CURRICULUM VITAE CLIFTON D. MCKEE, Ph.D.

PROFESSIONAL DATA

Contact Information

Johns Hopkins University Bloomberg School of Public Health Department of Epidemiology 615 N. Wolfe St., Room E6008 Baltimore, MD 21205

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Website: clifmckee.github.io

GitHub: clifmckee

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EDUCATION AND TRAINING

Degrees

Ph.D. / 2020 · Ecology · Colorado State University · Fort Collins, CO

Thesis: Evolutionary and ecological processes in microparasite communities of bats

Supervisor: Prof. Colleen Webb

M.S. / 2015 · Ecology · Colorado State University · Fort Collins, CO

Thesis: Spatial, demographic, and phylogenetic patterns of Bartonella diversity in bats

Supervisor: Prof. Colleen Webb

B.S. / 2011 · Ecology and Evolution · University of Pittsburgh · Pittsburgh, PA B.A. / 2011 · Environmental Studies · University of Pittsburgh · Pittsburgh, PA magna cum laude

Postdoctoral Training

2020–2022 · Johns Hopkins Bloomberg School of Public Health · Baltimore, MD

Member: Infectious Disease Dynamics · Bat One Health

Supervisor: Prof. Emily Gurley

PROFESSIONAL EXPERIENCE

Johns Hopkins University

Faculty Research Associate · Department of Epidemiology, Bloomberg School of Public Health (2023–present)

Postdoctoral Fellow · Department of Epidemiology, Bloomberg School of Public Health (2020–2022)

Centers for Disease Control and Prevention

Research Fellow · Bacterial Diseases Branch, Division of Vector-Borne Diseases (2015–2017)

PUBLICATIONS

Journal Articles (Peer Reviewed)

- *Corresponding author, †Equal contribution, ‡Student advisee
- [38] Niu Y[‡], **McKee CD**. Bat viral shedding: a review of seasonal patterns and risk factors. *In press at Vector-Borne and Zoonotic Diseases*. 2025. DOI: 10.1089/vbz.2024.0091.
- [37] Jackson J[‡], Shanta IS, **McKee C**, Luby SP, Haider N, Sharker Y, Plowright R, Hudson P, Gurley E. Identifying weather patterns affecting household date palm sap consumption in Bangladesh, 2013-2016. *PLOS ONE*. 2024; 19(11): e0313904. DOI: 10.1371/journal.pone.0313904.
- [36] McKee CD, Yu EX, Garcia A[‡], Jackson J[‡], Koyuncu A, Rose S, Azman AS, Lobner K, Sacks E, Van Kerkhove MD, Gurley ES. Superspreading of SARS-CoV-2: a systematic review and meta-analysis of event attack rates and individual transmission patterns. *Epidemiology & Infection*. 2024; 152: e121. DOI: 10.1017/S0950268824000955
- [35] McKee CD*, Peel AJ, Hayman DTS, Suu-Ire R, Ntiamoa-Baidu Y, Cunningham AA, Wood JLN, Webb CT, Kosoy MY. Ectoparasite and bacterial population genetics and community structure indicate extent of bat movement across an island chain. *Parasitology*. 2024; 151(7): 708–721. DOI: 10.1017/S0031182024000660
- [34] Mathis SM, Webber AE, León T, Murray EL, Sun M, White LA, Brooks LC, Green A, Hu AJ, Rosenfeld R, Shemetov D, ..., **McKee CD**, ..., Borchering RK. Evaluation of FluSight influenza forecasting in the 2021–22 and 2022–23 seasons with a new target laboratory-confirmed influenza hospitalizations. *Nature Communications*. 2024;15:6289. DOI: 10.1038/s41467-024-50601-9
- [33] Cortes-Azuero O, Lefrancq N, Nikolay B, **McKee C**, Cappelle J, Hul V, Ou TP, Hoem T, Lemey P, Rahman MZ, Islam A, Gurley ES, Duong V, Salje H. The genetic diversity of Nipah virus across spatial scales. *The Journal of Infectious Diseases*. 2024; jiae221. DOI: 10.1093/infdis/jiae221
- [32] Jung S-m, Loo SL, Howerton E, Contamin L, Smith CP, Carcelén EC, Yan K, Bents SJ, Levander J, Espino J, Lemaitre JC, ..., McKee CD, ..., Viboud C, Lessler J. Potential impact of annual vaccination with reformulated COVID-19 vaccines: lessons from the US COVID-19 scenario modeling hub. *PLOS Medicine*. 2024; 21(4): e1004387. DOI: 10.1371/journal.pmed.1004387

- [31] Lemaitre JC, Loo SL, Kaminsky J, Lee EC, **McKee C**, Smith C, Jung S-m, Sato K, Carcelen E, Hill A, Lessler J, Truelove S. *flepiMoP*: the evolution of a flexible infectious disease modeling pipeline during the COVID-19 pandemic. *Epidemics*. 2024; 47: 100753. DOI: 10.1016/j.epidem.-2024.100753
- [30] Howerton E, Contamin L, Mullany LC, Qin MM, Reich NG, Bents SJ, Borchering RK, Jung SM, Loo SL, Smith CP, Levander J, ..., McKee C, ..., Viboud C, Lessler J. Evaluation of the US COVID-19 Scenario Modeling Hub for informing pandemic response under uncertainty. *Nature Communications*. 2023; 14: 7260. DOI: 10.1038/s41467-023-42680-x
- [29] Fagre AC, Islam A, Reeves WK, Kading RC, Plowright RK, Gurley ES, **McKee CD***. Bartonella infection in fruit bats and bat flies, Bangladesh. Microbial Ecology. 2023; 86: 2910–2922. DOI: 10.1007/s00248-023-02293-9
- [28] Szentiványi T, **McKee C**, Jones G, Foster JT. Trends in bacterial pathogens of bats: global distribution and knowledge gaps. *Transboundary and Emerging Diseases*. 2023; 9285855. DOI: 10.1155/2023/9285855
- [27] Seidlova V, Straková P, Kejíková R, Nemcova M, Bartonička T, Salát J, Dufková L, Šikutová S, Mendel J, **McKee C**, Zukal J, Pikula J, Rudolf I. Detection of *Leptospira* species in bat cadavers, Czech and Slovak Republics. *Emerging Microbes & Infections*. 2022; 11(1): 2211–2213. DOI: 10.1080/22221751.2022.2117095
- [26] Kejíková R, **McKee C**, Straková P, Šikutová S, Mendel J, Rudolf I. First detection of *Bartonella* spp. in bat bugs *Cimex pipistrelli* (Hemiptera: Cimicidae), Central Europe. *Parasitology Research*. 2022; 121(11): 3341–3345. DOI: 10.1007/s00436-022-07668-4
- [25] Goodrich I, **McKee C**, Margos G, Kosoy M. Molecular characterization of a novel relapsing fever *Borrelia* species from the desert cottontail (*Sylvilagus audubonii*) in New Mexico, USA. *Journal of Wildlife Diseases*. 2022; 58(3): 646–651. DOI: 10.7589/JWD-D-21-00148
- [24] McKee CD*†, Islam A†, Rahman MZ, Khan SU, Rahman M, Satter SM, Islam A, Yinda CK, Epstein JH, Daszak P, Munster VJ. Nipah virus detection at bat roosts after spillover events, Bangladesh, 2012–2019. *Emerging Infectious Diseases*. 2022; 28(7): 1384–1392. DOI: 10.3201/eid-2807.212614
- [23] Rice BL[†], Lessler J[†], **McKee C**[†], Metcalf CJE[†]. Why do some coronaviruses become pandemic threats when others do not? *PLOS Biology.* 2022; 20(5): e3001652. DOI: 10.1371/journal.pbio.3001652
- [22] Ruiz-Aravena M[†], **McKee C**[†], Gamble A, Lunn T, Morris A, Snedden CE, Yinda CK, Port JR, Buchholz DW, Yeo YY, Faust C, ..., Munster VJ, Plowright RK. Ecology, evolution and spillover of coronaviruses from bats. *Nature Reviews Microbiology*. 2022; 20: 299–314. DOI: 10.1038/s41579-021-00652-2
- [21] Redd AD, Peetluk LS, Jarrett BA, Hanrahan C, Schwartz S, Rao A, Jaffe AE, Peer AD, Jones CB, Lutz CS, McKee CD, ..., Grabowski MK, Gurley ES, the Novel Coronavirus Research Compendium Team. Curating the evidence about COVID-19 for frontline public health and clinical care: the Novel Coronavirus Research Compendium. *Public Health Reports.* 2022; 137(2): 197–202.

DOI: 10.1177/00333549211058732

- [20] Islam A, McKee C, Ghosh PK, Abedin J, Epstein JH, Daszak P, Luby SP, Khan SU, Gurley ES. Seasonality of date palm sap feeding behavior by bats in Bangladesh. *EcoHealth*. 2021; 18: 359–371. DOI: 10.1007/s10393-021-01561-9
- [19] Zorrilla VO, Lozano ME, Espada LJ, Kosoy M, **McKee C**, Valdivia HO, Arevalo H, Troyes M, Stoops CA, Fisher ML, Vásquez GM. Comparison of sand fly trapping approaches for vector surveillance of *Leishmania* and *Bartonella* species in ecologically distinct, endemic regions of Peru. *PLOS Neglected Tropical Diseases*. 2021; 15(7): e0009517. DOI: 10.1371/journal.pntd.0009517
- [18] McKee CD*, Islam A, Luby SP, Salje H, Hudson PJ, Plowright RK, Gurley ES. The ecology of Nipah virus in Bangladesh: a nexus of land-use change and opportunistic feeding behavior in bats. *Viruses*. 2021; 13(2): 169. DOI: 10.3390/v13020169
- [17] **McKee C***, Bai Y, Webb C, Kosoy M. Bats are key hosts in the radiation of mammal-associated *Bartonella* bacteria. *Infection, Genetics and Evolution.* 2021; 89: 104719. DOI: 10.1016/j.meegid.2021.104719
- [16] Goodrich I, McKee C, Kosoy M. Trypanosoma (Herpetosoma) diversity in rodents and lagomorphs of New Mexico with a focus on epizootological aspects of infection in Southern Plains woodrats (Neotoma micropus). PLOS ONE. 2020; 15(12): e0244803. DOI: 10.1371/journal.pone.-0244803
- [15] Rudolf I, Blažejová H, Mendel J, Straková P, Šebesta O, Rettich F, Čabanová V, Miterpáková M, Betášová L, Peško J, Barbušinová E, **McKee C**, Osikowicz L, Šikutová S, Hubálek Z, Kosoy M. *Bartonella* species in medically important mosquitoes, Central Europe. *Parasitology Research*. 2020; 119(8): 2713–2717. DOI: 10.1007/s00436-020-06732-1
- [14] Goodrich I, **McKee C**, Kosoy M. Longitudinal study of bacterial infectious agents in a community of small mammals in New Mexico. *Vector-Borne and Zoonotic Diseases*. 2020; 20(7): 496–508. DOI: 10.1089/vbz.2019.2550
- [13] **McKee CD***, Krawczyk AI, Sándor AD, Görföl T, Földvári M, Földvári G, Dekeukeleire D, Haarsma A-J, Kosoy MY, Webb CT, Sprong H. Host phylogeny, geographic overlap, and roost sharing shape parasite communities in European bats. *Frontiers in Ecology and Evolution*. 2019; 7: 69. DOI: 10.3389/fevo.2019.00069
- [12] Bai Y, Osinubi MOV, Osikowicz L, **McKee C**, Vora NM, Rizzo MR, Recuenco S, Davis L, Niezgoda M, Ehimiyein AM, Kia GSN, Oyemakinde A, Adeniyi OS, Gbadegesin YH, Saliman OA, Ogunniyi A, Ogunkoya AB, Kosoy MY, Idanre Bat Festival Investigation Team. Human exposure to novel *Bartonella* species from contact with fruit bats. *Emerging Infectious Diseases*. 2018; 24(12): 2317–2323. DOI: 10.3201/eid2412.181204
- [11] Kellner A, Carver S, Scorza V, **McKee CD**, Lappin M, Crooks KR, VandeWoude S, Antolin MF. Transmission pathways and spillover of an erythrocytic bacterial pathogen from domestic cats to wild felids. *Ecology and Evolution*. 2018; 8(19): 9779–9792. DOI: 10.1002/ece3.4451
- [10] McKee CD^{*}, Osikowicz LM, Schwedhelm TR, Maes SE, Enscore RE, Gage KL, Kosoy MY. Acquisition of *Bartonella elizabethae* by experimentally exposed oriental rat fleas (*Xenop-*

- sylla cheopis; Siphonaptera, Pulicidae) and excretion of Bartonella DNA in flea feces. Journal of Medical Entomology 2018; 55(5): 1292–1298. DOI: 10.1093/jme/tjy085
- [9] Kosoy M^{\dagger}, McKee C^{\dagger}, Albayrak L, Fofanov Y. Genotyping of *Bartonella* bacteria and their animal hosts: current status and perspectives. *Parasitology*. 2018; 145(5): 543–562. DOI: 10.1017/S0031182017001263
- [8] Gorsich EE, McKee CD, Grear DA, Miller RS, Portacci K, Lindström T, Webb CT. Model-guided suggestions for targeted surveillance based on cattle shipments in the U.S. *Preventive Veterinary Medicine*. 2018; 150: 52–59. DOI: 10.1016/j.prevetmed.2017.12.004
- [7] McKee CD*, Osikowicz LM, Schwedhelm TR, Bai Y, Castle KT, Kosoy MY. Survey of parasitic bacteria in bat bugs, Colorado. *Journal of Medical Entomology.* 2018; 55(1): 237–241. DOI: 10.1093/jme/tjx155
- [6] McKee CD*, Kosoy MY, Bai Y, Osikowicz LM, Franka R, Gilbert AT, Boonmar S, Rupprecht CE, Peruski LF. Diversity and phylogenetic relationships among *Bartonella* strains from Thai bats. *PLOS ONE*. 2017; 12(7): e0181696. DOI: 10.1371/journal.pone.0181696
- [5] Urushadze L, Bai Y, Osikowicz L, **McKee C**, Sidamonidze K, Putkaradze D, Imnadze P, Kandaurov A, Kuzmin I, Kosoy M. Prevalence, diversity, and host associations of *Bartonella* strains in bats from Georgia (Caucasus). *PLOS Neglected Tropical Diseases*. 2017; 11(4): e0005428. DOI: 10.1371/journal.pntd.0005428
- [4] Bai Y, Urushadze L, Osikowicz L, **McKee C**, Kuzmin I, Kandaurov A, Babuadze G, Natradze I, Imnadze P, Kosoy M. Molecular survey of bacterial zoonotic agents in bats from the country of Georgia (Caucasus). *PLOS ONE*. 2017; 12(1): e0171175. DOI: 10.1371/journal.pone.0171175
- [3] McKee CD*, Hayman DTS, Kosoy MY, Webb CT. Phylogenetic and geographic patterns of bartonella host shifts among bat species. *Infection, Genetics and Evolution*. 2016; 44: 382–394. DOI: 10.1016/j.meegid.2016.07.033
- [2] Zinzow-Kramer WM, Horton BM, **McKee CD**, Michaud JM, Tharp GK, Thomas JW, Tuttle EM, Yi S, Maney DL. Genes located in a chromosomal inversion are correlated with territorial song in white-throated sparrows. *Genes, Brain and Behavior*. 2015; 14(8): 641–654. DOI: 10.1111/gbb.12252
- [1] Bai Y, Hayman DTS, **McKee CD**, Kosoy MY. Classification of *Bartonella* strains associated with straw-colored fruit bats (*Eidolon helvum*) across Africa using a multi-locus sequence typing platform. *PLOS Neglected Tropical Diseases*. 2015; 9(1): e0003478. DOI: 10.1371/journal.pntd.0003478

Articles, Editorials, and Other Publications (Not Peer Reviewed)

Deposited in a Pre-Print Server

[2] Stevens T, Zimmerman R, Albery G, Becker DJ, Kading R, Keiser CN, Khandelwal S, Kramer-Schadt S, Krut-Landau R, **McKee C**, Montecino-Latorre D, ..., Carlson CJ. A minimum data standard for wildlife disease studies. *EcoEvoRxiv.* 2024. DOI: 10.32942/x2tw4j.

[1] McKee CD*, Webb CT, Kosoy MY, Bai Y, Osikowicz LM, Suu-Ire R, Ntiamoa-Baidu Y, Cunningham AA, Wood JL, Hayman DT. Manipulating vector transmission reveals local processes in bacterial communities of bats. *bioRxiv.* 2021. DOI: 10.1101/2021.03.03.433743

HONORS AND AWARDS

Vice President for Research Fellowship · Colorado State University (2018)
Department of Biology Travel Award · Colorado State University (2014 & 2017)
Sharon E. and David E. Kabes Scholarship · Colorado State University (2015)
GDPE Research Grant · Colorado State University (2014)
Department of Biology Graduate Fellowship · Colorado State University (2013)
Phi Beta Kappa · University of Pittsburgh (2011)

EDITORIAL AND OTHER INVITED PEER REVIEW ACTIVITIES

Journal Peer Review Activities

Acta Chiropterologica \cdot Acta Tropica \cdot BMC Microbiology \cdot BMJ \cdot Current Zoology \cdot Ecology and Evolution \cdot Frontiers in Microbiology \cdot Frontiers in Veterinary Science \cdot Infection, Genetics and Evolution \cdot Journal of Animal Ecology \cdot Journal of Medical Entomology \cdot Journal of Zoological Systematics and Evolutionary Research \cdot Microbial Ecology \cdot Microbial Pathogenesis \cdot Molecular Ecology \cdot mSphere \cdot Nature Microbiology \cdot Parasitology Research \cdot Pathogens and Global Health \cdot PLOS Neglected Tropical Diseases \cdot PLOS ONE \cdot PNAS Nexus \cdot The European Zoological Journal \cdot The Lancet Planetary Health \cdot The Science of Nature \cdot Scientific Reports \cdot Systematic Biology \cdot Tropical Biomedicine \cdot Veterinary Microbiology \cdot Viruses \cdot Zoonoses and Public Health

Proposal Peer Reviews

Netherlands Organization for Health Research and Development (ZonMw), Infectious Disease Control Program (2023)

PROFESSIONAL ACTIVITIES

Society Membership

North American Society for Bat Research (2021–present) \cdot American Society of Tropical Medicine and Hygiene (2021–present) \cdot Society for the Study of Evolution (2019 & 2021) \cdot American Society of Naturalists (2019) \cdot Ecological Society of America (2014–2019)

PRACTICE ACTIVITIES

Software and Other Product Development

The Flexible Epidemic Modeling Pipeline (flepiMoP) · A software suite for simulating a wide range of compartmental models of infectious disease transmission [Available on GitHub]

USAMM R Shiny Visualization · Developed tool for visualizing estimated county-to-county animal shipments in the US · Presented to stakeholders at USDA APHIS (2015)

Other Practice Activities

Bat Virus Spillover Virus Compendium (Bat-Com) · Led team of researchers to curate and write public-facing summaries and assessments of research articles on the spillover of SARS-CoV, SARS-CoV-2, and related coronaviruses

2019 Novel Coronavirus Research Compendium (NCRC) \cdot Curated and wrote public-facing summaries and assessments of research articles on the Ecology & Spillover of SARS-CoV-2 and related coronaviruses (2020–2021)

Media Dissemination

Nipah virus: Of Fruit and Bats · Interview with This Podcast Will Kill You (2024)

Lurking in the Shadows · Feature article in The Scientist Magazine (2014)

Disease on the Wing · YouTube video produced by The Scientist Magazine (2014)

CURRICULUM VITAE CLIFTON D. MCKEE, Ph.D. PART II

TEACHING

Academic Advisees

Elias Chan · Master of Health Science, Epidemiology · Johns Hopkins Bloomberg School of Public Health (2024–present)

Yannan Niu · Master of Health Science, Epidemiology · Johns Hopkins Bloomberg School of Public Health (2023-2024) · Thesis: Bat viral shedding: a review of seasonal patterns and risk factors

Capstone Advisees

Kathy Brickman · Master of Public Health · Johns Hopkins Bloomberg School of Public Health (2024–present)

Juanita Losier · Master of Public Health · Johns Hopkins Bloomberg School of Public Health (2024-present)

Master's Thesis Reader

Andrés Garcia · Master of Health Science, Epidemiology · Johns Hopkins Bloomberg School of Public Health (2023–2024) · Thesis: Literature review on risk factors for transmission of *Orientia tsutsugamushi* in Thailand, Vietnam, and Malaysia

Jules Jackson · Master of Science, Epidemiology · Johns Hopkins Bloomberg School of Public Health (2022–2023) · Thesis: Identifying weather patterns affecting household date palm sap consumption in Bangladesh, 2013–2016

Doctoral Thesis Committee

Darshan Sreenivas · National Centre for Biological Sciences · Bengaluru, India (2024–present)

Final Oral Exam Participation

Laura MacKenzie · External Examiner for PhD viva · School of Biological Sciences, University of Aberdeen · Aberdeen, Scotland (2023)

Classroom Instruction: Instructor of Record

Johns Hopkins University

Introduction to R for Public Health Researchers (140.604.73/79) · Co-Instructors: Ava Hoffman, Carrie Wright · Winter/Summer Institutes (2023–present, enrollment: 16–35)

The One Health Approach to Epidemiology and Global Public Health (340.610.81) · Co-Instructor: Emily Gurley · Spring 4th Term (2023–present, enrollment: 35–44)

Classroom Instruction: Teaching Assistant

Johns Hopkins University

Topics in Infectious Disease Epidemiology (340.668.89) · Teaching Assistant (2022)

The One Health Approach to Epidemiology and Global Public Health (340.610.81) · Teaching Assistant (2021 & 2022)

Epidemiology of Infectious Diseases (340.627.01) · Teaching Assistant (2020 & 2021)

Colorado State University

Molecular and General Genetics (BZ 350) \cdot Teaching Assistant & Recitation Instructor (2014, 2018 & 2019)

Introduction to Evolution (BZ 220) · Teaching Assistant (2018)

Ecology (LIFE 320) · Teaching Assistant (2017)

Ornithology (BZ 335) · Teaching Assistant & Laboratory Instructor (2014 & 2015)

Attributes of Living Systems (LIFE 102) · Teaching Assistant & Laboratory Instructor (2013)

Other Teaching

Coursera \cdot Developing content for the Ecology and Environment module in an upcoming course on One Health Investigations of Outbreaks and Spillovers (2024–present)

Coordinator & Content Developer · Applied Modeling in Public Health Workshops · Johns Hopkins Bloomberg School of Public Health (2022–present)

Content Developer & Co-Instructor \cdot Zombiecology: Workshop on Disease Ecology \cdot National Science Olympiad \cdot Colorado State University (2018)

Teaching Excellence Recognition

Introduction to R for Public Health Researchers (140.604.79) · Summer Institute 2024 Introduction to R for Public Health Researchers (140.604.73) · Winter Institute 2024 Introduction to R for Public Health Researchers (140.604.79) · Summer Institute 2023

RESEARCH AND PRACTICE FUNDING

External Sponsored Grants and Contracts

Current Support

Project Title: Atlantic Coast Center for Infectious Disease Dynamics and Analytics

Dates: 09/12/2023 - 09/11/2028

Sponsoring Agency: CDC Center for Forecasting and Outbreak Analytics

Principal Investigator: Shaun Truelove

Main Grant Objective: Lead recruitment and logistics for Applied Modeling in Public Health workshop series; assist with workshop content; participate in model development for forecasts and

scenario projections of COVID-19, influenza, and RSV cases and hospitalizations

Role: Research Associate

Project Title: Ecology of Nipah Virus in Bangladesh (58-3022-3-029)

Dates: 09/01/2023 - 08/31/2026 Sponsoring Agency: USDA/ARS Principal Investigator: Emily Gurley

Main Grant Objective: Provide support for field work, data analysis, and writing manuscripts; draft protocols for approval, collate and analyze data coming from the field and the laboratory, and

lead development of manuscripts for publication

Role: Research Associate

Project Title: Safety and Healthcare Epidemiology Prevention Research Development (SHEP-

heRD) (75D30121F00005)

Dates: 09/03/2021 - 09/02/2024

Sponsoring Agency: Centers for Disease Control

Principal Investigator: Shaun Truelove

Main Grant Objective: Lead recruitment and logistics for Applied Modeling in Public Health workshop series; assist with workshop content; participate in model development for forecasts and scenario projections of COVID-19, influenza, and RSV cases and hospitalizations

Role: Research Associate

Project Title: Solving Opportunities for Spillover (SOS): Frequency and Mechanisms of Cross-species Transmission of Henipaviruses in Bangladesh (1U01AI168287-01A1)

Dates: 01/03/2023 - 01/02/2028Sponsoring Agency: NIH/NIAID Principal Investigator: Emily Gurley

Main Grant Objective: Provide support for field work, data analysis, and writing manuscripts; draft protocols for approval, collate and analyze data coming from the field and the laboratory, and

lead development of manuscripts for publication

Role: Research Associate

Past Support

Project Title: Preventing emergence and spillover of bat pathogens in high risk global hotspots

(G166-19-W7329)

Dates: 10/01/2018 - 09/30/2022

Sponsoring Agency: DARPA/Montana State University

Principal Investigator: Raina Plowright (Project Director), Emily Gurley (Site PI)

Main Grant Objective: Work with the team in Bangladesh to develop and implement protocols to sample bats, including identifying roosts best suited for sampling, ensuring protocols are aligned with broad aims of this grant, and that work is delivered on time; real-time analysis of virus

detection data

Role: Postdoctoral Fellow

Project Title: COVID-19 Technical Lead in the Implementation of the Strategic Preparedness

(202548990)

Dates: 08/03/2022 - 11/03/2022

Sponsoring Agency: World Health Organization

Principal Investigator: Emily Gurley

Main Grant Objective: Support COVID-19 technical lead to improve evidence-based guidance by

completing a systematic review of COVID-19 superspreading

Role: Postdoctoral Fellow

PRESENTATIONS

Invited Seminars

Invited Talks

Developing One Health surveillance systems for emerging zoonotic pathogens: moving towards pandemic prevention · University of Michigan School of Public Health · Ann Arbor, MI (2024)

Shadows on the wall: interpreting multiplex viral sero-surveillance in bats and domesticated animals in Bangladesh · Global Virus Network Annual Meeting · Durban, South Africa (2024)

Environmental and behavioral drivers of cross-species Nipah virus transmission in Bangladesh \cdot Emerging Infectious Diseases: Ecology and Evolution Workshop \cdot International Centre for Theoretical Sciences \cdot Bengaluru, India (2024)

Ecological stressors and pathogen shedding · Emerging Infectious Diseases: Ecology and Evolution Workshop · International Centre for Theoretical Sciences · Bengaluru, India (2024)

Investigating the ecology of Nipah and other bat-borne viruses at the human-animal interface in Bangladesh \cdot Uniformed Services University of the Health Sciences \cdot Bethesda, MD (2024)

Investigating Nipah virus spillover at the human-animal interface in Bangladesh · Cornell University College of Veterinary Medicine · Ithaca, NY (2023)

Global change, infectious disease, and public health: the need for ecological interventions \cdot WHO GOARN/RCCE Collective Service \cdot Virtual (2022)

Campus or Departmental Talks

Investigating the ecology of Nipah and other bat-borne viruses at the human-animal interface in Bangladesh · Johns Hopkins Bloomberg School of Public Health · Baltimore, MD (2023)

Bad roommates? Inferring bacterial interactions in coinfected individuals · Colorado State University Vice President Office of Research Symposium · Fort Collins, CO (2018)

Scientific Meetings

Oral Presentations

The Bat Virus Spillover Evidence Compendium (Bat-Com): What we know, and don't know, about the most important bat zoonoses \cdot American Society of Tropical Medicine and Hygiene \cdot New Orleans, LA (2024)

Empowering researchers through Publish-Review-Curate workflows \cdot Society for Scholarly Publishing \cdot Boston, MA (2024)

Superspreading of SARS-CoV-2: a systematic review and meta-analysis · American Society of Tropical Medicine and Hygiene · Chicago, IL (2023)

Comprehensive time tree analysis identifies bats as key to the radiation of mammal-associated Bartonella bacteria · Evolution · Providence, RI (2019)

Timing the diversification of a mammal parasite, $Bartonella \cdot Ecological Society of America \cdot New Orleans, LA (2018)$

Timing the diversification of a mammal parasite, Bartonella · Front Range Student Ecology Symposium · Fort Collins, CO (2018)

Long-term monitoring of *Bartonella* spp. bacteria in a captive colony of fruit bats and experimental evidence of bat flies as vectors of bartonella · Ecological Society of America · Portland, OR (2017)

Long-term monitoring of Bartonella spp. bacteria in a captive colony of fruit bats and experimental evidence of bat flies as vectors of bartonella · International Symposium on Infectious Diseases of Bats · Fort Collins, CO (2017)

Host phylogenetic distance and ectoparasite overlap predict Bartonella sharing in European bats · Front Range Student Ecology Symposium · Fort Collins, CO (2017)

Phylogeography of Bartonella bacteria in Eidolon spp. fruit bats across Africa · Ecological Society of America · Baltimore, MD (2015)

Phylogeography of *Bartonella* in *Eidolon* fruit bats across Africa · Front Range Student Ecology Symposium · Fort Collins, CO (2015)

Poster Presentations

The Bat Virus Spillover Evidence Compendium (Bat-Com): what do we know about viral spillover from bats after approximately 60 years of reported outbreaks? \cdot World One Health Congress \cdot Cape Town, South Africa (2024)

Nipah virus detection at bat roosts following spillover events in Bangladesh, 2012-2019 · International Conference on Emerging Infectious Diseases · Atlanta, GA (2022)

Manipulating vector transmission reveals local processes in bacterial communities of bats \cdot Ecology and Evolution of Infectious Disease \cdot Princeton, NJ (2019) \cdot **Poster award**

Dispersal of hosts and vectors predicts bacterial community structure across an island chain \cdot Front Range Student Ecology Symposium \cdot Fort Collins, CO (2019) \cdot **Poster award**

Timing the diversification of a mammal parasite, $Bartonella \cdot Colorado$ State University Graduate Student Showcase \cdot Fort Collins, CO (2017)

Linking patterns of bacterial parasite diversity across host and vector communities · Ecology and Evolution of Infectious Disease · Santa Barbara, CA (2017)

Phylogeography of Bartonella bacteria in Eidolon spp. fruit bats across Africa · Ecology and Evolution of Infectious Disease · Athens, GA (2015)

Large-scale patterns of Bartonella prevalence and diversity in African fruit bats · Ecological Society of America · Sacramento, CA (2014)

Large-scale patterns of Bartonella prevalence and diversity in African fruit bats · Ecology and Evolution of Infectious Disease · Fort Collins, CO (2014)

A novel method of isolating multi-locus sequence data for characterizing *Bartonella* diversity in bats · Front Range Student Ecology Symposium · Fort Collins, CO (2014)

ADDITIONAL INFORMATION

Personal Statement

I am an infectious disease ecologist interested in zoonotic pathogens and their dynamics within host populations and at the human-animal interface. Overall, my research sits firmly within One Health, seeking to understand disease emergence at the intersection between public health, animal ecology, and environmental change. I use a combination of fieldwork, statistical modeling, molecular genetics, and phylogenetics to investigate pathogen persistence in animal reservoirs, evolution of host specificity, and ecological drivers of pathogen spillover.

Keywords

disease ecology, epidemiology, data science, zoonoses, One Health, phylogenetics, evolution

Bioinformatics and Phylogenetics: Multiple sequence alignment (CLUSTAL, MAFFT), maximum likelihood (MEGA, IQ-Tree, RAxML) and Bayesian (MrBayes) phylogenetic inference, inference of population genetic structure (STRUCTURE) and recombination (ClonalFrame, SplitsTree), cophylogeny (ParaFit, PACo), ancestral state reconstruction and phylodynamics (BEAST)

Computing and Data Visualization: High proficiency in R for data manipulation, statistical analysis, and visualization (base and tidyverse, ggplot2, igraph), generating reports with R Markdown, and interactive web pages with Shiny; experience with git, LATEX, and Python (NumPy, Pandas, seaborn)

Modeling: Proficient in deterministic and stochastic models for infectious disease dynamics and hierarchical Bayesian modeling (JAGS)