

**BIOGRAPHICAL SKETCH**

Provide the following information for the Senior/key personnel and other significant contributors.  
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NAME: Ortlund, Eric Anthony

eRA COMMONS USER NAME (credential, e.g., agency login): EORTLUND

POSITION TITLE: Professor of Medicine

EDUCATION/TRAINING *(Begin with baccalaureate or other initial professional education, such as nursing, include postdoctoral training and residency training if applicable. Add/delete rows as necessary.)*

INSTITUTION AND LOCATION	DEGREE (if applicable)	Completion Date MM/YYYY	FIELD OF STUDY
University of South Carolina	B.S.	05/1997	Chemistry
University of South Carolina	Ph.D.	05/2002	Chemistry, Biochemistry
University of North Carolina – Chapel Hill	Post-Doc	06/2007	Biochemistry Structural Biology

**A. Personal Statement**

Our laboratory is keenly interested in macromolecular signaling within the context of normal and disease-state cellular functions. We leverage both structural and biochemical studies to assess protein/RNA/DNA function and enable structure-guided studies to assess function in the context of metabolism, stress, homeostasis and disease. We are excited to leverage our expertise to tackle fundamental questions centered SARS-Cov2 variant risk. I have a strong track record of collaboration, which has led to numerous contributions to the protein signaling community. Over the past 12 years, we have become interested not only in how ligands directly modulate gene expression, but how discrete lipid mediators and soluble metabolites drive signaling pathways leading to cellular- and organelle-specific effects, such as inflammation and changes in metabolism.

In addition to acting as PI of a laboratory in the Biochemistry Department, I serve as Director for Emory's Integrated Lipidomics Core. Lead and participate in several consortia efforts to generate a systems-level understanding of metabolism, immune function and disease. Our lab is part of the NIH funded Rapid Acceleration of Diagnostics, or RADx center. We are on the [Variant Task Force](#) and our object is to assess the threat of variants with respect to their ability to evade detection in rapid COVID tests and to evade detection by neutralizing antibodies. For this effort, we work closely with members of the Variant Task Force which includes members from Emory University and the University of Washington and works with the NIH, FDA, DOD, CDC, DOE, and BARDA. We are funded to map epitopes on the spike protein for commercial antibodies and our proposed project also involves antibodies derived from Indian donors provide through a collaboration with Dr. Rafi Ahmed (Emory University, National Academy of Sciences, Emory Vaccine Center).

Emory has made large investments to build infrastructure and expertise in cryoEM. We are leveraging these resources by taking advantage one-on-one training and cryoEm workshops available through our core (and nationally). We hold weekly meetings with the cryoEM core and other four labs on campus using these techniques and host bi-weekly presentations where students and trainees present their current research centered on single particle cryoEM or tomographic techniques. To achieve high-resolution, we desperately need access to the resources and expertise offered by the NCCAT!

Ongoing and recently completed projects that I would like to highlight include:

1R01DK115213 (MPI)

Ortlund (Contact PI); Calvert and Jui (mPI)

07/01/2017 - 06/30/2022 NIH

Targeting LRH-1 with Small Molecule Modulators

1U24DK112341-01 (MPI)  
Ortlund (contact mPI); Fernandez (MPI)  
09/01/2016 – 08/31/2022  
Georgia Comprehensive Metabolomics and Proteomics Unit for MoTrPAC

HHSN272201300018/HHSN27200009  
Rouphael/ Anderson (PI)  
09/01/2015 – 01/03/2023  
NIH Vaccine Treatments Evaluation Unit (VTEU); Consultation and 'omics testing of clinical samples

R01 DK103046  
Cohen (PI); Ortlund (MPI)  
06/01/2020 – 05/31/2025  
Them1-Mediated Metabolic Regulation and Pathogenic Role in NAFLD

#### Citations:

1. Suthar, M.S., Zimmerman, M., Kauffman, R., Mantus, G. Linderman, S, Hudson, W.H., Vanderheiden, A, Nyhoff, L., Davis, C., Adekunle, S., Affer, M., Sherman, M., Reynolds, S., Verkerke, H., Alter, D., Guarner, J., Bryksin, J., Horwath, M., Arthur, C., Saakadze, N., Smith, G., Edupuganti, S., Scherer, E., Hellmeister, K., Cheng, A., Morales, J., Neish, A., Stowell, S., Frank, F., **Ortlund, E.A.**, Anderson, E., Menachery, V., Rouphael, N., Mehta, A., Stephens, D., Ahmed, R., Roback, J., Wrammert, J., Rapid generation of neutralizing antibody responses in COVID-19 patients, *Cell Reports*, 2020, May 8:2020.05.03.20084442. PMCID: PMC7276302
2. Tillman, M.C., Imai, N., Li, Y., Khadka, M., Okafor, C.D., Juneja, P., Adhiyaman, A., Hagen, S.J., Cohen, D.E., **Ortlund, E.A.**, Allosteric regulation of Thioesterase Superfamily Member 1 lipid sensor domain by free fatty acids and lysophosphatidylcholine, *Proceedings of the National Academy of Sciences U S A*, 2020, 117(36):22080-22089, PMCID: PMC7486800
3. Xu, X., Wang, Y., Gutierrez, G.S., Damsker, J.M., Nagaraju, K., Hoffman, E.P., **Ortlund, E.A.**, "Disruption of a key ligand-H-bond network drives dissociative properties in Vamorolone for Duchenne Muscular Dystrophy Treatment", 2020, *PNAS*, Sep 29;117(39):24285-24293

## B. Positions, Scientific Appointments, and Honors

### Positions and Scientific Appointments

2002-02	Postdoctoral Fellow, Laboratory of Lukasz Lebioda; Dept of Chemistry, University of South Carolina, Columbia, SC
2003-07	Postdoctoral Fellow, Laboratory of Matthew Redinbo; Department of Chemistry, University of North Carolina, Chapel Hill; Lineberger Comprehensive Cancer Center, UNC
2007-2009-	Associate Professor; Department of Biochemistry, Emory University, Atlanta, GA
	Member, Executive Committee, Molecular Systems Pharmacology Graduate Program
2016-2018	Founder and Director of Emory Integrated Lipidomics Core
2019-	Founder and Director of Emory Integrated Metabolomics and Lipidomics Core
2010-14	PC4 and PC2 study section, American Heart Association
2012	RFA study section NIH Blood and Vascular Branch, National Heart, Lung, and Blood Institute, NIH
2012-13	French National Research Agency Grant Reviewer
2012-13	Foundation for Polish Science Grant Reviewer
2014	NIH Molecular and Cellular Endocrinology study section, NIH, <i>Ad hoc</i>
2012-	Editorial Board, ASPET, Molecular Pharmacology
2012-	Reviewer Board, Journal of Pediatric Biochemistry
2013-	Editorial Board, Nuclear Receptor Research
2013-	Editorial Board, Hepatology
2015	NIH LCMI Special Emphasis Panel/Scientific Review Group

2015	NIH Special Emphasis Panel ZRG1 EMNR-R(56), Cellular Mechanisms of Metabolism and Obesity
2016	Austrian Science Fund, study section
2016-	NIH Molecular and Cellular Endocrinology study section, Chair
2017	NIH ZRG1 EMNR A 07, Molecular and Cellular Endocrinology, Chair

### **Honors and awards**

2000	Bayer Corporation Award for Excellence in Chemistry, 2000
2002	College of Science and Mathematics Dissertation Fellowship
2004	Lineberger Comprehensive Center NIH NSRA Postdoctoral Fellowship
2013	American Crystallographic Association Margaret C. Etter Early Career Award
2013	Albert E. Levy Scientific Research Award for Junior Faculty
2013	Department of Biochemistry, Emory University Outstanding Teaching Award
2015	Awarded the W. M. Keck Foundation Medical Research Grant
2017	Postdoctoral Office "One in a hundred" Mentor of the Year Award
2017	"Special Recognition for Outstanding Research" Emory University School of Medicine

## **C. Contributions to Science**

### **1. Lipidomics and Lipid Signaling**

Leveraging our expertise in lipid structure, biochemistry and signaling we founded the Emory Integrated Lipidomics Core which uses cutting edge LC-MS/MS based approaches to identify and quantify lipids biological samples. We have collaborated with many groups at Emory and abroad to understand how lipids drive and are modulated both normal biology and disease. We have developed robust quantitative methods to identify and quantify bioactive lipids (e.g. oxylipins, cannabinoids), bile acids, short and long chain fatty acids, carnitines, CoAs and polyunsaturated lipids. We have also developed high-resolution methods for untargeted lipidomics leveraging scheduled MS/MS methods.

- a. Sanford, J.A., Nogiec, C.D., Lindholm, M.E., Adkins, J.N., Amar, D., Dasari, S., Drugan, J.K., Fernández, F.M., Radom-Aizik, S., Schenk, S., Snyder, M.P., Tracy, R.P., Vanderboom, P., Trappe, S., Walsh M.J.; Molecular Transducers of Physical Activity Consortium. Molecular Transducers of Physical Activity Consortium (MoTrPAC): Mapping the Dynamic Responses to Exercise, *Cell*, 2020, Jun 25;181(7):1464-1474 \***Ortlund, E.A.** part of MoTrPAC Consortium Research and Writing group
- b. Maner-Smith, K.M., Goll, J.B., Ford, D.A., Jensen, T.L. Khadka, M., Colucci, J.K., Gelber, C.E., Albert, C.J., Bosinger, S., Franke, S., Natrajan, M., Rouphael, N., Johnson, R., Zanz, P., Anderson, E.J., Hoft, D.F., Mulligan, M., **Ortlund, E.A.**, Alterations in the human plasma lipidome in response to Tularemia vaccination. *Vaccines*, 2020, Jul 24; 8(3):E414 PMID: PMC7564507
- c. Mhadka, M., Todor, Andrei, A., Maner-Smith, K.M., Colucci, J.K., Tran, ViLinh, T., Gaul, D.A., Anderson, E., Natrajan, M., Rouphael, N., Mulligan, M.J., McDonald, C., Suthar, M., Li, S., **Ortlund, E.A.**, The effect of anticoagulants, temperature and time on the human plasma metabolome and lipidome from healthy donors as determined by liquid chromatography-mass spectrometry. *Biomolecules*, 2019, 23:9(5)
- d. Cordy R.J., Lili L, Cabrera-Mora M, Chien J-T, Meyer E.V.S., Lapp S.A., Joyner C.J., Banton S., Tran V., Luvira V., Rungin S., Saeseu T., Rachaphaew N., Garcia A., Khadka M., Pakala S., Tharp G., MaHPIC Consortium, DeBarry J.D., Kissinger J.C., Bosinger S., Li S., **Ortlund, E.A.**, Jones D.P., Sattabongkot J., Uppal K., Barnwell J.W., Patrapuvich R., Moreno A., Galinski M.R., Distinct amino acid and lipid perturbations characterize acute versus chronic malaria. *JCI Insight*, 2019, ePub 10.1172/jci.insight.125156 PMID: PMC6538326

### **2. Phospholipid-regulated gene expression**

During my postdoctoral training, I tackled the characterization of liver receptor homologue -1 (LRH-1), an orphan nuclear receptor involved in glucose homeostasis, reverse cholesterol transport, lipid absorption, hormone synthesis, and cell proliferation. My structural and functional studies identified phospholipids as *bona fide* LRH-1 ligands, identifying LRH-1 as the first mammalian phospholipid-sensing transcription factor. We have continued to pursue this challenging project attempting to link phospholipid sensing with glucose, lipid, and bile acid homeostasis. We have recently established a collaboration with Dr. David Cohen (Harvard

Medical School/BWI) centered on the action of START domain containing proteins and their connection to phospholipid-driven gene regulation. Together, we have discovered that PC-TP stimulates LRH-1 transcriptional activity making PC-TP a critical player in the phosphatidylcholine signaling pathway that impacts glucose homeostasis. Thus, we are well positioned to elucidate the molecular mechanisms that drive the unique biology of phospholipid to control metabolism.

- a. **Ortlund EA**, Lee Y, Solomon IH, Hager JM, Safi R, Choi Y, Guan Z, Tripathy A, Raetz CR, McDonnell DP, Moore DD, Redinbo MR. Modulation of human nuclear receptor LRH-1 activity by phospholipids and SHP. *Nat Struct Mol Biol.* 2005;12:357-63. PMID: 15723037
- b. Lee JM, Lee YK, Mamrosh JL, Busby SA, Griffin PR, Pathak MC, **Ortlund EA**, Moore DD. A nuclear-receptor-dependent phosphatidylcholine pathway with antidiabetic effects. *Nature.* 2011;474:506-10 PMID: PMC3150801.
- c. Musille PM, Pathak MC, Lauer JL, Hudson WH, Griffin PR, **Ortlund EA**. Antidiabetic phospholipid-nuclear receptor complex reveals the mechanism for phospholipid-driven gene regulation. *Nat Struct Mol Biol.* 2012;19:532-7. PMID: PMC3960984.
- d. Musille PM, Pathak M, Lauer JL, Griffin PR, **Ortlund EA**. Divergent sequence tunes ligand sensitivity in phospholipid-regulated hormone receptors. *J Biol Chem.* 2013;288:20702-12 PMID: PMC3711333.

### 3. Phospholipid transport and signaling

Since 2006, I have studied how phospholipids are sensed and signal through interaction with the Sec14 family of proteins. This has largely been a collaborative effort between our lab and Dr. Vytas Bankaitis (Texas A&M Health Science Center). I was responsible for discovering the molecular mechanism driving both phosphatidylcholine and phosphatidylinositol recognition in the protein family. We were able to develop mutant variants with selective disruptions in their ability to transport either PC or PI and tie specific lipid shuttling abilities to cell functions such as proliferation, vesicle maturation, and protein synthesis. We have extensive experience in manipulating the phospholipid transfer ability of most known soluble phospholipid transport proteins in both yeast and mammals and are eager to apply this knowledge to understand the mechanisms that underlie exercise adaptation.

- a. Schaaf G, **Ortlund EA**, Tyeryar KR, Mousley CJ, Ile KE, Garrett TA, Ren J, Woolls MJ, Raetz CR, Redinbo MR, Bankaitis VA. Functional anatomy of phospholipid binding and regulation of phosphoinositide homeostasis by proteins of the sec14 superfamily. *Mol Cell.* 2008;29:191-206. PMID: 18243114.
- b. Schaaf G, Dynowski M, Mousley CJ, Shah SD, Yuan P, Winklbauer EM, de Campos MK, Trettin K, Quinones MC, Smirnova TI, Yanagisawa LL, **Ortlund EA**, Bankaitis VA. Resurrection of a Functional Phosphatidylinositol Transfer Protein from a Pseudo-Sec14 Scaffold by Directed Evolution. *Mol Biol Cell.* 2011;22:892-905. PMID: PMC3057712.
- c. Ren J, Pathak MC, Temple BRS, Lin C, Nile AH, Mousley CJ, Duncan MC, Eckert DM, Leiker TJ, Ivanova PT, Meyers MS, Murphy RC, Brown HA, Verdaasdonk J, Bloom KS, **Ortlund EA**, Bankaitis VA A Phosphatidylinositol Transfer Protein Integrates Phosphoinositide Signaling With Lipid Droplet Metabolism To Regulate a Developmental Program of Nutrient Stress-Induced Membrane Biogenesis. *Molecular and Cellular Biology.* 2014;25(5): 712-727. PMID: PMC3937096
- d. Flynn, A.R., Mays, S.G., **Ortlund, E.A.**, Jui, N.T., Development of Hybrid Phospholipid Mimics as Effective Agonists for Liver Receptor Homolog-1, *ACS Med Chem Lett.*, 2018, 9(10):1051-1056. PMID PMC6187417

### 4. Fatty acid signaling

We have studied the structure and mechanisms governing the function of soluble lipid transport proteins, which are able to bind specific lipids and deliver them to the nucleus for signaling through the nuclear hormone receptors PPAR $\beta/\delta$ . Our most recent contribution couples soluble lipid shuttling proteins to nuclear receptor-mediated gene activation. Our group identified a ligand-sensitive tertiary nuclear localization signal (NLS) on fatty acid binding protein 5 (FABP5). This molecular switch senses fatty acid type and oxidation status, allosterically relaying the information to the NLS. This serves as the driving force for fatty acid-specific nuclear translocation. Ultimately, we showed that FABPs provide a second level of regulatory control over nuclear receptor-mediated lipid signaling offering tremendous potential to modulate nuclear receptor

signaling in a tissue-specific way. Our most recent efforts center on the ability of FABPs to relay signals between tissue and cellular organelles to control metabolism and longevity.

- a. Armstrong E, Noy N, **Ortlund EA**. Structural basis for ligand regulation of the FABP5 – PPAR signaling pathway. *Journal of Biological Chemistry*. 2014; 289: 14941-14954. PMCID: PMC4031543.
- b. Folick A, Oakley HD, Yu Y, Armstrong EH, Kumari M, Sanor L, Moore DD, Zechner R, **Ortlund EA**, Wang MC. Lysosomal Signaling Molecules Regulate Longevity in *Caenorhabditis elegans*. *Science*. 2014;347(6217):83-6. PMCID: PMC4425353.
- c. Steensels, S., Qiao, J., Zhang, Y., Maner-Smith, K. M., Kika, N., Holman, C. D., Ivanova, A. A., Corey, K., **Ortlund, E. A.**, Ersoy, B. A., Acot9 promotes hepatic steatosis by trafficking short chain fatty acids towards lipogenesis and hepatic glucose production. *Hepatology*, 2020, June 4, Online ahead of print PMID: 32498134
- d. Alves-Bezerra, M., Li, Y., Acuna, M., Ivanovna, A.A., Corey, K.E. **Ortlund E.A.**, Cohen, D.E., Thioesterase Superfamily Member 2 Promotes Hepatic Triglyceride Secretion by Channeling Fatty Acids into the Glycerolipid Biosynthetic Pathway. *Hepatology*, 2019, Aug; 70(2):496-510. PMCID: PMC6551314

**Complete List of Published Works (80 total):**

<https://pubmed.ncbi.nlm.nih.gov/?term=ortlund&sort=date>

**BIOGRAPHICAL SKETCH**

Provide the following information for the Senior/key personnel and other significant contributors.  
Follow this format for each person. **DO NOT EXCEED FIVE PAGES.**

NAME: Patel, Anamika

eRA COMMONS USER NAME (credential, e.g., agency login): ANAMIKAPATEL

POSITION TITLE: Assistant Professor (Research Track), Department of Biochemistry

EDUCATION/TRAINING *(Begin with baccalaureate or other initial professional education, such as nursing, include postdoctoral training and residency training if applicable. Add/delete rows as necessary.)*

INSTITUTION AND LOCATION	DEGREE (if applicable)	Completion Date MM/YYYY	FIELD OF STUDY
Mohanlal Sukhadia University, India.	B.S.	07/1997	Chemistry, Botany, Zoology
Mohanlal Sukhadia University, India.	M.S.	07/1999	Chemistry
Central Salt & Marine Chemicals Research Institute, India	Ph.D.	11/2004	Chemistry
Technical University, Munich, Germany	Postdoctoral	10/2005	Biochemistry
Syracuse University, Syracuse, NY.	Postdoctoral	12/2009	Biochemistry

**A. Personal Statement**

I began my scientific training in the field of biochemistry and structural biology in 2001 and was intrigued by how you could visualize 3-D structures of proteins and link form to function. Since then, I've remained interested in studying the structure-function relationship of proteins, exclusively focusing on the field of transcriptional regulation. I possess specific training and expertise in X-ray crystallography, SAXS, cryo-EM, AUC, enzyme kinetics, and various biochemical and biophysical techniques required to study protein-protein and protein-DNA interactions. I started my postdoctoral training in Dr. Michael Cosgrove's laboratory at Syracuse University, where I was exposed to the interesting functions of chromatin modifying enzymes in transcriptional regulation and their significance in numerous diseases. During my stay in his lab, I worked on research projects that involved understanding the molecular mechanism of multiple histone H3 lysine 4 (H3K4) methylation catalyzed by the Mixed Lineage Leukemia-1 (MLL1) Core Complex, with an ultimate goal to provide a framework to develop a new class of therapies for MLL1-related leukemias. My work in his lab answered some long-standing questions in the field such as: how this five-protein complex gets assembled, and how the H3K4 tri-methylation activity of MLL1 is regulated by its complex components. These results led to a paradigm-shift in the field and suggested a novel mechanism used by this family of enzymes. I was also able to determine the structure of MLL1 in complex with one of its interacting proteins providing the basis for rational design of new class of drugs to target hyperactive form of MLL1 in some cases of leukemia.

After joining Department of Biochemistry at Emory University as an Assistant Professor (Research track), I have continued my research in the area of transcriptional regulation. I worked on a highly diverse class of transcription factors containing a tandem array of C2H2-type Zinc fingers in the group of Dr. Xiaodong Cheng at Emory. I solved several structures by X-ray crystallography and published few high-profile papers explaining how Zinc-fingers deviate from their conventional code of DNA recognition to achieve high sequence specificity without any gross penalties on DNA-binding. Recently, I shifted my focus towards cryo-EM due to two main reasons: first, my persistent interest in visualizing large transcriptional complexes and second, the technological advancement in cryo-EM now allow high-resolution structure determination. Driven by these interests, I undertook several research projects in Dr. Eric Ortlund's lab at Emory to study several protein complexes involved in transcription regulation to lipid transport with aim to determine their structures using cryo-EM. Given the fact that some of these complexes are resistant to crystallization, cryo-EM is the only available tool to derive structural information. I managed to produce good preliminary results for some of the protein complexes after several rounds of optimization that have laid a solid groundwork for current proposal. Over the past few years, I have attended a few workshops and training sessions to refine my skills in cryo-EM;

these points, in addition to the strong collaborative support we have, make us fully capable to carry out the proposed research.

## **B.Positions and Honors**

### **Positions and Employment**

**1999-2001:** Project Assistant at Central Salt & Marine Chemicals Research Institute, Bhavnagar, India.

**2006-2009:** Postdoctoral Research Fellow at Department of Biology, Syracuse University, Syracuse, NY.

**2010-2011:** Assistant Professor (Research Track), Department of Biology, Syracuse University, Syracuse, NY

**2012-2013:** Assistant Professor (Research Track), Department of Biochemistry and Molecular Biology, SUNY Upstate Medical University, Syracuse, NY

**2013-present:** Assistant Professor (Research Track), Department of Biochemistry, Emory University School of Medicine, Atlanta

### **Other Experience and Professional Memberships**

Member of American Association of Advancement of Science (AAAS)

Member of American Association of Cancer Research (AACR)

### **Honors**

2005: DAAD Short Term Research Fellow at Physics department, Technical University, Munich, Germany.

#### **1. C. Contributions to Science**

##### **Zinc Finger proteins and sequence specific DNA recognition.**

The C2H2 type zinc finger (ZF) proteins belong to the largest and most diverse family of transcription factors in human. They have well documented functional role in DNA binding. Despite being the largest family, the molecular targets and the biological function for the majority of them remains elusive. Based on structural knowledge of 2-3 tandem ZF, algorithms are derived to predict their DNA binding sequences. Yet, often they do not match with experimentally identified sequences, specifically in case of ZF array containing large number of tandem fingers. This suggesting the need for the structural studies of a tandem ZFs-DNA complex to enhance our understanding in their code of DNA recognition. My structure-function studies on two of these ZF proteins PRDM9 and ZFP568, each containing 13 and 11 tandem ZF, respectively, help to decipher the molecular basis of their DNA recognition. PRDM9 that directs the sequence specific binding to meiotic recombination hotspots and showed adaptability in their ZFs by flipping two differently positioned residue to the variations in DNA sequence and thus, allows them to respond to the variation in DNA sequences without any gross penalties in their binding affinities. My studies help to explain genetic data on how allelic variants of PRDM9 could influence their DNA binding targets and therefore, their meiotic recombination landscape. Along this line, I worked on a project in collaboration with Dr. Todd Macfarlan at NIH on a ZF protein ZFP568 that directly regulates the expression of a placental growth factor Igf2-P0 and is required for embryonic development in mice. I determined the structure of ZFP568 in complex with DNA, this is first structure available to date showing the largest number of ZFs in complex with DNA. The structures, I determined are of significant importance as they show deviations at several levels from the classical C2H2 type ZF DNA recognition code. The structures propose flexibility in the ZF by 2-,3-or 4-base pair specific recognition per zinc finger to enhance the binding capacity and also respond to the specific DNA shape. Collectively, these structures provide additional layers of adaptable binding ability between ZFs and DNA.

- a. **Anamika Patel**, Peng Yang, Matthew Tinkham, Mihika Pradhan, Ming-An Sun, Yixuan Wang, Don Hoang, Gernot Wolf, John R. Horton, Xing Zhang, Todd Macfarlan and Xiaodong Cheng. DNA Conformation Induces Adaptable Binding by Tandem Zinc Finger Proteins (2018). *Cell*, 147(1) 221-233. PMID:PMC5877318
- b. **Anamika Patel**, Xing Zhang, Robert Blumenthal and Xiaodong Cheng. Structural basis of human PR/SET domain 9 (PRDM9) allele C-specific recognition of its cognate DNA sequence (2017). *Journal of Biological Chemistry*, 292(39) 15994-16002. PMID:PMC5625032
- c. **Anamika Patel**, J.R. Horton, G.G. Wilson, X. Zhang, X. Cheng. Structural basis for human PRDM9 action at recombination hot spots (2016). *Genes & Development*. 2016. 30: 257-265. (This work was featured on the journal's cover.) PMID:PMC4743056

- d. **Anamika Patel**, H. Hashimoto, Xing Zhang, Xiaodong Cheng. Characterization of How DNA Modifications Affect DNA Binding by C2H2 Zinc Finger Proteins (2016). *Methods in Enzymology*, 573 387-403

## 2. **MLL1-WDR5-RbBP5-Ash2L-DPY30: a core complex responsible for histone H3 lysine 4 tri-methylation**

The disruption of Mixed Lineage leukemia-1 (MLL1) is frequently observed in acute lymphoblastic or acute myelogenous leukemia, which results in either increased or decreased MLL1's function. The MLL1 protein catalyzes histone H3 lysine 4 tri-methylation and regulates chromatin structure for active gene expression. The catalytic activity of MLL1 is tightly regulated by its core components. One question of significant importance in the field is that how MLL1's activity is regulated by its core components. My work showed that unlike other methyltransferase MLL1 core complex uses a novel mechanism involving two active sites to regulate its activity. We have identified a interaction motif in MLL1, which we named as "WIN" motif (also present in other family members: MLL2/3/4, SET1A and SET1B) required for binding with one of its regulatory subunit WDR5 and solved the first structure of MLL1 WIN motif in complex with WDR5. We also showed that targeting the interaction surface of MLL1-WDR5 successfully inhibits MLL1's activity and thus, provide a novel approach to target MLL1's activity in leukemia. As evidence of the high impact of this work, this research was published in several peer-reviewed journal articles as mentioned below. One article was pushed for accelerated publication and was featured on the cover of the Journal of Biological Chemistry. A second paper was highlighted as the paper of the week in the same journal.

- a. **Anamika. Patel**, V.E. Vought, V. Dharmarajan and M.S. Cosgrove (2011) A novel non-SET domain multi-subunit methyltransferase required for sequential nucleosomal histone H3 methylation by the MLL1 core complex. *Journal of Biological Chemistry* **286**(5): 3359-3369. PMID: PMC3030342
- b. **Anamika. Patel**, V. Dharmarajan, V.E. Vought and M.S. Cosgrove (2009) On the mechanism of multiple lysine methylation by the human Mixed Lineage Leukemia Protein-1 (MLL1) core complex. *Journal of Biological Chemistry* **284**(36): 2424-24256. PMID: PMC2782018  
*(selected as the paper of the week and spotlight the first author).*
- c. **Anamika. Patel**, V. Dharmarajan and M.S. Cosgrove (2008) Structure of WDR5 bound to Mixed Lineage Leukemia Protein-1 peptide. *Journal of Biological Chemistry* **283**(47): 32158-32161. PMID: 18829459. *(Accelerated Publication, Featured on the Journal Cover).*
- d. **Anamika. Patel**, V.E. Vought, V. Dharmarajan and M.S. Cosgrove (2008) A conserved arginine containing motif crucial for the assembly and enzymatic activity of the Mixed Lineage Leukemia protein-1 core complex. *Journal of Biological Chemistry* **283**(47): 32162-32175. PMID: 18829457.  
*(Featured on the Journal Cover).*

## 3. **Analytical Ultracentrifugation (AUC) to study protein-protein interaction and conformational switch**

I have my expertise in Analytical Ultracentrifugation to study Protein-Protein interactions and conformational changes in proteins in collaboration with various groups.

- a. S. Zhong, F. Hsu, C.J. Stefan, X. Wu, **Anamika. Patel**, M.S. Cosgrove and Y. Mao (2012) Allosteric activation of the phosphoinositide phosphatase sac1 by anionic phospholipids. *Biochemistry* **51**(15): 3170-7. PMID: PMC3329130
- b. M. Mbantenkhu, X. Wang, S. Wilkens, E. Hoffman, **Anamika. Patel**, M.S. Cosgrove and X.J. Chen (2011) Mgm101 - a DNA recombinase essential for the mitochondrial DNA maintenance. *Journal of Biological Chemistry* **286**(49): 42360-70. PMID: PMC3234957.
- c. T. Wollert, **Anamika. Patel**, Y.-L. Lee, V.E. Vought, M.S. Cosgrove, J.A. Mercer and G.M. Langford (2011) Myosin5a tail associates directly with Rab3A-containing compartments in neurons. *Journal of Biological Chemistry* **286**(16): 14352-14361. PMID: PMC3077635.



#### 4. Bili-proteins in cyanobacteria as photo-optical switch

My earlier work showed biochemical and structural characterization of photosynthetic bili-proteins in cyanobacteria from marine and fresh water environment. I showed that bili-proteins isolated from marine cyanobacterial species have higher thermal stability and antioxidant potential compared to fresh water species. We were able to determine for the first time a structure of a photo-optical switch of so-called E-isomer in bili-proteins.

- a. M. Schmidt, **Anamika. Patel**, Y. Zhao and W. Reuter (2007) Structural Basis for the Photochemistry of R-Phycocerythrocyanin. *Biochemistry* **46**: 416-423. PMID: 17209552
- b. L. Satyanarayana, C.G. Suresh, **Anamika. Patel**, S. Mishra and P.K. Ghosh (2005) X-ray crystallographic studies on C-phycocyanin from the cyanobacteria of different habitats: marine & fresh water. *Acta Crystallographica Section F: Structural Biology and crystallization communications* F61: 844-847. PMCID: PMC1978106.
- c. B.T. Paul, **Anamika. Patel**, G.S. Selvam, S. Mishra, P.K. Ghosh and R. Murugesan (2006) Photodynamic action of C-phycocyanins obtained from marine and fresh water cyanobacterial cultures: A comparative study using EPR spin trapping technique. *Free Radical research* **40**: 821-825. PMID: 17015260
- d. **Anamika. Patel**, S. Mishra, R. Pawar, S. Sonawane and P. K. Ghosh (2005) Purification and characterization of C-Phycocyanin from cyanobacterial species of marine and fresh water habitats. *Protein Expression and Purification* **40**: 248-255. PMID: 15766866

#### D. Additional Information: Research Support and/or Scholastic Performance

Ongoing

None

Completed

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DNA Methylation: Structures, Functions and Regulation

The goal of this grant was to determine central aspects, enzymatically and structurally, of (1) 5mC oxidation by Tet proteins, (2) modification-specific recognition by C2H2 zinc-finger and SRA-domain proteins, and (3) 5mC and 5hmC base excision by DNA glycosylases.