The Danish Alzheimer Intervention Study: Rationale, Study Design and Baseline Characteristics of the Cohort

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Key Words
Alzheimer’s disease • Dementia • Randomized controlled trial • Study design • Counselling • Psychosocial intervention • Support • Caregiving

Abstract
Background: There is a lack of appropriately designed trials investigating the efficacy of psychosocial interventions for patients with mild dementia and their family caregivers. This paper reports the rationale and design of the Danish Alzheimer Disease Intervention Study and baseline characteristics of the cohort. Methods: The study was a 1-year multicentre randomized controlled rater-blinded trial with randomization to follow-up and a multifaceted semitailored intervention programme or to follow-up only (with extension of follow-up to 3 years). The intervention included a counselling programme, teaching courses, written information and logbooks. The outcomes included clinical efficacy parameters, patient satisfaction and health economic consequences. Results: A total of 330 patients and their 330 caregivers were included during a period of 18 months. The majority (65.2%) of the caregivers were spouses. At inclusion the mean age of the patients and caregivers was 76.2 and 66.0 years, respectively. Conclusion: The study will explore the added value of a multifaceted intervention programme and contribute to the design of future interventions for patients with mild dementia and their caregivers.

Introduction
Alzheimer’s disease (AD) and other dementia disorders are characterized by progressive decline in cognitive, social and occupational function and are often associated with affective symptoms and behavioural disturbances. The majority of the patients are living in their own homes and require increasing assistance and supervision from caregivers, often a spouse. The role as a family caregiver for a patient with dementia is associated with a greater risk of developing stress and somatic and psychiatric health problems [1–3]. Several studies have indicated that counselling and psychosocial interventions for caregivers may have a significant positive effect in patients with moderate to severe AD [4–6] as well as in their caregivers [5, 7–16].
In general, however, results have been inconsistent, the quality of studies has been poor, and there is a need for appropriately designed trials [17]. For policy makers and service planners, who work with carer and patient support in dementia, good quality evidence is crucial for identifying who would benefit from the service and when. While there is an increasing awareness of the importance of early and continuous support and care, the majority of the patients have no or little access to diagnostic evaluation, treatment, and follow-up during the course of their disease [18, 19]. With increased awareness and better diagnostic methods many patients with progressing neurodegenerative dementia disorders are now diagnosed in the very early phase of their disease. Patients with very mild dementia often have nearly full autonomy and may request more attention and specific counselling programmes directed towards their own needs. Hence, it is necessary to develop and validate support programmes, which focus specifically on the needs of patients with mild dementia and their caregivers, and which include psychosocial support for the patient as well as for the caregiver. The aim of the Danish Alzheimer Disease Intervention Study (DAISY) was to evaluate and explore clinical effect, patient satisfaction and health economic consequences of a multifaceted and semitailored intervention programme in patients with mild dementia and their caregivers, in whom follow-up secured referral to conventional local support programmes where available and relevant.

We hypothesized that the multifaceted and semitailored intervention programme offered to the patients and their primary caregivers during the first year after the diagnosis might prevent the emergence of depressive symptoms and improve the quality of life in patients as well as in caregivers, and perhaps even stabilize the patients’ cognitive function for some time. This paper reviews current methodological issues in psychosocial interventions and reports the rationale and design of the DAISY study, and the baseline characteristics of the cohort.

**Material and Methods**

**Study Design**

The study design and reporting adhere to the Consolidated Standards for Reporting Trials Statement (www.consort-statement.org) [20]. The study design was compatible with the quality criteria and recommendations proposed by the Cochrane systematic review on support for carers of people with AD, which was available at the time of the initiation of the study [21] and later updated [22].

As shown in figure 1, the study was designed as a 1-year multicentre randomized controlled rater-blinded trial (RCT) with 1:1 randomization to follow-up only or to follow-up with the addition of the DAISY intervention, a multifaceted semitailored intervention programme, described in detail below. Two years after study start supplementary funding was obtained to extend the follow-up with a study visit 3 years after inclusion.

To randomize participants we used a central allocation process by a third party unaware of and concealed for the DAISY investigators. The allocation process was initiated after patient and caregiver identity and key baseline data had been registered in the central project office. The randomization was done using a computer programme (Stat-Direct version 2.3.7). We used a random block size algorithm to prevent imbalance between the allocation groups. Patients were stratified by centre, MMSE score (20–23; 24–30) and use (or nonuse) of symptomatic antidementia drugs. The assignment to group was done via letters from the central office to the local study coordinators. As described below, a range of primary and secondary quantitative outcome measures from standardized tests and scales, from interview data or from registry-based information were collected at baseline and at 6, 12 and 36 months. The randomization code and study results were not opened until after the last patient had completed the visit at 3 years. In the intervention group, patient and caregiver compliance
patients with probable AD who had vascular changes on cranial CT, which may contribute to their symptoms.

**Exclusion Criteria.** Patients with severe somatic or psychiatric comorbidity (including impaired hearing or vision), which would significantly impair their cooperation with the programme, were excluded. Patients participating in other intervention studies and patients living in a nursing home at baseline were also excluded.

**Primary Caregivers**

A primary caregiver was defined as the informal caregiver who was the main person responsible for the informal care to the patient and who had regular contacts (at least weekly) with the patient. For patients who had >1 primary caregiver, the invitation to join this study was given to only 1 person, at the patient’s choice.

**Baseline and Follow-Up Visits**

Patients and their primary caregivers participated together in all visits. The baseline visit took place prior to randomization and was performed by the local study coordinator in the local memory clinic. The follow-up visits were home visits performed at 6, 12 and 36 months by independent raters unaware of the randomization code. The raters were not involved in the intervention programme, and they were not employed in the same institutions as the study coordinators and counsellors. Patients and caregivers were instructed to try not to reveal which treatment arm they were in. The efficiency of concealment was checked using a questionnaire to the raters at the end of each follow-up visit. None of the raters visited the same patient-caregiver couple more than once.

**Drop-Outs**

For patients dropping out of the study, the date and reason for dropout was registered. If a patient dropped out for whatever reason, the caregiver was not allowed to continue in the study. If a primary caregiver dropped out, she/he was replaced with another primary caregiver if possible, in order to continue assessment of proxy-rated parameters (whereas scales rating caregiver health status were discontinued). If no replacement was possible, only the patient continued in the study, and proxy-rated assessments were discontinued.

**Roles and Training of Study Coordinators, Raters and Monitors**

In each of the 5 centres a study coordinator was responsible for recruitment, baseline assessments, implementation of the multifaceted and semitailored intervention, all counselling sessions and project coordination. The study coordinators were all designated specialist nurses with several years of experience in the field of dementia and in counselling. Prior to the initiation of the study the study coordinators participated in a 4-day centralized course, where they were instructed in the rationale and design of the study, in the intervention programme, in basic concepts of patient and caregiver intervention, and in the concepts and objectives of the specific counselling programme applied in this study. The course also contained formalized teaching of communication and counselling. Throughout the study intervention period, the study coordinators were offered supervision on a regular basis at 8 centralized follow-up sessions and at regular site visits by specialists from the coordinating study centre. The raters were nurses, therapists and other health professionals from the local municipalities or hospitals. They were responsible for follow-up of all patients and satisfaction with the intervention programme, and qualitative aspects of the intervention were assessed in separate sub-studies.

**Patients**

**Recruitment.** The study was conducted in 5 counties, representing rural and urban districts, in Denmark (fig. 2), and inclusion took place in 1 centre in each county during a period of 18 months. In each of the 5 counties information regarding the project was provided to all general practitioners, to private practice neurologists and psychiatrists, as well as to all hospital departments involved in diagnostic evaluation of patients with dementia. The local study coordinator and a designated physician in each of the local memory clinics assessed patients regarding inclusion criteria.

**Inclusion Criteria.** The study included community-dwelling patients with a clinical diagnosis of probable AD, mixed AD with vascular components or dementia with Lewy bodies (DLB), established within the past 12 months; age ≥50 years; mild dementia with MMSE score ≥20; a primary caregiver with close contact to the patient who was willing to participate in the study and the intervention; and informed consent from the patient and the caregiver. All patients met DSM-IV [23] for dementia and the NINCDS-ADRDA criteria for probable AD [24] or the McKeith criteria for DLB [25]. Participants classified as mixed AD were excluded. All patients met DSM-IV intervention; and informed consent from the patient and the caregiver. The local study coordinator and a designated physician in each of the local memory clinics assessed patients regarding inclusion criteria.

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and assessments at 6, 12 and 36 months. Both study coordinators and raters participated in centralized training sessions for the assessment of primary and secondary efficacy parameters. Central study monitors observed the progress of the study in each centre and checked case report forms for completeness and consistency throughout the study.

**The DAISY Intervention: A Multifaceted and Semitailored Intervention Programme**

The novel multifaceted and semitailored intervention programme was designed to comprise counselling, information and support to patients with mild dementia as well as to their primary caregivers during the initial months after a diagnosis had been established (fig. 3). The objective of the programme was to prevent the emergence of depressive symptoms and further impairment of health-related quality of life in patients as well as in caregivers, and perhaps even stabilize cognitive function in the patients for some time. The philosophy was to focus on positive resources, intact functions and retained skills and activities that the patients could still take part in. Administered by the local study coordinator in each centre, the programme was carried out in the intervention group only and initiated within the first month after inclusion. Having 5 key components, part of the intervention was tailored to the needs of the individual patient and/or caregiver, while other parts of the programme were based on group intervention. The participants were encouraged to take advantage of all components in the intervention programme as much as possible. Counselling sessions were planned on a running basis, while group-based teaching courses were scheduled to start when a sufficient number of participants had been allocated to the intervention arm. Thus, for logistical reasons the duration of the full intervention programme could vary from 8 to 12 months.

**Counselling Sessions**

The counselling aimed to prevent or reduce depressive symptoms, impairment of health-related quality of life and loss of social network. The counselling programme was based on constructivist principles [26]: the principles and methodology were anchored on the dynamics that characterizes the patient and the caregiver’s everyday life. The counselling was based on a philosophical approach in which each patient or caregiver was given the possibility of expressing his/her own life story and what is of personal importance and of great value to the individual. The counsellor offered the patient and the caregiver guidance with common decision-making, advice and activities that help the participants to construct a meaningful life.

At the initial counselling visit well-structured written notes were established. The notes were constructed to reflect the individual strengths and needs of the patient and the caregiver in 4 related areas: daily life, social network, pleasures and activities. The written notes were used to focus follow-up sessions, with the aim of improving coping strategies and to empower the patient and caregiver to focus on the positive factors and resources in their lives, according to the principles of self-validation. Self-validation is the process of recognizing and transcending the sense of self by various means to appreciate the unconditional value and meaning of our personal existence [27].

The counselling sessions were conducted by the local study coordinator in the home of the patient or in the hospital-based memory clinic. It was composed of: (a) 2 sessions with the patient and caregiver, (b) 2 sessions with the patient alone, (c) 2 sessions with the caregiver alone, and (d) 1 optional network session with the patient, caregiver and family network.

**Courses**

The aim of the courses was to provide patients and caregivers with basic information about dementia and its consequences and about living and coping with dementia. The courses also aimed to provide a forum for patients and caregivers to exchange experiences and coping strategies. Two parallel lines were aimed at patients and caregivers, respectively. The courses for patients took place simultaneously with the courses for their caregivers, but in a separate classroom. In each line the course programme included 5 scheduled sessions for groups of 12 participants. Each session included a standardized agenda with information on key topics related to dementia: general information about dementia, legal aspects, living with dementia, support to people with dementia and their caregivers, and a summary session with miscellaneous topics. The sessions were supported by written information produced specifically for patients and caregivers in this study. Furthermore, support group activities with open agendas were conducted at the end of each session. Here, the patients (or caregivers) could discuss topics and questions of their interest and get informal advice from other patients (or caregivers). All courses were coordinated by the local study coordinator, and local health professionals in the field of dementia were invited as teachers. Volunteers were invited to assist the teacher in the course sessions for patients.

**Information Folder**

A comprehensive information folder with structured information sheets about dementia and related aspects was developed specifically for this study and provided to all participants in the intervention group. The aim of the folders was to provide the participants with written information, which would support the information given at counselling sessions and teaching courses.
and also to serve as a reference guide later in the course of the dementia disease after completion of the intervention programme. There were separate and different folders for patients and for caregivers. The folders contained chapters on causes of dementia, diagnosis and treatment, legal aspects in relation to dementia, and sources and contact details for social support.

**Outreach Telephone Counselling**

The aim of outreach telephone counselling was to ensure regular contact and to follow up on issues discussed during the individual counselling sessions. At inclusion, patients and caregivers decided which one of them should receive the calls. During the intervention phase the study coordinator contacted the participants approximately 5–8 times with 3- to 4-week intervals. The calls focused on issues discussed at the individual sessions and education courses, but sometimes the conversations included other issues relevant to the individual participant.

**Logbook**

Patients and caregivers were each supplied with a logbook where they were free to write information and thoughts about their daily life. The logbooks aimed to encourage patients and caregivers to make notes about their daily life and prepare for the counselling sessions. Use of the logbooks in the counselling sessions was optional.

**Follow-Up Intervention**

Attempts were made to provide equal treatment for both intervention and control participants in all respects other than the add-on study intervention. At every study visit participants in both groups were interviewed about their current symptoms and daily life and informed about available support programmes (if any) in their local community, and they were free to participate in those during the study. Participation in other local support activities was registered for both groups. Identified special needs led to referral to local care facilities when available and relevant.

**Primary Outcome Parameters**

Since this is one of the first studies to examine the effect of support and counselling programmes in patients with very mild dementia, no consensus exists with respect to gold standards for documenting efficacy. The selection of primary and secondary efficacy parameters (listed in table 1) for this study was based on the specific aims of the intervention and on the outcome in previous similar intervention studies in patients with more advanced dementia. Although the study follow-up was extended to 3 years, the primary outcome parameters were defined for the patients and caregivers at the end of year 1. In the patients the primary outcomes were the emergence of depressive symptoms, proxy-rated health-related quality of life and global cognitive performance (change from baseline at 1 year). In the caregivers the primary outcomes were emergence of depressive symptoms and self-rated health-related quality of life (change from baseline at 1 year).

**Other Quantitative Outcome Parameters**

In addition, a wide range of secondary outcome variables were included (table 1). Two questionnaires, 1 for the patient and 1 for the caregiver, were used to assess additional health issues, social network, content, legal aspects, knowledge of dementia and driving. The caregiver questionnaire was completed by the caregiver, while the patient questionnaire was completed by the rater and based on an interview with the patient. Long-term follow-up data for mortality, morbidity, institutionalization and health care utilization will be drawn from national registries. The methods for assessing health care costs and private costs are described in a separate paper [28].

**Data Analysis**

Comparisons of patient and caregiver characteristics and study outcomes at baseline between randomization groups will be done by t tests for continuous variables and χ² tests for categorical variables. At the 6-, 12- and 36-month follow-up, the outcomes in the 2 randomization groups will be evaluated by the mean of their observed values and the mean change from baseline; comparison between randomization groups will be done by t tests in which subjects with missing values are omitted. Additionally, the difference in development of the primary outcomes over the follow-up period between randomization groups will be investigated in linear mixed models. Here, to adjust for possible bias because of differential dropout from the study, the assessment available at the 6-, 12- and 36-month follow-up will be weighted by the inverse of an estimate of the probability of staying in the study [29, 30]. These probabilities will be estimated from the data in logistic regression models for death and dropout with the dyad’s characteristics and the observed primary outcomes from previous visits as covariates. Difference in mortality and institutionalization between the randomization groups will be evaluated by a hazard ratio from a Cox regression model. Statistical significance will be assessed at a 5% level. Adjustment for multiple testing will be done by the Bonferroni method.

**Table 1. Primary and secondary efficacy parameters**

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<th>Primary efficacy parameters</th>
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<tr>
<td>Patient:</td>
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<tr>
<td>Depressive symptoms (Cornell’s depression scale) [40]</td>
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<tr>
<td>Proxy-rated health-related quality of life (EuroQoL VAS) [41]</td>
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<tr>
<td>MMSE [42]</td>
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<tr>
<td>Caregiver:</td>
</tr>
<tr>
<td>Depressive symptoms (Geriatric Depression Scale, GDS-30) [43]</td>
</tr>
<tr>
<td>Health-related quality of life (EuroQoL VAS) [41]</td>
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<tr>
<th>Secondary efficacy parameters</th>
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<tbody>
<tr>
<td>Health-related quality of life in patients (proxy-rated EuroQoL 5D and VAS) [41, 44]</td>
</tr>
<tr>
<td>Quality of life Alzheimer’s disease scale (QOL-AD) [41, 44]</td>
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<tr>
<td>Health-related quality of life in caregiver (EuroQoL 5D and VAS) [41]</td>
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<tr>
<td>Behavioral symptoms: Neuropsychiatric Inventory Questionnaire (NPI-Q) [45, 46]</td>
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<tr>
<td>Activities of daily living (ADCS-ADL) [47]</td>
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<tr>
<td>Insight scale [48]</td>
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<tr>
<td>Resource utilization in patient and caregiver (Resource Utilization in Dementia, RUD) [49]</td>
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<tr>
<td>Health care resource utilization and costs</td>
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<tr>
<td>Registry-based assessment of health care utilization and key social and health-related events from 12 months before inclusion and during 5 years of follow-up:</td>
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<tr>
<td>Time to nursing home placement and death</td>
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<tr>
<td>Number of hospital contacts in patient and caregiver</td>
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<td>Comorbidity and use of drugs in patient and caregiver</td>
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<th>Patient and caregiver knowledge and attitudes about key issues in dementia</th>
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<tr>
<td>Patient and caregiver content with intervention, public services and network</td>
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[51x105]
Power Calculations

Based on mean scores and score variations for MMSE, EuroQoL VAS, Cornell Depression Scale and GDS from previous studies in other similar AD populations [15, 31–33] we estimated that a group size of 165 was needed in order to detect a minimum effect size (defined as the difference between groups for mean change from baseline to 1 year in observed scores) of 1.2 (MMSE), 2.0 (Cornell), 1.8 (GDS), 6.0 (EuroQoL VAS) with type I error 5% and type II error 10%. The calculations were based on an estimated dropout rate of 20% at year 1. The minimum effect sizes were partly arbitrarily set, partly defined from outcomes in previous intervention studies [15, 31, 32].

Compliance

In the intervention group compliance was defined as the rate of adherence with the schedule for the major components of the multifaceted intervention programme. Thus, patients with satisfactory compliance were defined as patients who had participated with their caregivers in at least 3 counselling sessions (not including the optional network session) and in at least 3 teaching course sessions.

Evaluation of Participants’ Satisfaction with the Intervention Programme

In the intervention group evaluation forms were used at 12 months to quantitatively assess patient and caregiver satisfaction concerning each component in the add-on intervention programme and overall satisfaction with the programme.

Qualitative Study

A separate qualitative study used individual semistructured in-depth interviews of selected patients and caregivers from the intervention group in order to study the experienced outcome of the intervention in qualitative terms. The intervention group evaluation and the qualitative study were conducted and reported prior to the completion of follow-up in the RCT [34].

Ethics

The DAISY trial was conducted in accordance with the Helsinki declaration and evaluated by the local Scientific Ethics Committee [j. nr. (KF) 02-005/04]. All patients and caregivers gave informed consent to participation in the study, which was approved by the Danish Data Protection Agency (j. nr. 2003-41-3178) and registered in the Clinical Trial Database (www.controlled-trials.com/ISRCTN74848736).

Results

In total 330 patients (151 males and 179 females), of whom 30.9% lived alone, were included in the study during a period of 18 months. Their mean age was 76.2 years (range = 54–92), while the mean age of the 330 primary caregivers (110 males and 220 females) was 66.0 years (range = 22–90). The majority of the caregivers were spouses (65.2%). Of the 330 patients, 72.4% had been diagnosed as having AD, 25.9% had mixed AD, and 2.7% met the criteria for DLB. Most patients received antideater treatment with either a cholinesterase inhibitor (93.3%) or memantine (1%).

Discussion

To our knowledge, this prospective study is the largest randomized controlled trial to date to investigate the efficacy and cost-effectiveness of an intensive support and counselling programme for patients in the early phase of AD and their caregivers. With access to national registries for health and social care, the study offers a unique opportunity to follow the long-term outcome and benefits of the intervention.

It is well recognized that caring for a family member with dementia is a challenging task associated with impaired quality of life, and increased risk for depression and other health problems [1, 2], and that monitoring caregiver burden and quality of life is an important and integral task of dementia care. The beneficial effect of a wide range of interventions in order to enable caregiver coping, reduce caregiver burden and improve caregiver quality of life is well recognized from clinical practice and numerous previous studies. The most common interventions are training and education programmes, information-technology-based support, respite care, home care service, support groups and technical aids [5, 7–12]. Most, however, are directed towards caregivers to patients with moderate to severe dementia.

With advancing medicine and increased public awareness of dementia more patients are diagnosed in the early phase of dementia, when they may still have a relatively well preserved autonomy and often request individual counselling, information and training. Thus, a psychosocial intervention programmes in mild dementia must be adjusted to the special needs of caregivers and at the same time include interventions directed towards the patient.

The DAISY Intervention Programmes

The initiation of this study was based on experiences in our own memory clinic from counselling and teaching programmes for patients with mild dementia and their caregivers. We designed the DAISY intervention as a multifaceted and semitailored intervention programme, which aimed to prevent the emergence of depressive symptoms and impairment of health-related quality of life in patients as well as in caregivers, and perhaps even stabilize the functional status for some time. The programme comprised counselling, information and sup-
port to patients with mild dementia as well as to their primary caregivers during the initial months after a diagnosis had been established. Rather than focusing on compensation for lost functional abilities, the philosophy of the counselling programme was to focus on positive resources, intact functions and retained skills, and activities that the patients could still take part in. In order to ensure regular contacts and to follow up on issues discussed in the individual counselling sessions outreach telephone counselling was conducted at regular intervals. The aim of the courses was to provide patients and caregivers with basic information about dementia and its consequences and about living and coping with dementia. The courses also aimed to provide a forum for patients and caregivers to exchange experiences and coping strategies. The counselling and courses were supported by written information and log books to be kept by the patients and caregivers. A unique feature of the programme was the consistent two-lined design with activities and information specifically directed towards the patients as well as the caregivers. Another unique feature was the semiformatted design with some components adapted to the needs of the individual patient or caregiver and with other components common for all participants. The inclusion of the family network in one of the counselling sessions, at the discretion of the patient, aimed to ensure information to all, to prevent stigmatization, to identify important resources in the network and to enable the caregiver to recruit a larger network, when needed.

The intention of the programme was not only to investigate the short-term effect in the period where the intervention took place. By trying to empower both patients and caregivers to understand their symptoms and situations better, and by teaching coping strategies, the intervention and counselling was intended to have a long-term effect.

Follow-Up

For ethical reasons and in order not to leave any participants in the control group without any intervention except blinded rating, both groups received a standardized follow-up intervention. At every study visit participants in both groups were interviewed about their current symptoms and daily life and informed about available support programmes (if any) in their local community, and they were free to participate in those during the study. Any specific and urgent needs identified by the study raters were handled by referral to the general practitioner or local care programmes, when relevant. Thus, these attempts to provide equal treatment of both intervention and control participants in all respects other than the DAISY intervention left all participants with a service well above the level of usual care, as patients with dementia in Denmark, if diagnosed, are often left without any follow-up and without any systematic monitoring of caregiver needs [18, 19].

Selection of Patients

We included only patients with mild recently diagnosed dementia and only patients who had been offered a conventional diagnostic evaluation by a local specialist in dementia. By requiring appropriate diagnostic evaluation we wanted to prevent patients with questionable dementia and nonprogressive cognitive disorders from entering our programme. We did not allow patients with frontotemporal dementias in the programme, as they often have other needs. They may have difficulties in cooperating with the programme, and it may be difficult to mix patients with different disorders in the same sessions. Patients and caregivers were invited to the study regardless of their expected or expressed needs for intervention. Thus, the inclusion into the study was not restricted to patients and caregivers with active help-seeking behaviour. Participation in the study programme was quite demanding, particularly for those entering the DAISY intervention group. Adherence to the many counselling meetings and courses may be difficult for some, particularly for those with impaired somatic health status, busy caregivers, and for those living far away from the study centre. By the mere fact that a diagnosis and caregivers were required for entering the study, our population was selected. However, based on previously reported data on self-rated health and social performance [35, 36], the profile of patients and caregivers in our study was similar to that of typical drug trials and other clinical cohorts in mild AD [31, 32, 37].

Outcome Parameters

There was only little guidance from the literature for the selection of primary outcome parameters and for a priori sample size calculations. Our choices were based on the objectives of the DAISY intervention, on results in previous psychosocial intervention studies for caregivers to patients with more advanced dementia and on drug trials in mild AD. However, a wide range of additional quantitative outcome parameters were included. A unique feature of the study is the potential for follow-up of the cohort far beyond the actual study visits by using information in the systematic and comprehensive Danish health care registries.
Assessment of Efficacy in Psychosocial Interventions

In a recent systematic review aimed to assess the effectiveness of interventions based on information and support provision for informal caregivers of people with dementia in community settings 44 randomized studies were included according to assessment of quality and relevance [17]. Overall the methodological quality of the studies was poor. For instance, only 4 of the studies included an adequate randomization process and concealment of allocation, and a priori sample size calculations were rare. Only few studies were blinded. The majority of the studies had a maximum follow-up of 12 months, and many reported only the positive results. The reviewers concluded that the lack of adherence to best practice in trial-based studies on effectiveness was overwhelming. There was a significant but very small overall effect on depressive symptoms in the caregivers, which should be interpreted with caution. The meta-analysis did not identify any significant effect in other outcome parameters. The authors concluded that there is a pressing need to ensure that supportive interventions at the development stage are accompanied by good quality randomized evaluation in which outcomes that are important for clinicians and carers are measured. Subsequently, at least 1 recent RCT on psychosocial intervention for family carers was unable to identify any effect on primary outcome variables [38].

Thus, there is a great need for randomized controlled studies, which takes into account these methodological considerations, and also for studies directed towards patients with mild dementia. The DAISY study meets all quality criteria established by the Cochrane group [21, 22].

Summary of Strengths and Limitations

In summary, the strengths of the present study are related to the quality of the rater-blinded RCT design, the focus on patients with mild dementia and their caregivers, the inclusion of patients as well as caregivers in the intervention programme, the multifaceted and semitailored design of the intervention, the long-term follow-up and the unique potentials for registry-based follow-up. In addition, the study collected also qualitative data, as there is a risk that important aspects of supportive interventions, such as perceptions and attitudes concerning the programme or staff members, adverse effects or the positive aspects of sharing experiences with others, may be overlooked in quantitatively designed studies. Therefore, experience must also be gained from qualitative studies, which may help design future quantitative studies [34, 39].

Conclusion

There are also limitations associated with the study. The study includes a broad range of patients with mild dementia and not only those with identified needs for psychosocial intervention, which could potentially lead to unexpected adverse effects of the intensive support programme. On the other hand, based on the inclusion criteria the patient group is selected with a high representation of patients with a supportive social network and may not adequately reflect the average population of AD patients. Because there is no established consensus on primary outcomes, the study is to some extent also explorative and will contribute to the design of future studies.

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