OIF Virtual Young Investigators Symposium
Basic, Translational and Clinical Aspects of OI and Other Rare Bone Disorders

Abstract Submission Form
Deadline: October 1, 2020

The Osteogenesis Imperfecta Foundation (OIF) is seeking abstract submissions from young investigators working in the field of osteogenesis imperfecta (OI) and/or rare bone disease research to be presented at the 2020 OIF Virtual Young Investigator Symposium on Tuesday, November 17 from 5pm-8pm EST.

This virtual one-day meeting is being offered as a pre-meeting event for the 2020 International Conference on Children’s Bone Health’s Virtual Forum and is open to US and international participants. Investigators are welcome to submit abstracts to both meetings. Submissions for the Young Investigator Symposium will be reviewed by a scientific review committee and selected abstracts will be invited to present a 10-15 minute overview of their work with 5 minutes for questions from the audience. Investigators are encouraged to share novel findings. For more information about the ICCBH Virtual Forum visit www.iccbh.org

Eligibility and review process: The OIF will accept abstracts from medical and graduate students, post-doctoral fellows and young faculty (within 10 years of their last graduate degree) working both in the US and abroad. A scientific review committee, led by Dr. Morello, will evaluate and select abstracts for presentation. Investigators are encouraged to share unpublished research. Only presentation titles will be published. Areas of research include, but are not limited to: Osteogenesis imperfecta (OI), Paget’s disease of bone, Generalized Arterial Calcification of Infancy (GACI), Hypophosphatemia, Autosomal Recessive Hypophosphatemic Rickets (ARHR2), Osteopetrosis, Fibrodysplasia Ossificans Progressiva (FOP), Fibrous Dysplasia and McCune Albright Syndrome, Hypophosphatasia (HPP), Gorham-Stout Disease, Melorheostosis, Jansen’s disease, X-Linked Hypophosphatemia (XLH), Multiple Hereditary Exostoses (MHE), and Multicentric Carpotarsal Osteolysis Syndrome (MCTO).

Selected investigators will be invited to present a 10-15 minute overview of their work with 5 minutes for questions from the audience. Meeting registration is open to medical professionals interested in OI and rare bone disease research.

How to apply: Please complete and return this application to Jenny Stup at jstup@oif.org by October 1, 2020. For questions, contact Erika Carter at ecarter@oif.org or 301-947-0083.

The OI Foundation is committed to supporting research to help improve the quality of life for people living with OI. This often times includes engaging groups throughout the rare bone disease community, such as the Rare Bone Disease Alliance (RBDA www.rbdalliance.org). The OI Foundation is pleased to open this meeting to the entire rare bone disease research community. To learn more about the work of the OIF please visit www.oif.org.
OSTEOGENESIS IMPERFECTA FOUNDATION ABSTRACT SUBMISSION FORM

TITLE OF PRESENTATION:

APPLICANT INFORMATION

Name (Last, First, M.I.)

Address:

City:  State:  Zip Code:  Country:

Email address:  Daytime Phone:

Please list education and professional training in chronological order beginning with college and ending with current affiliation:

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<th>Institution/Location</th>
<th>Dates of Attendance</th>
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Please explain your proposed presentation in no more than **500 words**. Be sure to include other contributors to the work.
Please describe your previous research experience in the space provided below.
Please list publications that you have authored or to which you have contributed. Please separate peer-reviewed publications from others.