

CORRECTION



Correction: Myhre syndrome in adulthood: clinical variability and emerging genotype-phenotype correlations

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There was an error in the labeling of Table 1.

The values for codons 500 and 496 were switched. In the previous version, codon 500 was shown as 14/24 and codon 496 as 9/24, whereas this values should be reversed. This mistake is not present in the text, where the values are shown correctly.

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