**A close up of a name

AI-generated content may be incorrect.**

**PRE-APPLICATION: Google PhD Fellowship Program 2025**

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| **Student Name:** | Megan Darrell | | **Banner ID:** | | 810446601 |
| **Thesis Mentor:** | | Sophie Molholm | |  | |

**Describe the desired impact your research will make on the field and society, and why this is important to you (350 word-limit)**

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| Entering college, I was torn between studying neuroscience—captivated by the brain’s unparalleled complexity—and bioinformatics, an innovative field that required logical rigor and creativity. As I delved deeper into research, I quickly realized the two disciplines exist not in competition, but in parallel—and that the most meaningful advances in healthcare would come from integrating them.  In my first research role, I helped develop a convolutional neural network to non-invasively measure naturalistic gait in neurological disease models, and was immediately captured by the ability of AI-driven tools to reveal profound insights into brain function and behavior. Eager to explore the power of bioinformatics to drive medical discovery, I’ve since applied novel AI methodologies to rigorously analyze data across a wide range of clinical domains—from identifying transcriptomic subtypes in a rare pediatric autoimmune condition at the NIH, to my most recent work, which has uncovered critical healthcare disparities in large clinical trials. These prior experiences have been the driving force behind my current research, as they reveal the immense potential of novel informatics approaches to extract meaningful insights from complex health data—insights that would remain hidden using traditional analyses.    Currently, my work centers on autism spectrum disorder (ASD)—a condition characterized by significant clinical and biological heterogeneity, which poses profound challenges for both diagnosis and treatment. In my proposal, we leverage a large, diverse dataset from children in the Bronx—a population historically excluded from research—that integrates rigorous clinical assessments and EEG from eight behavioral paradigms. Instead of relying on traditional analysis methods—which risk oversimplifying the complex neural and behavioral heterogeneity in ASD—we propose an innovative alternative: cluster-based identification of neural subgroups within ASD.  This methodology—applied to a uniquely rich clinical dataset—offers a rare and powerful opportunity to identify functional neural mechanisms that drive phenotypic variation in ASD, a challenge that has long hindered progress in the field. Above all, I am deeply motivated by the hope that this work moves us closer to personalized, effective care for individuals with autism—while demonstrating how data-driven methods can reveal insights into the brain’s complexity. |