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8	MULTIFOCAL MOTOR NEUROPATHY PRESENTING AS PSEUDODYSTONIA
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1 Concord and Royal North Shore Hospitals, 2 Sydney, New South Wales, Australia 3 E-mail: y.con@bigpond.com P: +61 2 9747 2688 4 F: +61 2 9744 5276 5 6 7 **Email Addresses of Authors** Dr Nidhi Garg: nidhi.garg@sydney.edu.au 8 9 Associate Professor Robert Heard: robert.heard@sydney.edu.au 10 Associate Professor Lynette Kiers: lynette.kiers@mh.org.au Professor Richard Gerraty: richard.gerraty@monash.edu 11 12 Professor Con Yiannikas: y.con@bigbond.com 13 14 **Running Title** MMN presenting as pseudodystonia 15 16 Keywords 17 18 Multifocal motor neuropathy, cramping, dystonia, peripheral nerve hyperexcitability, conduction block 19 20 21 22 **ABSTRACT** 23 24 Multifocal motor neuropathy is an immune-mediated neuropathy. The clinical presentation is 25 typically dominated by wasting and weakness. We describe four cases presenting with prominent 26 cramping resembling a primary movement disorder. All cases had features of focal motor conduction 27 block on neurophysiological studies. The involuntary movements resolved in all four patients 28 following treatment with intravenous immunoglobulin. The cases presented highlight an unusual 29 presentation of multifocal motor neuropathy and emphasise that peripheral nerve pathology can 30 present with movement disorders mimicking central nervous system disease. Furthermore, the 31 movement disorder appears to be particularly sensitive to standard therapy. 32 33 **INTRODUCTION** Multifocal motor neuropathy (MMN) is a rare immune-mediated neuropathy, typically presenting 34 35 with asymmetrical, and slowly progressive, distal upper limb wasting and weakness without sensory

1 abnormalities¹. A definitive diagnosis requires demonstration of focal motor conduction block on 2 neurophysiological studies with normal sensory nerve conduction across the lesion². The disorder is 3 associated with GM1 antibodies in approximately 50% of cases but the pathogenic role of such antibodies along with the mechanism of conduction block remains unclear¹. Cramp, fasciculation and 4 hypertrophy are well-recognized clinical features but prominent spasm and twitching mimicking a 5 6 primary movement disorder has not been described. We present four cases of MMN presenting with 7 unusual focal movements, which we describe as pseudodystonia. In addition, we discuss the spectrum 8 of involuntary movements that may be associated with peripheral nerve pathology along with possible 9 mechanisms. 10 **CASE REPORT** 11 12 Clinical characteristics of presented cases are summarized in Table 1. 13 14 Case 1 15 A 26-year-old female presented with a 3-month history of involuntary movement of the right hand 16 17 with recurrent flexor spasm of the wrist and fingers with intermittent jerking movements (Video 1). 18 The movements occurred both at rest and with action. There was possible weakness of thumb 19 abduction, but assessment of strength was limited due to continuous movements of the hand. There 20 were no sensory abnormalities. Neurophysiological studies revealed focal motor conduction block in the forearm segment of the right median nerve (Figure 1A). Sensory and other motor studies were 21 22 normal. Needle electromyography demonstrated fasciculations and non-rhythmic grouped discharges 23 of normal motor units in abductor pollicis brevis (APB) and flexor pollicis longus (FPL). Anti-24 ganglioside antibodies were negative. A diagnosis of possible MMN was made and she received 25 treatment with intravenous immunoglobulin (IVIg). Involuntary movements and neurophysiological abnormalities resolved completely within three months of treatment (Video 1 and Figure 1B). 26 27 Case 2 28 A 35-year-old female presented with a 12-month history of weakness of the right hand associated with 29 painful cramping, twitching and spasm (Video 2), which would often wake her from sleep. The 30 31 cramping preceded the weakness by several years. In addition, there was weakness of right ankle 32 dorsiflexion and right toe extension. The sensory examination was normal. Neurophysiological 33 studies revealed proximal conduction block involving the median nerves bilaterally and the right ulnar 34 nerve (Figure 1E). The lower limb compound muscle action potential amplitudes were preserved

suggesting conduction block proximal to the fibular head on the basis of neurogenic recruitment and

motor cortex stimulation studies. Sensory studies including somatosensory evoked responses were

normal. Needle electromyography showed fasciculation, myokymic discharges as well as cramp

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discharges in APB, FPL and ulnar finger flexors. Serum was positive for anti-GM1 antibody. IVIg
was commenced with partial improvement in muscle power and resolution of involuntary movements,
although the patient experiences mild muscle twitching in the week before her next IVIg dose is due.

5 Case 3

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- 6 A 51-year-old female presented with a 2-year history of cramps in the right hand, particularly
- 7 affecting the thumb and index finger, which would often curl down. In addition, there was occasional
- 8 cramping of the left hand. Clinical examination revealed weakness of APB, FPL, abductor digiti
- 9 minimi (ADM) and first dorsal interossei (FDI) bilaterally with no sensory abnormalities.
- Neurophysiology revealed focal motor conduction block in the forearm segment of the median nerves
- bilaterally (Figure 1C and 1D) and borderline conduction block of the right ulnar nerve above the
- 12 elbow. Sensory studies including somatosensory evoked responses were normal. Needle
- 13 electromyography demonstrated neurogenic recruitment with fasciculations and high-frequency
- discharges in APB and FDI without evidence of active denervation. Anti-ganglioside antibodies were
- negative at the time of diagnosis, but a repeat sample three years later was positive for anti-GM1 IgM
- antibodies. Cramping and weakness improved after commencement of IVIg, but recurred when IVIg
- dosing frequency was increased from four- to five-weekly.

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- 19 <u>Case 4</u>
- A 59-year-old male presented with a 6-month history of dystonic posturing of the left hand with ulnar
- 21 deviation of digits III-V and the wrist. On examination, there was moderate weakness of left FDI and
- 22 ADM. The sensory examination was normal. Nerve conduction studies demonstrated reduced
- 23 persistence of left median F-wave responses and absent left ulnar F-wave responses with normal
- 24 compound muscle action potential amplitudes and sensory studies. Needle electromyography revealed
- 25 fasciculation potentials in APB and ADM and markedly reduced recruitment in ulnar-innervated
- 26 muscles as well as a mild reduction in recruitment in APB. In the context of normal compound muscle
- 27 action potential amplitudes and significant weakness of FDI and ADM, the findings were in keeping
- with proximal conduction block. Serum was positive for anti-GM1 IgM antibodies. Pseudodystonia
- and muscle power improved following treatment with IVIg.

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DISCUSSION

- While movement disorders are primarily classified as disorders of the central nervous system, the four
- cases described highlight that peripheral nerve pathology may also be associated with prominent
- 34 involuntary movements which may dominate the clinical presentation and be difficult to differentiate
- 35 from central disease processes. Involuntary movements can be seen in a variety of peripheral
- disorders including immune-mediated neuropathies, peripheral nerve hyperexcitability syndromes,
- 37 following peripheral trauma and post-irradiation plexopathy.

Although the precise mechanisms for abnormal movements in peripheral nerve disease remain elusive, they are likely to be multifactorial and to vary between different disease processes. Such movements may result from activation of peripheral nerve fibers causing spontaneous impulse generation due to motor fiber hyperexcitability which may manifest as cramping, focal spasms, or twitching with semirhythmical movements. Furthermore, loss of or disruption to afferent pathways due to abnormal sensory input may secondarily leads to abnormal movements such as tremor and pseudoathetosis.

 In the cases presented, abnormal movements were unusual and striking, and may have been confused with dystonia or other central disease processes, such as chorea or athetosis. The phenomenology in Case 1 with the appearance of slow writhing movements, could easily have been mistaken for athetosis. However, when the phenomenology is examined closely, it is apparent that there is an element of sustained posturing in all cases, which is not consistent with chorea or athetosis. Furthermore, in all cases, components of the movements are segmental and in the distribution of one or more peripheral nerves with demonstrable motor conduction block on neurophysiological studies, although they progress in Cases 1 and 2 into widespread muscle involvement and fist formation. For example, Video 1 (00:45 – 01:15) demonstrates initial movements involving FPL, APB, and flexor digitorum profundus (II) prior to flexion of digits III-V followed by fist closure and wrist flexion. This phenomenology suggests initial median, followed by ulnar and subsequent widespread muscle involvement. Although ulnar nerve conduction studies were normal, there may have been proximal pathology involving the ulnar nerve not demonstrable on standard nerve conduction studies.

Other features which may support a peripheral generator and argue against dystonia include the absence of specific triggers (such as writing in the case of writer's cramp), the presence of movements at rest and lack of improvement with sensory tricks. Furthermore, in Case 2, movements occurred during sleep. This would provide an argument against dystonia, which typically disappears during sleep³. This feature was not reported in the other cases, but is an important aspect to elucidate in the clinical assessment as it may refine the differential diagnosis.

The presence of electrophysiological manifestations of peripheral nerve hyperexcitability (such as fasciculations, myokymia, continuous motor unit activity and cramp potentials) and evidence of conduction block in all cases along with the presence of segmental movements on examination supports a peripheral process underlying the primary aetiology. However, both videos demonstrate prominent cramping with synchronized movements of fist closure and wrist flexion. The mechanism of sustained contraction and synchronized activation of multiple muscles is uncertain. Evidence suggests that there is substantial shared central input to the motoneuron pools innervating the finger

1 muscles and hence central modulation through peripheral feedback may play a role. Furthermore, the 2 interdependence of finger movements is affected by biomechanical factors, such as tendinous 3 interconnections, which may also be a contributing factor⁴. 4 Although it is accepted that involuntary movements may occur in MMN, why the peripheral nerve 5 6 becomes hyperexcitable is unclear and the precise source of ectopic activity remains undefined. The 7 most plausible hypothesis stems from results of nerve excitability studies which suggest 8 hyperpolarisation in the nerve just distal to the site of conduction block. It has been proposed that 9 depolarisation at the lesion site leads to sodium influx into the distal axon resulting in overactivity of 10 the Na+/K⁺ pump with juxtaposed lengths of depolarised and hyperpolarised axonal membrane leading to ectopic discharges⁵. 11 12 13 Abnormal movements in the form of focal dystonia have also been described following peripheral trauma and nerve injury and may be related to central modulation⁶ and reorganization following 14 interruption to afferent sensory input. "Jumpy stump" is a term that has been used to describe jerking 15 movements of a stump developing after limb amputation. Post-traumatic tremor has also been 16 reported with neurophysiological studies in one case supporting a peripheral generator⁷. 17 18 19 Tremor in association with peripheral neuropathy ("neuropathic tremor") has been associated with 20 hereditary and immune-mediated neuropathies, including Guillain-Barré syndrome, chronic 21 inflammatory demyelinating polyneuropathy, MMN and in particular, IgM paraproteinaemic neuropathy, in which tremor occurs in up to 80% of patients⁸. The underlying pathophysiology is 22 likely to be driven by delayed and desynchronized sensory input, with impairment of central 23 modulation through the cerebellum and its pathways^{8,9}. Furthermore, in certain subgroups of chronic 24 inflammatory demyelinating polyneuropathy, the cerebellum may be a direct immune target 25 contributing to tremor¹⁰. 26 27 28 Pseudoathetosis may be seen in peripheral neuropathies and ganglionopathies, as well as posterior 29 column pathology. In contrast to tremor which is likely to result from slowed sensory conduction with desynchronisation, pseudoathetosis represents a more severe form of sensory dysfunction with 30 31 deafferentation and loss of sensory input due to neuronal loss or conduction block and results from 32 failure to maintain sustained motor output and muscle contraction. 33 34 Tremor associated with inflammatory and paraproteinaemic neuropathies tends to be refractory to immunomodulatory treatment although it may fluctuate with disease activity⁸. In contrast, the focal 35 36 movement disorders associated with the four cases of MMN presented in this series were exquisitely

sensitive to IVIg therapy supporting a rapidly reversible functional mechanism such as	antibody-
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2 mediated ion channel blockade.

CONCLUSION

- 5 Although movement disorders are usually of central origin, peripheral nerve disorders should be
- 6 included in the differential diagnosis. Assessment for MMN should be considered, particularly when
- 7 movements are focal, segmented and in the distribution of one or more peripheral nerves. Our
 - experience suggests that pseudodystonia associated with MMN is highly responsive to IVIg therapy.

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1	Dr N. Garg: Contributed to the design and conceptualization of the manuscript, analysis
2	and interpretation of data, and writing of the first draft and revising the manuscript.
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4	Professor R. Heard: Contributed to the analysis and interpretation of data and review
5	and critique of the manuscript.
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7	Associate Professor L. Kiers: Contributed to the analysis and interpretation of data
8	and review and critique of the manuscript.
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10	Professor R. Gerraty: Contributed to the analysis and interpretation of data and review
11	and critique of the manuscript.
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13	Professor C. Yiannikas: Contributed to the design and conceptualization of the
14	manuscript, analysis and interpretation of data, and writing of the first draft and review
15	and critique of the manuscript.
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26	N. Garg and R.N.S. Heard declare that there are no additional disclosures to report. L. Kiers
27	holds stock ownership with CSL and serves on scientific advisory boards for Baxter
28	Immunoglobulin, CSL Intragam 10 steering committee, National Blood Authority
29	Immunoglobulin Advisory Committee, Australia. R. Gerraty serves on the scientific advisory
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L5	REFERENCES

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- 1. Vlam L, van der Pol WL, Cats EA, Straver DC, Piepers S, Franssen H, van den Berg LH. 16 Multifocal motor neuropathy: diagnosis, pathogenesis and treatment strategies. Nat Rev 17 Neurol 2011; 8(1): 48-58. 18
 - 2. Joint Task Force of the EFNS and the PNS. European Federation of Neurological Societies/Peripheral Nerve Society guideline of management of multifocal motor neuropathy. Report of a joint task force of the European Federation of Neurological Societies and the Peripheral Nerve Society – first revision. J Peripher Nerv Syst 2010; 15(4): 295-301.
 - 3. Häger-Ross C, Schieber MH. Quantifying the independence of human finger movements: comparisons of digits, hands, and movement frequencies. J Neurosci 2000; 20(22): 8542-50.
 - 4. Evatt ML, Freeman A, Factor S. Adult-onset dystonia. Handb Clin Neurol 2011; 100: 481-511.
- 5. Kiernan MC, Guglielmi JM, Kaji R, Murray NM, Bostock H. Evidence for axonal membrane 27 28 hyperpolarisation in multifocal motor neuropathy with conduction block. Brain 2002; 125(Pt3): 664-675. 29
- 6. van Rooijen DE, Geraedts EJ, Marinus J, Jankovic J, van Hilten JJ. Peripheral trauma and 30 movement disorders: a systematic review of reported cases. J Neurol Neurosurg Psychiatry 31 32 2011; 82(8): 892-898.
- 7. Costa J, Henriques R, Barraso C, Ferreira J, Atalaia A, de Carvalho M. Upper limb tremor 33 induced by peripheral nerve injury. Neurology 2006; 67(10): 1884-1886. 34
- 8. Saifee TA, Schwingenschuh P, Reilly MM, et al. Tremor in inflammatory neuropathies. J 35 Neurol Neurosurg Psychiatry 2013; 84(11): 1282-1287. 36

- 1 9. Schwingenschuk P, Saifee TA, Katschnig-Winter P, et al. Cerebellar learning distinguishes 2 inflammatory neuropathy with and without tremor. Neurology 2013; 80(20): 1867-1873. 3 10. Querol L, Nogales-Gadea G, Rojas-Garcia R, et al. Neurofascin IgG4 antibodies in CIDP 4 associate with disabling tremor and poor response to IVIg. Neurology 2014; 82(10): 879-886. 5 6 Figure 1. Neurophysiology 7 Figure legend: Nerve conduction studies. Case 1: focal motor conduction block in the 8 9 forearm segment of the median nerve (A) with resolution following treatment with intravenous immunoglobulin (B). Case 3: bilateral conduction block in the forearm segment 10 of the median nerves (C and D). Case 2: proximal conduction block between Erb's point and 11 the axilla (E). 12 W: wrist; E: elbow; A: axilla; EP: Erb's point; IVIg: intravenous immunoglobulin; APB: abductor 13 14 pollicis brevis; a = amplitude; A = negative peak area; d = negative peak duration 15 16 Video 1 Video legend: Case 1: Flexor movements start in the thumb and finger in a median nerve 17 18 distribution and then progressively involve all the fingers and wrist in a cramping fashion with 19 later ulnar nerve involvement. In the second part of the video, involuntary movements have 20 dramatically resolved following intravenous immunoglobulin therapy. 21 22 Video 2 23 Video legend: Case 2: Video demonstrating semi-rhythmic twitching of digits with abduction and flexion movements. This is followed by flexor spasm of the wrist and fingers of 24 the hand. The patient tries to extend the fingers to reduce the amount of pain she is 25 26 experiencing. 27 **Table 1. Clinical Characteristics**
- 29 RUL: right upper limb; LUL: left upper limb; RLL: right lower limb; IVIg: intravenous

30 immunoglobulin