## **Research Summary**

(As of 12-31-2024)

Dr. Jizhong Zhou is a George Lynn Cross Research Professor in School of Biological Sciences, School of Civil Engineering and Environmental Sciences, School of Computer Science, and is the Director of the Institute for Environmental Genomics, at the University of Oklahoma (OU). He is a Visiting Senior Scientist in the Earth and Environmental Sciences at Lawrence Berkeley National Laboratory, and an Adjunct Professor of School of Environment at Tsinghua University, Beijing, China. Dr. Zhou has distinguished himself as an international leader in **genomics-enabled environmental sciences**, especially in areas related to: (i) climate change biology, (ii) environmental remediation, (iii) theoretical ecology, (iv) genomic technologies, and (v) computational technologies.

Dr. Zhou has made several ground-breaking discoveries related to climate change biology. First, he discovered that microorganisms play central roles in regulating soil carbon (C) dynamics through three primary feedback mechanisms<sup>1,2</sup>. This was the **first** study revealing functional differences in microbial communities due to warming that may affect projections of climate warming. These results represent critical findings in global change biology, and was ranked by Faculty 1000 as among the top 2% of published articles in biology and medicine. More recently, he discovered that climate warming leads to divergent succession of grassland bacterial and fungal communities<sup>3</sup>, but convergent in archaeal community<sup>4</sup>, accelerates microbial temporal scaling<sup>4,5</sup>, network complexity and stability<sup>6</sup>, and positive soil priming<sup>7</sup>, but reduces microbial diversity<sup>4,8</sup>. Also, using integrated genomics technologies with isotope and process analysis, he demonstrated the *in situ* vulnerability of tundra soil C to climate warming and the importance of microbes in mediating such vulnerability<sup>9</sup>. This study was highlighted as a breakthrough by *The* Washington Post, Department of Energy, and ASM Microbe because northern permafrost stores more than 30% of the global soil organic C, and is a wildcard that could alter the future trajectory of global climate change. In addition, he demonstrated that long-term field exposure of a grassland ecosystem to elevated CO<sub>2</sub> dramatically altered the composition and structure of the belowground microbial community<sup>10</sup>, which was the first comprehensive study at the whole community level to document the responses of soil microbial communities to elevated CO<sub>2</sub>. These results were highlighted by Science as a convincing demonstration of the frontier of integrative biodiversity studies (Naeem et al. 2012. Science: 336: 1401-1406). Recently, he and his colleagues demonstrated that elevated CO<sub>2</sub>-stimulated soil respiration is enhanced by N limimation, and the underlying microbial mechanisms<sup>11</sup>, which implies that the heightened release of CO<sub>2</sub> back to the atmosphere from soil may be pervasive given that fact that N is generally limited in nature. All of these studies have made conceptual advances in climate change biology, and lead to a paradigm shift by demonstrating the urgent needs of incorporating microbial functional traits into global change modeling because whether microbially mediated feedback is positive or negative depends on which functional guilds are affected by climate changes and to what magnitude. Because of his outstanding contributions in research related to climate change microbiology, environmental remediation, and theoretical ecology, he received the 2022 Soil Science Research Award from Soil Science Society of America for recognizing outstanding research contributions in soil science, and Distinguished Scientist Award by Southeastern Universities Research Association (SURA) in 2024 for recognizing scientists who are performing world-leading research.

Dr. Zhou has made several pioneering discoveries in the area of **environmental remediation**. <u>First</u>, over the last two decades, he has pioneered studies on groundwater microbiomes in response to extreme heavy metal contamination by illustrating: (a) the impact of heavy metal contamination on the functional community structure<sup>12,13</sup>, (b) crucial roles of horizontal gene transfer in driving the evolution of microbial communities in response to contamination<sup>14</sup>, (c) microbial mechanisms underlying successful *in situ* bioremediation to reduce high uranium contamination to below drinking water standards<sup>15-21</sup>, (d) community assembly processes governing the succession of microbial communities to substrate

amendment<sup>22-26</sup>, and (e) artificial intelligence-assisted predictions of groundwater geochemistry. contamination and functions with omics data<sup>27,28</sup>. Second, using GeoChip and other technologies, he and his colleagues demonstrated a broad capacity and high potential for intrinsic bioremediation of the Deep Water Horizon oil plume in the Gulf of Mexico<sup>29</sup>, which was described by President Barack Obama in a prime-time address as "the worst environmental disaster America has ever faced". A subsequent publication with GeoChip data alone in *The ISME Journal*<sup>30</sup> was specially highlighted by the European Commission's Science for Environment Policy. These studies also provide a shining example of how GeoChip technology is used in an integrated synergistic fashion to address complex emergent environmental problems, a cornerstone of basic science. Third, he has established a Global Water Microbiome Consortium (GWMC). With a systematic global-sampling effort, he and his colleagues examined global diversity and biogeography of microbiomes from ~1200 activated sludge samples collected from 269 wastewater treatments (WWTPs) in 23 countries across 6 continents<sup>31</sup>. This is the **first** comprehensive global scale study to address various theoretical questions in engineered systems. The results demonstrated extremely high microbiome diversity of global wastewater treatment plants, with a small global core strongly linked to system's performance. The findings of this study highlight how little we know of the world's microbiome, even in one of the most common and well-controlled systems in the built environment, and have important implications for microbial ecology and wastewater treatment processes. In addition, Dr. Zhou is an international leader in studying functional genomics to assess microbial responses to environmental stresses – a core of microbial ecology - via his central role in the first comprehensive book on Microbial Functional Genomics<sup>32</sup>. He pioneered the elucidation of microbial cellular responses to contaminants and various stressors, energy metabolism and regulatory networks in several environmentally important microorganisms, including Shewanella, Desulfovibrio, Rhodanobacter, as well as Clostridium and Thermoanaerobacter. Many new insights were revealed using integrated functional genomics/systems biology approaches, especially with sulfate-reducing microorganisms as demonstrated by his invited synthesis in *Nature Reviews Microbiology*<sup>33</sup>. Primiarly because of his leaderships in environmental and functional genomics, he received received U.S. Congressional recognition with the U.S. Department of Energy's Ernest Orlando Lawrence Award in 2014 – one of the highest honors presented by the U.S. government in science, and the 2022 ISME-IWA BioCluster Grand Prize Award for recognizing the importance and impact of interdisciplinary research at the interface of microbial ecology and water/wastewater treatment.

Dr. Zhou has also made several meritorious contributions to the field of theoretical ecology. First, in contrast to the traditional view of microbial assembly, several of his pioneering studies have explicitly demonstrated the importance of stochastic processes underlying microbial community assembly - a central but poorly understood topic in ecology<sup>34</sup>. He was the **first** to discern unusual microbial diversity patterns in soils and, through mathematical modeling analysis, revealed that spatial isolation (i.e. dispersal limitation) was the major underlying mechanism<sup>35</sup>, and to discover that divergent microbial communities with different functions can be formed under identical environmental conditions<sup>36</sup>. A novel framework was also recently developed to conceptualize the relationships between community assembly processes and environmental stresses<sup>37</sup>. He provided the **first** evidence that microbial community **succession** is stochastic rather than deterministic<sup>22</sup>, and the relative importance of stochastic assembly processes in shaping community structure decreased as stress increased<sup>37</sup>. In addition, he was the first to document that the responses of grassland soil microbial communities to climate change are primarily governed by stochastic processes<sup>5,38</sup>, and the microbiome assembly in global WWTPs is also more stochastic<sup>31</sup>, in complete contrast to conventional thought because WWTPs are well-controlled engineered ecosystems. Such findings have important implications for studying ecosystem functioning, biodiversity preservation, and ecosystem management. Second, using integrated genomic technologies, his studies convincingly demonstrated that species area relationships (SARs), a universal law in ecology, generally hold for microorganisms, but that microorganisms have substantially lower spatial turnovers (z values, < 0.1) due to their unique biology<sup>39-41</sup>. His recent study on the temporal scaling of soil bacteria and fungi provides explicit evidence for the existence of species-time relationships (STRs), another possible universal law in biology<sup>5</sup>. This study is the **first** to demonstrate that there is phylogenetic-time relationships (PTRs), and that warming accelerates both STRs and PTRs. <u>Third</u>, contrary to a common belief, he and his colleagues demonstrated that taxonomic and phylogenetic diversity of soil bacteria, fungi, and nitrogen fixing bacteria all are better predicted by temperature than pH<sup>42</sup>. This was the **first** study in microbial ecology showing that the metabolic theory of ecology (MTE) is a powerful framework for predicting broad large-scale microbial biodiversity, but further theoretical modelling development is needed to account for the unique characteristics of microorganisms. Primiarly due to his outstanding contributions in research related to climate change biology, environmental remediation, and theoretical ecology, he received the 2019 ASM Award for Environmental Research - for recognizing an outstanding scientist with distinguished research achievements in microbial ecology and environmental microbiology.

Dr. Zhou has made several pioneering advances in the area of **genomic technologies**, particularly in the development of various innovative technologies for microbial analyses, which were crucial for the early development of this emerging field. First, his GeoChip<sup>18,43-45</sup> is a revolutionary, high throughput technology for addressing a central goal in microbial ecology - linking community structure to function. Over last 25 years, he has solved various challenging problems in developing chip-based technologies for environmental applications to enable scientists to address research questions that formerly they could not in terms of specificity, sensitivity, quantitation and reproducibility 44,46-57. The most recent generation of GeoChip (Version 5.0) contains 180K probes targeting ~370K functional genes in ~1,500 gene categories important to different microbial processes, including carbon, nitrogen, phosphorus, and sulfur cycling, energy processing, metal resistance and reduction, organic contaminant degradation, stress responses, antibiotic resistance, pathogenesis, viral-host interactions from broad, functionally divergent taxonomic groups, such as bacteria, archaea, fungi, protists, and viruses (mainly phages)<sup>44</sup>. The GeoChip 5 is highly specific, sensitive, and quantitative based on both computational and experimental assays<sup>44</sup>. The GeoChip technology was highlighted in the NRC's report on the New Science of Metagenomics (2007), and in the roadmap for NSF's National Ecological Observatory Network (NEON, 2009), and won a R&D 100 Award in 2009 as one of the 100 most innovative scientific and technological breakthroughs that year. Using GeoChip technology, he has addressed various fundamental questions in ecological theory, climate change, environmental remediation, and human health, as evident by numerous publications (>100) in Nature indexed journals, including Science<sup>29</sup>, Nature Climate Change<sup>1,6,9</sup>, PNAS<sup>11,22,39,58-60</sup>, Nature Communications<sup>2,61</sup>, Ecology Letters<sup>10</sup>, The ISME Journal<sup>30,62,63</sup>, Environmental Science & Technology<sup>24,26</sup>, and Water Research<sup>64,65</sup>, and other prestigious journals, e.g., Nature Ecology & Evolution<sup>66</sup>, mBio<sup>28,67</sup>, and Global Change Biology  $^{68-70}$ . The GeoChip paper in the Inaugural issue of ISME  $J^{18}$  is among the 5 top-cited articles in the first 10 years of the journal. In brief, GeoChip has become a powerful technology enabling comprehensive visualization of the functional structure of microbial communities important to energy, global change, ecosystem management and allows microbiologists to address research questions which could not otherwise be undertaken. Second, his methods for effectively extracting high quality, high molecular weight DNA from environmental samples have been very influential and widely used by other scientists<sup>71,72</sup>. The paper<sup>71</sup> describing these methods has been cited >4,000 times and was among the 20 most cited papers in Applied and Environmental Microbiology (a leading microbiology journal) since 2008. These methods, because they are capable of recovering high molecular weight DNA from environmental samples, have provided a foundation for contemporary metagenomics and molecular microbial ecology. Third, he was the first to discern the problems of heteroduplexes and PCR-induced mutations in 16S rRNA gene-based cloning and developed approaches to minimize such artifacts<sup>73</sup>. The cloning-based method was very important in early molecular microbial ecology studies in the 1990s. Fourth, he was the first to discover that sequencing-based metagenomic technologies are poorly reproducible and quantitative 74.75, which are mainly due to the effects of random sampling processes<sup>57,76</sup>, and developed mathematical approaches to predict and minimize the artifacts associated with random sampling <sup>76</sup>, for which the paper describing these approaches was selected as an Editor's Pick. As an mBio's Editor commented, this study would make a fundamental contribution for designing and interpreting microbiome studies. Since most ecological studies involve random sampling, these results and guidance have had a wide impact. His

perspective on metagenomics technologies for ecological applications<sup>57</sup> has also been influential in the field of microbiology. All these studies have been **instrumental** in shaping molecular microbial ecology during its great expansion over the last two decades. <u>Finally</u>, he developed a novel Cas9 nickase-based tool for editing bacterial genomes<sup>77</sup>. This study was highlighted by AEM as an Editor's Pick, and the tool has since been used by many other microbiologists for their studies. <u>Primarily due to his pioneering advances in omics-based technologies for environmental studies and microbial functional genomics, he recieved Presidential Early Career Award for Scientists and Engineers in 2001 - the highest honor for young scientists and engineers in US.</u>

Dr. Zhou has also made several pioneering advances in the area of **computational technology**. First, based on the random matrix theory (RMT), a relatively young mathematical theory proposed in 1960s, he developed a novel, sensitive, and robust approach with mathematically defined thresholds for predicting network interactions in microbial communities<sup>78-80</sup>, overcoming a great challenge in network biology. The network pipeline<sup>81</sup> has been widely used by microbial ecologists with little computational background, as evidenced by the  $\sim 135,000$  networks constructed by  $\sim 9,000$  users from > 70 countries. This early pathbreaking work represents paradigm-shifting, transformative research in microbial ecology. Prior to this, most microbial biodiversity studies ignored the "interactions" and associated functions among different microorganisms. More recently, he has developed a novel approach for inferring direct dependences in association networks, called iDIRECT (Inference of Direct and Indirect Relationship with Effective Copula-based Transitivity), by overcoming/ameliorating several mathematical problems (e.g. underdetermination, self-looping and interaction strength overflow) in network ecology<sup>82</sup>. Second, he has developed various novel mathematical approaches to quantitatively estimate the relative importance of deterministic vs stochastic processes in governing community assembly – a major bottleneck in ecology. His most recent innovation provides an effective and robust tool for quantitatively assessing ecological stochasticity<sup>58</sup>. Another novel general framework for quantitatively inferring community assembly mechanisms by phylogenetic-bin-based null model analysis (iCAMP)<sup>38</sup> is capable of reliably estimating the relative roles of selection, dispersal, and "drift". These new general tools are not only crucial for microbial ecology, but also for macrobial ecology. By the end of 2024, the R packages of NST and iCAMP have been downloaded for >32,000 and >31,000 times, respectively. Plus, the web-based galaxy pipelines have also been used by many users (iCAMP by ~110 users for > 1,100 times; and NST by ~80 users for > 1,500 times). These tools and pipelines have greatly enabled non-computational researchers in applying sophisticated mathematical and statistical methods to explore frontier ecological questions. In addition, he has pioneered the development of novel mathematical frameworks to incorporate massive genomics information into Earth system models<sup>2,11,83,84</sup> – an urgent issue with the greatest challenge in global change research. He was the first to demonstrate that incorporating microbial functional traits into ecosystem models can substantially reduce model uncertainty and improve modeling prediction in response to longterm experimental warming<sup>2</sup>, and elevated CO<sub>2</sub>, and/or nitrogen deposition<sup>11,84</sup>. These groundbreaking studies have been instrumental in steering microbial ecology in new research directions.

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