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Associations between unmet needs for daytime activities and company and scores on the Neuropsychiatric Inventory-Questionnaire in people with dementia: A longitudinal study

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Associations between unmet needs for daytime activities and company and scores on the Neuropsychiatric Inventory-Questionnaire in people with dementia: A longitudinal study

Abstract:

Objectives: To examine prospectively the association between unmet needs for daytime activities and company and behavioural and psychological symptoms of dementia.

Methods: We included 451 people with mild or moderate dementia, from eight European countries, who were assessed three times over twelve months. Unmet needs were measured with the Camberwell Assessment of Need for the Elderly. Three sub-syndromes of the Neuropsychiatric Inventory-Questionnaire were regressed, one-by-one, against unmet needs for daytime activities and company, adjusting for demographic and clinical-functional covariates.

Results: Unmet needs for daytime activities were associated with more affective symptoms at baseline, six and twelve months, mean 0.74 ($p<0.001$), 0.76 ($p<0.001$) and 0.78 ($p=0.001$) points higher score respectively, and with more psychotic symptoms at baseline (mean 0.39 points, $p=0.007$) and at six months follow-up (mean 0.31 points, $p=0.006$).

Unmet needs for company were associated with more affective symptoms at baseline, six and twelve months, mean 0.44 ($p=0.033$), 0.67 ($p<0.001$) and 0.91 ($p<0.001$) points higher score respectively, and with more psychotic symptoms at baseline (mean 0.40 points, $p=0.005$) and at six months (mean 0.35 points, $p=0.002$) follow-up.

Conclusion: Interventions to reduce unmet needs for daytime activities and company could reduce affective and psychotic symptoms in people with dementia.

Keywords: dementia, needs assessment, daytime activities, company, BPSD

Introduction

For people with dementia, thorough assessments of individual needs are important for efficiently delivering high-quality health and social services that are individually tailored (van der Roest et al., 2009; Curnow et al., 2019). These assessments should include the perspective of the person with dementia, as his or her perceptions of unmet and met needs may differ from those of informal caregivers or health care professionals. Studies have shown that people with dementia generally report fewer unmet needs than researchers and their informal caregivers report them to have (van der Roest et al., 2009; Kerpershoek et al., 2017).

Studies investigating unmet needs in home-dwelling people with dementia by use of the widely used Camberwell Assessment of Need for the Elderly (CANE), found that daytime activities and company were two of the most commonly reported areas of unmet need (van der Roest et al., 2009; Miranda-Castillo et al., 2010; Mazurek et al., 2019). The item daytime activities includes social, work, leisure and learning activities, and the item company is described as social contact. Other unmet needs frequently reported by people with dementia, as well as their caregivers, include needs related to memory problems, information, and psychological distress (van der Roest et al., 2009; Miranda-Castillo et al., 2010; Curnow et al., 2019). In a large European cohort study including people with dementia from eight countries, daytime activities and company were again two of the items that both people with dementia and caregivers most frequently reported as unmet needs (Kerpershoek et al., 2017).

Identifying and seeking to meet unmet needs of people with dementia is important because unmet needs have been found to be associated with a lower health-related quality of life (Hoe et al., 2006; Miranda-Castillo et al., 2010; Kerpershoek et al., 2017; Handels et al., 2018; Janssen et al., 2018). Miranda-Castillo et al. (2010) suggest that unmet needs mediate the relationship between behavioural and psychological symptoms of dementia (BPSD) and quality of life. BPSD is a term referring to a heterogeneous range of phenomena, considered to be highly prevalent and occur in the majority of people with dementia over the course of the disease (Kales et al., 2015). The term BPSD has lately been raised as controversial, and there is an ongoing discussion to find a more psychosocial term that reflects the multiple causes of behaviour in dementia care (Cunningham et al., 2019; Wolverson et al., 2019). Though we acknowledge the importance of this debate, we do not aim to take a stand in it. We have chosen to use the term BPSD in this manuscript, as this is the term most widely used in our references. BPSD have been cited as major risk factors for higher caregiver burden, greater functional impairment, more rapid cognitive decline, poorer quality of life and nursing home admission (Kales et al., 2015; Wergeland et al., 2015). The grouping of BPSD into sub-

syndromes has been suggested as a more effective strategy for examining interventions than to report on each of the symptoms individually (van der Linde et al., 2014). Symptom groups commonly used are affective symptoms, psychosis, hyperactivity and euphoria (van der Linde et al., 2014).

Unmet needs are widely considered to be one of the contributory factors of BPSD (Cohen-Mansfield et al., 2015; Kales et al., 2015; Black et al., 2019; Cunningham et al., 2019). Many stakeholders in fact claim that BPSD are better considered as responses to unmet needs and suggest that the term ‘unmet needs’ might be used instead of BPSD (Wolverson et al., 2019). The links between unmet needs and BPSD may indicate that unmet needs should always be assessed, preferably with a standardized measure such as the CANE, in order to understand BPSD. These links may further indicate that meeting unmet needs should be a first choice to prevent and treat BPSD. From a research perspective, few studies have included a measurement of specific unmet needs when examining possible associations with BPSD in home-dwelling people with dementia. Thus, the aim of the current study was to examine prospectively over twelve months the association between unmet needs for daytime activities and company and the severity of different BPSD sub-syndromes.

Methods:

The Access to Timely Formal Care (Actifcare) study was an EU Joint Programme – Neurodegenerative Disease Research (JPND) project where access to and uptake of formal community care services were explored in the following eight European countries: Germany, Ireland, Italy, the Netherlands, Norway, Portugal, Sweden, and the United Kingdom. This study included data from the Actifcare prospective cohort study, a longitudinal study following people with dementia and their informal caregivers. Details about the Actifcare project and its cohort study can be found in the protocol paper (Kerpershoek et al., 2016).

Before the initiation of the cohort study, a joint training session for the data collectors from all eight countries was carried out in order to coordinate data collection and ensure consistency and a mutual understanding of how to complete the measures.

Participants:

In the Actifcare study, 451 dyads of people with dementia and their informal caregivers were included at baseline. For the present study, only data describing the people with dementia, not the informal caregivers, were included. Inclusion criteria were being home-dwelling and having a diagnosis of mild to moderate dementia indicated by a Clinical Dementia Rating

scale (CDR) score of 1 or 2 or a score on the Mini Mental State Examination (MMSE) of 24 or lower. To be included, the participants should not have been receiving formal personal care related to dementia at baseline but should be believed by a health care professional to require such care within one year. A subjective risk estimate was used to estimate need for additional assistance, based on available sources such as psychologists, general practitioners, memory clinic staff members, and other health care or social care professionals. These sources differed between countries and participants depending on where the participants were recruited from. Data were collected at baseline, six and twelve months.

Measures:

Outcome measure: BPSD (collected at baseline, six and twelve months)

BPSD were measured using the brief version of the Neuropsychiatric Inventory-Questionnaire (NPI-Q) addressing the severity of the following twelve symptoms: delusions, hallucinations, agitation/aggression, depression/dysphoria, anxiety, elation/euphoria, apathy/indifference, disinhibition, irritability/lability, motor disturbance, night-time behaviours and appetite/eating, each on a scale from 0 to 3 with 3 indicating more severe symptoms (Kaufer et al., 2000). The NPI-Q was completed by the informal caregiver.

Main exposure variables: Needs for daytime activities and company (collected at baseline, six and twelve months)

Needs were measured using the Camberwell Assessment of Need for the Elderly (CANE) scale (Reynolds et al., 2000). The CANE is an interview-based questionnaire designed to map the needs of older people ('needs present'; if answered with 'yes', then 'met' or 'unmet') and amount of help ('received' and 'needed') in relation to 24 items that address psychological, physical and environmental domains (Orrell & Hancock, 2004). The two items 'daytime activities' and 'company' were selected for this study for which only data on whether needs were present and, if so, met or unmet, were used. In the Actifcare study, needs of the person with dementia were reported by themselves, the caregiver, and the researcher; based on an overall perspective from extensive interviews with the person with dementia and the caregiver. In this study, we wanted to include the perspective of the person with dementia along with all other information. We therefore used scores for needs assessed by the researcher which are based on the reports from the person with dementia and the informal caregiver, together with all other information available to the researcher. The categories 'no

need' and 'met need' were collapsed into one category and compared to 'unmet need'. The needs variables were treated as time-dependent covariates in the analyses.

Covariates (collected at baseline, six and twelve months)

Level of dementia was measured with the Clinical Dementia Rating scale (CDR) (Hughes et al., 1982). Six domains of cognitive and functional performance are characterized using a scale of 0–3, where 0 indicates normal function and 3 indicates severe decline. The CDR was completed by the researchers after each interview based on all available data, and the sum of boxes scores, where the six item scores are added up (0–18 points) were used for this study (O'Bryant et al., 2008). Comorbidity was measured using the Charlson Comorbidity Index (Charlson et al., 1987), where higher scores indicate more comorbidities. Quan et al. have suggested updated weights of the contribution of chronic comorbidities of this index as a result of advances in medical treatment (Quan et al., 2011), and these updated weights were applied for each of the Charlson Comorbidity Index item scores before a sum score was produced for use in the analyses. Instrumental activities of daily living (IADL) were measured with Lawton and Brody's IADL scale, ranging from 0 to 8 with a lower score indicating a higher level of dependence (Lawton & Brody, 1969). Living situation was divided into two categories: 1) living alone and 2) living with someone.

Covariates (collected at baseline only)

The participants were from different European regions and, grouped in line with Handels et al. (Handels et al., 2018): North (Sweden and Norway), Middle (the Netherlands, Germany, UK and Ireland) and South (Portugal and Italy). Furthermore, all participants had a diagnosis of dementia meeting the DSM-IV criteria (American Psychiatric Association, 2000) following an assessment by a clinical professional. When an aetiological dementia diagnosis was available, this was recorded using the following categories: Alzheimer's disease (AD), Vascular dementia (VaD), mixed AD and VaD, Lewy body dementia (LBD) or 'other' dementia. Education of person with dementia was used in the analyses as a continuous variable of years of full-time education.

Statistics

The twelve BPSD symptoms assessed with the NPI-Q are quite different and using a sum score in analyses is not a preferred solution as two different participants with the same sum score may have significantly different clinical presentation. To identify clusters and group the

symptoms measured by the NPI-Q, a principal component analysis (PCA) was performed initially for the NPI-Q. We kept all items regardless of initial correlation and used varimax rotation and an eigenvalue greater than 1.0. The PCA resulted in three factors (see table 1) that were used in the analyses: agitation (agitation, euphoria, disinhibition, irritability and motor disturbance), affective (depression, anxiety, apathy and appetite) and psychotic (delusions, hallucinations and night-time behaviours). The three items anxiety, appetite and delusions each loaded on two factors. These items were placed in the factor on which they loaded most heavily, which was also the factor in which they are commonly found to fit (van der Linde et al., 2014).

<insert table 1 here>

To describe the proportion of the participants with clinically relevant levels of BPSD at baseline, we have chosen to categorise the sum score in each NPI-Q sub-syndrome into 3 groups: no/not significant, mild/moderate and severe symptoms. There is no common agreement on cut-offs for clinically relevant symptoms using the NPI-Q, and we have used a cut-off between no/not significant and mild/moderate which is in line with similar cut-offs used for the NPI (Lyketsos et al., 2002; Aalten et al., 2007). The difference in proportion of clinically relevant symptoms between participants with no/met need and unmet need is investigated with Chi-square analyses using the following two categories: mild, moderate and severe symptoms vs no/not significant symptoms.

Linear mixed models with random intercepts and slopes were used, with the three NPI-Q sub-syndromes as the dependent variables (one-by-one) and unmet needs vs met needs/ no needs for daytime activities or company as independent variables. The CDR, Charlson Comorbidity Index, IADL and a time variable (coded as 0 for baseline, 1 for six months and 2 for 12 months) were all treated as time-dependent covariates in the analyses. Because a linear time variable had equally good fit as the more complex three level dummy variable in a likelihood ratio test, the simpler continuous linear variable was preferred. The other variables were all treated as fixed time-invariant variables (dementia diagnosis, region, baseline age measured on the continuous scale). First six unadjusted linear mixed models were used, then six adjusted models where age, sex, CDR, region, Charlson Comorbidity Index, IADL, diagnosis, and living together/alone were added to the model. An interaction term (needs by time) was added to test whether differences changed over time. The inclusion of both the random intercept and slope improved the fit of the models significantly as revealed by a likelihood ratio test, and thus both terms were included.

Statistical analyses were performed using IBM SPSS Statistics version 25 and Stata version 16.0.

Ethical considerations:

Ethical approval was obtained separately in each of the participating countries. Written informed consent was obtained from participants or, for people with dementia with reduced ability to consent, from an informal caregiver/ legal representative.

Results:

Data from between 425 and 437 (depending on which CANE item and which NPI-Q sub-syndrome was being analysed) participants were sufficiently complete to be used for baseline analyses. The mean age of the participants at baseline was 78 years (SD 7.85), and 55% were female. The mean CDR sum of boxes score was 7.1 (SD 2.43), indicating mild dementia. A total of 28.9% had unmet needs for daytime activities, and 27.3% had unmet needs for company. For other characteristics of the participants, see table 2.

<insert table 2 here>

Table 3 shows proportions of participants with clinically significant BPSD symptoms at baseline. A larger proportion of the participants with unmet needs both for daytime activity and company had mild to moderate symptoms of affective and psychotic symptoms, compared to participants with no need or met need. Few participants had severe symptoms.

<insert table 3 here>

Daytime activities

Participants with unmet needs for daytime activities had higher scores on the NPI-Q affective items with a mean of 0.74 (95% Confidence Interval [CI] 0.34, 1.14, $p<0.001$), 0.76 (95% CI 0.46, 1.06, $p<0.001$) and 0.78 (95% CI 0.32, 1.24, $p=0.001$) points higher at baseline, six months and twelve months, respectively (Table 4, mixed model, adjusted). The SDs for NPI-Q affective scores for the reference groups (no/met need) were 2.46, 2.35 and 2.47 at baseline, 6 months, and 12 months, respectively. Thus, the effect sizes for the differences in scores on affective symptoms corresponded to 0.30, 0.32 and 0.32 SDs at baseline, six months and twelve months respectively. Unmet needs for daytime activities were also associated with more severe symptoms on the psychotic factor of the NPI-Q at baseline (mean of 0.39 points higher, 95% CI 0.10, 0.67, $p=0.007$) and at the six-month follow-up (mean of 0.31 points

higher, 95% CI 0.09, 0.52, $p=0.006$). These effect sizes for psychotic symptoms corresponded to 0.23 SD at baseline and 0.19 SD at six months. The differences in the NPI-Q affective and psychotic items between the groups with no/met and unmet needs did not change over time (interaction terms unmet needs*time were not significant; $p=0.935$ for affective items, $p=0.500$ for psychotic items) see figure 1. Scores on the agitation factor of the NPI-Q were not associated with unmet needs for daytime activities.

<insert figure 1 here>

Company

Participants with unmet needs for company had higher scores on the NPI-Q affective items with a mean of 0.44 (95% CI 0.04, 0.84, $p=0.033$), 0.67 (95% CI 0.35, 0.99, $p<0.001$), and 0.91 (95% CI 0.41, 1.41, $p<0.001$) points higher at baseline, six months and twelve months, respectively (Table 4, mixed model, adjusted). These effect sizes for the differences in scores on affective symptoms corresponded to 0.18, 0.29 and 0.36 SDs at baseline, six months and twelve months respectively. Unmet needs for company were associated with more severe symptoms on the psychotic factor of the NPI-Q at baseline (mean of 0.40 points higher, 95% CI 0.12, 0.69, $p=0.005$) and at the six-month follow-up (mean of 0.35 points higher, 95% CI 0.12, 0.58, $p=0.002$). These effect sizes for difference in scores on psychotic symptoms corresponded to 0.24 SD at baseline and 0.21 SD at six months. The differences in the NPI-Q affective and psychotic items between the groups with no/met and unmet needs did not change significantly over time (interaction terms unmet needs*time were not significant; $p=0.170$ for affective items, $p=0.694$ for psychotic items), even though there was a tendency towards a larger difference in scores over time for affective symptoms (see Figure 2). Scores on the agitation factor of the NPI-Q were not associated with unmet needs for company.

<insert figure 2 here>

<insert table 4 here>

Discussion

In this longitudinal study, we found that unmet needs for daytime activities and for company were associated with more affective and psychotic symptoms over twelve months. We also found a lack of association between agitation symptoms and unmet needs for daytime activities and company.

The differences we have found are significant, but the effect sizes are small. This is,

however, on a group level. As seen in table 3, a larger proportion of the participants with unmet needs had clinically significant BPSD at baseline compared to those with no needs or met needs. On an individual level, the presence of clinically significant symptoms may make a large impact on the life of a person with dementia, as well as on their caregivers, and even a small reduction of symptoms may improve their everyday lives.

The association between unmet needs for daytime activities and company and affective and psychotic symptoms

According to previous studies, unmet needs are, in general, associated with BPSD (Miranda-Castillo et al., 2010). In the Unmet Needs Model, Cohen-Mansfield et al. described BPSD (‘problem behaviours’) as a result of unmet needs stemming from a decreased ability of people with dementia to communicate those needs and to provide for themselves (Cohen-Mansfield et al., 2015). They focussed mainly on agitation in nursing home residents when describing the model, which is a setting that likely includes people with more severe dementia than the participants in our study. Yet the principle that behaviour is need-driven may also apply to community-dwelling people in a mild or moderate phase of dementia and to other symptoms such as affective and psychotic symptoms.

Apathy, depression, and anxiety (all included in our affective factor) are the most prevalent BPSD, and anxiety and depression are common in an early stage of dementia (Kales et al., 2015). The participants in our study were in a mild or moderate stage of dementia where affective symptoms are common. They could be starting to experience a decrease in their ability to meet their own needs for daytime activities and social life due to ADL impairments. Impairment in ADL has been found to be associated with a higher number of unmet needs (Eichler et al., 2016). Experiencing loss of function may contribute to affective symptoms because one may lose one’s sense of autonomy or feel less valued. Company and daytime activities are both considered to be social needs, and unmet social needs have been found to be associated with higher levels of depression along with unmet psychological needs (Alltag et al., 2018).

Depressive symptoms are described as being intensified or maintained by the absence of positive feelings resulting from participation in enjoyable and meaningful activities (Orgeta et al., 2017). Furthermore, having depressive symptoms, anxiety or apathy may lead to not taking the initiative to be active and to meet people even if it would be beneficial, thereby resulting in unmet needs for daytime activities and company.

To our knowledge, no previous studies have found an association between psychotic

symptoms and unmet needs for daytime activities and company. Psychotic symptoms in dementia may share similarities with symptoms of schizophrenia, where reduced social activity and interest, loss of motivation and reduced productive activity are often present (Cipriani et al., 2020). It might be that people with dementia who experience psychotic symptoms are withdrawing from activities and from social interaction since their symptoms make it difficult for them to function in some kinds of activities and social settings. Delusions may make it difficult to trust others and to communicate in a relevant way. Further, psychotic symptoms may make it harder for caregivers to fulfil needs in people with dementia. It may also be that unmet needs for daytime activities and company contribute to psychotic symptoms in people with dementia, e.g. due to lack of interaction with other people. On the other hand, too much stimuli may add to psychotic symptoms.

The lack of association between agitation and unmet needs for daytime activities and company

In this study, no associations were found between the agitation symptom cluster and unmet needs for daytime activities or company. Agitation itself is a heterogeneous term and is often used to describe diverse symptoms such as pacing, hoarding, making disruptive sounds, asking repetitive questions and becoming upset easily (Kales et al., 2015). In our PCA, the items included in the agitation factor were agitation, euphoria, disinhibition, irritability, and motor disturbance. Even if these items loaded on the same factor, they may have less in common than the items in the affective or psychotic factor. Van der Linde et al. found that studies using PCA on the NPI-Q generally suggest the following symptom groups: 1) affective symptoms, 2) psychosis, 3) hyperactivity and 4) euphoria (van der Linde et al., 2014). Our agitation factor includes both 3 and 4, indicating that this factor may be our most heterogeneous.

The need for daytime activities and company

In research on needs among people with dementia, unmet needs for daytime activities and company are frequently found (van der Roest et al., 2009; Miranda-Castillo et al., 2010; Kerpershoek et al., 2017). Even if professional support was frequently provided for company and daytime activities, unmet needs were still reported in these areas (van der Roest et al., 2009). Involvement in meaningful activities has been found to be important for people with dementia because it gives them feelings of enjoyment and pleasure, connection and belonging as well as autonomy and identity (Phinney et al., 2007). Daytime activities and company are

connected to the essential psychological needs for occupation, inclusion and attachment in person-centred care (Kitwood, 1997), and their importance may be explained by the association between occupation, health and well-being (Christiansen & Townsend, 2011).

Studies have shown that social isolation and reduced access to their usual activities may increase the risk of mental health problems in older adults (Armitage & Nellums, 2020), and it is likely that people with dementia are at particular risk. Assessing at an early stage of the disease, as well as reassessing regularly, whether needs for daytime activities and company are met and providing these if needed may prevent or reduce BPSD and enhance quality of life in people with dementia.

Implications for post-diagnostic support to reduce unmet needs

Although we have found associations between unmet needs for daytime activities and company and affective and psychotic symptoms, the direction here may be discussed. It may be that unmet needs for daytime activities and company contribute to affective and psychotic symptoms; it is also possible that these symptoms contribute to unmet needs; or it could go both ways as a downward spiral. However, this implies that if post-diagnostic support can reduce either unmet needs or affective and psychotic symptoms, this could affect the other part of the equation.

Assessing the needs of people with dementia is useful both for helping to identify interventions and services that should be tailored to each individual and planning the provision of health care on a macro level (Reynolds et al., 2000; Curnow et al., 2019). An assessment of unmet needs should be carried out as early as possible in the process of dementia and updated regularly. Sometimes it takes a while to establish a diagnosis, but interventions to target unmet needs do not have to await the diagnosis. With the assessment of symptoms and functioning and post-diagnostic support assigned to the same municipal dementia-resource team, as provided by the Norwegian model, post-diagnostic support can even include pre-diagnostic support. Moreover, it can be individually tailored and be initiated as soon as symptoms and/or needs become known (Michelet et al., 2020).

Enabling people with dementia to engage in meaningful activities as part of their everyday lives should be part of post-diagnostic support (Gitlin et al., 2009; Kales et al., 2015; Orgeta et al., 2017; Lobbia et al., 2018). Person-centred care includes the promotion of social participation and meaningful activities, and these are important components of several psychosocial interventions for people with dementia. Evidence of efficacy has been found for a variety of such interventions delivered to home-dwelling people with mild to moderate

dementia; however, the use of such interventions remains low (Keogh et al., 2019). Informal caregivers play a crucial role in several of the interventions. In this study, we address caregivers only as partners in the provision of interventions, even though several of the interventions may also have an effect on caregivers' health and well-being.

Adult day services such as day care for people with dementia may serve to meet the needs for daytime activities and company, given that the service is age appropriate and individually tailored (Strandenaes et al., 2017; Strandenæs et al., 2019). In a review, attending adult day care was found to increase social engagement for people with dementia through participation in activities with peers with whom they feel safe and comfortable. Further, participants who attended adult day care exhibited significantly less depression and fewer behavioural issues compared to participants who did not attend (Du Preez et al., 2018).

Group interventions targeting activities, such as behavioural activation and Cognitive Stimulation Therapy (CST), have been found to offer several positive effects including reducing anxiety and depressive symptoms, improving quality of life and communication, reducing problematic behavioural symptoms, and increasing scores on ADL for community-dwelling people with dementia (Orgeta et al., 2017; Lobbia et al., 2018). Caregivers being taught to use activities individually tailored to the capabilities and interests of people with dementia in the Tailored Activity Program (TAP) report reduced behavioural symptoms (Gitlin et al., 2009).

Strengths and limitations

The strength of this study is that the data were from a large cohort study with participants from eight countries across Europe and may, therefore, be representative of a larger group of people with dementia. However, this heterogeneity could also be a limitation as the recruitment of participants differed, including the sources used in estimating that need for additional assistance would likely be required within one year. The sample studied was a convenience sample. There might also have been heterogeneity among the researchers collecting data as the perceptions of different symptoms and use of the measures may differ across researchers in different countries. However, joint training was conducted, and there were meetings and regular contact within the project group to coordinate the data collection for consistency and improved inter-rater reliability.

Furthermore, in the analyses, the twelve NPI-Q items were reduced to three factors following a PCA. This may have resulted in the loss of some of the details in the data. Having three NPI-Q factors is, however, comparable to other studies that have used the NPI-Q

(Truzzi et al., 2013). The NPI-scores are not based on direct observations or on the view of the person with dementia, but on proxy information, from an informal caregiver. This may be a limitation because proxy information could be influenced by caregiver distress or relationship quality.

From the needs assessment (CANE) data, the researchers' assessments were used rather than those of the people with dementia. The researchers did consider the scores from the people with dementia and the informal caregiver as well as other available information, but there is always a risk that the perspective of the people with dementia was not given enough weight in these scorings.

Conclusion

In this study, we found that unmet needs for daytime activities and for company were associated with more affective and psychotic symptoms but not with more symptoms of agitation. This is in line with previous findings and may serve to elaborate the importance of structured and repeated assessment of needs and a proactive approach toward fulfilling unmet needs for daytime activities and company for people with dementia. Psychosocial interventions in post-diagnostic support creating meaningful occupations and addressing social needs may reduce unmet needs for daytime activities and company and, thereby, reduce affective and psychotic symptoms.

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Consortium members: Maastricht University (NL): Marjolein de Vugt, Claire Wolfs, Ron Handels, Liselot Kerpershoek. Martin-Luther University Halle-Wittenberg (DE): Gabriele Meyer (WP2 leader), Astrid Stephan, Anja Bieber, Anja Broda, Gabriele Bartoszek. Bangor University (UK): Bob Woods (WP3 leader), Hannah Jelley. Nottingham University (UK): Martin Orrell. Karolinska Institutet (SE): Anders Wimo (WP4 leader), Anders Sköldunger, Britt-Marie Sjölund. Oslo University Hospital (NO): Knut Engedal, Geir Selbaek (WP5 leader), Mona Michelet, Janne Rosvik, Siren Eriksen. Dublin City University (IE): Kate Irving (WP6 leader), Louise Hopper, Rachael Joyce. CEDOC, Nova Medical School/Faculdade de Ciências Médicas, Universidade Nova de Lisboa (PT): Manuel

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Disclosure statement

Geir Selbæk is a member of the advisory board of Biogen in Norway. The authors report no other conflict of interest.

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Table 1. Principal component analysis of the Neuropsychiatric Inventory-Questionnaire (NPI-Q), Varimax rotation with Kaiser Normalization

Item	Component		
	1	2	3
Disinhibition	0.73		
Agitation/aggression	0.73		
Irritability/lability	0.69		
Elation/euphoria	0.46		
Motor disturbance	0.31		
Depression/dysphoria		0.77	
Apathy/indifference		0.69	
Anxiety		0.49	0.38
Appetite/eating	0.33	0.40	
Hallucinations			0.78
Night time behaviours			0.72
Delusions	0.40		0.53

Table 2. Characteristics of the participants at baseline

Age - Mean (SD), n = 451	77.77 (7.85)
Sex, female, n = 451	246 (54.5%)
Living alone, n = 451	88 (19.5%)
Education, years of full time education - Mean (SD), n = 449	9.82 (4.48)
Region, n = 451	
North (Sweden and Norway)	110 (24.4%)
Middle (UK, Ireland, the Netherlands and Germany)	222 (49.2%)
South (Portugal and Italy)	119 (26.4%)
Diagnosis, n = 451	
AD	218 (48.3%)
VaD	53 (11.8%)
Mixed	56 (12.4%)
LBD	6 (1.3%)
Other	27 (6.0%)
Unspecified dementia	91 (20.2%)
CANE daytime activities - with unmet needs, n = 450	130 (28.9%)
CANE company - with unmet needs, n = 450	123 (27.3%)
NPIQ agitation - Mean (SD), maximum 15 points, n = 439	2.93 (2.77)
NPIQ affective - Mean (SD), maximum 12 points, n = 436	3.37 (2.60)
NPIQ psychosis - Mean (SD), maximum 9 points, n = 444	1.46 (1.84)
Charlson Comorbidity Index, updated weights - Median (IQR), n = 441	2 (2, 3)
Clinical Dementia Rating, Sum of boxes - Mean (SD), n = 448	7.06 (2.43)
Instrumental Activities of Daily Living - Mean (SD), n = 445	3.45 (1.99)

Table 3. Proportions of participants with clinically significant symptoms at baseline, per subsyndrome, classified as no/not significant symptoms - mild/moderate symptoms – severe symptoms, grouped by no/met need and unmet need for daytime activities and company

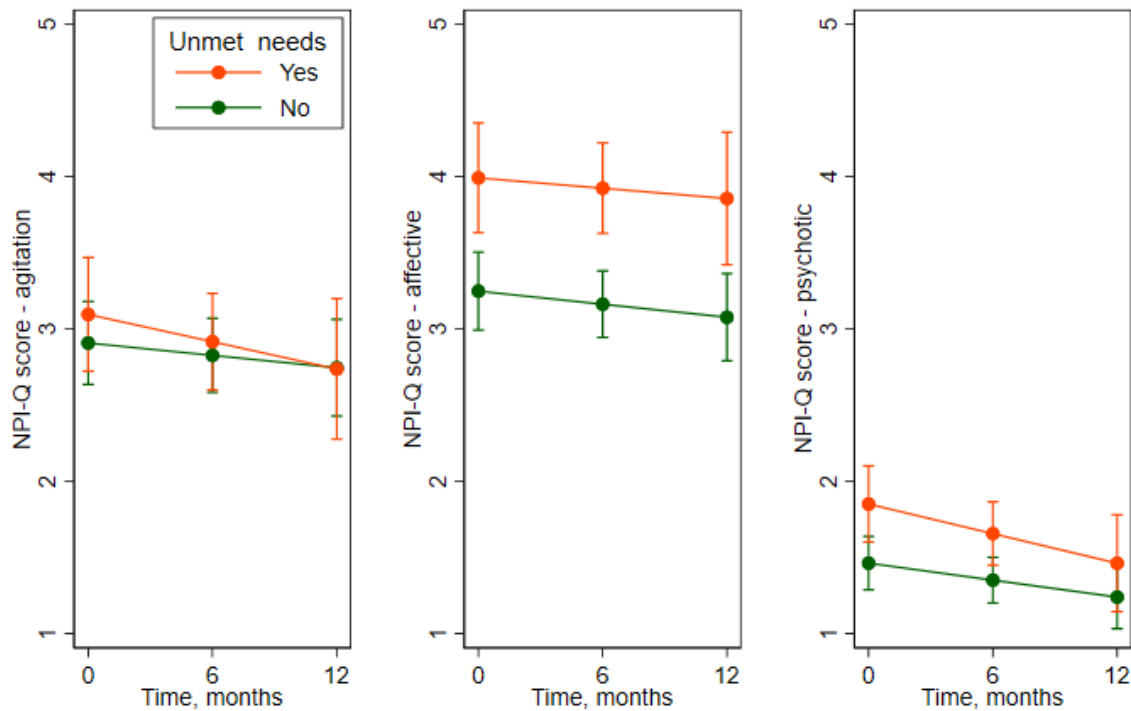
		Daytime activities		Pearson Chi-Square*	Company		Pearson Chi-Square*
		No need/ Met need N (%) n=304-308	Unmet need N (%) n=125-129		No need/ Met need N (%) n= 306-310	Unmet need N (%) n= 120 -123	
Agitation Maximum score: 15	Score 0-4 no/not significant	235 (77.3%)	91 (71.1%)	p= 0.212	237 (77.5%)	87 (71.3%)	p= 0.225
	Score 5-10 Mild/ moderate	67 (22.0%)	32 (25.0%)		66 (21.6%)	31 (25.4%)	
	Score 11-15 Severe	2 (0.7%)	5 (3.9%)		3 (1.0%)	4 (3.3%)	
Affective Maximum score: 12	Score 0-3 no/not significant	189 (62.2%)	49 (39.2%)	p< 0.001	186 (60.8%)	50 (41.7%)	p= 0.001
	Score 4-8 Mild/ moderate	110 (36.2%)	67 (53.6%)		116 (37.9%)	60 (50.0%)	
	Score 9-12 Severe	5 (1.6%)	9 (7.2%)		4 (1.3%)	10 (8.3%)	
Psychosis Maximum score: 9	Score 0-2 no/not significant	250 (81.2%)	82 (63.6%)	p< 0.001	249 (80.3%)	81 (65.9%)	p= 0.002
	Score 3-6 Mild/ moderate	54 (17.5%)	42 (32.6%)		58 (18.7%)	36 (29.3%)	
	Score 7-9 Severe	4 (1.3%)	5 (3.9%)		3 (1.0%)	6 (4.9%)	

* For the Chi-square analyses the scores for mild/moderate and severe clinically significant symptoms have been collapsed and compared to the scores for no clinically significant symptoms

Table 4. Mean difference in NPI-Q sub-syndromes between groups: no need/met need vs unmet need, concerning daytime activities and company

			Mixed model - Unadjusted		Mixed model - Adjusted	
Variable	n	Visit	Difference – mean (95% CI)	p-value	Difference – mean (95% CI)	p-value
Daytime activities						
NPI-Q - agitation	432	Baseline	0.23 (-0.20, 0.66)	0.286	0.19 (-0.22, 0.59)	0.366
	376	6 months	-0.14 (-0.57, 0.29)	0.536	0.09 (-0.22, 0.40)	0.569
	332	12 months	0.39 (-0.15, 0.92)	0.154	-0.01 (-0.49, 0.47)	0.972
NPI- Q - affective	429	Baseline	0.76 (0.33, 1.18)	0.001	0.74 (0.34, 1.14)	<0.001
	372	6 months	0.80 (0.36, 1.24)	<0.001	0.76 (0.46, 1.06)	<0.001
	330	12 months	1.08 (0.58, 1.58)	<0.001	0.78 (0.32, 1.24)	0.001
NPI-Q - psychotic	437	Baseline	0.44 (0.14, 0.74)	0.004	0.39 (0.10, 0.67)	0.007
	380	6 months	0.26 (-0.05, 0.56)	0.100	0.31 (0.09, 0.52)	0.006
	342	12 months	0.37 (-0.00, 0.75)	0.052	0.22 (-0.12, 0.57)	0.205
Company						
NPI-Q - agitation	428	Baseline	0.11 (-0.33, 0.54)	0.636	0.21 (-0.19, 0.62)	0.304
	372	6 months	0.06 (-0.38, 0.50)	0.779	0.19 (-0.14, 0.51)	0.256
	327	12 months	0.46 (-0.12, 1.04)	0.119	0.16 (-0.35, 0.68)	0.538
NPI-Q - affective	426	Baseline	0.33 (-0.10, 0.77)	0.128	0.44 (0.04, 0.84)	0.033
	368	6 months	0.66 (0.21, 1.11)	0.004	0.67 (0.35, 0.99)	<0.001
	325	12 months	0.99 (0.43, 1.54)	<0.001	0.91 (0.41, 1.41)	<0.001
NPI-Q - psychotic	433	Baseline	0.26 (-0.04, 0.57)	0.090	0.40 (0.12, 0.69)	0.005
	376	6 months	0.27 (-0.05, 0.58)	0.094	0.35 (0.12, 0.58)	0.002
	336	12 months	0.40 (-0.00, 0.81)	0.052	0.30 (-0.07, 0.67)	0.114

Daytime activities

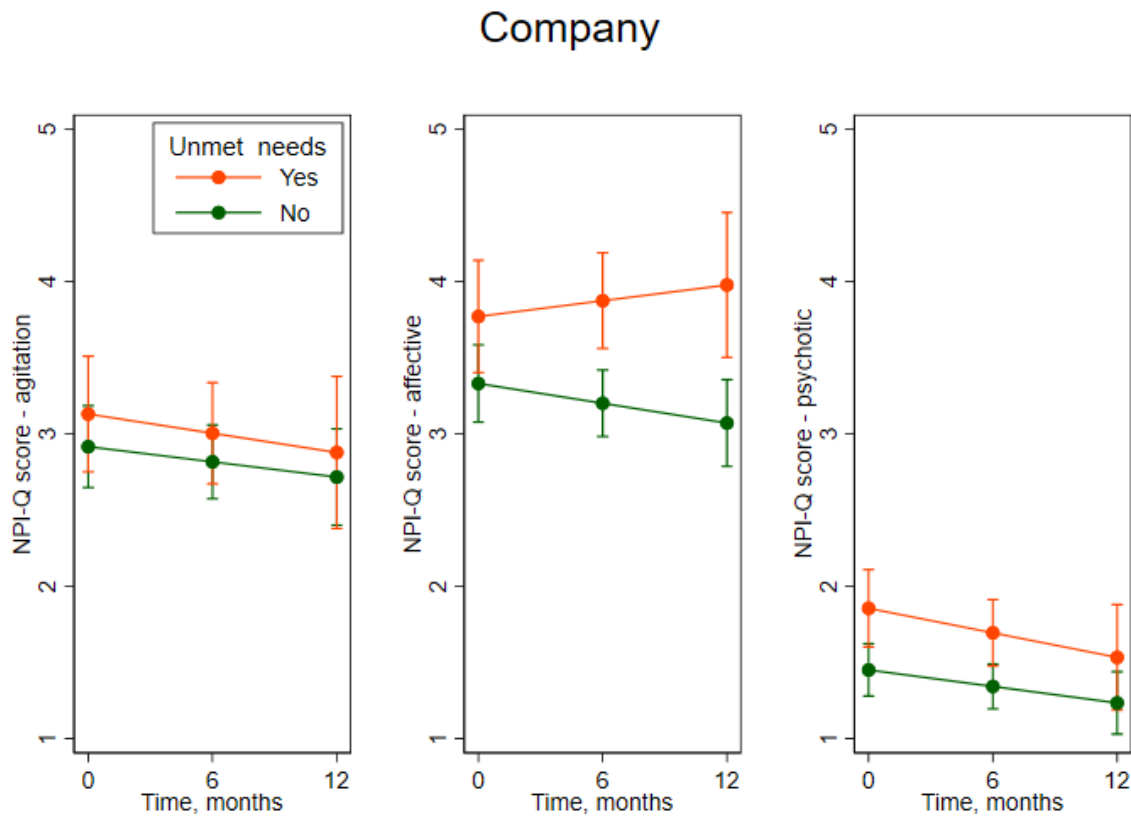


NPI-Q agitation sub-syndrome= agitation, euphoria, disinhibition, irritability and motor disturbance – max score= 15

NPI-Q affective sub-syndrome= depression, anxiety, apathy and appetite – max score= 12

NPI-Q psychotic sub-syndrome= delusions, hallucinations and night-time behaviours – max score= 9

Figure 1. Mean Neuropsychiatric Inventory-Questionnaire (NPI-Q) scores over time for those with met needs for daytime activities versus those with unmet needs. Vertical lines are 95% confidence intervals. Estimated in mixed regression where needs was modelled as a time dependent covariate, adjusted by age, sex, living alone/ with someone, region, diagnosis, education, Charlson Comorbidity Index, Clinical Dementia Rating and Instrumental Activities of Daily Living



NPI-Q agitation sub-syndrome= agitation, euphoria, disinhibition, irritability and motor disturbance – max score= 15

NPI-Q affective sub-syndrome= depression, anxiety, apathy and appetite – max score= 12

NPI-Q psychotic sub-syndrome= delusions, hallucinations and night-time behaviours – max score= 9

Figure 2. Mean Neuropsychiatric Inventory-Questionnaire (NPI-Q) scores over time for those with met needs for company versus those with unmet needs. Vertical lines are 95% confidence intervals. Estimated in mixed regression where needs was modelled as a time dependent covariate, adjusted by age, sex, living alone/ with someone, region, diagnosis, education, Charlson Comorbidity Index, Clinical Dementia Rating and Instrumental Activities of Daily Living

Figure captions:

Figure 1:

(Footnote:)NPI-Q agitation sub-syndrome= agitation, euphoria, disinhibition, irritability and motor disturbance – max score= 15

NPI-Q affective sub-syndrome= depression, anxiety, apathy and appetite – max score= 12

NPI-Q psychotic sub-syndrome= delusions, hallucinations and night-time behaviours – max score= 9

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Figure 2:

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