Two Adult Siblings With Myotonic Dystrophy Type 1 With Different Phenotypes Presenting With Chronic Respiratory Insufficiency and Sleep Apnea Syndrome

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Myotonic dystrophy type 1 (DM1) is an autosomal dominant disease characterized by progressive muscle weakness and myotonia along with multiple organ system involvements. Overall, DM1 patients show reduced life expectancy, mainly due to respiratory or cardiac abnormalities. Chronic respiratory impairment is associated with increased morbidity in DM1. The main ventilatory dysfunction etiology in DM1 is complex, consisting of both peripheral respiratory dysfunction and central respiratory drive dysfunction as well as upper airway muscle dysfunction leading to obstructive sleep apnea syndrome (SAS) and aspiration. Advancements in early diagnosis of DM1 and management with non-invasive therapeutic tools have improved life expectancy for DM1 patients. We present herein two siblings with DM1, a thin elder brother and an obese younger sister with visceral fat accumulation. Although neither had voluntary symptoms related to respiratory dysfunction, their apnea-hypopnea indices revealed severe SAS and subsequent arterial blood gases studies showed hypercapnia as well as hypoxia, suggesting central nervous system involvement with peripheral respiratory dysfunction. Non-invasive positive pressure ventilation during sleep was started following pulmonary assessment. Respiratory function should be assessed in DM1 patients, even those free of respiratory symptoms, because respiratory muscle weakness occurs in a high percentage of these patients and will shorten their lives.

Keywords: myotonic dystrophy type 1, siblings, different phenotypes, respiratory insufficiency, sleep apnea syndrome
INTRODUCTION

Myotonic dystrophy type 1 (DM1) is the most common type of muscular dystrophy in adults, caused by a trinucleotide cytosine-thymine-guanine (CTG) repeat expansion in the dystrophia myotonica protein kinase (DMPK) gene on chromosome 19q13.3, and represents multiple organ symptoms including respiratory dysfunction (1). Because of respiratory dysfunction in DM1, besides a restrictive ventilatory pattern due to respiratory muscle weakness, involvement of the central nervous system (CNS) has recently been reported (2–5). Sleep apnea syndrome (SAS) is a prevalent disorder particularly among obese patients including those with metabolic syndrome (MetS) which is associated with risks of cardiovascular morbidity and mortality, type 2 diabetes, and all-cause mortality and are also reportedly associated with DM1, i.e., 63.6% of DM1 patients who underwent polysomnography had obstructive SAS due to weakness of the expiratory muscles, or hemidiaphragm palsy secondary to phrenic nerve dysfunction (3–7). We present herein siblings with DM1, an older brother and a younger sister with severe SAS accompanied by hypercapnia, suggesting that both probably had central respiratory dysfunction, although no voluntary symptoms were noted.

CASE REPORT

Case 1 (A 44-Year-Old Male)

A 44-year-old Japanese male, an office worker, was transported to our emergency department with a complaint of temporary loss of consciousness. He had experienced fatigue for 3 years and had fallen easily in his daily life starting 2 months before admission. He was feverish and had a cough and phlegm for several days before admission. On the way to work, he felt drowsy and fell down the stairs of the train station, necessitating transport to our hospital by ambulance. The peripheral blood analysis noted about 600 repeats in the elder brother and about 900 repeats in the younger brother and sister (Figure 1). Chest computed tomography (CT) showed severe infiltration in the upper posterior fields of both lungs as well as food debris in the esophagus. He was diagnosed as having aspiration pneumonia. His past medical history included diabetes mellitus that had been treated by a local doctor; glycated hemoglobin was approximately 7%. He had experienced ileus six times since age 30 years. He was not married. His father had died of dilated cardiomyopathy at the age of 70 and had also been thin. On the other hand, his mother was healthy but his maternal uncle was diabetic. His younger sister (Case 2) also had mild muscle weakness of the four extremities, handgrip myotonia, and diffuse muscle atrophy. Electromyography performed to test the biceps brachii muscle and femoral quadriceps muscle on the left side revealed frequent myotonic discharges. Because DM1 was strongly suspected, we recommended that, after providing informed consent during genetic counseling, he undergo genetic testing for DM1 together with his younger sister. The number of CTG repeats in the DMPK gene was abnormally expanded to about 600 repeats in the elder brother and about 900 repeats in the younger sister and sister.

Abbreviations: ABG, arterial blood gas; AH1, apnea-hypopnea index; CNS, central nervous system; CT, computed tomography; CTG, cytosine-thymine-guanine; DM1, myotonic dystrophy type 1; DMPK, dystrophia myotonica protein kinase; IGT, impaired glucose tolerance; MRI, magnetic resonance imaging; NIV, non-invasive therapeutic tools; NPPV, non-invasive positive pressure ventilation; SAS, sleep apnea syndrome.

TABLE 1 | Laboratory values and arterial blood gas of Case 1.

|   | a | b |
|---|---|---|
| (A) LABORATORY VALUES OF CASE 1 |   |   |
| WBC (×10^9/l) | 6,400 | 6,100 |
| RBC (×10^12/l) | 389 | 465 |
| Hb (g/dl) | 12.3 | 14.1 |
| PLT (×10^9/l) | 13 | 19 |
| CRP (mg/dl) | 8.9 | 0.1 |
| Total protein (g/dl) | 5.7 | 6.9 |
| Total bilirubin (mg/dl) | 1.3 | 0.6 |
| AST (GOT) (U/l) | 243 | 24 |
| ALT (GPT) (U/l) | 202 | 21 |
| Alkaline phosphatase (U/l) | 579 | 293 |
| Gamma-glutamyl transferase (U/l) | 193 | 71 |
| Creatine kinase (U/l) | 160 | 191 |
| Blood urea nitrogen (mg/dl) | 8.2 | 14.4 |
| Creatinine (mg/dl) | 0.5 | 0.6 |
| Sodium (mmol/l) | 142 | 146 |
| Potassium (mmol/l) | 4.1 | 4.9 |
| Chloride (mmol/l) | 104 | 104 |
| Uric acid (mg/dl) | 4.2 | 4.8 |
| Total cholesterol (mg/dl) | 189 | 204 |
| HDL-C (mg/dl) | 65 | 77 |
| Triglyceride (mg/dl) | 64 | 145 |
| Glucose (mg/dl) | 146 | 113 |
| Glycated hemoglobin (%) | 7.0 | 7.3 |
| (B) ARTERIAL BLOOD GAS VALUES OF CASE 1 |   |   |
| pH | 7.4 | 7.4 |
| PaCO₂ (mmHg) | 44.0 | 48.3 |
| PaO₂ (mmHg) | 170.9 | 81.5 |
| HCO₃⁻ (mmol/l) | 25.3 | 29.2 |
| BE (mmol/l) | −0.5 | 3.6 |
| SaO₂ (%) | 99 | 96 |

Numbers in bold type lie outside of the reference range.

a, Data obtained on admission to an emergency department with 10 L/min of oxygen from a mask with a reservoir.
b, Data obtained at outpatient care 3 months after discharge, before starting NPPV.
ALT, alanine aminotransferase; AST, aspartate aminotransferase; CRP, C-reactive protein; HDL-C, high-density lipoprotein cholesterol.
the younger sister, in contrast to those of healthy individuals who have 5 to 37 repeats (9), thereby confirming the diagnosis of DM1 in these siblings (Figure 2).

Case 2 (A 41-Year-Old Woman)

His younger sister was a 41-year-old woman, a housewife. She was obese according to the Japanese criteria (11), i.e., she was 156 cm in height, weighing 69.6 kg (102 cm waist circumference and a 164.2 cm² visceral fat area calculated employing abdominal CT). Slightly impaired fasting glucose levels had been pointed out on her medical check-ups since 3 years earlier, but she had never consulted a doctor. She had noted repeated falls and injuries for the past 5–6 years. She had married at the age of 37 and had undergone infertility treatment. When she was admitted for detailed examination to our hospital, she showed distal muscle weakness of the four extremities as described above, handgrip myotonia, and distal muscle atrophy, which were consistent with the characteristics of DM1. The laboratory data showed elevated fasting glucose with dyslipidemia, which met the Japanese criteria for metabolic syndrome (MetS) (Table 2A) (12), and an oral glucose tolerance test using 75 gram glucose showed impaired glucose tolerance (IGT) (Table 2B), followed by starting treatment with diet and exercise. Based on the presence of MetS, a sleep study was conducted using an apnomonitor Type 4 (SAS-2100, NIHON KOHDEN, Tokyo, Japan) (Table 3) (6, 7, 13). The apnea-hypopnea index (AHI) was defined as the average number of apnea and hypopnea episodes per hour of recording time in bed. The younger sister’s AHI was high at 55.9, which suggested that she had severe SAS, as defined by an AHI of 30 or more, though no snoring, apneic episodes, nocturnal dyspnea or signs of daytime somnolence were observed (14, 15). Her arterial blood gas (ABG) showed hypoxia accompanied by hypercapnia and her lung function test, using spirometry (CHESTAC-8900, Chest, Tokyo, Japan), revealed a decreased percent predicted vital capacity (VC) of 67.9% while Forced Expiratory Volume in 1 s (FEV1)/Forced Vital Capacity (FVC) (Gaensler index) was adequate at 87.9% (Table 3) (16).

We then also analyzed the respiratory function of the elder brother (Case 1), although, like his sister, he had not complained of voluntary symptoms suggestive of SAS. His AHI examined using an apnomonitor Type 4 was 55.1 and his ABG showed CO₂ retention without hypoxemia (Table 1B). His percent predicted VC was decreased, at 51.9%, while the Gaensler index was sufficient at 93.9%, similar to his sister’s values (Table 3). His aspiration pneumonia was regarded as possibly being due to DM1, which can cause expiratory muscle weakness. The indications for non-invasive positive pressure ventilation (NPPV) were assessed by an experienced pulmonologist. Based on chronic alveolar hypoventilation (patients with decreased percent predicted VC are at high risk), diurnal hypercapnia (PaCO₂ ≥ 45 mmHg) according to the ABG, evidence of nocturnal hypoventilation on nocturnal pulse-oximetry, defined as SpO₂ < 92% more than 4 times during the entire sleep time or >4% of the entire sleep time as well as AHI ≥ 10 events/h, these two siblings were started on NPPV. The NPPV consisted of 6 cmH₂O of inspiratory positive airway pressure, 4 cmH₂O
of expiratory positive airway pressure and a 12 bpm back-up rate in spontaneous/timed mode during sleep (4, 5, 16–18). These two siblings have been followed at outpatient care by the same pulmonologist and there has been no apparent worsening of symptoms since starting NPPV therapy. Both patients have shown good compliance. The ABG of the brother revealed slight elevation of PaCO$_2$, that is, pH 7.4, PaCO$_2$ 51.5 mmHg, PaO$_2$ 76.9 mmHg, HCO$_3^-$ 28.5 mmol/l, BE 2.1 mmol/l, and SaO$_2$ 95%.

Because the sister’s arteries are very thin, making it difficult for us to determine her ABG, spirometry was performed 3 months after initiating NPPV therapy and revealed almost no interval changes as compared to the baseline data before starting NPPV, with the direct value of VC being 1.9 L, VC 68.3% pred, direct value of FVC 1.84 L, FVC 66.2% pred, direct value of FEV1 1.52 L, FEV1 59.4% pred, and FEV1% (Gaensler index; FEV1/FVC) 101.8%.

As to other complications, the brother showed negative results on brain magnetic resonance imaging (MRI), echocardiography, abdominal ultrasound and, except for the diabetes, on endocrinologic examinations (8). The sister underwent brain MRI which showed localized white matter hypointense lesions in the right temporoparietal lobe, a common finding in DM1 (19), which may have accounted for her hypoventilation (5). Other studies including echocardiography, abdominal ultrasound, and endocrinological examinations yielded essentially normal results except for cataracts and IGT.

**DISCUSSION**

Patients with DM1 typically present with distally predominant muscular atrophy often resulting in an emaciated appearance. At the same time, MetS is significantly more common in DM1 patients than in the general population, which is considered to be due to a sedentary lifestyle related to muscle weakness (20). As mentioned above, body types differed markedly between our DM1 siblings, though both had respiratory dysfunction. Our Case 2 presented with MetS, prompting us to investigate her for sleep disorders. These investigations yielded evidence suggestive of severe SAS, while ABG and lung function tests detected type 2 respiratory failure, indicating that the main cause of her respiratory disorder was likely to be DM1 rather than the concomitant MetS (5, 6).

The DM1 phenotype is known to show extremely wide variability among affected patients, with some being asymptomatic while others have severe congenital manifestations (1, 8, 9). Among these symptoms and features, respiratory muscle weakness occurs in a high percentage of patients with DM1 in an early stage of the disease, i.e., the overall prevalence of ventilatory restriction is reportedly 36% (2, 21), and chronic respiratory failure can develop during its clinical course (5, 22, 23). Respiratory dysfunction in DM1 is predominantly a restrictive ventilatory pattern associated with alveolar hypoventilation, chronic hypercapnia, and sleep disturbance in the form of sleep apnea and sleep-related disordered breathing. Central respiratory drive is known to be significantly involved in the alveolar hypoventilation in DM1 (2–5), but the precise mechanism in the CNS has yet to be elucidated. Furthermore, there is no consensus in the literature regarding the relationship between CTG repeat length and the severity of respiratory dysfunction (3, 5). Sleep apnea may increase morbidity and mortality by leading to daytime respiratory failure, with expiratory muscle dysfunction often manifesting sooner than that of inspiratory muscles, possibly leading to early recurrent...
TABLE 2 | Laboratory values of Case 2.  

| (A) LABORATORY VALUES OF CASE 2 OBTAINED ON ADMISSION |  
|--------------------------------------------------------|  
| WBC (/l) | 7,100  
| RBC (×10^4 /ml) | 465  
| Hb (g/dl) | 12.6  
| Pt (×10^4 /ml) | 19  
| AST (GOT) (U/l) | 32  
| ALT (GPT) (U/l) | 37  
| Alkaline phosphatase (U/l) | 399  
| Gamma-glutamyl transferase (U/l) | 91  
| Creatine kinase (U/l) | 160  
| Blood urea nitrogen (mg/dl) | 11.1  
| Creatinine (mg/dl) | 0.4  
| Sodium (mmol/l) | 146  
| Potassium (mmol/l) | 4.5  
| Chloride (mmol/l) | 109  
| Uric acid (mg/dl) | 3.7  
| Total cholesterol (mg/dl) | 254  
| HDL-C (mg/dl) | 60  
| Triglyceride (mg/dl) | 188  
| Glucose (mg/dl) | 129  
| Glycated Hemoglobin (%) | 6.2  

| (B) DATA OBTAINED BY 75 GRAM OGTT TEST THE DAY AFTER ADMISSION OF CASE 2 |  
|---------------------------------------------------------------|  
| Glucose (mg/dl) & Insulin (mU/ml) |  
| Fasting | 90 & 6.5  
| OGTT Post-30 min | 151 & 66.6  
| OGTT Post-60 min | 162 & 50.3  
| OGTT Post-90 min | 175 & 47.9  
| OGTT Post-120 min | 176 & 56.9  

Numbers in bold type lie outside of the reference range.  

ALT, alanine aminotransferase; AST, aspartate aminotransferase; HDL-C, high-density lipoprotein cholesterol.  

OGTT, oral glucose tolerance test.  

pneumonia due to a weak cough and insufficient airway clearance (5, 8). Despite restrictive lung syndrome progressing slowly (24), problems and complications related to the respiratory system still represent the main cause of reduced life expectancy in DM1, i.e., respiratory problems are reportedly among the leading causes of death in adult DM1 patients (51–76%) with most of these patients dying between 50 and 60 years of age (5, 25). Furthermore, since DM1 affects the CNS, studies have pointed out that an abnormal sensitivity of the central respiratory drive to chemical blood changes, particularly of \([\text{CO}_2]\), might contribute to the pathogenesis of respiratory dysfunction in DM1 (2, 26). Because we could not examine the causes of chronic respiratory insufficiency of our patients in detail, we can rule out neither respiratory muscle weakness nor CNS involvement as potential causes of respiratory insufficiency. Furthermore, DM1, though not necessarily by itself, can contribute, along with other coexisting sleep breathing disorders, to the conditions of patients like ours. However, we consider the main cause to be DM1 in our cases, based on the results of whole body screening examinations showing no signs of coexisting sleep breathing disorders such as obstructive pulmonary diseases, emphysematous lung lesions or intestinal lesions suggestive of restrictive lung diseases.

The development of chronic, progressive respiratory failure can have prognostic effects, particularly in terms of survival, in many neuromuscular diseases including DM1. Early detection of respiratory dysfunction in addition to early diagnosis of DM1 is crucial. Therapeutic management employing non-invasive therapeutic tools such as NPPV may increase the life expectancies of DM1 patients (3, 4).

CONCLUSION

Respiratory system function should be examined in DM1 patients, even those with minimal or no symptoms, because respiratory muscle weakness occurs in a high percentage of DM1 patients, and chronic respiratory failure may develop (5, 22, 23).

DATA AVAILABILITY

All datasets generated and analyzed for this study are included in the manuscript and/or the supplementary files.

ETHICS STATEMENT

Both subjects gave written informed consent for publication of the information and images related to this case report. As this
is a case report without experimental intervention, no formal research ethics approval was required.

**AUTHOR CONTRIBUTIONS**

GK, MK, ST, JI, TNi, and KM substantially contributed to the acquisition, analysis, or interpretation of the data obtained. KO, HI, and MF drafted the manuscript or revised it critically for important intellectual content. TNa and YS provided approval for publication of the content. All of the authors agreed to be accountable for all aspects of this study in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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