formation and metastasis.

24. DE023105 (Yang) Role: co-l 6/1/2014- 5/31/2019 0.5 calendar

NIH DC to Qu Lab: \$64,543

Regulation of Skeletal Development and Homeostasis by Ift Protein

The goal is to dissect the molecular mechanism of IFT80 interactions that confers cilia formation and OB differentiation and function by characterizing IFT80 structural domains, interacting proteins and their functions.

23. AG048388 (Yang) Role: co-l 8/1/2014- 5/30/2019 0.5 calendar

NIH DC to Qu Lab: \$72,513

Function of Regulator of G Protein Signaling in Aging Skeleton

The goal of this project is to discover the role and mechanism of RGS12 in OC differentiation and activation in pathologic age condition, and provide new and more effective therapeutic targets to age-associated osteoporosis and other bone diseases.

22.AR066101(Yang) Role: Co-I 5/1/2014-4/30/2019 0.6 calendar

NIH DC to Qu Lab: \$71,320

Role of Rgs12, A Regulator of G Protein Signaling, In Bone Remodeling

Qu's role is to develop proteomics strategies for characterization of RGS12-related pathways and sub-proteomes.

Completed

21. HD075363 (Feltri) Role: co-l 9/28/2012- 9/1/2017 0.5 calendar

NIH DC to Qu Lab: \$28,500

Subcellular Domains of Myelinating-Glia: Capturing Axonal Contact

The goal is to study the proteomes of subcellular domains of neuron development.

20. CHE-1412405 (Hevel) Role: Co-I 8/1/14-7/31/17 0.6 calendar

NSF DC to Qu Lab : \$41,521 Collaborative Research: Protein Arginine Methylation

The goal of this grant is to investigate the specific PRMT specificity and activity on methylating arginine residues in cells.

19. SUNY Cooperate fund (multiple) Role: Co-l 6/1/2014 - 5/30/2015 0 calendar

Center for Hearing and Deafness DC to Qu Lab: \$12,500

Brain Network: Membrane Permeable Transcriptional Regulators for Retinal Repair
The goal is to provide a seed grant to explore the relationship of Vitamin D level and children diabetes.

18. Roche Research Grant (Balthasar) Role: Co-l 6/1/2014 - 5/30/2016 0 calendar

Hoffmann-La Roche Inc DC to Qu Lab: \$73,500

Investigation of the Utility of LC/MS for Characterization of the Plasma and Tissue PK of A Novel Series of Anti-CEA Monocolonal Antibodies.

The goal is to provide a seed grant to explore the relationship of Vitamin D level and children diabetes.

17. EY025061(Zhang) Role: co-l 12/2/2014- 12/1/2016 0.5 calendar

NIH DC to Qu Lab: \$34,507

Study of the ER-Mitochondria Interface as A New Target in Diabetic Retinopathy
The overall goal of this pilot study is to establish a role of MAM in retinal cell metabolism in diabetes

16. HL103411(Neelamegham) Role: Co-l 7/1/2011-6/30/2016 0.6 calendar

NIH DC to Qu Lab: \$101,370

Systems Biology of Glycosylation

Qu's role is to develop on a dual-enzyme-digestion and nano-LC/CID/HCD/ETD method for more efficient fragmentation of glycosylated proteins in complex biological systems, and to participate in the bioinformatics efforts to elucidate the complex sugar structure.

15. Al060260 (Read) Role: Co-l 7/1/2010-6/30/2015 0.6 calendar

NIH DC to Qu Lab: \$74,800 Protein Arginine Methylation in Trypanosomes

Qu's role is to develop and employ a dual-enzyme/activation, high-resolution SCX fractionation and nano-LC/MS strategy for the global identification of methylation proteins in the proteomes of trypanosomes.

14. DA023223 (Morris) Role: Co-l 7/1/2012-6/30/2017 0.6 calendar

NIH DC to Qu Lab: \$18,110

Gamma-hydroxybutyrate: Toxicokinetics, Toxicodynamics and Treatment Strategies Qu's role is to develop an ultra-sensitive and accurate targeted nano-LC/MS strategy for the quantification of multiple drug transporters in animal models.

13. GM073646 (Blanco) Role: Co-l 7/1/2010-6/30/2014 0.6 calendar

NIH DC to Qu Lab: \$11,240
Pharmacogenetics of Human Carbonyl Reductases

Qu's role is to employ a nano-LC/MS method developed by Qu lab in 2008 to quantify CBR1 and CBR3 in human tissues.

12. HL61610 (Canty) Role: Co-l 9/1/06 - 8/31/15 1.2 calendar

NIH DC to Qu Lab: \$54,750

Metabolic Adaptation and Functional Recovery of Hibernating Myocardium

Qu's role is to discover the tissue biomarkers for hibernating myocardium and remodeling.

11. R21 HD075363 (Feltri) Role: Co-I 9/1/2012-8/31/2014 0.6 calendar

NIH DC to Qu Lab: \$26,877

Subcellular Domains of Myelinating-Glia: Capturing Axonal Contact

Qu's role is to discover the biomarkers that are responsible for the neuron cell differentiation and the development of pseudopods, by the development and optimization of an ion-current-based method.

10. 1051350 (Yu) Role: Co-l 4/1/2011-3/31/2015 0.6 calendar

NSF DC to Qu Lab: \$16,500

The Role of Protein Arginine Methylation in the Co-transcriptional Recruitment of premRNA Splicing Factors

Qu's role is the use of CID/ETD nano-LC/MS methods to determine the methylproteins and localize the exact methylation sites on key proteins pulled-down by TAP procedures.

9. R21 DA027528(Multi) Role: Co-Pl 9/1/2009-8/31/2011 1.2 calendar

NIH DC to Qu Lab : \$50,421

Peripheral Biomarkers of Cocaine Dependence and Relapse

Qu's role is to design and execute the proteomics studies for the discovery of brain biomarkers for cocaine addiction and withdrawn.

8. R03 CA139562 (Mojica) Role: Co-l 4/1/2009-3/31/2011 1.2 calendar

NIH DC to Qu Lab : \$56,700

Identification of Colon Cancer Protein Biomarkers in the Blood

Qu's role is to design and execute the proteomics studies for proteomics comparison of the normal and cancerous epithelial cells enriched from clinical samples.

7. NS045630 (Feltri) Role: sub-award PI 8/1/2011-7/31/2012 0 calendar

NIH DC to Qu Lab : \$12,000 Laminin Receptors and Signals in Schwann Cells

Qu's role is to develop a method to discover the biomarkers for the neuron cell differentiation.

6. Al085569 (Schwartz) Role: sub-award Pl 7/1/2009-6/30/2012 0.6 calendar

NIH DC to Qu Lab : \$56.526

Integration of Clinical, Genomic And Proteomic Data Using A Bioinformatic Approach Qu's role is to develop a proteomics strategy to compare the PBMC proteomes from NP and LTNP HIV patients.

5. EY007361 (Fliesler) Role: sub-award PI 3/15/2010-12/31/2010 0 calendar

NIH DC to Qu Lab: \$18,000

Isoprenoid Metabolism in the Retina.

Qu's role is to develop a proteomics and bioinformatics method to elucidate the mechanisms of retina degeneration in a SLOS model.

4.1S10RR024521(Straubinger) Role: Co-l 4/1/2009 – 3/31/2010 0 calendar

NIH/NCRR DC to Qu Lab : \$0

High Performance Computational System to Support LCMS/Proteomics Analysis

Funds the purchase of a state-of-the-art computational cluster to accelerate proteomics analysis and provide mass storage for large datasets, for the proteomics facility Qu is currently running.

3. GM073646 (Blanco) Role: Co-l 3/1/2005-2/28/2010 0 calendar

NIH DC to Qu Lab : \$9,000

Pharmacogenetics of Human Carbonyl Reductases

Qu's role was to develop a highly sensitive and reliable method for the quantification of CBR enzymes in livers.

2. 1S10RR021221(Straubinger) Role: Co-l 04/01/2005-03/31/2006 0 calendar

NIH DC to Qu Lab: \$ 0

LC/Quadrupole Ion Trap Mass Spectroscopy System

Funds a state-of-the-art ion-trap LC/Linear Trap Quadrupole instrument for peptide sequencing and drug metabolite characterization.

1. 1S10RR023650(Straubinger) Role: Co-l 04/01/2007- 03/31/2008 0 calendar

NIH DC to Qu Lab: \$0

High Sensitivity Liquid Chromatography Tandem Mass Spectrometry System

Funds a state-of-the-art ultra-sensitive LC/MS instrument for drug and proteomic analysis.