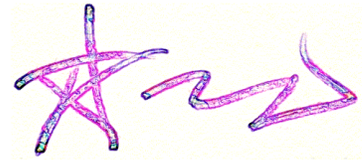


EuqriC Science Communications

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August 9th, 2019.

To whom it may concern,

The following paper, “**A case report of the diagnosis of Laron Syndrome using monoplex-PCR technology with a whole-genome amplification template**” by *Adina Neumann, Miguel Ángel Alcántara Ortigoza, Ariadna González-del Ángel, Felipe Camargo and Esther López-Bayghen*, has been thoroughly revised, edited and proofread by our team of professional science journalists, researchers and editors, with more than 15 years of experience in the Life Sciences area. We have ensured that the paper meets the quality standards of your Journal, minimizing all errors to speed the publication process, including peer review, and guaranteeing that the information stated in this paper possesses the meaning that the authors originally intended.

Should you have any questions, please feel free to contact us.

Best regards,

A handwritten signature in black ink, appearing to read 'Jacqueline Robledo', is positioned above a horizontal line.

Jacqueline Robledo, M. Sc.
Managing Editor,
EuqriC Science Communications