


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| | | |
|---|---|---|
|  | Адрес | Контакты |
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Образование

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| 2014–2018 | Московская обл., г.Долгопрудный | МФТИ (ГУ) | аспирантура |
| 2007–2014 | Московская обл., г.Долгопрудный | МФТИ (ГУ) | Бакалавриат, Магистратура |

Преподавание

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|-------------------|-------------------|-----------------------|---|
| 2021– наст.вр. | Россия, Москва | РНИМУ им. Пирогова | Практикум по биоинформатике, магистерские программы "Медицинская Биоинформатика" и "Молекулярная Иммунология" |
| 2020– 2022 | Россия, Москва | МГУ им.Ломоносова | Практикум по структурной биологии (расчет структуры белка методом ЯМР, работа в программе PyMol), магистратура "Структурная Биология" |

Работа в ИБХ

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| 2025–наст.вр. | Научный сотрудник |
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Владение языками

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Гранты и проекты

| | |
|---------------|---|
| 2021– 2023 | Структура и метаболизм люциферина биolumинесцентных энхитреид <i>Henlea</i> sp. |
| 2019– 2022 | Структурная биология мембранных белков для создания новых лекарственных и диагностических средств |

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10. **Myshkin MY**, Dubinnyi MA, Kulbatskii DS, Lyukmanova EN, Kirpichnikov MP, Shenkarev ZO (2019). CombLabel: rational design of optimized sequence-specific combinatorial labeling schemes. Application to backbone assignment of membrane proteins with low stability. *J Biomol NMR* 73 (10-11), 531–544, [10.1007/s10858-019-00259-z](https://doi.org/10.1007/s10858-019-00259-z)
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