Cognitive Change in Donepezil Treated Patients with Vascular or Mixed Dementia

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ABSTRACT: Background: Vascular dementia (VaD) and mixed Alzheimer’s disease (AD/VaD) are common. How best to monitor treatment is not clear. Our objective was to compare responsiveness and construct validity of change scores, following donepezil treatment, of the standardized Mini-Mental State Examination (sMMSE) and other measures potentially usable in primary care.

Methods: A six-month, outcome measurement study. The Disability Assessment for Dementia (DAD), CLOX-1 and 2, Phonetic Fluency, a short Neuropsychiatric Inventory, (the NPI-Q), Clinical Global Impression (CGI) and the SymptomGuide™ (SG) were measured. Construct validity was tested by correlating change scores, and responsiveness by calculating standardized response means (SRMs).

Results: Of 148 treated patients, 116 completed. The mean sMMSE increased by 0.7 (95% Confidence Interval (CI) = -0.005, 1.41; p=0.06; SRM= 0.15). There was no statistically significant difference in the DAD. The NPI-Q (-1.4; 95% CI = -2.08, -0.72; p<0.01; SRM=0.24), CLOX-1 (0.9; 95% CI = 0.19, 1.61; p<0.01; SRM=0.21), CLOX-2 (0.9; 95% CI = 0.17, 1.63; p=0.03; SRM=0.26), Phonetic Fluency (0.9; 95% CI = 0.19, 1.61; p=0.02; SRM=0.21) and SG (0.35; 95% CI = 0.20,0.51; p<0.01; SRM=0.28) each detected significant improvement. The CGI suggested improvement in 74 completers (64%) – mostly “minimal” (44/116, 38%) – while 21/116 (18%) were worse. Change scores at 24 weeks were at best modestly correlated with each other (range -0.22 to 0.30).

Discussion: Different measures showed different responsiveness, in a setting in which the mean treatment effect seems to have been small, but clinically detectable. Patient-centered and executive function measures might be useful in vascular and mixed dementia.

RÉSUMÉ: Changerments cognitifs chez les patients atteints de démence vasculaire ou mixte traités par le donépézil. Contexte : La démence vasculaire (DV) et la maladie d’Alzheimer mixte (MA/DV) sont des maladies fréquentes. Toutefois, on ne connaît pas quelle est la meilleure façon de faire le suivi de ces patients au cours du traitement. Le but de notre étude était de comparer la réactivité et la validité conceptuelle des taux de changement observés au mini examen de l’état mental (MMSE) et à d’autres instruments de mesure utilisables dans les soins de première ligne, suite au traitement par le donépézil. Méthode : Nous avons effectué une étude des résultats obtenus après six mois de traitement. Nous avons utilisé l’échelle d’évaluation de l’invalidité associée à la démence (DAD), les tests du cadran de l’horloge CLOX-1 et 2, le test de la fluence verbale phonologique (TFVP), un inventaire neuropsychiatrique abrégé, le NPI-Q, le test d’impression clinique globale CGI et le Guide des symptômes (GS). la validité conceptuelle a été évaluée au moyen des corrélations des scores de changement et la réponse au traitement a été évaluée par le calcul des moyennes des réponses standardisées (MRS). Résultats : Parmi les 148 patients traités, 116 ont complété les évaluations. La moyenne au MMSE a augmenté de 0,7 (IC à 95% : -0,005 à 1,41; p = 0,06; MRS = 0,15). Nous n’avons pas noté de différence significative au point de vue statistique pour le DAD. Le NPI-Q (-1,4; IC à 95% : -2,08 à -0,72; p < 0,01; MRS = 0,24), le CLOX-1 (0,9; IC à 95% : 0,19 à 1,61; p < 0,01; MRS = 0,21), le CLOX-2 (0,9; IC à 95% : 0,17 à 1,63; p = 0,03; MRS = 0,26), le TFVP (0,0 IC à 95% : 0,19 à 1,61; p = 0,02; MRS = 0,21) et GS (0,35; IC à 95% : 0,20 à 0,51; p < 0,01; MRS = 0,28) ont tous détecté une amélioration significative. Le CGI suggérait une amélioration chez 74 sujets (64%) et cette amélioration était minime (44/116, 38%), alors que 21/116 étaient pires (18). Les scores de changement à 24 semaines étaient au mieux corréllés faiblement entre eux (écart de -0,22 à 0,30).

Discussion : Différentes mesures ont montré une réactivité différente dans le contexte d’un effet moyen du traitement qui semble avoir été faible mais détectable au point de vue clinique. Des mesures centrées sur le patient et des mesures de la fonction exécutive pourraient être utiles dans la démence vasculaire et la démence mixte.
extent to which vascular lesions give rise to dementia in the face of traditional AD markers is disputed.5,9

The treatment of VaD and mixed AD/VaD with cholinesterase inhibitors, although generally recommended,10-12 can be controversial. Sceptics point to the small size of the treatment effects in the two major clinical trials of donepezil in VaD.13,14 More recent work, which attempted to learn from the two trials, likewise was inconclusive. While patients with VaD who were treated with donepezil showed improvement on a neuropsychological test battery — Vascular-Alzheimer Disease Assessment Scale-Cognitive Subscale15 — this was not reflected in the clinical assessment.16 In consequence, the 4th Canadian Consensus Conference on the Diagnosis and Treatment of Dementia recommended treatment of patients with possible Alzheimer’s disease and a cerebrovascular component, but not pure vascular dementia.17

Although it might simply be that donepezil does not much help people with vascular and mixed dementia,18 the small treatment effect might also reflect the insensitivity of commonly used outcome measures to detect change, especially if subtle effects on executive function are present.1,11,19 For this reason, there was interest in evaluating both commonly employed clinical trial outcome measures, as well as measures that might better detect executive function and thereby be more sensitive to clinically important changes in VCI. Our objectives were: to evaluate, in an open label clinical trial, whether the standardized Mini-Mental State Examination (sMMSE) captured any impact of donepezil on general cognition in primary care patients with clinico-radiographically defined VaD and mixed AD/VaD, and; to evaluate the responsiveness and construct validity of measures of executive dysfunction, behaviour, and activities of daily living (ADLs). Finally, as with any compound, we studied the safety and tolerability of donepezil in subjects with these types of dementia.

2. METHODS

2.1 Patients, setting, sample size and selection criteria

This was a six-month, open-label study, conducted between June 2005 and April 2008 at 30 primary care clinics across Canada. Treatment was with 5 mg/d donepezil for six weeks, to be increased to 10 mg/d for the following weeks; adjustable dosing was permitted. Patients were first screened and then examined at a separate baseline visit within one week, and then at 6, 12, and 24 weeks post baseline.

We expected a mean improvement on the sMMSE of 0.73 points at six months, with a standard deviation of 3.5,20,21 A two-sided α = 0.05 t-test required 205 patients for 85% power to detect a significant improvement from baseline; with an anticipated drop-out rate of approximately 20%, the plan was to enrol 260 patients in the study.

To be eligible for inclusion, patients had to meet both clinical and radiographic criteria. The clinical inclusion criteria, in addition to written informed consent by the patient and a primary caregiver, were: 50 years-of-age or older, with a reliable caregiver; VaD and/or mixed AD/VaD diagnosed using DSM-IV-TR criteria;22 mild-moderate severity, operationalized in people with a Hachinski Ischemia Score > 423 and a Functional Assessment Staging Tool (FAST)24 score of 4 or 5. Radiographic inclusion criteria specified lesions compatible with a diagnosis of cerebrovascular pathology: e.g., the presence of one strategic (e.g., thalamic) or more than one infarct, two or more lacunes, or significant leukoaraiosis (e.g., extending > 3 mm from the ventricles).25 Patients were excluded if they had recently participated in other studies, were clinically unstable (including a systolic blood pressure > 160 mmHg and/or diastolic blood pressure > 95 mmHg) or had a history of intolerance or hypersensitivity to donepezil.

2.2 Measures

Data were recorded on patient and caregiver demographics, the patient’s medical history, general and neurological examination, vital signs, electrocardiogram (ECG), serum chemistry, hematology and urinalysis and concomitant medications. Vascular risk factors included hypertension, diabetes, hypercholesterolemia, transient ischemic attack (TIA), stroke, coronary artery disease, atrial fibrillation, smoking and body mass index. The six-item Hamilton Depression Scale (HAM_D-6) was used for screening.26

The primary efficacy measure was the 30-point sMMSE (a higher score is better).27 Other measures were chosen for their familiarity and/or potential feasibility for use in Canadian primary care settings. Activities of daily living were measured using the Disability Assessment for Dementia (DAD), which includes assessment of planning, initiation and execution of basic and instrumental activities (a higher score is better).28 Executive function was assessed using two variants of the clock drawing tests, CLOX-1 and CLOX-2 (a higher score is better)29 and the Phonetic letter fluency test (a higher score is better).30 In the CLOX test, two clocks are drawn, one free hand (CLOX 1) and one copied (CLOX 2). Each CLOX test was scored separately. Phonetic letter fluency is a measure of how many words a subject can generate in one minute. The total number of words constituted the score. To assess behavior we used a brief form of the Neuropsychiatric Inventory, the NPI-Q.31 The brief NPI-Q is used to measure severity of behavioral manifestations of dementia together with the level of distress that each symptom causes the main caregiver. The 10-item NPI-Q is based on: delusions, hallucinations, agitation/aggression, depression/dysphoria, anxiety, elation/euphoria, apathy/indifference, disinhibition, irritability/lability, and aberrant motor behavior. The 12-item version (used here) adds sleep and appetite/eating disorders. These scores are each rated as 0 (absent); 1 (mild); 2 (moderate) or; 3 (severe) giving a range from 0-36. The NPI-Q also measures the caregiver distress associated with each symptom, ranging from 0 (no distress) to 5 (very severe), so that the NPI-Q distress (NPI-Q-D) score ranges from 0 to 60. Lastly, global functioning was measured using the Clinical Global Impression (CGI). The CGI measures a physician’s global impression of a subject’s clinical condition, at baseline in terms of severity (CGI-S), and at follow-up as interval change from baseline (CGI-I). At baseline, the subject was rated on a scale ranging from 1 (normal, not at all ill) to 7 (among the most extremely ill patients). At follow-up visits, the subject was rated on a 7-point scale ranging from 1 (very much improved) to 7 (very much worse).

In this study, the SymptomGuide™ (SG) was introduced to assess individual responses to treatment.32 The SG provides descriptions of common dementia symptoms. In the version used
here, 9–12 descriptors were provided for each of 27 symptoms, in a paper format. (There are descriptors for 60 symptoms in the online format: http://www.dementiaguide.com/symptomguide/) Caregivers completed the SG in an interview, facilitated by the CGI rater, where the caregivers chose from the list of symptoms those that were the most troubling to them. Next they ranked the symptoms, with the least troublesome ranked as 1, and each other is ranked higher and their frequency of occurrence. The SG score is the average frequency of the symptoms in the individual with the weights represented by their ranks. As some troublesome symptoms can occur very frequently (e.g., many times a day in looking for a missing object, whereas others can occur less frequently, e.g., refusing to attend a regular weekly outing), the data were represented on a log scale to accommodate the large range, with the differences in the log scores between the visits represented graphically as the ratio of the last score to the baseline score. The SG method explicitly incorporates patient and caregiver input, which aids the understanding of clinical meaningfulness.33

Adverse events (AEs) that were observed, reported by the patients/caregivers, or elicited by direct questioning, were recorded as to severity, seriousness, cause and outcome.

2.3 Analysis

The efficacy analyses was based on the full analysis set sample, defined as all patients who received at least one dose of donepezil and who had baseline and at least one post-baseline assessment of efficacy. Missing data were imputed using the last observation carried forward (LOCF) endpoint, defined as the last assessment recorded post baseline. Note that both LOCF and observed case (OC) analyses were conducted. Neither method is without controversy,34,35 but given the emphasis here on detecting observed treatment effects, OC analyses are reported. The least squares (LS) mean and standard error were presented for variables analyzed with the Mixed Model. The covariates that were fitted as fixed effects in all the primary and secondary analyses were Center, Week, and Baseline score. The term subject was fitted as a random effects term. In addition to descriptive statistics, scores were compared between the baseline and 24 week visits, for patients who completed the study. Analysis of variance was used to compare continuous scores and chi-square for categorical variables. For the SG, which must combine frequencies across a large range, the non-parametric Mann-Whitney U test was used to analyse change. The CGI is a change score, so that the comparison measure is no change from baseline (or a score of “4”), where “1” = very much better, and “7” = very much worse). For all tests, a significant difference was set at p<0.05.

To assess the clinical detectability of the various measures, standardized response means were calculated (where the mean before/after difference is divided by the standard deviation of the change).36 Interpretation is as with other standardized effect sizes. Here of most relevance is that the threshold for clinical detection is the Standard Response Mean (SRM) of 0.2; SRMs in the range of 0.2-0.4 are said to be small,37 which is common in trials of cholinesterase inhibitors in dementia.38 To assess their construct validity, the change scores were correlated (Pearson and Spearman correlation coefficients (when appropriate) were used) with each other and compared against the global clinical judgments.

2.4 Ethics

The ethics committees of each participating centre approved the protocol. A central ethics review was used for practices that were not affiliated with an academic centre. In addition, the protocol was approved by the Research Ethics Committee of the Capital District Health Authority, Halifax, NS and the Sunnybrook Health Science Centre for the sub-studies on the SymptomGuideTM and imaging, respectively.

3. RESULTS

3.1 Baseline clinical characteristics

Due to challenges with patient recruitment for this study, enrolment was stopped when 200 patients had been screened rather than extend the recruitment period even further to achieve the originally targeted 260 patients. Of the 200 screened patients, 148 received treatment and took at least one dose of donepezil.
From these, 137 had at least one post-dose efficacy measure and 116 completed the study, for a drop-out rate of 21.6% (Figure 1). Patients who discontinued were slightly older, but otherwise were of similar demographic and clinical characteristics, including the proportion with VaD versus mixed AD/VaD. Overall, about half of the patients (77; 52%) had a diagnosis of VaD, with a mean duration of illness of 2.1 years. The 71 patients (48%) with mixed AD/VaD had a mean illness duration of 1.8 years. Ten patients had incomplete data at the final visit, although CGI data were available on all patients.

The patients were predominantly elderly (87% were aged 65 years or older) with a slight female preponderance (55%), although the male/female split was almost even in VaD, compared with more women (42; 62%) in mixed AD/VaD. Otherwise, the diagnostic subgroups had the same mean body mass index (BMI) (27.2). Clinically, the study population consisted mainly of people with mild dementia. The mean baseline sMMSE score was 23.4, with VaD patients having a higher mean baseline sMMSE (24.8) than those with mixed AD/VaD (22.0). By definition, the Hachinski Ischemia Scale (HIS) scores were higher in VaD than in mixed AD/VaD; overall, patients represented a range of ischemic injury, with most having a moderate ischemic burden (mean HIS=8.0 ± standard deviation (SD) 5.0). Otherwise, HAM-D-6 (mean 4.3 ± 0.5) and CGI-S (3.4 ± 0.4) were consistent with a group with mild-moderate dementia, as specified in the inclusion criteria. All patients reported at least one co-morbid illness. Vascular co-morbidities were the most common (total reported = 305), especially hypertension (present in 105 people), dyslipidemias (59) heart disease (43) and diabetes (30). Seventy-three psychiatric co-morbidities were reported, chiefly anxiety (17) and depression (28). Patients were using one to two drugs on average at baseline; the most common drug classes were antithrombotic agents (39), analgesics (38), and anti-hypertensives (37).

The majority of the caregivers were women (Table 1). Most patients were cared for by a spouse and most were retired.

### 3.2 Efficacy measures

Regarding the primary efficacy measure, although the sMMSE scores increased slightly at Week 24 (LS mean change = 0.72, Table 2), this difference was not statistically significant (p=0.06, Table 2) and the effect size was small (SRM = 0.15, AD – Alzheimer’s disease; SD – standard deviation; VaD – vascular dementia

<table>
<thead>
<tr>
<th>Measure</th>
<th>Primary Diagnosis</th>
<th>Baseline</th>
<th>Week 12</th>
<th>Week 24</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>sMMSE</td>
<td>All Subjects</td>
<td>23.4 (4.5;10-30)</td>
<td>23.98 (4.6;10-30)</td>
<td>24.12 (4.7;11-30)</td>
<td>0.06</td>
</tr>
<tr>
<td>DAD</td>
<td>All Subjects</td>
<td>80.1 (18.6;25-100)</td>
<td>81.4 (20.3;10-106)</td>
<td>78.9 (23.6;9-5-100)</td>
<td>0.90</td>
</tr>
<tr>
<td>NPI-Q</td>
<td>All Subjects</td>
<td>6.8 (5.4;0-28)</td>
<td>5.4 (4.7;0-23)</td>
<td>5.4 (5.1;0-30)</td>
<td>0.01</td>
</tr>
<tr>
<td>NPI-Q-D</td>
<td>All Subjects</td>
<td>7.4 (6.8;0-30)</td>
<td>6.1 (6.6;0-35)</td>
<td>5.8 (6.9;0-41)</td>
<td>0.11</td>
</tr>
<tr>
<td>CLOX -1</td>
<td>All Subjects</td>
<td>8.7 (3.9;0-15)</td>
<td>8.7 (4;10-15)</td>
<td>9.6 (3;9;5-15)</td>
<td>0.006</td>
</tr>
<tr>
<td>CLOX -2</td>
<td>All Subjects</td>
<td>11.2 (3.2;9-12)</td>
<td>11.5 (3;4;9-15)</td>
<td>12.1 (2;9;6-15)</td>
<td>0.03</td>
</tr>
<tr>
<td>DAD</td>
<td>VaD</td>
<td>11.6 (3.1;0-15)</td>
<td>11.8 (3;3;9-15)</td>
<td>12.6 (2;4;4-15)</td>
<td>0.12</td>
</tr>
<tr>
<td></td>
<td>Mixed AD/VaD</td>
<td>11.6 (3.4;0-15)</td>
<td>11.2 (3;6;0-15)</td>
<td>12.6 (2;4;4-15)</td>
<td>0.03</td>
</tr>
<tr>
<td>Phonic</td>
<td>All Subjects</td>
<td>7.4 (3.7;0-17)</td>
<td>8.1 (4;1;0-18)</td>
<td>8.3 (4;2;0-21)</td>
<td>0.01</td>
</tr>
<tr>
<td></td>
<td>VaD</td>
<td>8.2 (3.5;1-17)</td>
<td>9.1 (3;9;1-17)</td>
<td>8.9 (4;1;2-21)</td>
<td>0.39</td>
</tr>
<tr>
<td></td>
<td>Mixed AD/VaD</td>
<td>6.6 (3;7;0-16)</td>
<td>7.2 (4;2;0-18)</td>
<td>7.6 (4;1;0-17)</td>
<td>0.05</td>
</tr>
</tbody>
</table>

CLOX-1 - clock drawing (hands free); CLOX-2 - clock drawing (copied); DAD - Disability Assessment for Dementa; NPI-Q - brief Neuropsychiatric Inventory; NPI-Q-D - brief Neuropsychiatric Inventory – Distress; sMMSe - standardised Mini-Mental State Examination; VaD - vascular dementia
Apart from ClOX-2 and Phonetic Fluency (Table 2), there was no difference in results between the mixed AD/vaD patients and the vaD patients (data not shown). The percentage of patients with no response (a change from baseline of zero or less) on the sMMSE decreased from 47.5% at Week 12 to 40.1% at Week 24. After six months of treatment with donepezil, the DAD showed no statistically significant difference (Table 2) with a very small effect size (SRM = -0.05, Table 3). Likewise, the NPI-Q-D scores were not significantly better.

Other tests were more responsive. The ClOX-1 and ClOX-2 tests both showed statistically significantly better performance (Table 2), although with small effect sizes (Table 3). The NPI-Q similarly recorded significantly fewer disturbances (Table 2) with very small effect sizes (SRM = -0.24, Table 3). The Phonetic Fluency test showed statistically significant increases after six months (Table 2) with small, yet detectable effect sizes (SRM = 0.21, Table 3).

The SG showed statistically significant improvement with donepezil treatment (Figure 2). For example, it shows some improvement (with the ratio up to two times) in 46 patients, with worsening occurring in 14 patients. The effect size of the differences were small (SRM = -0.28, Table 3). The mean change in the CGI was 3.32 (95% confidence interval = 3.10-3.54). Similarly, the most common CGI-I response was improvement, mostly “minimal” improvement (Figure 3). The Spearman correlation between the changes in the SG (expressed by the ratio SG at the last week to the baseline visit) and CGI was 0.32 (p=0.001). Although overall the executive function and judgment based measures each point in this direction (Table 3) the individual correlation coefficients between measures was small (range -0.22 to 0.30).

### Table 3: Comparative responsiveness of outcome measures (mean, standard deviation, range and sample size)

<table>
<thead>
<tr>
<th>Measure</th>
<th>Baseline</th>
<th>Week 24</th>
<th>SRM</th>
</tr>
</thead>
<tbody>
<tr>
<td>sMMSE</td>
<td>23.4 (4.5;10-30)</td>
<td>N=137</td>
<td>24.12 (4.7;11-30) N=114</td>
</tr>
<tr>
<td>DAD</td>
<td>80.1 (18.6;25-100)</td>
<td>N=137</td>
<td>78.9 (23.6;9.5-100) N=114</td>
</tr>
<tr>
<td>NPI-Q</td>
<td>6.8 (5.4;0-28)</td>
<td>N=137</td>
<td>5.4 (5.1;0-30) N=114</td>
</tr>
<tr>
<td>NPI-Q-D</td>
<td>7.4 (6.8;0-30)</td>
<td>N=137</td>
<td>5.8 (6.9;0-41) N=114</td>
</tr>
<tr>
<td>ClOX-1</td>
<td>8.7 (3.9;0-15)</td>
<td>N=137</td>
<td>9.6 (3.9;0-15) N=111</td>
</tr>
<tr>
<td>ClOX-2</td>
<td>11.2 (3.2;0-12)</td>
<td>N=131</td>
<td>12.1 (2.9;0-15) N=106</td>
</tr>
<tr>
<td>Phonetic Fluency</td>
<td>7.4 (3.7;0-17)</td>
<td>N=136</td>
<td>8.3 (4.2;0-21) N=113</td>
</tr>
<tr>
<td>Symptom Guide</td>
<td>-2.7 (1.5;5.22)</td>
<td>N=128</td>
<td>-2.9 (1.7;6.3-2) N=106</td>
</tr>
</tbody>
</table>

CLOX-1 - clock drawing (hands free); CLOX-2 - clock drawing (copied); DAD - Disability Assessment for Dementia; NPI-Q - brief Neuropsychiatric Inventory; NPI-Q-D - brief Neuropsychiatric Inventory – Distress; sMMSe - standardised Mini-Mental State Examination; SRM - Standardized Response Mean

### Figure 2: Change in Symptom Guide (SG) score from baseline visit to last visit

### Figure 3: Proportion of patients by Clinical Global Impression at Week 24

3.3 Safety and adverse effects

Three hundred and sixty-three adverse events were recorded in the 148 patients, of which 130 were felt to be related to treatment (Table 4a). Adverse events (AE) were defined as any...
Table 4a: Overview of adverse events (AE)

<table>
<thead>
<tr>
<th>Number of Patients</th>
<th>All Causality</th>
<th>Treatment-Related</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients evaluable for AEs</td>
<td>148</td>
<td>148</td>
</tr>
<tr>
<td>Patients with AEs</td>
<td>110 (74.3)</td>
<td>62 (41.9)</td>
</tr>
<tr>
<td>Patients with serious AEs</td>
<td>24 (16.2)</td>
<td>2 (1.4)</td>
</tr>
<tr>
<td>Patients with severe AEs</td>
<td>22 (14.9)</td>
<td>4 (2.7)</td>
</tr>
<tr>
<td>Patients discontinued due to AEs</td>
<td>20 (13.5)</td>
<td>12 (8.1)</td>
</tr>
<tr>
<td>Patients with dose reduced or temporary discontinuations due to AEs</td>
<td>24 (16.2)</td>
<td>18 (12.2)</td>
</tr>
</tbody>
</table>

Untoward medical occurrence in a patient administered an investigational product or medical device. Serious adverse events (SAE) were defined as any untoward medical occurrence at any dose that results in death, is life threatening, requires inpatient hospitalization or prolongation of existing hospitalization, results in significant or persistent disability/incapacity or results in congenial abnormality or birth defect. All adverse events were rated for severity (mild, moderate, severe) by the investigating physician. During the course of this study, two patients (1.4%) died. One patient had two strokes, one in the active treatment period and one during the post treatment period. The second patient died following a worsening of general health status. None of these three events or two deaths was considered by the investigator as related to study drug. Fewer than half of the patients (41.9%) reported treatment-related AEs (Table 4a). Few patients reported treatment-related serious and severe AEs (1.4% and 2.7%, respectively). Twenty patients (13.5%) permanently discontinued the study due to an AE (all causalities), whereas 24 patients (16.2%) had to either reduce their dosage or temporarily discontinue due to AEs.

The most common treatment-related AEs (Table 4b) were gastrointestinal disorders, reported in 36 patients (24%), chiefly nausea (14.9%) and diarrhea (8.1%). Nervous system disorders were reported by 16 patients (10.8%) and psychiatric disorders by 9.5% of patients.

Table 4b: Frequency and severity of treatment-related adverse events

<table>
<thead>
<tr>
<th>System Organ Class Preferred Term</th>
<th>N (%)</th>
<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gastrointestinal disorders</td>
<td>36 (24.3)</td>
<td>15</td>
<td>21</td>
<td>0</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>12 (8.1)</td>
<td>4</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Nausea</td>
<td>22 (14.9)</td>
<td>13</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>Vomiting</td>
<td>7 (4.7)</td>
<td>5</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>General disorders</td>
<td>7 (4.7)</td>
<td>3</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Metabolism and nutrition disorders</td>
<td>7 (4.7)</td>
<td>3</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Nervous system disorders</td>
<td>16 (10.8)</td>
<td>8</td>
<td>8</td>
<td>0</td>
</tr>
<tr>
<td>Dizziness</td>
<td>8 (5.4)</td>
<td>5</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Psychiatric disorders</td>
<td>14 (9.5)</td>
<td>6</td>
<td>6</td>
<td>2</td>
</tr>
<tr>
<td>Insomnia</td>
<td>8 (5.4)</td>
<td>5</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>

4. DISCUSSION

In this six-month study, several measures that might feasibly be employed in primary care (such as the sMMSE, CLOX tests and SG) of patients with VaD and mixed AD/VaD showed the varying interpretations of change following treatment with donepezil. On the one hand, there was no statistically significant change in the sMMSE or activities of daily living; change in behaviour seemed to vary depending on the measure. On the other hand, there appears to have been some overall clinical benefit, as suggested by both individualized tests (CGI and the SG) and by those standardized tests that measured executive function (CLOX-1, CLOX-2, and Phonetic Fluency). This latter observation is potentially clinically important, given the role of executive dysfunction across the VCI spectrum.

The data must be interpreted with caution due to recruitment difficulties the sample size was less than aimed for. With 116 completers, we estimate a 34% chance of type II error (missing a significant difference) in the sMMSE. In addition, generalizability may not be assured given that the VCI construct was operationalized here for two subcomponents (VaD and mixed AD/VaD) which may not be widely used in all primary care settings. We underscore that the open-label design does not allow us to attribute change (or lack of it) to treatment with donepezil. An open-label trial, however, is appropriate for the evaluation of measurement issues, where the design can be viewed as a setting in which patients and treatment have been held constant, but measures varied. In consequence, we have been careful to keep inferences to that – i.e. measurement, not efficacy.

Of note, most patients did not deteriorate over the six months, so the magnitude of change is mostly in the “clinically detectable, but small” range.

Other factors associated with participation in a clinical trial – e.g., selection, vascular risk factor control, the placebo response, learning effects, chance – might account for the stability seen in most patients. Whether the response is maintained, worsened or enhanced over a longer period is not clear. With regard to safety, all enrolled subjects were analyzed for AEs, vital signs data, and ECG data. Laboratory parameters were analyzed for subjects who received at least one dose of donepezil and who had baseline and at least one post-baseline laboratory assessment (n=125 [84.5%]). Based on the advanced age and underlying medical condition of the
subjects enrolled in this study, the reported AEs and abnormal laboratory values were not unexpected.

Several lines of evidence, in addition to SRM estimates, point to clinical change on average having been detectable, but small. Here, the SG showed an effect size of -0.28 (between the baseline and week 24 visits). The SG’s summative intensity score (combining symptom frequency and the rank of its importance) for all symptoms corresponds to approximately 1.7 times a day for their occurrence at baseline, and about 1.2 times a day at week 24. This amounts to about an average 33% decrease in symptom frequency per individual, which on its face would seem to be both detectable and, on average, clinically meaningful, even if small. This analysis of a symptom-based approach to measuring treatment effects is in accord with a review which concluded that, despite their being central to the interpretation of trial results and to decisions regarding whether to employ trial findings in clinical practice, patient- and caregiver-centered measures of clinical significance have not been adequately studied33. Here, inspection of the distribution of the scores suggests that while most patients showed only a small change, the average effect reflects that a few patients showed a very large change – typically in the significant diminution of common and troubling symptoms. Likewise, a few patients showed important worsening, although on average more patients showed large improvement rather than large decline. While it might be that such stability at six months is better than expected, the clinical trial experience in VaD often shows placebo patients with little decline at six months11,16,22.

These data lend support to the contention that some part of the controversy over the effects of cholinesterase inhibitor treatment reflects how outcomes are measured. Many clinicians and caregiver advocacy groups have felt persuaded that they can see meaningful benefits from treatment. But if what they see has little chance to be captured by standard clinical trial instruments, there is no way for such evidence to be adjudicated40. Given how many new drugs for dementia are being tested, it remains useful to measure benefits which are likely to be both important and clinically meaningful, and which might capture unexpected benefits arising from novel approaches to treatment. Despite the intensity of effort and resources being put into them, as argued elsewhere32 it is doubtful that the emphasis on biomarkers as outcome measures will meet the need to demonstrate clinically meaningful change without, at the very least, being judged against clinically meaningful outcomes. For this reason, there remains a role for studies which address how to detect changes that might be associated with drug treatment, especially where clinical experience suggests that current measures are missing useful clinical information41-44. In Canada, this may have particular relevance to provincial formularies, many of which will only reimburse the cost of dementia treatment to people whose Mini-Mental State Examination test scores improve. Further work on which symptoms are most responsive, which are most resistant to change, or how any such patterns might be localized, are intriguing questions that are motivating further inquiries.

ACKNOWLEDGMENTS

Kenneth Rockwood is supported by the Dalhousie Medical Research Foundation as the Kathryn Allen Weldon Professor of Alzheimer Research. Sandra Black is supported by the Neuroscience Research Program and the Department of Medicine, Sunnybrook Health Sciences Centre, University of Toronto as the Brill Professor of Neurology at the University of Toronto.

The authors would like to acknowledge Isabelle Lussier, PhD and David A Gold, PhD for the initial development of this study, Marc-André Bédard, PhD for his assistance during the conduct of the study, and Kalliopi Dimos, BSc for the management of the conduct of this study, as well as An Zeng PhD and the late Yikang Xu, PhD for statistical support and review of earlier drafts of this manuscript.

ROLE OF THE AUTHORS

KR: Was a paid consultant to Pfizer Canada for this study and, as such, developed the research protocol with Pfizer Canada, trained participating physicians, wrote the first draft of the manuscript and revised subsequent drafts. KR had access to all of the data. The SG data reside fully with KR and he was responsible for their analyses.

SEB: Was a paid consultant to Pfizer Canada for this study and, as such, developed the research protocol with Pfizer Canada, wrote imaging protocol, trained participating physicians and reviewed and contributed to the manuscript. SEB had access to all of the data. The imaging data resides fully with SEB and she was responsible for their analysis.

AM: Supervised the analyses and designed analyses for the SG.

MR: Conducted analyses on SG data, held in-house at DGI Clinical Inc.

ID: Supervised the development of the statistical analysis plan, analyses of the data and interpretation of the data. ID reviewed and contributed to the manuscript.

CONFLICT OF INTEREST

Kenneth Rockwood was paid as consultant to this project. In addition, KR is President, Chief Scientific Officer and a shareholder in DGI Clinical Inc. (formerly DementiaGuide Inc.), which owns the SymptomGuide™ that was employed as an outcome measure in this trial. In the last five years, KR has also sat on advisory boards for Pfizer Canada, Glaxo SmithKline and Eli Lilly & Co, and given a talk sponsored by Bristol Myers Squib.

Sandra E. Black was paid as consultant to this project. In addition, SB receives honoraria from CME and ad hoc consulting from Pfizer, Janssen-Ortho, Novartis Pharmaceuticals, Lundbeck and Myriad Pharmaceuticals and operating funds from Pfizer Inc., Janssen Ortho, Novartis Pharmaceuticals, Myriad Pharmaceuticals, Sanofi-Aventis, Astra-Zeneca, Boehringer Ingelheim and Nova Nordisk.

Isabelle Defoy is an employee of Pfizer Canada Inc. This study was sponsored and managed by Pfizer Canada Inc.

REFERENCES


