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Outcome of paediatric acute flaccid myelitis associated with enterovirus D68

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Paper for DMCN



Paediatric cluster of acute flaccid myelitis in Scotland associated with enterovirus D68 (EV-D68) infection and outcome at 18 months.

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Keywords:	EV-D68, AFM in Children, AFM and enterovirus, Acute Flaccid Myelitis, Acute Flaccid Paralysis

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[1 Case Series; 2 Figures, 1 Table, 2 Appendix online-only]

Outcome of paediatric acute flaccid myelitis associated with enterovirus D68 (EV-D68): a case series

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ABBREVIATIONS

AFM Acute flaccid myelitis

EV-D68 Enterovirus D68

NPA Nasopharyngeal aspirate

[Abstract]

 Enterovirus D68 (EV-D68) is an emerging infection associated with acute flaccid myelitis (AFM). Cases of AFM associated with EV-D68 infection have increased in recent years and the evidence for a causal link is growing. However, our understanding of the epidemiology, clinical features, prognosis, and neurological sequelae of EV-D68 requires ongoing surveillance and investigation. We report five cases of AFM in previously typically developing children (2–6y) from South East Scotland during September and October 2016 after infection with EV-D68 (all detected in the nasopharyngeal aspirates). All cases presented with significant neurological symptoms, which were severe in two cases requiring intensive care support because of respiratory paralysis. At 18 months follow-up, two cases remain ventilator-dependent with other cases requiring ongoing community rehabilitation. These cases represent one of the largest reported paediatric cluster of AFM associated with EV-D68 in Europe. The epidemiology and clinical information add to the knowledge base and the 18 months outcome will help clinicians to counsel families.

What this paper adds:

- Nasopharyngeal aspirate is more sensitive for viral isolation and isolated in all cases.
- Clinical outcome at 18 months after enterovirus D68 with acute flaccid myelitis provides information on extent of recovery and level of disability.

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Case Series

[main text]

Acute flaccid myelitis (AFM) is a clinical syndrome, characterized by weakness in one or more limbs with or without respiratory and bulbar muscle weakness, with no specific treatment. There are a number of viruses associated with AFM including polio, enterovirus A71, and flaviviruses.^{1–3} The European region was declared polio-free in 2002.^{4,5} Enterovirus A71 has been linked with outbreaks of AFM recently.^{6,7} Other non-polio enteroviruses have also been associated with AFM but more recently cases of AFM associated with Enterovirus D68 (EV-D68) have been reported.^{8–11}

Since its discovery in 1962, EV-D68 has been associated with sporadic cases of respiratory disease and minor outbreaks worldwide.¹² However, in 2014, the USA declared a nationwide outbreak and reported cases worldwide of over 2000 confirmed cases of EV-D68.^{10,13} There was an increased incidence of AFM during these EV-D68 outbreaks and the evidence for causality has increased significantly.^{9,14,15} In the UK, a cluster of neurological illness associated with EV-D68 was reported in South Wales in 2016.¹¹ There is a paucity of information available about the long-term prognosis and recovery from AFM associated with EV-D68 infection.

We report a cluster of AFM associated with EV-D68 in children with their clinical presentations, public health investigations, diagnosis, management, and outcome at 18 months. This case series will strengthen the existing literature; the clinical outcome at 18 months could help clinicians and families to target interventions.

METHOD

After one case of AFM associated with EV-D68 infection presented to the Royal Hospital for Sick Children Edinburgh on September 10th, 2016, a multidisciplinary incident management team was convened to investigate and set up monitoring for further possible cases. A clinical alert was sent to paediatric services throughout Scotland to be aware of potential cases of AFM associated with EV-D68 infection and to report all possible cases to the incident management team.

The following case definitions for AFM associated with EV-D68 infection were used for identifying subsequent cases after September 10th, 2016.

Possible Case: Person presenting with AFM in Scotland from September 10th, 2016 without other identified cause.

Confirmed Case: Person presenting with AFM in Scotland from September 10th, 2016 with laboratory confirmed EV-D68

Parents of cases were interviewed using a trawling questionnaire (Appendix S2, online supporting information) based on recall and covered exposures within 4 weeks before onset of

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symptoms. All confirmed cases were managed by paediatric neurology and intensive care specialists at the Royal Hospital for Sick Children Edinburgh.

Clinical investigations included brain and spine magnetic resonance imaging (MRI), electrophysiological studies, cerebrospinal fluid analysis, and virology testing (stool samples, throat swabs and nasopharyngeal aspirates [NPA] for a panel of viruses including EV-D6) using real time polymerase chain reaction. Stool specimens were all tested for the presence of polio virus. Fairly extensive neuroinflammatory investigations were carried out in the first two cases and not done in subsequent cases as the diagnosis became more apparent.

All children were regularly followed up by paediatric neurology, respiratory, physiotherapy, occupational therapy, speech and language therapy, psychology, and dietetic teams over the following 18 months. Data on this follow-up period were extracted from clinical records.

Parents gave written consent for the use of anonymised data for investigation and dissemination. NHS Lothian Caldicott Guardian's approval was obtained for data storage and dissemination.

RESULTS

Clinical presentation and epidemiology

All confirmed cases were residents within the South East Scotland region. Figure 1 displays the chronology of onset of prodromal and neurological symptoms for this cluster. From the questionnaires, there were no common sources (food, environmental exposures, recent travel, and previous direct contact with each other) identified. All children had received age-appropriate vaccinations, including inactivated polio vaccine. Their prodromal illness, clinical presentation, neurological weakness, investigations, treatment, and progress have been detailed in Table I. All presented with asymmetric flaccid weakness of varying severity, severe pain, three with bulbar involvement, and the two severely affected cases required intubation and respiratory support.

Investigations

All confirmed cases had EV-D68 detected from NPA samples after their admission. Two tested positive for EV-D68 on throat swab. However, in one case repeated throat swabs were negative and only confirmed on NPA. Viral typing confirmed EV-D68 B3 lineage. Faecal testing for polio was negative in all cases.

All children had a brain and spinal MRI and demonstrated an abnormal high T2 signal in the spinal cord grey matter (Fig. 2). Three of these cases showed high T2 signal particularly in the cervical spinal cord, with one case having entire cord involvement. Four of the children also showed signal abnormality in the dorsal pons and medulla. Cerebrospinal fluid results showed an elevated white cell count with a predominantly lymphocytic picture, negative for EV-D68 and negative for other viruses or bacteria.

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Treatment

One case did not receive any treatment other than supportive care and made excellent recovery. No significant improvement in clinical symptoms were observed after use of any treatment. Treatment with antihypertensives was commenced in the two most severely affected cases to control hypertension. All five children required significant input from a multidisciplinary therapy team to manage limb and truncal weakness. Gabapentin was used to control pain with good effect. Clinical psychologists were involved in supporting the children and their families.

Clinical outcome at 18 months

Clinical progress at 18 months post admission is summarized in Table I. Recovery for affected cases was most significant in the first 12 months; however, they continued to show improvement even beyond this time. One child had recovered almost fully with only mild lower limb weakness and a mild gait abnormality persisting. Two of the most severely affected required long-term home ventilation and were able to manage short periods without ventilation during the day while awake. Both children can feed orally with varying degrees of improvement in bulbar function. The autonomic dysfunction suffered during the early part of their admission resolved fully and antihypertensive therapy stopped. Limb pain lasted 6 months.

Persisting limb weakness and loss of function was predominantly in the lower limbs for two children and in the upper limbs for three children. Despite improvements, there has been persisting proximal limb weakness with muscle wasting in one or more proximal limbs in all cases apart from one. These four cases have been referred for evaluation for nerve transfer. We could not reliably use any standardized functional scoring system because of their unique pattern of weakness and adaptation. Unlike other motor disorders there seems to be persisting regeneration in some muscle groups.

DISCUSSION

The confirmed cases in this cluster were linked in time, place, and person and presented to hospital with significant neurological symptoms within a 2-week period. No clear hypothesis for a common environmental exposure was identified in this cluster based on recall of parents or guardians. After the presentation of this cluster we retrospectively retested all throat swabs and NPAs for EV-D68 which had previously tested positive for enterovirus. This retrospective testing found 59 samples between July and October 2016 were positive for EV-D68. These were in a select population of those presenting to hospital, mostly children with respiratory symptoms; however, these positive samples indicate wider circulation of EV-D68 in South East Scotland during this time. The baseline clinical data of those children reported to have EV-D68 without AFM did not identify any potential risk factors for AFM apart from the fact that the AFM cohort were much younger.

The age of presentation, pattern of limb weakness, bulbar involvement, cerebrospinal fluid results, MRI findings, and nerve conduction studies are like other reported case series.^{8–} ^{11,13} Two cases with autonomic involvement with severe hypertension and evidence of end organ damage during acute presentation had not been reported before. There was no

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association between steroid treatment and subsequent hypertension. One case developed worsening respiratory function after the use of general anaesthetic during lumbar puncture. It is not possible to determine whether this was exacerbated by use of anaesthesia, but we advise caution with use of general anaesthetic for suspected cases in future. We wanted to carry out follow-up MRI studies and we delayed this to avoid general anaesthesia. NPA was found to be the investigation which most consistently identified EV-D68 infection even when the throat swab was negative.

Different treatments (intravenous immunoglobulin, steroids, etc.) have been used in acute settings and the benefits of these treatments are not well understood. Although the numbers are small one case in our cohort made almost full recovery without any treatment. Further studies are needed to establish the benefits of these treatments.

Long-term outcome is variable and in our cohort all children can walk, talk, and feed with varying degrees of persistent neurological deficits at 18 months follow-up. Permanent proximal limb weakness with muscle wasting was present in four cases with varying severity. There seems to be continued improvement even at 18 months and rehabilitation in community settings with psychological support should be provided for future affected cases.

Recent developments have shown that infection with EV-D68 can cause AFM in murine models and the virus has been isolated from the spinal cord of infected mice.¹³ The short duration of prodromal illnesses before onset of acute neurology favours pathogenesis through direct destruction of the nerve by the virus. Direct or indirect association of viruses with neuroinflammatory disorders (N-methyl-D-aspartate, multiple sclerosis, Guillain-Barré syndrome, etc.) are rare but have been increasingly recognized. It is important to try to understand the susceptibility in those affected. This is a case series and hence there are limitations. There is an urgent need for a coordinated, cooperative, international approach to monitoring this emerging infection globally particularly as cases of AFM associated with EV-D68 infection emerge in other regions out with North America and Europe.^{16–18}

CONCLUSIONS

EV-D68 associated AFM present with short prodromal illness, asymmetric limb weakness, bulbar involvement, severe pain, and some autonomic involvement in previously well children, confirmed by MRI and EV-68 viral isolation. This condition has a devastating effect on the physical and mental health of the child and family and there is a lack of any prevention or treatment for complications. There are significant resource implications for health care services; not only in the provision of intensive care but also for long-term home ventilation and community rehabilitation. Further research should be targeted at the prevention of infection as well as understanding and preventing serious sequelae of AFM aiming to minimize complications. To achieve these goals, a global coordinated response from researchers, clinicians, virologists, and public health is required to understand and ultimately prevent this infection and its potential severe neurological sequelae. It is imperative that we act now as outbreaks of EV-D68 and reports of cases of AFM associated with infection become more common.

ACKNOWLEDGMENTS

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We would like to thank all the parents for their continued effort increase awareness of this condition and consenting to the publication of this report. We would also like to thank NHS Lothian and Fife Public Health Departments for all their assistance with this investigation. This research received no specific grant from any funding agency in the public, commercial or not-for-profit sectors. JS is an NHS Research Scotland (NRS) Fellow and acknowledges the financial support of NRS through NHS Lothian. The authors have stated that they had no interests that might be perceived as posing a conflict or bias.

SUPPORTING INFORMATION

The following additional material may be found online:

Appendix S1: NHS Lothian EV-D68 associated AFM study group.

Appendix S2: Parent interview trawling questionnaire.

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Clinical details

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Case 4

Case 5

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 Table I: Characteristics of confirmed cases of EV-D68 with associated AFM and clinical features at 18 months follow-up

Case 3

Case 2

Age at onset	2у	5у	4y	6у	2у
Sex	Male	Male	Female	Female	Male
Medical history	Mild asthma	None	None	None	CMPI
Prodromal symptoms (duration in days)	Fever, earache, reduced appetite (5)	Fever, headache, malaise, pyrexia (4)	Fever, headache, vomiting (2)	Fever, cough, headache, reduced appetite (1)	Fever, coryza (7)
Asymmetric severe flaccid limb weakness	Yes	Yes	Yes	Yes	Yes
Limb weakness	UL>LL	LL>UL	UL>LL	UL>LL	LL>UL
Cranial nerve involvement	Yes	No	Yes	Yes	Yes
Bulbar symptoms	Yes	No	Yes	Yes	No
Reduced or absent reflexes	Yes	Yes	Yes	Yes	Yes
Autonomic symptoms	Yes	No	No	Yes	No
Severe pain	Yes	Yes	Yes	Yes	Yes
MRI cord abnormality (location)	Yes (C2–C7)	Yes (C2–C7)	Yes (movement artefact)	Yes (C2–C7)	Yes (entire cord, maximum T1–T2)
MRI abnormality of pons and medulla	Yes	No	No	Yes	Yes
EV-D68 polymerase chain reaction	NPA (+) Throat swab (-) CSF (-) Stool (-)	NPA (+) Throat swab (-) Stool (-)	NPA (+) Throat swab (-) Stool (-)	NPA (+) Throat swab (+) Stool (-)	NPA (+) Throat swat (+) CSF (-) Stool (-)
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(left biceps) Nerve conduction studies	insertional activity, frequent fibrillations. No voluntary activity Normal sensory	performed Not performed	performed Not performed	insertional activity, widespread fibrillations. Two discrete complex repetitive discharges. No spontaneous activity Normal sensory	performed Not performed
studies	Reduced CMAP	periorited	-	Early CMAP	penomed
Treatment	IVIG, Steroids	None	IVIG	IVIG	IVIG, steroid
Respiratory support	Yes (long-term invasive ventilation)	No	No	Yes (long-term invasive ventilation)	No
Intensive care support	Prolonged	No	No	Prolonged	Short term
Hospital stay (days)	376	67	125	278	62
		At 18mo follo	ow-up		
Mobility	Walking short distances	Walking short distances.	Almost normal mobility.	Walking short distances.	Almost normal mobility.
Speech	Normal	Normal	Normal	Almost normal	Normal
Swallow	Normal	Normal	Normal	Almost normal	Normal
Weakness	Significant proximal upper limb (asymmetric) with muscle	Significant lower limb proximal weakness (asymmetric)	Mild distal unilateral lower limb weakness affecting gait	Significant proximal upper limb (asymmetric) with muscle wasting	Significant proximal upper limi (asymmetric) with muscle wasting
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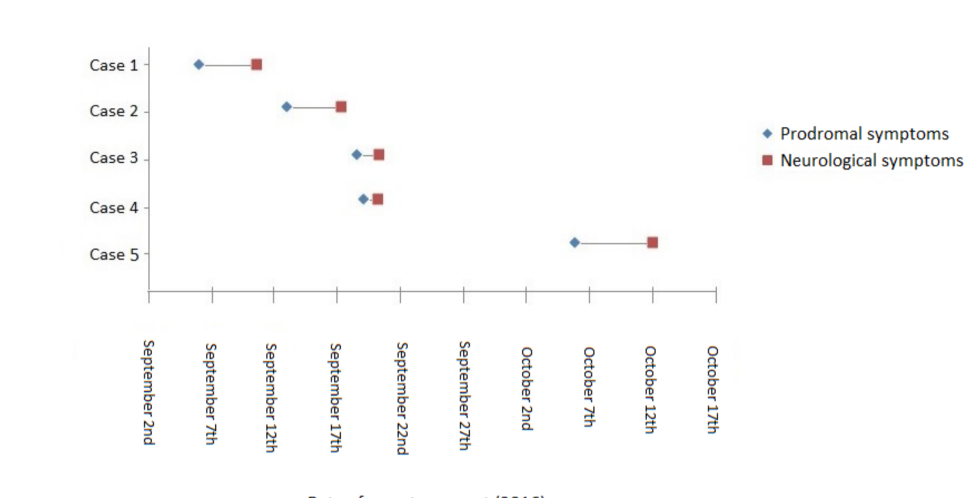
Scoliosis	Yes	Yes	No	Yes	No
Head tilt	Yes	No	No	Yes	Yes
Breathing	Tracheostomy and ventilatory support during sleep	Normal	Normal	Tracheostomy and ventilatory support during sleep	Normal

EV-D68, enterovirus D68; AFM, acute flaccid myelitis; CMPI, cow's milk protein intolerance; UL, upper limb; LL, lower limb; MRI, magnetic resonance imaging; NPA, nasopharyngeal aspirate; CSF, cerebrospinal fluid; CMAP, compound muscle action potential; IVIg, intravenous immunoglobulin. [Figure legends]

Figure 1: Timeline of symptom onset or confirmed cases of acute flaccid myelitis associated with enterovirus D68 infection

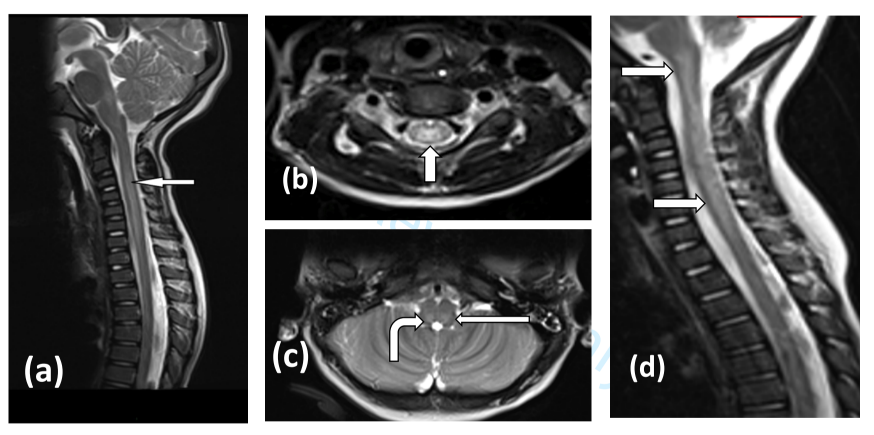
Figure 2: Magnetic resonance imaging (MRI) of the spinal cord and brainstem in acute flaccid myelitis caused by enterovirus D68. The saggital T2-weighted sequences (a,d) showing longitudinal hyperintense signal and the Axial T2-weighted sequences (b,c) showing hyperintensity in spinal cord grey matter (b) and dorsal brainstem (c).

 Paper for DMCN



Date of symptom onset (2016)

Figure 2. MRI spine and brainstem in AFM due to EV D68- Scotland



Magnetic resonance imaging (MRI) of the spinal cord and brainstem in acute flaccid myelitis caused by enterovirus D68. The saggital T2-weighted sequences (a,d) showing longitudinal hyperintense signal and the Axial T2-weighted sequences (b,c) showing hyperintensity in spinal cord grey matter (b) and dorsal brainstem (c).

Paper	for	DMCN
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DOB / /

Patient Name			DOB//
NHS Lothian Acute Neuro	logic Illness with Limb W	/eakness in Children: F	orm
Form to be completed by or in conjunction Part 1A Patient details	n with a physician who provided care a	to the patient during the neurologic	c illness.
1 .Today's date///	dd/mm/yyyy) 2.Name of person com	oleting form:	
3. HPT Board			
4. Name of physician who can provide ac	lditional clinical/lab information, if nee	eded	
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Past Medical History			
13 . Was patient admitted to a hospital?			
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14. Date of admission to first hospital			
Details of general ward	HDU or ICU ad	mission \Box yes \Box no & dates	_//
Ventilated 🛛 yes 🖾 no Date ventila	ation started///	Date came off ventilator/	
Outcome			
outtome			
15. Confirmed Enterovirus D68 🛛 yes	□no Sample		
16. Date of discharge from hospital	// Discharge to hom	e □yes, another hospital □yes	
Details:			
Clinical status two months after onset: [1At home 🗆 still in hospital 🗖 improv	ed Onot improved Oventilated	□Died· /
			//
Comments :			
	Mac Keith Pre	SS	

1	Case no					
1 2	Patient NameDOB//					
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5 6	Explain-unusual illness in the UK and we would like to gather more information in order to understand the illness better					
7 8 9	Family history:					
10	Details of who does the case live with (name/age)					
11 12						
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17	Details of pets in household					
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19 20	Anyone in the household with similar symptoms of intercurrent illness? yes no unknown add details here and here	tails/dates				
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25	Details of any contact with anyone else with similar symptoms					
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30	Details of nursery playgroup childminder school attended					
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34	Details of attendance at any health care waiting rooms eg GP, dentist, A&E					
35 36						
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39 40	Are they up to date according to UK childhood schedule? yes no unknown					
41	44 How many dama of instituted wells we size (IDM) and decomposited to have been reasized by					
42	44. How many doses of inactivated polio vaccine (IPV) are documented to have been received by					
43 44	the patient before the onset of weakness? & when	doses	□unknown			
45	45. How many doses of oral polio vaccine (OPV) are documented to have been received by the					
46 47						
47	patient before the onset of weakness? & when	doses	□unknown			
49	46. If you do not have documentation of the <i>type</i> of polio vaccine received:					
50 51	What is total number of documented polio vaccine doses received before onset of weakness?	doses	□unknown			
52 53	47. Details of last vaccination received & when?					
54	48. Flu vaccine details what when?					
55 56						
57						
58						

1		se no tient Name		DOB / /			
2 3		TRAVEL IN UK AND ABROAD in 4 weeks pre-ons	set of limb w		Yes	No	NS
4 5 5 7	1.	Spent any nights away from home WITHIN THE UK? (Holida friends or relatives, hotels, campsites, etc)	ays or busines	ss trips; staying at			
3 9 10 11		IF YES: Please state dates and From: / / / where		To: <u>/</u>	/		<u> </u>
12 13 14		Address visited, including postcode:					
5 6 7 8	2.	Spent any nights away from home OUTSIDE THE UK (Hol. friends or relatives, hotels, campsites, etc)	idays or busi	ness trips; staying at			
19 20 21		IF YES: Please state dates and country/resort		To: <u>/</u>	/		
22 23		Country and resort:		_			
24 25	Env	vironmental exposures: in the 4 weeks before onset, has the	patient:		Yes	No	Unk
26 27 28 29	Swa	vam or played in a swimming pool, indoor or outdoor? If Yes, o	details:				
80 81 82 83 84	Tak	ken part in water sports including sailing, canoeing, windsurfir	ng, fishing? If	Yes, details:			
5 6 7 8 9	cou	ken part in any outdoor activity that brought them into contact urses in fields or open land, including hill-walking, mountain-b tails:					
1 2 3 4	Any	y insect bites (including tick bites) ? If Yes, details:					
.5 .6 .7 .8	Woi	orked/played in a garden or allotment (including usual home g	ardening)? If	Yes, details:			
50 51 52 53 54		ed or had contact with any household chemical for cleaning (e s, details:	e.g. bleach, c	leaning sprays)? If			
55 56 57 58 59		ed or had contact with any garden chemicals, such as weedki tails:		ides? If Yes,			
50		Mac Keith Press					<u> </u>

1	Case no Patient Name DOB / ,	,		
2				
3 4 5 6	Food and drink exposures: In the 4 weeks before onset, has the patient:	Yes	No	Unk
7 8 9 10 11	Eaten any meals/ snacks bought from fast-food outlets? Fast-food outlets include any restaurant, stall or shop where food is paid for before it is eaten, such as sandwich bars, canteens, burger bars, kebab shops, fish and chip shows, hot dog stands, food outlets at markets			
12 13	Eaten any meals or snacks from any other restaurants, cafes, pubs or hotels?			
14 15 16	Eaten any meals or snacks at any function or gathering, e.g. at a party, reception, barbecue, picnic, etc?			
17 18 19 20	Eat any meals/snacks bought from grocers, bakers, supermarkets or delicatessens which were consumed away from the premises?			
21 22	Eaten any fish or shellfish?			
23 24	Eaten any tinned food?			
25 26	Eaten any new, unusual or imported foods? (e.g. wild berries, teas)			
27 28 29	Used any Chinese or herbal medicines etc (details)			
31 32 33 34 35 37 33 30 40 41 42 44 44 50 51 22 54 55 55 55				
56 57				
58 59 60	Mac Keith Press			

		14 days prior to onset of initial il	iness DOO_/_/_/
	ant to take this away and		
ay & date	Morning	Afternoon	Evening
		14	

Paper for DMCN

	Case no	
1	Patient Name	DOB / /
2		
3		
4		
5		
6	PART 2 Virology testing: details completed	on//2016 .Name of Dr completing this bit of the form:
7		
8		ID
9	73. Was CSF tested for the following	
10		Date of specimen collection//
11	pathogens?	
12		Enterovirus DCD: Desitivo Negativo Negativo
13		Enterovirus PCR: Positive Negative Not done
14		If positive: type: 🛛 Not typed
15		
16		Herpes Simplex Virus PCR: Positive Negative Not done
17		
18		Cytomegalovirus PCR: Positive Negative Not done
19		
20		Varicella Zoster Virus PCR: Positive Negative Not done
21		
22		Other pathogen identified: specify:
23		
24		Type of test:
25		
26		
27		
28		
29	74. Was a respiratory tract specimen tested	
30	for the following nother and	Date of specimen collection//
31	for the following pathogens?	
32		Enterovirus/rhinovirus PCR: Positive Negative Not done
33	Page 4 of 6	
34		If positive: type:
35		
36	Page 4 of 6	Adenovirus PCR: Positive Negative Not done
37		
38		If positive: type:
39		
40		Influenza virus PCR: 🗆 Positive 📄 Negative 📄 Not done
41		If positive: type:
42		
43		Other pathogen identified: specify:
44		
45		Type of test:
46 47		
47 48		
40 49		
50	75. Was a stool specimen tested for the	
51	••••••••••••••••••••••••••••••••••••••	Date of specimen collection// Date of specimen collection//
52	following pathogens?	
53		
54		Enterovirus PCR: Positive Negative Not done
55		
56		If positive: type:
57		
58		Poliovirus PCR: Positive Negative Not done
59		

60

	Patient Name	DOB_/_/ Poliovirus culture:
		Other pathogen identified: specify:
		Type of test:
	76. Was serum tested for the following	Date of specimen collection// Date of specimen collection//
	pathogens?	
		West Nile Virus: Positive Negative Not done
,		If positive, test type: 🗆 IgM 🖾 PCR
;		Other pathogen identified: specify:
		Type of test:
)	77. Describe any other laboratory finding(s)	considered to be significant
} 		
+ 5		
5		
7		
3		
9		
) 1		
2		
2 3 1		
2 3 1 5		
2 8 1 5 5		considered to be significant
2 6 7		
2 8 8 8 8 8 8 9		
2 3 4 5 5 7 3 9 9 2 3 4 5		
234557390 234557		
2 3 1		

Paper for DMCN

1	Case no Patient Name							_D(ОВ/	'/_			
2 3 4 5 6	PART 3 Clinical Neurological details completed on /	/2016	5 .N	ame c	of Dr o	comp	oleting	this	s bit o	f the	forr	n:	
7	17. Signs/symptoms/condition at <u>ANY</u> time during the illness:												
8 9 10	Date of onset of limb weakness:///												
11 12		R	ight	Arm		Left A	Arm		Right I	eg		Left	Leg
13	18. Since neurologic illness onset, which limbs have been acutely weak?												
14 15	[indicate yes(y), no (n), unknown (u) for each limb]	Y	Ν	U	Y	Ν	U	Y	Ν	U	Y	N	U
16 17	19. Date of neurologic exam (recorded at worst weakness thus far)						1	,					
18 19	(dd/mm/yyyy)				-		.//	'		_			
20 21	20 . Reflexes in the affected limb(s): (recorded at worst weakness thus far)	A	refle	exic/hy	oorefle	exic (C)-1) □ N	lorm	nal (2)	🗆 Нур	erre	flexic	(3-4+)
22 23	21. Any sensory loss/numbness in the affected limb(s), at any time during						Y N	ι	U				
24 25	the illness? (paresthesias should not be considered here)												
26 27	22 . Any pain or burning in the affected limb(s)? (at any time during illness)	Y	N	U	Y	N	U	Y	N	U	Y	N	U
28									Yes	No		Unk	nown
29 30	23. Sensory level on the torso (ie, reduced sensation below a certain level of	the tor	so)?	' (at an	y time	durir	ng illness	5)					
31 32	24. At any time during the illness, please check if the patient had any of the f	ollowir	ng cr	anial n	erve si	gns:							
33 34	Diplopia/double vision (If yes, circle the cranial nerve involved if kn	own: 3	3 / -	4 / 6)									
35 36	□Loss of sensation in face □ Facial droop □Hearing loss	□ Dysp	bhag	jia		Dysart	hria:						
37 38	25 . Any pain or burning in neck or back? (at any time during illness)												
39 40	26 . Bowel or bladder incontinence? (at any time during illness)												
41 42	27 .Cardiovascular instability (e.g, labile blood pressure, alternating tachy/bra	adycar	dia)	? (at ar	iy time	e durii	ng illness	5)					
43	28. Change in mental status (e.g, confused, disoriented, encephalopathic)? (at any	time	e during	; illnes	s)							
44 45	29 . Seizure(s)? (at any time during illness)												
46 47	30. Received care in ICU because of neurologic condition? (at any time during	g illnes	s)										
48 49	31 . Received invasive ventilatory support (e.g, intubation, tracheostomy) bec	ause o	f ne	urologi	cal cor	ditio	n?						
50 51													
52													
53 54													
55													
56 57													

Page	23	of	25
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	Case no Patient Name				DOB//
,	Within the 4-week period BEFORE onset of				
	limb weakness, did patient:	Yes	No	Unkn	
	32 . Have a respiratory illness?				33 . If yes, date of onset// & details of symptom
	34 . Have a fever, measured by parent				
	or provider and \geq 38.0°C?	0	~		35 . If yes, date of onset//
	36. Receive oral, IM or IV steroids?			2	
	37. Receive any other systemic				38. If yes, list:
	Immunosuppressant(s)?			14	
					2
	41 . Does patient have any underlying illnesses? Past Medical History				42. If yes, list
4	43. On the day of onset of limb weakness, did				
	patient have a fever? (see definition above)				
		1			1

1	Case no								-		
2	Patient Name								L	DOB//	
3 4											
5	Neuroradiographic findin	ngs: (Indi	cate based on	<u>most abnormal</u> st	udy)						
6 7	MRI of spinal cord 4	17. Date o	of study	//	_ (mm/dd	/уууу)					
8 9	48. Levels imaged:	□cervica	I □thoracic	□lumbosacral	□unknov	vn					
10 11	49. Gadolinium used?	□yes	□no	□unknown							
12 13 14 15	50. Location of lesions:	□cervi □conu □unkn	s E	∃thoracic cord ∃cauda equina			ected (if a	applicable): 5 2		c:	
16 17 18 19	For cervical and thoracic cord lesions		at areas of spi re affected?	inal cord	□predo □both e		gray mati fected	ter	□predo □unkno	minantly white wn	natter
20 21		54 . Wa	s there cord e	dema?	□yes	□no	□unkr	nown			
22											
23 24 25 26	For cervical, thoracic cord or conus lesions	55 . Did GAI	any lesions ei)?	nhance with	□yes	□no	□unkr	nown			
27											
28 29	For an describe la forme	EC Did	41 - 1								
30	For cauda equina lesions		the ventra l nance with GA		□yes	□no	□unkn	own			
31 32			the dorsal ne nance with GA		□yes	□no	Dunkn	own		ID	
33 34 35		62. Date □ yes		_//]unknown	(dd/mr	n/yyyy)	0				
36 37 38	59 . Any supratentorial (i.e, le cortical, subcortical, basal gator thalamic) lesions		□yes □no	o 🗆 unknown			2				
39 40			60 .If yes, inc	licate location(s)	□cortex	□su	bcortex	□basal g	anglia	□thalamus	□unknown
41 42				d any lesions with GAD?	□yes	□no	□unkr	nown			
43	62. Any brainstem lesions?		□yes □no	o 🛛 unknown							
44 45			63 . If yes, in	dicate location:	□midbra	ain 🗆	lpons	□medul	la	□unknown	
46 47				d any lesions with GAD?	□yes	□no	□unkr	nown			
48	65. Any cranial nerve lesions	s?	□yes □no								
49 50			66 . If yes, in CN(s):	dicate which	CN	∏unilat	eral 🗆 b	ilateral	CN	□unilateral	Dbilateral
50					CN	-	eral □b		CN	Dunilateral	
52				d any lesions with GAD?	□yes	□no	□unkr	00WP			
53 54 55	68. Any lesions affecting the cerebellum?										

69. Was an EMG done? Uyes no unknown If yes, date ___/__/___ (*mm/dd/yyyy*)

70. If yes, was there evidence of acute motor neuropathy, motor neuronopathy, motor nerve or anterior horn cell involvement? Uyes 🗆 no 🗆 unkn

Za. CSF Image: CSF	arliest and the		umbar punctu	re performed?	□yes □no	□unknow	n If yes, com	plete 72 (<i>If i</i>	more than 2 CSF e	examinations,
Puncture WBC/mm3 neutrophils lymphocytes monocytes eosinophils RBC/mm3 Glucose mg/dl Protein m 2a. CSF om LP1 Image: CSF Image: CS		en most abnorm	al)							
puncture WBC/mm3 neutrophils lymphocytes monocytes eosinophils RBC/mm3 Glucose mg/dl Protein m 2a. CSF		Date of lumbar		%	%	%	%			
om LP1 CSF		puncture	WBC/mm3					RBC/mm3	Glucose mg/dl	Protein mg/
2b. CSF	72a. CSF									
om LP2	from LP1									
1. Any other significant details of clinical illness?	72b. CSF									
1. Any other significant details of clinical illness?	from LP2									
tor Review Only	71. Any other s	significant details	of clinical illne	ess?						
	,	0								