Successful institutional care for behavioral and psychological symptoms of dementia in a repopulated area after the 2011 Fukushima disaster: A case report

Yoshitaka Nishikawa1,2,3,4 | Hidehito Niimura2,5 | Akihiko Ozaki6,7 | Yuko Kimura2 | Tomohiro Morita8 | Toyoaki Sawano9,10 | Hiroaki Saito11 | Masaharu Tsubokura3,8,10

1Kawauchi Intensive Care Homes for the Elderly, Futaba-gun, Japan
2Kawauchi Village National Health Insurance Clinic, Futaba-gun, Japan
3Department of Internal Medicine, Hirata Central Hospital, Ishikawa-gun, Japan
4Department of Health Informatics, School of Public Health, Kyoto University, Kyoto, Japan
5Department of Neuropsychiatry, Keio University School of Medicine, Tokyo, Japan
6Department of Breast Surgery, Jyoban Hospital of Tokiwa Foundation, Iwaki, Japan
7Department of Epidemiology and Biostatistics, Teikyo University Graduate School of Public Health, Tokyo, Japan
8Department of Internal Medicine, Soma Central Hospital, Soma, Japan
9Department of Surgery, Minamisoma Municipal General Hospital, Minamisoma, Japan
10Department of Public Health, Fukushima Medical University School of Medicine, Fukushima, Japan
11Department of Gastroenterology, Sendai Kousei Hospital, Sendai, Japan

Correspondence
Yoshitaka Nishikawa, Kawauchi Intensive Care Homes for the Elderly, Futaba-gun, Japan.
Email: ynishikawa-tky@umin.ac.jp

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Key Clinical Message
Caregiving in a long-term facility played a key role in improvements of this patient’s behavioral and psychological symptoms of dementia, which also led to a reduced caregiver burden on her family members. Considering the global population aging trend, the lesson from this case may apply to other settings beyond disasters.

Keywords
dementia, Fukushima nuclear accident, long-term care, nursing homes, social capital

1 INTRODUCTION

In March 2011, Fukushima, Japan, experienced a nuclear disaster. An 81-year-old woman was diagnosed as behavioral and psychological symptoms of dementia (BPSD) during a long-term evacuation in June 2012; thereafter, her cognitive ability further worsened further. Yet, admission to a long-term care facility in her home area alleviated her BPSD.
BPSD impair the dementia patients’ daily lives and can impose a significant caregiver burden. It is therefore important for BPSD patients to receive formal long-term care. In limited-resource areas without abundant informal family-caregiving resources, there is a particular need for formal caregiving for those with BPSD. However, there is little information on the provision of care for patients with BPSD in limited-resource areas.

In March 2011, an earthquake and tsunami struck northeastern Japan, triggering the Fukushima Daiichi Nuclear Power Plant (FDNPP) accident, which forced the evacuation of residents living in surrounding areas. Kawauchi Village, Fukushima, lies in a mountainous area 10-30 km southwest of the FDNPP (Figure 1). Even before the disaster, this area had a clinic but no hospital facilities providing intensive long-term care. Because radiation exposure in Kawauchi Village was within the acceptable level for living, the former residents started to repopulate from April 2012. The resident’s limited access to health care after the disaster was previously described. The average age of the local population remaining after this disaster was older because many younger residents did not return out of fear of radiation exposure. This major disaster also created social and environmental challenges. As the mass evacuation changed the residents’ living and social environments, it also affected those with chronic diseases, such as diabetes, dyslipidemia, and impaired motor function. Along with several chronic diseases, this disaster also exacerbated symptoms in those with dementia. It has been reported that changes to circumstances, such as evacuation, could affect patients’ mental and cognitive conditions. Both cognitive functions and BPSD deteriorated among patients who experienced this disaster. A previous study has also shown an association between housing damage and cognitive decline, because of new onset of depression and disruption of social contacts. In response to the increasing demand for care for the aging population in this impacted village, a long-term care facility was established in November 2015.

The case of Kawauchi Village involves rapid changes in the age distribution of the local population combined with limited age-care resources; it is thus representative of other areas affected by nuclear disaster. Here, we report on a patient who developed dementia after the Fukushima nuclear disaster and who received long-term care in a new facility established in the repopulated area. This case emphasizes the importance of providing formal care for patients with BPSD, and their family members, in limited-resource areas.

2 | CASE REPORT

An 81-year-old woman with dementia was admitted to the long-term care facility in Kawauchi Village after the 2011 Fukushima disaster. Before the disaster, she had lived in...
Kawauchi Village with her husband and their son’s family. She had experienced a stroke about twenty years ago without neurological sequelae and had no other remarkable past medical history. On March 11, 2011, the Great East Japan Earthquake struck the area, triggering the FDNPP accident. The patient and her husband evacuated to a shelter apart from other family members because of the mandatory evacuation order. After moving to temporary housing in Koriyama, she lived alone with her husband, while some relatives lived in temporary housing nearby.

In June 2012, she presented with memory loss. Physical examination revealed no neurological findings. Her symptoms were stable and were followed up in an outpatient clinic. A cognitive function test had not been performed before the disaster, but the patient and her family did not notice dementia symptoms at that time. Symptoms including agitation, irritability, aggression, and personality change emerged in October 2012, particularly during attendance at daycare activities. At that time, the patient’s score on the Mini-Mental State Examination (MMSE) was 14 out of 30, indicating dementia. At first, she was suspected of having Pick disease because of the personality change. Brain computed tomography revealed hippocampal atrophy without frontotemporal lobar degeneration. Brain magnetic resonance imaging (MRI) revealed a previous putamen lacunar infarction, which could not completely rule out vascular dementia. Brain MRI also showed hippocampal-dominant atrophy that was 14.6 times as atrophic as other parts of the brain. The neurological changes accompanied by aggressive behavior were consistent with severe BPSD. Together with clinical symptoms and the results of MRI, the diagnosis was confirmed as Alzheimer’s disease.

The patient was treated with memantine 10 mg/d, but the BPSD, such as undressing, hiding or stealing objects, agitation, and aggression, continued to worsen. Although the family members made their best effort to provide informal care to the patient, as the patient and her husband lived apart from the family members in the temporary housing, the family’s support alone could not cover the entire care of the patient. In February 2015, on order of the designated psychiatrist, she was mandatorily hospitalized for medical care and protection. At the time of admission, the patient’s cognitive function could not be measured. Although its use is controversial in dementia patients, percutaneous endoscopic gastrostomy was performed because she suddenly began to refuse oral intake in September 2015; this decision was based on the family’s values and preferences. The patient’s BPSD, including agitation and aggression, were slightly alleviated and she was transferred to the nonpsychiatric ward. In January 2016, she was moved to the long-term care facility in Kawauchi Village with her husband.

At the time of the patient’s admission to the long-term care facility, she was immobilized and needed constant care. She could not understand any explanations or instructions and her cognitive function could not be measured. Although she could not understand explanations, she could swallow and exhibited only mild dysphagia in a modified water-swallowing function test (MWST) and scored 3b (swallowed successfully but with wet hoarseness). She was started on oral intake of semisolid foods in addition to gastronomy feeding.

She was admitted to the facility with her husband, and they spent the daytimes together while she received care. Some caregivers in the facility had been raised in Kawauchi Village and had known the patient long before her admission. These caregivers conversed with the patient about life in Kawauchi and their mutual acquaintances. Her son and relatives living nearby continued to visit the facility several times a week and took the patient in a wheelchair for walks around the facility grounds. Caregivers respected the patient’s motivation to continue her daily activities such as dressing, bathing, and eating. Her swallowing function improved to an MWST score of 4 (swallowed successfully without choking or wet hoarseness) in June 2016, when she ate all meals orally without a gastronomy feeding tube.

The patient’s BPSD improved sufficiently that the gastronomy feeding was successfully removed in March 2017. She still exhibited cognitive impairment (MMSE: 13) but without severe BPSD by December 2017. She ate meals orally, used a wheelchair independently, and lived with her husband in the long-term care facility in the repopulated village.

3 | DISCUSSION

This patient’s BPSD improved after her admission to a long-term care facility in a repopulated village following the 2011 Fukushima disaster. Despite development of dementia and further deterioration after the 2011 Fukushima disaster, she was able to receive enhanced support in the facility from caregivers already known to the patient and the patient’s relatives who lived nearby. The caregiver burden experienced by family members was thus offset by the facility.

This case report showed that medication along with long-term care in a facility located in the patient’s home area helped to manage the patient’s BPSD and improve her eating function. First, she received care provided with respect and dignity by caregivers, which might have helped alleviate the BPSD, as previously noted in Japanese research. Some of the institutional caregivers were originally from the patient’s village and conversed with her about topics related to the village and their mutual acquaintances, which seemed to provide mental support. In addition, family members visited the facility frequently because it was located in the repopulated village. Furthermore, the patient spent daytimes with her husband in the facility, as a previous report suggested that group migration mitigated the loss of cognitive function even after a disaster. Building social capital along with antipsychopathic treatment has
benefits in terms of improving the symptoms of BPSD.\textsuperscript{21} Examples of heightened social capital that may contribute to the alleviation of BPSD symptoms include conversations with caregivers who know the patient well, admission with a family member, frequent visits by relatives who live nearby, and familiar landscapes. In this case, the long-term care facility played an important role in providing care and aggregating social capital.

This case also suggested that institutional caregiving helped to complement the informal caregiving by family members in the repopulated village after the 2011 Fukushima disaster. Caregiver burden refers to the physical and mental stress of caregivers caused by informal caregiving, which is reportedly associated with health issues such as sleep deprivation, depression, and suicide.\textsuperscript{22} After the 2011 Fukushima disaster, the affected areas faced an increasing need for caregiving because of depopulation and an increase in the aging population.\textsuperscript{7,23} However, because of these demographic changes, it may have been difficult for the patients’ family to provide all of the care needed.\textsuperscript{24,25} In Kawauchi Village, the percentage of residents aged over 65 years has increased from 34.0\% before the disaster in 2011 to 37.9\% in 2015; and this percentage was higher than Japan’s national average of 26.7\% in 2015.\textsuperscript{26} Similar to the demand experienced by Kawauchi Village, demand for long-term care is starting to exceed the supply because of increasing life expectancy and declining fertility rates, particularly in limited-resource areas. In such areas, there is a need to alleviate caregiver burden experienced by families. Institutional caregiving is a potential solution for decreasing informal care burden on family caregivers in an aging society.

In conclusion, we report on a patient developed dementia and BPSD after the Fukushima nuclear disaster and experienced an alleviation of her BPSD after admission to a long-term care facility with her husband. Caregiving in the facility also played an important role in reducing caregiver burden on her family members living in the area after the disaster.

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ETHICS STATEMENT
For this type of study, ethics approval by an institutional review board is not required. Written informed consent was obtained from the patient.

CONFLICT OF INTEREST
None declared.

AUTHOR’S CONTRIBUTION
YN: drafted the article. HN, AO, YK, TM, TS, HS, RK, and MT: performed the critical revision of the article for important intellectual content. All the authors involved in the interpretation of the case, and conception and design, and final approval of the article.

ORCID
Yoshitaka Nishikawa http://orcid.org/0000-0003-3313-1990

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