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Dear Prof. Smit, Prof. Curtin, Prof. Marzari and other members of the selection committee,

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My name is Yury Polyachenko, I am a 3rd-year undergraduate student at Moscow Institute of Physics and Technology (MIPT). I have been interested in computer simulations since the 9th grade of high school, where for 3 years I was studying spiral structure formation in galaxies both implementing C++ and Matlab code and running simulations using Agama package. Later on, as a freshman at MIPT, I joined Prof. Genri Norman's lab of Computational Condensed Matter Physics and Life Systems. As my 1st project in this lab, I created a molecular dynamics (MD) package from scratch with an ultimate goal to test and improve MKT equations (see CV).

Now, I have been working with Prof. Norman for over 2 years. I am currently investigating a discontinuity phenomenon in a stable-metastable phase transition in the Lennard-Jones system using LAMMPS and Matlab. I discovered a property which exhibits an anticipated disruption at the transition point, therefore enables detection of a phase transition as a possible application. Throughout my research, I delivered 2 oral reports at Russian national conferences and currently we are preparing our results for submission to PRL.

This past summer I attended CECAM SISSA MD school, where I had an opportunity to strengthen my knowledge in enhanced sampling, reversibility, polymer and protein dynamics. I was immensely excited about prof. Mark Tuckerman's lectures on applying ML and MD to the drug design. This experience spurred me on to continue exploring this field more. Therefore, in the fall term, I completed several courses on MD, HPC and ML at MIPT, being ranked as the top 1 in my class. I believe EPFL, being a very applied-oriented university and a major player in CECAM, provides an excellent opportunity to challenge myself with non-trivial problems related to my background and interests this summer. Therefore, I am interested in joining several scientific groups in which I believe I can fully realize my scientific potential.

Laboratory of Molecular Simulation ___

This lab offers many interesting projects, yet the one that intrigues me the most is the "Self-diffusion from a potential energy field" project since it is closely related to my research experience. As my first educational project on MD was about self-diffusion, I was greatly excited to learn how this lab's members pointed out ways to use TuTraSt for the real-world applications such as e.g. ion batteries and even for a very general and fundamental problem of studying chemical bonds in a molecule via electron density analysis. Moreover, in SISSA I had a practice on rare events and enhanced sampling where I learned about physical ideas close to ones behind the TuTraSt. I really liked the fusion of rigour and power in the idea of "compensating" the potential landscape of a system to make it better sample more extreme aka rare regions of configurational space. Finally, the lab's tests for CH4 were conducted in LAMMPS, and it is mentioned that to speed up computations authors want to translate the code from Matlab to Python and C++. I have been working in LAMMPS in my current project and I consider myself proficient in all these 3 languages (see CV and git). So my computer skills will certainly be relevant in this project if I was to work on it.

Laboratory for Multiscale Mechanics Modeling lammm_

Many recent lab's publications have a molecular approach so my experience will be relevant. Furthermore, I am looking forward to mastering multiscale methods. I am particularly interested in the projects on neural network potentials, since I it combines the best from both DFT and classical potentials approaches. At MIPT I had a course "ML in condensed matter physics" where I became acquainted with basic ideas on using NN potentials. My final project for the course was to find a reasonable systematically improvable set of features and construct an ML potential for a crystal structure. In the end, I came up with an idea close to Moment Tensor Potentials. I believe there is a way to optimize and improve the methods currently used in the lab's research. I am interested in implementing the concept of "learning on the fly" which takes the idea of balancing the DFT and potential dynamics even further than traditional NN potentials. Thus it achieves accuracy close to the pure DFT with orders of magnitude less computational time.

Theory and Simulation of Materials THEOS ______

This lab has a list of Masters Projects and they say those are adaptable for Bachelor Projects. At MIPT I have completed 1 semester of Landau-Lifshitz quantum mechanics course and received 10/10 grade. By this summer, I will complete the Landau-Lifshitz course on quantum mechanics. Therefore I think by that time I will be qualified to work on the adapted versions of offered masters projects since for most of them proper QM background is essential. Furthermore, having completed basic QM courses I will be looking forward to mastering DFT and other computational QM techniques and I believe THEOS lab to be a great place to do so. Besides QM, I have experience with Python and Jupyter, so I can work on the "Visualization plugins for the Materials Cloud platform" for which Python is the main demand.

To sum up, I am extremely interested in joining EPFL as a summer intern. The program would provide an excellent opportunity to develop both professionally and academically, which I believe would prove critical when applying for a doctoral degree program at EPFL. I thank you for your time and sincerely appreciate your consideration of my application. Please do not hesitate to contact me at .

Sincerely,

Polyachenko Yury