Caregiver Burden is Reduced by Personalized Social Support for Non-Dependent Old Persons: a Longitudinal Study of 876 Old Persons and Their Caregivers

Sylvie Arlotto (✉ sylvie.arlotto@ap-hm.fr)  
Assistance Publique Hopitaux de Marseille  
https://orcid.org/0000-0002-2548-8001

Stéphanie Gentile  
Aix-Marseille Universite

Anne Claire Durand  
Assistance Publique Hopitaux de Marseille

Sylvie Bonin-Guillaume  
Aix-Marseille Universite

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Abstract

Background. Informal care provided by family caregivers to old persons is associated with a high risk of burden and poor health status. This study aimed to analyze the impact of Personalized Social Support (PSS) for non-dependent old persons living in the community on caregiver burden, satisfaction, and frailty.

Methods. This non-interventional longitudinal study was performed in the southeast of France: old persons asking for PSS (>70 years of age, with no disability and no severe chronic disease, living at home) and their caregivers were included with a 6-month follow-up. Eligible dyads were visited at home by social workers. Caregiver burden was assessed with Mini-Zarit and frailty status with FiND (Frail Non-Disabled).

Results. 876 dyads were eligible for PSS. Old persons were 82.2 ± 5.8 years old and 77.6% were women. Most caregivers were women (64.5%). Most caregivers were the children of the old person (61%), the rest were mostly spouses. Nearly 64% of old persons were frail and 38% were highly dependent. Follow-up was conducted for 686 dyads (78.3%). Of these, only 569 had PSS. The PSS was mainly for housework and meal preparation. At the time of follow-up, 53% of the caregivers for whom the old person had PSS had less difficulty helping their old person. Two-thirds (61.5%) of caregivers were fully satisfied with the PSS. Whether or not they had received the PSS, 73% of caregivers had reduced burden at 6 months, only 6% still had a high burden, 17% were less frail, and 18% felt healthier than at the time of inclusion.

Conclusions. Our study highlights that non-dependent old person’s caregivers also experience burden. Implementing social support for activities of daily living had a major impact on their burden but not on their level of frailty. This means that the determinants of caregiver frailty are more complex and further studies are needed.

1 Background

In the coming years, the population of old persons will increase considerably. In France, the number of persons aged 75 and over will double between 2013 and 2060, to reach nearly 13 million (17% of the general population) (1). In this context, promoting quality of life, well-being and dignity is a challenge, especially for the most dependent old persons (2). Recent data have shown that about 20% of the European population over 65 years of age is to some extent dependent and, for the most part, assisted by a caregiver: mainly women family members (3). These caregivers are essential to relay care for the old person and will be increasingly involved in the coming years (4). The economic value of this unpaid informal care was estimated to be between 6.1 and 8.3 billion euros per year in France (5).

In France, between 8 and 11 million persons aged 16 years or more regularly help one or more persons in their entourage for health or disability reasons (i.e. nearly one out of six persons in this age group) (6). In 2015, 3 million persons aged 60 and over and living at home regularly received assistance for a health problem or disability (7). The caregiver’s profile varies according to the age of the person being cared for.
With advancing age, the massive help provided by spouses (80% when assisted persons live in a couple) decreases in favor of help provided by children: 55% of old persons aged 70 to 74 receive help from their spouse and 32% from their children. From the age of 75, this situation is reversed: 32% receive help from their spouses and 52% from their children (6).

The type of care provided by caregivers varies greatly, from treatment and care management to shopping assistance and driving out-of-town. When the help provided exceeds the physical and mental capacities of the caregiver, it becomes a chronic stressor generating what is called caregiver's burden (3).

For health workers, it is important to correctly identify the main caregiver, assess the level of stress associated with caregiving, and create a partnership with the caregiver (8). Chronic stress has a negative effect on the caregiver's mental and perceived health (depression, anxiety, psychosomatic and immunological disorders, cardiovascular problems...) and increases the risk of chronic health problems (heart disease, high blood pressure...) and the development of new diseases. In addition, the stress and emotional distress of caregivers has serious consequences for them, in particular their progressive inability to fulfill their role as caregivers, resulting in early institutionalization or unplanned hospitalization of the old person (9) and the risk of abuse or mistreatment for the care recipient, particularly those with dementia and or disruptive behavioral symptoms (10–14). Because of these consequences for caregivers and at a second level for the care recipient, it is necessary to introduce interventions and support for the caregiver (3,15–17).

Two main factors contribute to reducing the burden of primary caregivers. The first is to receive specific help from another family member (18). The second is to benefit from assistance of formal systems (6). In France, the Caisse d'Assurance Retraite et de Santé Au Travail (CARSAT: national administration for retirement and health at work) is responsible for pension management and health and safety at work, and provides assistance to vulnerable persons. This national institution is represented in all French regions. Southern CARSAT (for the Provence Alpes Côte d'Azur and Corsica regions) has implemented Personalized Social Support (PSS), a formal service developed to provide social support to retired beneficiaries who are not yet dependent but who are at risk of becoming so (19). After assessment by a social worker, support is proposed to old persons who request it, who are not dependent (Group Iso Resources 5 and 6 of the AGGIR grid (20)), and who do not receive other assistance for dependency. However, CARSAT does not fully cover the costs associated with the support: the level of CARSAT coverage is correlated with the income level of the beneficiary. The higher the income, the lower the share of CARSAT coverage. Thus, the financial contributions of old persons vary from 10% to 73% of the PSS cost. The impact of this PSS has never been assessed for the beneficiary or for the caregiver.

The main objective was to analyze the impact of PSS implemented for old persons on the burden, perceived health, and frailty of their caregiver. The second objective was to describe the main characteristics of PSS.

2 Methods
1.1 Type of study

A non-interventional observational longitudinal study was conducted in southeastern France in the PACA region (5 million inhabitants, 1 million persons aged 65 years and over) between 2016 and 2017 among 876 dyads: persons aged 70 and over, CARSAT beneficiaries, and their caregivers.

1.2 Study population

All beneficiaries were non-dependent, without serious chronic illness, living at home, who applied for CARSAT Southeast assistance, and whose self-appointed caregiver was eligible. After information and consent, the dyads (old person and identified caregiver) who agreed to participate in the study were consecutively included between 1 April 2016 and 30 June 2017 and were followed for 6 months (T6). If a follow-up after 6 months was not possible, the dyad was excluded from the study.

1.3 Data collection procedure

At the time of inclusion (T0), the old person was assessed at home by a social worker, according to the usual process, during 60 minutes. Social workers had been trained in the concept of frailty and the modalities of the study. The caregiver was invited to be present at the time of assessment of the old person and to complete a self-assessment questionnaire with the help of the social worker if necessary. When not present, the caregiver was assessed during a phone call by a trained Clinical Research Associate recruited specifically for the study. The 6-month follow-up (T6) was done through a telephone call from the Clinical Research Associate. Dyads were considered "unreachable" after 10 unanswered phone calls. The variables collected are presented in Table 1

1.4 Assessment

1.4.1 Mini-Zarit Scale

There were seven questions on a three-point Likert scale (9,21). The items were scored 0 (never), 0.5 (sometimes), and 1 (almost always). The total score results from the sum of the responses and ranges from 0 to 7. Four levels of burden were defined:

- 0 ≥ no burden ≥1
- 1.5 ≥ light burden ≤3
- 3.5 ≥ moderate burden ≤ 5
- 5.5 ≥ severe burden ≤7
1.4.2 FiND questionnaire

The FiND questionnaire consists of five questions. Two questions (A and B) were specifically aimed at identifying individuals with mobility disability. The difficulty to walk 400 meters (A) or the difficulty to climb 10 stairs (B) revealed the presence of mobility disability. Three additional questions (items C–E) were aimed at assessing signs, symptoms, or conditions commonly considered as components of the frailty syndrome: weight loss (item C), exhaustion (item D), and sedentary behavior (item E). Then the caregiver was classified in three categories: disabled, frail, or robust. Subjects reporting none of these 5 items were classified as robust (22).

1.4.3 Autonomy Gerontology Iso-Resource Groups (AGGIR)

This scale is based on the observation of the activities performed or not by the person (20). It measures the level of need for assistance in daily activities. There are six Iso Resources Groups (GIR), from 1 to 6 with decreasing dependency. Group 6 corresponds to independent subjects.

1.4.4 Frailty Group Iso-Resource Evaluation (FRAGIRE)

This grid consists of 17 questions (23). The result is calculated according to an algorithm that produces a score from 0 to 100. This score reflects 3 levels of frailty

- Low frailty: scores < 40
- Intermediate frailty: 40 ≥ score < 60
- High frailty: score ≥ 60

1.5 Data analysis

Statistical analysis was conducted using SPSS Statistics version 20 software. All variables were examined through classical descriptive analysis. Qualitative variables were described by their frequencies and percentages and quantitative variables by their mean and standard deviation (±), minimum, median, and maximum.

A univariate analysis was performed to verify that the population at T6 was comparable to the population at T0. The variable «getting help», recorded as yes/no/yes + nursing, was examined by univariate analyses. The associations between qualitative variables were measured by the Chi 2 test and the exact Fischer test for small numbers. A Student Test or Analysis of Variance (ANOVA) was performed for the quantitative variables.

3 Results
1.1 Characteristics of the dyads included in the study

The study flowchart is presented in Figure 1. Of the 876 dyads interviewed at the time of inclusion, the follow-up was completed for 686 of them (78.3%). For the other 190 dyads, for which follow-up was not possible, the most frequent reasons were unreachable patient and/or caregivers (12.1% of the cohort), withdrawal (7.5%) or death of the old person (2%).

The characteristics of the remaining 686 dyads are summarized in Table 2. The population at follow-up had the same characteristics as the total population at the time of inclusion. The average age of the old person was 82.3 years (± 5.9); 11% were over 90 years old. The great majority were women (78%) and half lived alone (53%). The old persons were not dependent and were classified in GiR 5 or 6, respectively 48% and 52%. According to the FRAGIRE scale, 94% were frail; 48% of them were at high risk of frailty.

In 66% of cases, caregivers were women with an average age of 62.7 years (± 13.6). Most of the time, caregivers were children or stepchildren (62%); spouses represented 27%. The others (11%) were friends, neighbors, grandchildren, and siblings. Among the other caregivers, 71.7% were family members. In 75% of the cases, the caregivers were a couple and had children, 35% of whom were still living with them. Almost half (48%) of caregivers had a high level of education and almost 40% of them were employed. For 48% of them, incomes were low.

At the time of inclusion, the support provided by caregivers had an impact on their life, particularly on their outings and their availability to have free time for a few days. In addition, 92.5% of caregivers experienced difficulties in providing support. Many reasons were reported, including poor health, lack of time, and negative impact on their other family obligations (for example, the time available for children). One in five caregivers also reported a lack of material or financial resources. Finally, 17% of them indicated that they had a poor relationship with their old person.

In addition to CARSAT’s support, one in four dyads (27%) received intervention from another professional, in most cases (88%) a health professional.

1.2 Characteristics of the personalized social support (PSS)

PSS was mainly for household chores (93.6%); 17.3% received shopping assistance, and 16.1% received ironing assistance. Other types of support included escort during outings (6.7%), meal preparation assistance (6.2%), grooming assistance (2.3%), meal assistance (1.9%), management assistance (1.9%), and dressing assistance (0.9%)(Table 3).

One in five old persons received two supports, and 10% more than two. The most frequent help association was household chores and ironing (29%) or shopping (27%) or meal preparation (5%) or outings assistance (5%).
1.3 Level of satisfaction with the PSS

Most caregivers, as well as the old persons, were very satisfied with the quality of PSS (Table 3). Almost two-thirds of old persons as well as their caregivers felt that their needs were fully met. The main cause of dissatisfaction was the number of hours received, which was considered insufficient by two-thirds of the dyads. One out of five felt that they should have had other support.

Nearly 9% complained of dysfunctions in the organization of the support plan, such as delays, absences, and frequent changes of service providers. But these problems were resolved quickly in half of the cases (Table 3).

1.4 Refusal of the PSS

For about one-fifth of dyads, PSS was not implemented at the time of the follow-up. This was mainly due to the old person's refusal to receive someone at home (29.9%), price considered too high (19.9%), and proposed support considered not adequate (8.5%) or could not be provided (6.8%). In addition, 10% preferred other support systems than PSS. And 11% were still awaiting implementation. Finally, 16% reported having other reasons for refusing PSS, including institutionalization.

The demographic characteristics of the dyads who did not receive PSS were similar to those of the dyads who received it. However, caregivers who did not have PSS had a poorer relationship with their old person and were less satisfied with their relationship with the health professional.

1.5 Impact of the PSS

Most dyads received PSS (569 with PSS versus 117 without PSS), which reduced the power of statistical comparisons and thus the ability to prove the impact of the PSS intervention. Among the caregivers who received PSS, a minority (n = 38) reported that they had reduced the number of hours they spent with their old person. For more than 80% of caregivers who received PSS or did not receive it, the time spent with their old person remained unchanged from the inclusion time, averaging 7.5 ± 7.0 hours and 3.8 ± 2.5 visits per week.

In addition, among caregivers who received support, obtaining support for a specific task was associated with a reduction of the time spent on that task. This was true for the household chores: 74% of caregivers who provided home help reported that they had reduced the time spent on this task by obtaining specific household help, while the decrease in time spent on household chores was only 22% for those who had not received specific help. Time reduction was also significant for ironing assistance (55.6% vs 29.1%, p < 0.05), outings assistance (31.8% vs 8.7%, p < 0.05), shopping assistance (56.7% vs 10.1%, p < 0.05), and meal preparation assistance (36.4% vs 8%, p < 0.05) (Figure 2)
Figure 3 shows the degrees of change observed for burden, perceived difficulties, self-rated health, and frailty at the 6-month follow-up. At follow-up, 53% of caregivers experienced fewer difficulties in caring for their old person. This improvement was proportional to the support provided: it was significant for those who had a nursing assistance in addition to PSS compared to those who had only PSS and those who had no assistance at all (62.7% and 53.6% respectively vs 39.1%, p < 0.05).

The burden was reduced for 73% of caregivers, whether or not they received PSS. The analysis of the mini-Zarit items revealed that caregivers who received PSS obtained better results on the item concerning the impact on their daily life (6.3 % vs. 11.8% for those without PSS, p < 0.05).

At follow-up, less than one in five caregivers felt their health status was better than at the time of inclusion. In addition, according to the FiND questionnaire, 17% of them were less frail, but 12% appeared frailer. The improvement in frailty was not linked to the implementation of PSS, even if the items constituting the FiND were analyzed separately. However, the improvement in frailty was greater for spouses than for other categories of caregivers (23% vs. 16% for child caregivers and 6% for other caregivers, p<0.05). Caregivers living with the old persons were more likely to have improved their frailty compared to caregivers not living with them (24.5% versus 12.9, p = 0.000). They were, for the most part, spouses, and therefore older and frailer than other caregivers.

At follow-up, 15% of the old persons reported better health status, while 31% reported poorer health. These changes in health status were not related to the implementation of PSS. Concerning the level of caregiver involvement, those caregivers who performed the most tasks (> 3 tasks) were more likely to have reduced their level of burden (90.1% vs. 84.3%, p = .008), less likely to have difficulty caring for the old person (59% vs. 42%, p = .001), and more likely to have improved their level of frailty (19.2% vs. 9.4%, p = .013). Caregivers with a moderate/severe burden at inclusion (37.9%) were more likely to have increased their level of burden (94.8% versus 84.2% who decreased by at least one level or more, p = 0.000) and were more likely to have less difficulty caring for their old person (65.3% versus 44.4%, p = 0.000).

4 Discussion

No studies, to our knowledge, have been conducted on caregivers of a non-dependent old-age population without serious chronic disease living at home. Most studies have focused on caregivers of dependent populations (24). Our results show that the population of caregivers of non-dependent old persons has a profile similar to that of caregivers of dependent old persons (6,19) and persons with disabling illness (11,25–27) in terms of socio-demographic characteristics and of level of burden, which also has a negative impact on their health(24–26,11)(19,26–28). Our results also show that the caregiver's burden appears well in advance of the recipient's dependency. Moreover, the number of hours and tasks carried out are as numerous as those carried out by the caregivers of dependent persons (19).

PSS was implemented for 83% of old persons. Most dyads were satisfied and considered that it covered their needs. To the extent that almost all participants benefited from PPS, the measurement of impact
was limited due to the size of the two subgroups (569 with PSS versus 117 without PSS). Paradoxically, results showed that, at follow-up, the burden level had improved for 73% of caregivers, whether they had PSS or had merely been consulted about needing PSS.

According to a literature review, the burden includes several component: a psychological component related to the difficulties experienced and an emotional component related to the relationship with the old person, and finally a physical component related to the type and number of tasks performed (29). That the burden improved for all caregivers suggests that the consultation of social workers with the caregivers to identify the needs of the old person for the implementation of PSS had a positive impact on two of the three components of burden, the psychological and emotional components. Indeed, the consultation could be felt by caregivers as a space to talk, a time to be listened to, which allowed them to express their difficulties and which reduced their feeling of loneliness. This consultation probably reassured them, and they were able to project themselves towards the possibility of obtaining future support if it became necessary (30).

Low social support or social isolation, and bad relationship between caregiver and the care recipients are known to increase the risk of burden (19,31) and abuse in community-dwelling old persons. Social support is a protective factor against abuse of old patients (32). This study did not aim at assessing mistreatment or abuse of the old persons. Yet we can hypothesize that this evaluation for PSS, accepted or not, prevented some mistreatment, abuse, or early unplanned hospitalizations particularly for caregivers at high risk, i.e, frail, high burden and high number of daily tasks. (13).

The perceived health and the frailty level improved only for one in five caregivers; this was unrelated to obtaining PSS (33). The fact that the burden, perceived health, and frailty improved independently of the implementation of PSS shows that the impact of a dedicated consultation is very positive overall, as shown in the bibliography (34).

As expected, obtaining PSS reduced the amount of time the caregiver spent performing the tasks. The results show that obtaining specific help reduces not only the time spent on the targeted task but also the time spent on other tasks. This is particularly true for caregivers who were ironing or cleaning before obtaining PSS; many (30% and 20% respectively) reported reducing the time spent on this task, even if the assistance provided by PSS was not specific to the task. It is likely that the professionals involved did more than the targeted service and that the old persons were able to focus on fewer tasks but more effectively, which indirectly relieves the caregiver. This could also explain the improvement in the difficulties of caregivers, which was correlated with the implementation of PSS and affected 56% of caregivers with PSS compared to 39% of those without it (p < 0.05).

However, except in 38 cases, the number of hours and weekly visits to the old person did not decrease since obtaining the PSS. Several explanations are possible. First, data were collected only for caregivers who did not live with the old person, which may have biased the positive impact of the PSS. We found that spouses were the caregivers who did the most work, but the number of hours was not collected for them, so it is possible that the effect of PSS may not be highlighted. Second, the needs may have
increased during the 6 months or they were not fully covered by CARSAT, and therefore the impact on the amount of time may be moderated. Finally, maintaining the number of hours devoted to assistance may not have the same qualitative significance. Thanks to PSS, the caregiver spends less time performing domestic tasks but spends more time on the relationship with the old person.

Among the biases in selection, the first is related to the financial contribution requested from the old person. Our population is precarious: the number of caregivers close to the poverty line is much higher than that of the general population of the PACA region: 17% in PACA against 49.9% for our study (35). However, some caregivers may have underestimated their incomes because they feared that CARSAT would refuse the social support requested or that the contribution requested would be too high. A second potential selection bias is that we included old persons who asked for help with daily living tasks. Thus, the status of the old persons included in our study is in an intermediate level of autonomy. Our results showed that only 5% of the old persons were assessed as at low risk of frailty according to the FRAGIRE grid. A third limitation regarding the measurement bias on the health status of caregivers is that we did not collect objective health elements, such as walking speed. As a result, comparisons of pre- and post-PSS health status were based solely on perceived health.

The originality of this study is its target: caregivers of old persons without serious chronic diseases, autonomous, and living at home. No studies, to our knowledge, have targeted these caregivers, although in terms of frequency, they are certainly the most numerous and constitute a real public health challenge, especially in the future.

Moreover, while most dependency studies have focused on psychological or medical support activities, our study concerns the impact of social assistance exclusively. The study is based on a large sample and a small proportion of persons lost to follow-up. All these points constitute a good level of internal validity.

These results seem to underline the importance of supporting natural caregivers, and the benefit of a dedicated consultation. These consultations should target, first of all, caregivers who have many tasks and/or a high level of burden.

There are many organizations founded and managed by informal caregivers, whose purpose is to provide information and support to peers (6). Unfortunately, they are not always easily accessible. They require that caregivers be aware of their role as caregivers, and that they have sufficient awareness to detect not only the needs of the person they support, but also their own needs and/or difficulties. In addition, these organizations are often focused on a particular pathology, which may exclude caregivers of old persons with no particular pathology.

The first action would therefore be to systematically identify during routine medical consultations whether a person is a caregiver in order to advise and refer them to dedicated organizations and associations. Action should be taken before physical and/or psychological disorders appear. We could consider setting up a specific consultation for them, as is done in the context of dependent old persons. Most of these types of consultations are medicalized or for psychological care. We believe that these
consultations could first identify the needs of caregivers and make a situational diagnosis that would make it possible to offer personalized support.

5 Conclusion

Our results show that the population of caregivers of non-dependent old persons is similar to that of caregivers of dependent old persons or persons suffering from serious diseases in terms of socio-demographic characteristics but also in terms of burden level, perceived health status, and risk of frailty. The evaluation of the implementation of a social assistance plan to improve home support for old persons shows that the intervention of a social worker in the home, whether or not the assistance plan is officially granted, improves the burden on caregivers, but does not quantitatively change the caregiver’s investment in the old person.

Health professionals and/or social workers should identify caregivers in order to offer care adapted to their needs and expectations and not only through the needs of the person being cared for. The implementation of dedicated consultations could help caregivers discover the limits of their role and to evaluate the difficulties they encounter so that they can be supported.

6 Abbreviations

PSS: Personalized Social Support
CARSAT: Caisse d'Assurance Retraite et de Santé Au Travail
PACA: Provence Alpes Côte d'Azur
CNIL: Commission Nationale de l'Informatique et des Libertés
CRA: Clinical Research Associate
AGGIR: Autonomy Gerontology Iso-Resource Groups
GIR: Iso Resources Groups
ANOVA: Analysis of Variance
FRAGIRE: Frailty Group Iso-Resource Evaluation

7 Declarations

1.1 Ethics approval and consent to participate

This study was done in accordance with the bioethics laws at the time of data collection and did not require submission to the Protection of Persons Committee. The study was conducted in compliance with
the IRB. The approach with the study population was consistent with CARSAT's routine professional practice.

Only data collection concerning caregivers was carried out specifically for this study. Prior to the inclusion to the study, every subject received an information letter. At the beginning of the assessment they were then informed again verbally to obtain a non opposition to participate to the study. The subject was free to withdraw from the study at any time. All nominative data were managed exclusively by CARSAT, which is authorized by the Commission Nationale de l'Informatique et des Libertés (CNIL) (n2005-38) to deal with the protection of personal data.

1.2 Consent for publication
Not applicable.

1.3 Availability of data and material
The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

1.4 Competing interests
The authors declare that they have no competing interests

1.5 Funding
This study was funded by CARSAT Southeast, as part of the implementation of social support for autonomous old persons. The funder (CARSAT) agreed with the design of the study. Health worker of the CARSAT partly participated to the data collection (old subjects and caregivers at inclusion) on a routine practice. The CARSAT did not participate to the data analysis and interpretation, neither to the writing of the manuscript.

1.6 Authors' contributions
ACD, SBG and SG developed the protocol. SA and ACD participated in the data collection. SA and SG were responsible for data analysis. SA, SBG and SG interpreted the data. All authors contributed to the writing and revision and approved the final manuscript.

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Tables

Table 1. Variables collected from the dyad at the time of inclusion and at follow-up
Variables collected from caregivers at the time of inclusion (T0)

| Sociodemographic data |
|-----------------------|
| Age, gender, family status (single/married) and number of children living at home |

The relationship with the old person

Spouse caregivers (spouses, partners, and ex-spouses)

Child caregivers (children and stepchildren)

Other caregivers (no family relationship and other family members)

The level of education

(no diploma, Youth training, BTEC First Diploma, A levels, higher education level), which has been grouped into 2 categories:

< A level (UK)

≥ A level

Professional activity grouped into two categories:

Caregivers who are working

Inactive caregivers (includes retired, unemployed and disabled)

Monthly income of the tax household

< 740€, between 740 and 1200€, between 1201 and 2200€, between 2201 and 4400€, > 4400€

Income: the variable was created from 3 variables: monthly income, family status, and number of children living at home

low income for single caregivers with incomes below 1200€ and caregivers living alone or in couple with dependent children and with incomes below 2200€ (36)

not low income

Activities performed for the old persons at the time of inclusion

Number of hours and days per week spent with the old person

Number and type of tasks performed with the old person:

administrative tasks, activities of daily living (transportation, household chores, ironing, meal preparation, shopping) and nursing (meal assistance, treatment assistance, getting up, going to bed, dressing, grooming)

The consequences of the assistance provided at the time of inclusion

Perceived difficulties in fulfilling their role as caregivers (5 points on the Likert scale from "not at all" to "a lot")
Difficulty in fulfilling their role as caregivers due to their own health status, family obligations, lack of material or financial resources, lack of specialized institutions or services, lack of dialogue with professionals or support services, lack of time, lack of expertise and lack of information.

Feel a negative impact of the help on relationships with other family members, on the day's outings, on leaving for a few days (5-point Likert scale from "never" to "always"). This variable was recorded as binary, "impact or not".

The quality of the caregiver's relationship with the old person (5-point Likert scale from "very good" to "very difficult"). This variable was recorded as binary, "good or bad relationship".

Caregiver Burden: Mini-Zarit Scale (ZBI) (9,21,37)

Caregiver's frailty: the FiND questionnaire (22)

Health status and health perceived by the caregiver. (38)

Perceived health measured by a question: "How do you rate your health?" (5 points on the Likert scale from "excellent to "bad")

**Variables collected from old persons at the time of inclusion**

Sociodemographic data: age, gender, and family status

The level of dependency (20) by the Autonomy Gerontology Iso-Resource Groups (AGGIR) scale

Frailty of the old person (23) by the Frailty Group Iso-Resource Evaluation (FRAGIRE)

**Variables collected from caregivers at follow-up**

Number and type of tasks performed with the old person:

management tasks, activities of daily life (transportation, household chores, ironing, meal preparation, shopping) and nursing (meal assistance, treatment assistance, getting up, going to bed, dressing, grooming);

Perceives more difficulties to fulfill their role as caregivers (5-point Likert scale from "not at all" to "a lot")

Impact on perceived difficulties: this variable was calculated from the “perceived difficulties” variable at inclusion and follow-up. It represents the change in perceived difficulties and is divided into 5 levels (from significant improvement to significant deterioration).

Caregiver Burden: Mini-Zarit Scale (9,21,37)

Impact on burden: this variable was calculated from the "Mini-Zarit" variable at inclusion and follow-up. It represents the change in burden level and is divided into 5 levels (from significant improvement to significant deterioration).

Caregiver's frailty: the FiND questionnaire (22)

Impact on frailty: this variable was calculated from the "FiND" variable at inclusion and follow-up. It represents the change in frailty level and is divided into 5 levels (from significant improvement to significant deterioration).

Health status and health perceived by the caregiver.
Perceived health measured by a question: "How do you rate your health?" (5 points on the Likert scale from "excellent to "bad")

Impact on perceived health: this variable was calculated from the "perceived health" variable at inclusion and follow-up. It represents the change in perceived health and is divided into 5 levels (from significant improvement to significant deterioration).

**Variables collected from caregivers and old persons at follow-up**

Receiving help (yes/not)

*if receiving help*

what help (many possible answers):

- household chores
- ironing
- shopping
- transportation
- meal preparation
- meal assistance
- grooming assistance
- dressing assistance
- securing the house (shower, stairs...)
- remote assistance
- artistic activities
- physical activities

satisfaction (5-point Likert scale), recoded in binary (totally satisfied or not)

if not satisfied, why not?

quality of aid (5-point Likert scale), recoded in binary (very good quality or not)

having experienced a problem with the performance of the service

which one

rapid resolution of the problem
if not receiving help

reasons for the refusal (economic, change of opinion, proposed aid different from the aid requested...)

Other professional care support (nurse, physiotherapist, ...)

Table 2. Characteristics of dyads with or without Personalized Social Support (PSS) at inclusion and at follow-up
|                          | Total     | PSS       | No PSS    | p     |
|--------------------------|-----------|-----------|-----------|-------|
|                          | N = 686   | n = 569   | n = 117   |       |
| **Caregiver characteristics** |           |           |           |       |
| Age. mean ± SD           | 62.7 ± 13.6 | 62.9 ± 13.3 | 61.5 ± 14.7 | .310  |
| Women % (n)              | 66 (453)  | 67 (381)  | 61.5 (72) | .259  |
| Single % (n)             | 21.9 (149)| 20.8 (117)| 27.4 (32) | .118  |
| Working % (n)            | 39.8 (273)| 401 (228) | 3805 (45) | .746  |
| High school education % (n) | 48.4 (326)| 49.4 (276)| 43.9 (50) | .283  |
| Low income % (n)         | 48.2 (299)| 49.5 (293)| 42.2 (46) | .166  |
| **Relationship Caregiver – Old Person** |           |           |           |       |
| Spouse caregivers % (n)  | 27.4 (188)| 27.6 (157)| 26.5 (31) | .906  |
| Child caregivers % (n)   | 61.5 (422)| 61.2 (348)| 63.2 (74) |       |
| Other caregivers % (n)   | 11.1 (76) | 11.2 (64) | 10.3 (12) |       |
| **Caregivers limited in the assistance they provided at T0 by** |           |           |           |       |
| own health status % (n)  | 45.1 (301)| 45.7 (253)| 42.5 (48) | .535  |
| family obligations % (n) | 30.8 (205)| 31.6 (175)| 26.5 (30) | .285  |
| lack of material/financial resources % (n) | 19 (127) | 18.1 (100) | 23.9 (27) | .149  |
| lack of specialized institutions/services % (n) | 7.8 (52) | 7 (39) | 11.5 (13) | .107  |
| lack of dialogue with professionals % (n) | 6.4 (43) | 5.6 (31) | 10.6 (12) | .048  |
| lack of time % (n)       | 45 (300)  | 44.8 (248)| 46 (52)   | .807  |
| lack of expertise % (n)  | 15.2 (101)| 14.1 (78) | 20.4 (23) | .091  |
| lack of information about the disease % (n) | 4.1 (27) | 3.6 (20) | 6.2 (7) | .207  |
| **Caregivers for whom the assistance provided has had an impact on** |           |           |           |       |
| family Life % (n)        | 27.3 (186)| 26.5 (150)| 31 (36)   | .323  |
| the day's outings % (n)  | 47.7 (324)| 47.4 (267)| 49.1 (57) | .737  |
| leaving for a few days % (n) | 48.1 (327)| 49 (276) | 43.6 (51) | .284  |
having a difficult relationship with the old person % (n) | 17.4 (118) | 15.9 (90) | 24.6 (28) | .026

| At follow-up | Reduced number of hours per week % (n) | 55.9 (38) | 57.9 (33) | 45.5 (5) | .447
| Fewer difficulties % (n) | 53 (350) | 55.7 (307) | 39.1 (43) | .001
| Burden improvement % (n) | 73.3 (439) | 73.5 (371) | 72.3 (68) | .821
| Self-rated health improvement | 17.7 (118) | 17.1 (95) | 20.4 (23) | .411
| Frailty improvement | 16.9 (108) | 16.4 (88) | 19.4 (20) | .462

Table 3_ Characteristics of Personalized Social Support (PSS) and satisfaction
| PSS received % (n)                      |
|----------------------------------------|
| Household chores                       | 96.3 (545) |
| Shopping assistance                    | 17.3 (98)  |
| Ironing assistance                     | 16.1 (91)  |
| Outing assistance                      | 6.7 (38)   |
| Meal preparation                       | 6.2 (35)   |
| Grooming assistance                    | 2.3 (13)   |
| Meal assistance                        | 1.9 (11)   |
| Management                             | 1.9 (11)   |
| Dressing assistance                    | 0.9 (5)    |
| Securing the house                     | 0.9 (5)    |
| Remote assistance                      | 0.5 (3)    |
| Artistic activities                    | 0.5 (3)    |
| Physical activities                    | 0.2 (1)    |
| Dyads totally satisfied with PSS % (n) | 61.5 (340) |

| If not satisfied. Why not? % (n)       |
|----------------------------------------|
| Need more time                         | 70.5 (67)  |
| Need for other assistance              | 20 (19)    |
| Too expensive                          | 2.1 (2)    |
| Other reasons                          | 7.4 (7)    |
| Very good quality of PSS% (n)          | 64 (355)   |
| Having experienced a problem with the performance of the service % (n) | 8.9 (48) |
| Changes in work hours, delays. absences.... % (n) | 62.2 (28) |
| Quality of service, frequent change of service provider.... % (n) | 22.2 (10) |
| Other (delay. don't know....) % (n)    | 15.6 (7)   |
| Rapid resolution of the problem % (n)  | 47.2 (17)  |

**Figures**
Figure 1

Flow-chart showing participation in the study, from inclusion to follow-up.

876 dyads included
18 old persons deceased
66 dyads refused follow up
106 dyads unreachable

686 dyads analyzed at follow-up

569 dyads with PSS
117 dyads without PSS
Figure 2

Impact of PSS on burden, self-rated health, and frailty.

* This image shows how the time spent on a task, such as household chores, decreases as a result of obtaining household-specific assistance (74%), but also as a result of obtaining assistance other than for the household (22%).
Figure 3

Impact of PSS on the time spent on tasks.