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SHORT REPORT

National audit of cognitive assessment in people with pwMND A national audit of cognitive assessment in people with motor neurone disease (pwMND) in Scotland

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Abstract
Cognitive and behavioral abnormalities are recognized as an integral part of Motor Neurone Disease (MND) and occur at all stages of the disease. The early detection of cognitive and behavioral symptoms in MND is critical. Such symptoms are only reported when we explicitly ask, evaluate, document, and assess. In the National Institute for Health and Care Excellence (NICE) MND guideline (2016), formal cognitive and behavioral assessment is incorporated in MND management and is fundamental to providing appropriate care to pwMND. Cognition is explicitly stated in 14 separate recommendations in the guidelines. The NICE guidelines therefore constitute pre-defined standards which we audited. This audit highlights that health professionals increasingly recognize the significance of cognitive screening in MND and follow more structured approaches in implementing this compared to previous years.

Keywords: Motor neurone disease, cognition, cognitive assessment

Background
Cognitive and behavioral abnormalities are recognized as part of Motor Neurone Disease (MND). Approximately 15% of people with MND (pwMND) fulfill diagnostic criteria for fronto-temporal dementia (typically behavioral variant), while milder changes in behavior and cognition affect a further 35% of pwMND (1,2). Cognitive and behavioral impairments have important functional implications for pwMND and their families: they increase caregiver burden, reduce survival, and impact care planning (3). These impairments may not be readily apparent at clinical interview, hence the need to assess using standardized measures. In the National Institute for Health and Care Excellence (NICE) MND guideline (2016), cognitive and behavioral assessment is incorporated in MND management and is fundamental to providing appropriate care to pwMND (Table 1) (4,5,6).

The Edinburgh Cognitive and behavioral ALS screen (ECAS) has been specifically developed and validated (7,8). In Scotland, assessment of pwMND is provided either via neuropsychology or members of the multidisciplinary team (MDT), usually the clinical nurse specialist. Furthermore, Scotland benefits from an integrated national healthcare team for MND closely aligned to the Scottish MND register (relaunched in 2015 as Clinical Audit Research and Evaluation of MND – CARE-MND) (9). This is an electronic platform for prospective, population-based research.

We conducted an audit to analyze “real-world” experience of cognitive assessment in pwMND in Scotland. Using the NICE guidelines, we measured whether we follow best practice.

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Objectives

To audit cognitive assessment in pwMND in Scotland against two predetermined standards by NICE:

- Following MND diagnosis, cognitive screening is recommended in pwMND.
- Care planning should be adapted for people with cognitive impairment and behavior change.

Methods

During 2016, several educational activities occurred within Scotland and the UK to encourage cognitive and behavioral screening of MND as part of routine assessment. At the end of 2016, we presented the preliminary results throughout Scotland and identified obstacles to screening.

Data were captured from the CARE-MND Register. Two time periods were audited:


Results

(1) Between January 2015 and December 2016, 393 new cases were captured (Table 2). ECAS was undertaken in 36% (n = 140). Almost 33% (n = 131) did not have any form of cognitive assessment. No data were recorded in the remaining cases (31%; n = 122); this is likely attributed to the following: there is no funded clinical neuropsychology in some areas; long-term staff sickness; in some regions, cognition was considered satisfactory and therefore cognitive screening was not offered.

Total ECAS scores were available in 67% (n = 94) and 50% had cognitive impairment (defined by the cutoff total ECAS scores) (Figure 1(a)). There was a non-significant trend toward fewer interventions (non-invasive ventilation, gastrostomy, riluzole) in cognitively impaired patients (Figure 1(b)).

(2) In 2017, data were available from 193 incident cases (Table 2). Around 57% (n = 110) underwent cognitive screening, with ECAS performed in 86% (n = 95), and other forms of cognitive assessments in the rest. Total ECAS scores were available in 85% (n = 80); 54% of those had cognitive impairment (Figure 1(a)). Comparing those with and without cognitive impairment on treatment, there was no statistically significant difference (Figure 1(b)).

Discussion

This audit highlights that health professionals increasingly recognize the significance of cognitive screening in MND and follow more structured approaches compared to previous years. Cognitive testing increased by 21% and ECAS was the most commonly performed assessment. Of those assessed, 56% were cognitively impaired, consistent with previous population studies.

During the second cycle, cognitive assessments were not recorded for 11.4%. The reasons include:
(1) delay in data input in the CARE-MND platform, (2) ECAS was not collected because it was too soon after diagnosis. We offer pwMND follow-up appointments with an MDT member 4–6 weeks post-diagnosis. This delay prevents misconstruing emotional overburdening with cognitive impairment.

Regarding treatment, there was a non-significant trend toward fewer interventions in cognitively impaired patients and other groups have also identified this (10). Those with cognitive change or FTD may have difficulty coping with NIV (11). Larger cohorts are required confirming the association between interventions and cognitive impairment.

### Table 2. Patient demographics.

<table>
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<tr>
<th></th>
<th>Audit phase 1</th>
<th>Audit phase 2</th>
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<tbody>
<tr>
<td>Median age of onset</td>
<td>67 (Median Abs deviation = 8)</td>
<td>66 (Median Abs deviation = 8.5)</td>
</tr>
<tr>
<td>% males, % females</td>
<td>63%, 37%</td>
<td>61%, 39%</td>
</tr>
<tr>
<td>Median disease duration (in months)</td>
<td>27 (median Abs deviation = 13)</td>
<td>25 (Median Abs deviation = 13)</td>
</tr>
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</table>

Figure 1. (a) Implementation of cognitive assessments Phase 1 & 2. In the UK, the cutoff for the ECAS total is 105. A score AT or BELOW 105 suggests that a person may have cognitive impairment. “Pending” were the cases where ECAS was undertaken but the data were not yet interpreted or inputted to the CARE-MND platform. (b) Interventions by cognitive status.
impairment in MND. Specifically, more data are needed on NIV compliance, the role of patients, carers and physicians in the decision-making process, and end-of-life practices in pwMND with cognitive change. Our future work involves investigating the profile of supportive/palliative interventions in pwMND with mild or moderate cognitive impairment, behavioral changes and the frontotemporal syndrome. Further studies should elucidate how to ensure appropriate interventions are not inadvertently denied.

Additional pathways should be developed for cognitive/behavioral screening for pwMND such as: (1) Masterclasses/training days to enhance health care professionals’ knowledge in ECAS; (2) Establishment of dedicated ECAS clinics or incorporating psychologists into clinic; (3) Ongoing support and access to neuropsychology services. Neuropsychological intervention helps the MDT manage the particularly complex cases.

Finally, the online register has contributed to accurate and effective data capturing. CARE-MND Platform is a national resource with an average case ascertainment coverage of 99% (7). Centralizing data from across Scotland are unique and beneficial for patient care. Ongoing multidisciplinary collaboration is essential for mastering data capture, attaining compliance with evidence-based recommendations and continuing quality improvement.

Acknowledgements

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Declaration of interest

Abrahams S. is the author of the Edinburgh Cognitive and Behavioral ALS Screen (ECAS). The authors report no conflicts of interest. The authors alone are responsible for the content and writing of this article.

Data sharing statement

Requested for access to data should be addressed to the corresponding author and the CARE-MND Consortium.

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