

Bibliography

Bibliography of congenital muscular dystrophies: Series IV (2000) with Appendix II

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2. Introduction

This is a collection of articles that deal with congenital muscular dystrophies and/or cobblestone lissencephaly

which appeared in world literature in 2000, and is the 4th series of supplementary bibliography on the subjects.

In 1997, a comprehensive bibliography, comprising 423 original articles, 60 book chapters, 207 proceedings reports and 157 abstracts, was issued as a chapter of the book ‘Congenital muscular dystrophies’ (CMD) [1]. Keeping this as a basis, newly appearing articles on CMD in literature have been collected, compiled and published as a supplementary bibliography on CMD once every year for the last 3 years: Series I (1997) [2], Series II (1998) [3] and Series III (1999) [4].

Furthermore, those articles on CMD which were omitted from the previous list were gathered and published as an appendix to Series III (1999) [4]. We present here Series IV (2000) with Appendix II as the supplementary bibliography on CMD.

Please note the compiling policy which restricted the subject matter to only CMD; apparently related conditions such as congenital myotonic dystrophy, congenital non-dystrophic (non-progressive) myopathies, and type 1 lissencephaly/cortical dysplasias are excluded.

A copy of any article quoted in the Bibliography Series I–IV is available from the compiler (Y. Fukuyama) on request.

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3. Part A: Series IV – literature in 2000

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3.2.1. II-1 Fukuyama CMD

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3.3.1. III-1 Merosin deficiency

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3.3.3. III-3 General or unspecified or dealing with both types

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