Correction: Clinical and genetic characterization of chanarin-dorfman syndrome patients: first report of large deletions in the ABHD5 gene

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Correction
Following the publication of this article [1], it was clarified that the clinical follow-up of one of CDS family described in the manuscript was performed by Dr. Amalia Sertedaki and Talia Kakourou. The authorship of the article has been changed accordingly. The submitting authors would like to apologise to Amalia Sertedaki and Talia Kakourou for this error and they would like to thank Catherine Dacou-Voutetakis for underlining the problem.

Authors’ contributions
CR carried out the molecular genetic studies and the interpretation of the results. RAC and LM made substantial contributions to interpretation of data and participated in manuscript preparation. AS, TK, SME, DP, AC and RC were involved in the clinical evaluation of patients and manuscript revision. DT made substantial contributions to conception, analysis and interpretation of data and drafted the manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

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Reference
1. Redaelli C, Coleman RA, Moro L, Dacou-Voutetakis C, Elsayed SM, Prati D, Colli A, Mela D, Colombo R, Tavian D: Clinical and genetic characterization of Chanarin-Dorfman Syndrome patients: first report of large deletions in the ABHD5 gene. Orphanet J Rare Dis 2010, 5:33.

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