

Dr. Giorgia Mandrile



Dr. Giorgia Mandrile is a medical geneticist at the San Luigi Gonzaga Hospital (Orbassano – Italy). She trained in University of Torino and in her PhD studied primary hyperoxalurias genetics. She is involved in basic and clinical research on PH, mainly focused on genotype-phenotype correlations and PH epidemiology. She is also involved in several advisory boards for PH. Currently, she is the Italian section of the European Primary Hyperoxaluria online database.

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