6-11-2013

Correlations among Coping Skills and Life Satisfaction in Ethnic Older Caregivers

Didiana de la Osa
Florida International University, didiana.delaosa@yahoo.com

DOI: 10.25148/etd.FI13080510
Follow this and additional works at: https://digitalcommons.fiu.edu/etd

Part of the Psychology Commons

Recommended Citation

https://digitalcommons.fiu.edu/etd/899

This work is brought to you for free and open access by the University Graduate School at FIU Digital Commons. It has been accepted for inclusion in FIU Electronic Theses and Dissertations by an authorized administrator of FIU Digital Commons. For more information, please contact dcc@fiu.edu.
CORRELATIONS AMONG COPING SKILLS AND LIFE SATISFACTION IN
ETHNIC OLDER CAREGIVERS

A thesis submitted in partial fulfillment of the
requirements for the degree of
MASTER OF SCIENCE
in
PSYCHOLOGY
by
Didiana de la Osa

2013
To: Dean Kenneth G. Furton  
College of Arts and Sciences

This thesis, written by Didiana de la Osa, and entitled Correlations among Coping Skills and Life Satisfaction in Ethnic Older Caregivers, having been approved in respect to style and intellectual content, is referred to you for judgment.

We have read this thesis and recommend that it be approved.

_______________________________________
Dionne Stephens

_______________________________________
Lisa Arango

_______________________________________
Mary Levitt

_______________________________________
Leslie Frazier, Major Professor

Date of Defense: June 11, 2013

The thesis of Didiana de la Osa is approved.

_______________________________________
Dean Kenneth G. Furton  
College of Arts and Sciences

_______________________________________
Dean Lakshmi N. Reddi  
University Graduate School

Florida International University, 2013
The purpose of the present study is to extend our current understanding of the effects of caregiver burden on life satisfaction by examining whether or not there are ethnic differences in coping strategies used to manage caregiving. Several specific hypotheses were tested in order to determine the linkages among age, gender, ethnicity (i.e., familism, filial piety), caregiver burden, coping with caregiving, and life satisfaction. A total of 103 Hispanic and Non-Hispanic White participants ages 60 and older were included in this study (mean age was 67.42; 16.5% male; 83.5 % female; 52.4% Hispanic; 47.6% Non-Hispanic White). The results suggest that demographics and certain coping skills can influence levels of life satisfaction and burden experienced by caregivers. The findings from this study shed light on how to structure effective psychoeducational interventions, facilitate adaptive coping, reduce burden, and improve life satisfaction for older adult caregivers.
# TABLE OF CONTENTS

<table>
<thead>
<tr>
<th>CHAPTER</th>
<th>PAGE</th>
</tr>
</thead>
<tbody>
<tr>
<td>I. INTRODUCTION</td>
<td>1</td>
</tr>
<tr>
<td>II. LITERATURE REVIEW</td>
<td>4</td>
</tr>
<tr>
<td>Coping Skills</td>
<td>5</td>
</tr>
<tr>
<td>Life Satisfaction</td>
<td>17</td>
</tr>
<tr>
<td>III. METHODOLOGY</td>
<td>31</td>
</tr>
<tr>
<td>Participants</td>
<td>31</td>
</tr>
<tr>
<td>Measures</td>
<td>31</td>
</tr>
<tr>
<td>Procedure</td>
<td>35</td>
</tr>
<tr>
<td>Data Analytic Plan</td>
<td>35</td>
</tr>
<tr>
<td>IV. RESULTS</td>
<td>39</td>
</tr>
<tr>
<td>Descriptive Data</td>
<td>39</td>
</tr>
<tr>
<td>Coping and Depression</td>
<td>40</td>
</tr>
<tr>
<td>Demographic Differences in Burden</td>
<td>44</td>
</tr>
<tr>
<td>Demographic Differences in Coping</td>
<td>45</td>
</tr>
<tr>
<td>Demographic Differences in Life Satisfaction</td>
<td>50</td>
</tr>
<tr>
<td>The Role of Familism and Filial Piety</td>
<td>51</td>
</tr>
<tr>
<td>Coping as a Moderator for Demographics and Life Satisfaction</td>
<td>52</td>
</tr>
<tr>
<td>Coping as a Moderator for Burden and Life Satisfaction</td>
<td>55</td>
</tr>
<tr>
<td>V. DISCUSSION</td>
<td>57</td>
</tr>
<tr>
<td>Outcomes of Caregiving</td>
<td>58</td>
</tr>
<tr>
<td>Limitations</td>
<td>65</td>
</tr>
<tr>
<td>Potential Implications</td>
<td>66</td>
</tr>
<tr>
<td>REFERENCES</td>
<td>76</td>
</tr>
<tr>
<td>APPENDICES</td>
<td>87</td>
</tr>
</tbody>
</table>
CHAPTER I

Introduction

The interaction between ethnicity, caregiver burden, and coping styles in the elderly is a multidimensional and elaborate issue that is still not well understood. However, it is imperative that the interactions among these variables be investigated because the aging population in the United States is increasing in diversity which may impact what we know about effective coping for late life issues such as caregiving. The influences of psychological and sociocultural factors on the coping skills that caregivers employ has been explored (Campbell, Converse, & Rodgers, 1976; Chakrabarti & Gill, 2002; Hooker, Frazier, & Monahan, 1994; Lewinsohn, Redner, & Seeley, 1991; Shimazu, Shimazu, & Odara, 2005). More specifically, the role of personality (Chakrabarti, et al., 2002; Hooker, et al., 1994; Hooker, Monahan, Bowman, Frazier, & Shifren, 1998; Lewinsohn, et al., 1991), culture (Caspi & Elder, 1986; Greer & Brown, 2011; Greer & Chwalisz, 2007; Outten, et al., 2009; Utsey, Ponterotto, Reynolds, & Cancelli, 2000), relationship with care-recipient (Lawrence, Tennstedt, & Assman, 1998; McClendon, Smyth, & Neundorfer, 2004; Quayhagen & Quayhagen, 1988), economic status (Borg & Hallberg, 2006; Jang, Chiriboga, Kim, & Phillips, 2008; Rice, Near & Hunt, 1980), role strain (Barusch & Spaid, 1991; Bonebright, Clay, & Ankenmann, 2000; Choo, Low, Karina, Poi, Ebenezer, & Prince, 2003; Collins & Jones, 1997), as well as occupational stress (Bonebright, Clay, & Ankenmann, 2000; Gueritalt-Chalvin, Kalichman, Demi, & Peterson, 2000), and gender (Collins, et al., 1997; Hooker, Manoogian-O’Dell, Monahan, Frazier, & Shifren, 2000; Kristensson Ekwall & Rahm Hallberg, 2006) have all been shown to increase the experience of stress (Collins, et al., 1997; Gueritalt-Chalvin, et al.,
decrease life satisfaction (Arango-Lasprilla, Olivera Plaza, Drew, Perdomo Romero, Arango Pizarro, Francis, & Kreutzer, 2010; Borg, et al., 2006; Haley, LaMonde, Han, Burton, & Schonwetter, 2003), influence depression (Chang, Noonan, & Tennstedt, 1998; Collins, et al., 1997; Haley, et al., 2003; Hooker, et al., 1998; Lawrence, et al., 1998), and increase co-morbid conditions in caregivers (Hooker, et al., 1998; Haley, et al., 2003; Lazarus & Folkman, 1984). The proposed study intends to expand on previous research focusing specifically on how the burden of caregiving affects life satisfaction in an ethnically diverse sample of older adults. The mediational role of coping will be investigated in varied ethnic groups to examine its impact on outcomes of caregiver satisfaction beyond the known effects of demographics, such as gender and age. The knowledge of preferred coping styles in ethnic older caregivers is necessary to better understand the relationship between caregiving and life satisfaction in this population.

The present study aims to investigate the influence that demographic variables, such as age, gender, ethnicity, and caregiver burden, may have on the evaluation of aging caregivers’ life satisfaction. Elderly female caregivers are expected to experience lower levels of life satisfaction than their male counterparts as a consequence of their more frequent use of ineffective coping strategies, such as escape and avoidance, (Lutzky & Knight, 1994; Matud, 2004; Ptacek, Smith, & Dodge, 1994), which have been shown to increase levels of distress (McClendon, et al., 2004). Furthermore, being of a certain ethnicity may have an effect on the coping strategies employed by caregivers in this study. Employing maladaptive coping strategies can exacerbate caregiver stress and, in turn, lower life satisfaction. Knowing what resources are more successful among certain
ethnicities can aid efforts of intervention and ultimately improve their satisfaction with life.
Coping Skills

The literature on stress and coping has grown extensively over the last 30 years. Yet, to date the most widely accepted theory of the stress and coping process is that of R. Lazarus and S. Folkman, (1984). The Transactional Theory of Stress and Coping (Lazarus & Folkman, 1984), articulates the process of stress and coping as a result of how individuals interact with their environment (Folkman & Lazarus, 1980). The first phase of coping with stress is known as the primary appraisal process, in which the individual determines whether the stressor presents a threat or challenge (Lazarus & Folkman, 1984). How the stressor is assessed determines how the individual reacts on cognitive, physical, and psychological levels. The primary appraisal process also influences the next phase, the secondary appraisal process (Lazarus & Folkman, 1984). Secondary appraisal is a cognitive evaluation of whether or not the resources in the individual's reservoir are sufficient to manage the stressor (Lazarus & Folkman, 1984). The secondary appraisal process can mediate the primary appraisal process and is critical in deciding the effects of the stressor and the coping strategies used to manage it (Lazarus & Folkman, 1984). To summarize, the Transactional Model of Stress suggests that when an individual experiences an event it is immediately appraised in terms of the effects of the stressor, simultaneously and in conjunction with the primary appraisal process, a secondary appraisal process determines how the individual will cope with the stressor. Both processes are integral to adjustment to the stressor.
According to Lazarus & Folkman, (1984), individuals may engage in a variety of coping strategies to manage immediate and chronic stress. Coping is conceptualized as the thoughts and actions; or emotions, cognitions, and behaviors, which an individual engages in to reduce, ameliorate, or adjust to the stressor. There are a wide array of behavioral responses individuals engage in to manage distress, from self-blame, positive reappraisal, action-oriented strategies, to strategies such as turning to religion, substance use, and seeking social support (Carver, 1997; Chakrabarti & Gill, 2002). However, Lazarus & Folkman, (1984), and others (Chakrabarti & Gill, 2002; Folkman, 1984) have stated that these strategies can be divided into two conceptually different forms of coping. One form entails active and effortful strategies designed to control, change or eliminate the stressor, known as problem-focused coping (Folkman & Lazarus, 1980). The second form of coping focuses on managing emotions and reactions brought about by the stressor, known as emotional-focused coping (Folkman & Lazarus, 1980).

Research abounds on problem-focused and emotion-focused coping (Carver, 1997; Cheng, 2001; Cheng & Cheung, 2005; Folkman & Lazarus, 1980; Lazarus & Folkman, 1984). In addition, a large body of research exists exploring the coping strategies of spousal caregivers of Alzheimer's or Parkinson's disease patients (Choo, et al., 2003; Hooker, et al., 1994; Hooker, et al., 2000; McClendon, et al., 2004; Morano, 2003). Studies show that people, in general, use a combination of both problem-focused and emotion-focused coping strategies in order to deal with the stressors they encounter (Cheng, 2001; Cheng, et al., 2005; Folkman & Lazarus, 1980). Indeed the flexible use of coping styles has been found to be more effective in reducing the negative impact a stressor may have on an individual (Cheng, 2001; Frazier, 2000). However, simply
because a person has a vast array of coping strategies does not ensure effective coping. Research has shown that those who successfully employ coping strategies do so, not by applying them in a random fashion, but rather by tailoring them to the particular stressor (Cheng, 2001; Frazier, 2000).

**Measuring coping.** To gain more conceptual clarity and more focused assessment of coping strategies some researchers have argued that assessing coping as a dispositional process as well as contextually sensitive process is more informative (Carver, Scheier, & Weintraub, 1989). There are several measures available that assess both maladaptive and adaptive coping skills as well as problem-focused and emotion-focused coping strategies (Carver, 1997; Carver, et al., 1989; Endler & Parker, 1990; Lazarus & Folkman, 1984). Carver, et al., (1989), however, argue that distinguishing between the two kinds of coping is not sufficient. The authors posited that one may employ dispositional coping styles, defined as a preferred set of coping strategies used across circumstances, or contextually sensitive coping styles, where one utilizes coping methods tailored to the specific stressor. Thus, Carver, et al., (1989), created a theoretically driven measure of functional and non-functional coping. The instructions of the scale can be tailored to measure either dispositional or situational strategies of coping.

The COPE consists of 15 sub-scales that measure different coping strategies:

- Active Coping
- Planning
- Suppression of Competing Activities
- Restraint Coping
- Seeking Social Support for Instrumental Reasons
- Seeking Social Support for Emotional Reasons
- Focusing on Venting of Emotions
- Behavioral Disengagement
- Mental Disengagement
- Positive Reinterpretation and Growth
- Denial
- Acceptance
- Humor
- Substance Abuse
- and Turning to Religion (Carver, et al., 1989). Active coping is
defined as the participants’ efforts to actively remove the stressor from their lives, while the use of Planning is related to cognitions employed to aide in coping with the stressor. In addition, Suppression of Competing Activities entails putting other projects aside to better handle the stressor, so as not to distract oneself or occupy one's time on other activities. Moreover, Restraint coping reflects holding back and not acting prematurely in order to effectively deal with the stressor. Using Emotional Support is defined as seeking sympathy, moral support, or compassion, whereas Using Instrumental Support is seen as seeking advice or assistance. Venting of emotions is described as the tendency to focus on the stressor and the distress it causes and express those emotions, while Behavioral Disengagement is said to be the actions taken to avoid the stressor and to reduce the effort employed in dealing with the stressor. Mental disengagement is said to occur when behavioral disengagement fails and entails mentally distracting one's self from the stressor or efforts used to cope with it. Positive reinterpretation is a coping strategy aimed at managing the distress caused by a stressor, rather than the stressor itself. Furthermore, Denial is a form of appraisal that diminishes perceived distress caused by the stressor. Conversely, Acceptance is said to be a functional coping skill in which the person accepts the reality of a stressor and accommodates it into their lives. Humor is coping strategy used to make light of the situation, while Substance Use is a coping skill used in order to disengage oneself from the stressor. Finally, Turning to religion might serve as a source of emotional support, may help positive reappraisal take place, or may be a method of active coping (Carver, et al., 1989).

**Aging and coping.** Differences in coping styles across age groups have been documented in the literature (Aldwin, Sutton, Chiara, & Spiro, 1996; Brändstadter &
Research has found that the elderly tend to use more passive coping strategies, such as acceptance, compared to their younger counterparts (Birkeland & Natvig, 2009; Folkman, et al., 1987), which is unfortunate since proactive measures of coping have been found to improve functional ability and independence in everyday activities among the elderly (Fiksenbaum, Greenglass, & Eaton, 2006). These discrepancies in coping styles remained stable and consistent even when the different age groups were coping with similar stressors (Folkman, et al., 1987). In addition, those in the older age groups were less likely to use escapism, hostility and instrumental methods as coping mechanisms (Aldwin, et al., 1996). However, the results of the coping resources employed were not found to differ between younger participants, middle-aged participants and the elderly participants (Aldwin, et al., 1996; Hamarat, Thompson, Steele, Matheny, & Simons, 2002). Thus, although coping strategies vary by age groups, with the elderly using more passive methods, there exist comparable outcomes across age groups in the effectiveness of coping strategies employed in handling perceived stressors.

Some researchers suggest that differences in coping across age groups are a sign of adaptive flexibility in coping (Brändstadter, et al., 1990; Brändstadter, et al., 1997; Brändstadter, et al., 1993). Active measures of coping to achieve goals in youth are beneficial and preferable, however, more accepting or accommodating methods of coping assist in adapting to the deterioration and losses faced in old age, allowing the elderly to maintain their satisfaction with life even in the face of disability (Brändstadter, et al., 1990; Brändstadter, et al., 1997; Brändstadter, et al., 1993). Agren, (1998), found that
the oldest of old cope with challenges by redirecting their focus away from activities which are difficult to engage in due to functional disabilities and toward those activities that are still feasible. The use of adaptive and flexible coping may explain why some researchers find no differences in life satisfaction across age groups (Hamarat, et al., 2002).

However, the role of aging can add a significant amount of stress to those engaging in caregiving tasks in later life (Roberto, 1995). Physical deterioration, illness, functional disability, and lack of patience can pose a hindrance to caregivers. Indeed, caregivers have been shown to perceive aging as adversely affecting their functionality (Minnes, Woodford, & Passey, 2007). Research has shown that although they are faced with additional stressors, aging caregivers are more reluctant to seek out services and help to share the responsibilities of caregiving tasks than younger caregivers (Hayden & Heller, 1997). Furthermore, problem-focused coping strategies, such as problem-solving and planning are said to reduce burden and actually bring about positive gains in the specific subset of the elderly caregivers (Ribeiro & Paul, 2008). As a result of the previously mentioned patterns of coping employed by the elderly, this can put aging caregivers at a disadvantage in coping with their caregiving roles.

**Caregiver burden and coping with caregiving.** Caregiving is stressful and has an impact on caregivers' mental health. Being a caregiver has been correlated with melancholy mood, social isolation, family conflict, drug and alcohol use, poor health, and impaired immune system functioning (Choo, et al., 2003). Caregivers with high levels of stress are at greater risk for abusive or neglectful behavior towards those receiving their care, perhaps because they lack effective channels through which to express their
emotional distress. Moreover, variables related to patient and caregiver characteristics can interact to affect caregiver burden (Choo, et al., 2003). In the patient, factors such as behavior problems, functional impairment and severity, and chronicity of illness may lead to increases in caregiver burden experienced. Caregiver variables such as age, education level, health status, social support and relationship with the care receiver impact level of burden as well. Research has shown that these influences are more pronounced in elderly female caregivers (Choo, et al., 2003). The influence of ethnicity has also been shown to play an important role in how caregivers handle their roles (Choo, et al., 2003). More expansive social networks may reduce caregiver burden by diffusing the responsibilities of caregiving tasks and reducing feelings of isolation associated with the caregiving role (Guarnaccia & Parra, 1996). However, some research states that ethnic caregivers may not feel they receive enough social support (Haley, et al., 1996). Moreover, assistance in the caregiving role has been correlated with lower levels of burden, however, paid help did not alleviate stress (Guarnaccia, et al., 1996; McCabe, Yeh, Lau, Garland, & Hough, 2003).

Spousal caregivers have been found to experience greater burden than other subsets of caregivers (George & Gwyther, 1986; Neal, Ingerson-Dayton, & Starrels, 1997; Schulz, O’Brien, Bookwala, & Fleissner, 1995). Wives have been shown to experience more distress than husbands when taking care of a disabled spouse. Conversely, husband caregivers have actually reported an improvement in the relationship with their wives after they became caregivers (Fitting, Rabins, Lucas, & Eastham, 1986). Female spousal caregivers felt the greatest reward when they felt companionship from their husbands (Raschick & Ingersoll-Dayton, 2004), which is the
case in some disease contexts such as Parkinson's Disease, but not in the cases of other
disease contests, such as Alzheimer's disease (Hooker, et al., 2000). Adult sons caring
for their parents also experience improvement in the relationship and experience higher
levels of gratification than spousal caregivers (Hinrichsen, Hernandez, & Pollack, 1992).
Hinrichsen, et al. (1992), postulate this may be a consequence of adult children caregivers
perceiving their caregiving responsibilities as an opportunity to return the nurturance
their parents offered them.

Social support has been shown to reduce the impact of burden on caregivers of
patients with psychosis (Grandon, al., 2008; Quayhagen, et al., 1988). In fact, as much as
5% of such burden is attributed to lack of social/affective support (Grandon, al., 2008).
In addition, emotional over involvement may lead to heightened levels of distress and
burden when caregivers feel the care receiver's symptoms are out of one's control. It has
also been found that emotion-focused methods of coping are less effective in alleviating
perceived burden than problem-focused methods (Kristensson Ekwall & Rahm Hallberg,
2006; McClendon, et al., 2004) and may actually increase the levels of stress suffered
(Grandon, et al., 2008). Paradoxically, emotion-focused coping strategies are the most
commonly used among caregivers with lower levels of burden (Chakrabarti, et al., 2002).
However, higher levels of perceived stress prompted the use of problem-focused coping
strategies (Chakrabarti, et al., 2002).

Research shows that successful interventions for caregivers included improving
reframing skills, problem solving skills and seeking social support (McClendon, et al.,
2004; Quayhagen, et al., 1988). Acceptance was found to reduce depression, increase
positive affect and reduce any negative symptoms. The previously mentioned coping
skills were also linked to longer survival time for care recipients (McClendon, et al., 2004). Wishfulness and fantasy were found to be correlated with poor mental health, leading to greater depression and anxiety and. Similarly, escape and distancing were found to increase emotional distress (McClendon, et al., 2004).

Thus, the experience of caregiver burden is best described as the adverse effects caregiving responsibilities have on one's physical and emotional health, social life, and financial status (Zarit, Reever, & Bach-Peterson, 1980). Moreover, the experience of caregiver burden may be influenced by factors, such as demographics and culture (Choo, et al., 2003), relationship with the care recipient (George, et al., 1986; Hinrichsen, et al., 1992; Neal, et al., 1997; Raschick, et al., 2004; Schulz et al., 1995) and coping strategies employed (Grandon, et al., 2008; Kristensson Ekwall, et al., 2006, McClendon, et al., 2004; Quayhagen, et al., 1988). Caregivers who use more emotion-focused coping strategies have been known to fare worse than those who employ problem-focused coping methods (Baker & Robertson, 2008; McClendon, et al., 2004; Quayhagen, et al., 1988; Rose, Strauss, Neundorfer, Smyth, & Stuckey, 1997; Saad, et al., 1995).

Acceptance is an exception to the negative effect emotion-focused coping strategies have on caregivers since use of this specific coping method has led to decreased levels of burden (Baker et al., 2008; Rose, et al., 1997; Saad, et al., 1995).

**Gender and coping.** Men and women have been found to use different coping strategies when faced with stressors (Folkman, et al., 1980; Lutzky, et al., 1994; Matud, 2004; Ptacek, et al., 1994; Tamres, Janicki, & Hegelson, 2002; Yeh, Huang, Chou, & Wan, 2009). Although the differences are minor, women utilize more emotion-focused coping styles (Endler & Parker, 1990), such as avoidance (Lutzky, et al., 1994; Matud,
2004; Ptacek, et al., 1994) and seeking social support (Lutzky, et al., 1994; Ptacek, et al., 1994; Tamres, et al., 2002; Yeh, et al., 2009). Men, on the other hand, have been shown to employ more problem-focused coping styles when facing stressors (Endler & Parker, 1990; Folkman & Lazarus, 1980; Ptacek, et al., 1994). Although both sexes use a combination of emotion-focused and problem-focused coping strategies, men are somewhat more likely to utilize instrumental methods of coping with a stressor (Ptacek, et al., 1994).

A preference for certain coping strategies is also seen across genders in caregivers (Navaie-Waliser, Spriggs, & Feldman, 2002). Female caregivers were more likely than their male counterparts to engage in emotion-focused coping strategies, such as turning to religion (Navaie-Waliser, et al., 2002) and wishful thinking (Rose, et al., 1997). In addition, female caregivers were more likely than male caregivers to reallocate time from other activities, such as employment and leisure, to caregiving tasks (Navaie-Waliser, et al., 2002), which may be detrimental since these actions could lead to social isolation and financial strain. In fact, the tendency to forego employment and social activities may provide some explanation as to why women report more difficulties than men in caregiving (Navaie-Waliser, et al., 2002) as well as higher levels of distress (Adams, Aranda, Kemp, & Takagi, 2002; Navaie-Waliser, et al., 2002).

However, the literature shows that male caregivers who employ more emotion-focused strategies, with the exception of acceptance, report levels of distress similar to female caregivers (Rose, et al., 1997). Similarly, female caregivers who employed more problem-focused coping skills experienced less depression and caregiver burden (Essex, Seltzer, & Krauss, 1999). Rose, et al., (1997), reported that although there are
differences in coping styles across genders, distress may be a function of the coping strategies used by the caregiver, rather than a result of gender.

**Ethnicity and coping.** Like gender, ethnicity may be differentially associated with caregiver burden and caregiver coping. However, there appears to be a lack of consensus in the literature concerning disparities between coping strategies used by non-Hispanic Whites and other ethnicities. Some research suggests that the coping styles employed are similar across ethnicities (Barber, 2002; McCallum, Longmire, & Knight, 2007), while other studies report there indeed exist discrepancies (Knight & McCallum, 1998; Kosberg, Kaufman, Burgio, Leeper, & Sun, 2007; Pinquart & Sorensen, 2005; Sun, Kosberg, Leeper, Kaufman, & Burgio, 2010). Researchers who have found variations in coping styles suggest certain ethnic groups have higher rates of using emotion-focused coping than non-Hispanic Whites (Knight, et al., 1998; Kosberg, et al., 2007; Pinquart, et al., 2005; Sun, et al., 2010; Trail Ross, & Aday, 2006; Wykle, & Segall, 1991). Non-Hispanic Whites have been found to seek assistance from support groups and receive other forms of help from mental health professionals more often than other ethnicities (Montoro-Rodriguez, & Gallagher-Thompson, 2009).

Caregivers who identify as African American have been shown to use different coping styles than Non-Hispanic White caregivers (Knight, et al., 1998; Kosberg, et al., 2007; Pinquart, et al., 2005; Sun, et al., 2010; Trail Ross, et al., 2006; Wykle, et al., 1991). One of the coping methods used more frequently by this group is the use of religion (Haley, et al., 1996; Kosberg, et al., 2007; Pinquart, et al., 2005; Sun, et al., 2010; Wykle, et al., 1991). For this population, religion acts as a window to access spiritual comfort and emotional support from the congregation (Sun, et al., 2010). In
fact, African American caregivers have been found to turn to religiosity more so than both Hispanic and Non-Hispanic White caregivers (Morano, & King, 2005), which is beneficial since higher levels of religiosity has been shown to lower levels of distress. Social support is another coping strategy seen more frequently in African Americans than in non-Hispanic Whites (Pinquart, et al., 2005; Wykle, et al., 1991). Not only was this group more likely to receive more social contact from others, but they also reported more informal support in their caregiving tasks (Pinquart, et al., 2005). For many African American caregivers, the caregiving role was shared with others and responsibilities of the caregiver were dispersed. Avoidance of the stressors and positive reappraisal of the caregiving role as a rewarding experience were also strategies common among African American caregivers (McCallum, et al., 2007; Pinquart, et al., 2005; Trail Ross, et al., 2006).

Another ethnic difference in coping is that, Hispanic caregivers tend to engage in active coping less than Non-Hispanic White caregivers (Montoro-Rodriguez, et al., 2009; Pinquart, et al., 2005). Female Hispanics have been shown to engage in avoidance and escape coping strategies in order to handle the stress of the caregiver role (Montoro-Rodriguez, et al., 2009; Sander, et al., 2007). However, Hispanics as a group report acceptance of the caregiver role and its responsibilities more frequently than their Non-Hispanic White counterparts (Sander, et al., 2007). Informal support from loved ones in which caregiving tasks are shared (Cox, & Monk, 1990; Pinquart, et al., 2005) and religious coping which provides comfort and another avenue for social contact and support (Navaie-Waliser, et al., 2001) were also used by Hispanics in efforts to alleviate the stress of caregiving.
Interestingly, Asian caregivers also engage in more emotion-focused coping strategies than Non-Hispanic White caregivers (Pinquart, et al., 2005). Research has shown that Asians relied heavily on informal support with the caregiving responsibilities much like African Americans and Hispanics did and were the ethnic group least likely to turn to formal caregiving, such as paid help (Kim & Knight, 2008, Pinquart, et al., 2005). Asian caregivers also engaged in cognitive coping strategies (Kim, et al., 2008) to buffer the impact of caregiving stressors, such as avoidance and distancing themselves from the stressors (Lee & Sung, 1998).

**Variables that affect coping.** Taken together, research has shown that demographic factors such as age, gender, and ethnicity can influence both the primary and secondary appraisal process described in the Transactional Theory of Stress and Coping. The impact these variables have on the primary appraisal process is demonstrated by group differences in evaluations of caregiving. Women tend to report less rewards from caregiving responsibilities than men (Fitting, et al., 1986; Hinrichsen, et al., 1992). Because of their own functional decline, the elderly tend to view caregiving as more burdensome than younger caregivers (Minnes, et al., 2007). In addition, African Americans, Hispanics, and Asians view their caregiving roles as a normative experience which they do not have to endure alone (Guarnaccia & Parra, 1996).

The secondary appraisal process is also influenced by the demographic variables mentioned previously. Women have been shown to employ more emotion-focused methods of coping, such as using religion or wishful thinking (Navaie-Waliser, et al., 2002; Rose, et al., 1997) than male caregivers. Furthermore, aging caregivers use more passive methods of coping than their younger counterparts (Hayden & Heller, 1997).
Similarly, certain ethnicities tend to employ emotion-focused coping strategies, such as turning to religion and avoidance to handle the effects of caregiving, more so than Non-Hispanic Whites (Knight, et al., 1998; Kosberg, et al., 2007; Pinquart, et al., 2005; Sun, et al., 2010; Trail Ross, et al., 2006; Wykle, et al., 1991). The above mentioned influences of demographic variables on coping strategies used by caregivers are a focus of this study. The impact of gender and ethnicity on coping styles employed by elderly caregivers will be examined.

**Life satisfaction**

Life satisfaction is important within the context of caregiving because it is affected by the particular burden of caregiving as well as the efficacy of coping. Life satisfaction can be conceptualized as the happiness and contentment one feels in his or her life as an overall synthesis of the many individual domains that compromise one's life (Lewinsohn, et al., 1991). However, it is not merely an emotional state, but an evaluative process in which life is assessed in the context of subjective criteria, such as what constitutes success (Caspi & Elder, 1986; Diener, Emmons, Larsen, & Griffin, 1985; Lewinsohn, et al., 1991; Mallard, Lance, & Michalos, 1997; Meadow, et al., 1992; Michalos, 1986). Life satisfaction as an evaluative process is a dynamic and multifaceted assessment of the overall conditions of one’s life (Caspi et al., 1986; Diener, 1984; Mallard, et al., 1997; Michalos, 1986).

Findings show that elevated life satisfaction has been linked to general well being and positive effects on mood and emotions (Pasupuleti, Allen, Lambert, & Cluse-Tolar, 2009). Those with high levels of life satisfaction report higher self-esteem and feeling good about their lives (Lewinsohn, et al., 1991). They tend to be happier and have more
social and cultural interactions (Griffin & McKenna, 1998). There is also a propensity to be more efficient and productive in handling problems that may arise in the workplace as well as demonstrating more creativity and open-mindedness (Pasupuleti, et al., 2009).

**Measuring life satisfaction.** Life satisfaction, because it is the self-reported outcome of a highly subjective evaluative process, can be difficult to measure (Caspi & Elder, 1986; Diener, Emmons, Larsen, & Griffin, 1985; Lewinsohn, et al., 1991; Mallard, Lance, & Michalos, 1997; Meadow, et al., 1992; Michalos, 1986). In the present study, life satisfaction is conceptualized in accordance with Multiple Discrepancies Theory (MDT) which states that net satisfaction is a result of the divergence between one’s current state and one’s ideal state (Michalos, 1986). Specifically, satisfaction is "a linear function of the discrepancy between what one has and 1) what one wants, 2) what others have, 3) the best one has had in the past, 4) what one expected to have three years ago, 5) what one expects to have in five years, 6) what one deserves, and 7) what one needs" (Mallard, et al., 1997, p. 260; Michalos, 1986). Given that certain comparisons are made between one's current state and one's ideal state, the congruence of these determine the levels of satisfaction one experiences (Mallard, Lance, & Michalos, 1997; Meadow, et al., 1992; Michalos, 1986). That is, if one feels that they currently have what they desire, that it is as good as what others have and what they have had in the past, and that they want to maintain it in the future, they will have increased life satisfaction. Since caregiving is a situation that creates stress, it is likely that life satisfaction may be compromised because it may exacerbate discrepancies in the previously mentioned domains.
Furthermore, life satisfaction is influenced by Top Down (TD), Bottom Up (BU) or Bidirectional processing (BD) (Diener, 1984; Deiner, Suh, Lucas, & Smith, 1999; Mallard, et al., 1997). Top Down processing states one has a propensity to experience things positively or negatively and this influences interactions one engages in (Diener, 1984; Mallard, et al., 1997). Proponents of the TD perspective argue personality traits, such as high self-esteem (Campbell, et al., 1976), and sociability (Diener, 1984), have a positive impact on assessment and perception of life and one's satisfaction with it. The TD framework proposes overall life satisfaction influences satisfaction in individual life facets or sub-domains (Diener, 1984; Mallard, et al., 1997). In contrast, BU processing suggests external events and demographics, such as age, marriage, standard of living, work, school and leisure activities, have a direct effect on life satisfaction (Diener, 1984; Deiner, et al., 1999; Mallard, et al., 1997). The BU processing model conceptualizes overall satisfaction to be a result of the combination of satisfaction experienced in individual domains of one's life. A BD view of overall life satisfaction posits that the influence of both BU and TD processes are present in one's evaluation of life satisfaction (Gerhart, 1987; Mallard, et al., 1997). In this view, both satisfaction in individual aspects of life and the personality traits one expresses interact to influence the assessment of one's happiness.

Bottom Up processing was chosen in the present study to conceptualize life satisfaction because it considers demographics and the assessment of and satisfaction with life facets, such as participants age, gender, ethnicity, level of caregiver burden, and appraisal of the caregiving role, and how all these sub-domains come together to form an experience of overall life satisfaction (Diener, 1984; Diener, et al., 1999; Mallard, et al.,
BU processing fits well with another model used here to conceptualize life satisfaction, MDT, which states that perceptions of discrepancies between one's present state and one's ideal state in life domains, such as the one's suggested by the BU processing model, and with life as a whole influence the experience of overall satisfaction (Mallard, et al., 1997; Michalos, 1986). Therefore, in the present study life satisfaction will be determined by the participant's assessment of the degree of closeness between their present state and ideal state, their evaluation of condition of their lives, as well as their report of subjective satisfaction. The degree to which the participants feel they have accomplished important goals in their lives and the desire to change aspects of their lives will also be used to assess their degree of life satisfaction.

**Caregivers and life satisfaction.** There is a large body of literature which shows that life satisfaction is affected negatively by the caregiver role (Arango-Lasprilla, et al., 2010; Borg & Hallberg, 2006; Haley, et al., 1996; Haley, et al., 2003). Lawton, Moss, Kleban, Glicksman, and Rovine (1991), proposed a two-factor model which argues that caregiving can be both a source of satisfaction as well as exhaustion. The authors argue that the commitment to caring for someone can be a positive and fulfilling experience, but may also be a burden at times because of role strain the depletion of caregiver’s resources (Borg, et al., 2006; Lawton, et al., 1991; Morano, 2003). Satisfaction with caregiving responsibilities has been correlated with increased positive affect but has not been shown to efficiently alleviate burden or negative affect (Lawton, et al., 1991; Wilson-Genderson, Pruchno, & Cartwright, 2009). In addition, caregiver burden has been correlated with negative affect and has been known to reduce positive affect (Lawton, et al., 1991; Wilson-Genderson, et al., 2009).
Haley, et al., (2003) present the Stress Process Model which specifies variables which affect caregiver outcomes. These variables include primary stressors, those directly related to caregiving, and secondary stressors, such as poor health (Haley, et al., 2003). Protective factors are also described in The Stress Process Model. These include positive appraisals of the caregiving situation, adaptive coping responses, and positive social interactions (Haley, et al., 2003). These factors work together to impact the levels of depression and life satisfaction felt by caregivers. Those with higher levels of negative appraisal of caregiver duties and ineffective coping tend to experience increased depression and lowered life satisfaction (Haley, et al., 2003).

According to the Stress-Process Model, one factor that has a buffering effect on life satisfaction is the relationship the caregiver shares with the care recipient (Lawrence, et al., 1998). The quality of the relationship can mediate the effect problem behaviors have on the caregiver's levels of depression and life satisfaction (Lawrence, et al., 1998). When problem behaviors occur, the relationship between the caregiver and the care-recipient suffers, which may lead the caregiver to experience elevated levels of depression and a decreased experience of life satisfaction (Lawrence, et al., 1998). The relationship may also serve as a moderator in this interaction. When the quality of the relationship was high, caregivers experienced higher levels of distress when the care receiver deteriorated and functional ability declined (Lawrence, et al., 1998).

Haley, et al., (2003) applied the Stress-Process Model in a study that examined risk factors and protective factors in caregivers and how these factors predicted depression and life satisfaction. The authors found that objective caregiver strains, such as duration of caregiving, or severity of patient symptoms, were only moderate predictors
of the caregiver’s levels of depression or life satisfaction. Other factors, however, were found to be better predictors of caregiver’s state of being, for example negative social interactions were found to be correlated with higher ratings of depression (Haley, et al., 2003). Conversely, greater numbers of social interactions were caregivers felt content with the social support they received were correlated with a better state of being (Borg, et al., 2006; Haley, et al., 2003; Morano, 2003; Waldron-Perrine, et al., 2009) and good caregiver health was associated with greater life satisfaction (Haley, et al., 2003). Furthermore, the subjective appraisal a caregiver had of their required tasks, along with the benefits they felt were received from caregiving were also more closely associated with life satisfaction and depression than objective stressors (Haley, et al., 2003; Morano, 2003). Those who evaluated their responsibilities as less stressful and found caregiving to be rewarding and fulfilling reported higher life satisfaction and lower depression (Haley, et al., 2003).

**Aging and life satisfaction.** Beyond the effects of caregiver burden, the aging process may create an added burden to caregivers because of possible deterioration and reduction in functionality, thus impacting life satisfaction. Aging caregivers are at an increased risk for reduction in life satisfaction and increase in depression compared to noncaregivers (Haley, LaMonde, Han, Narramore, & Schonwetter, 2001). The manner in which their caregiving tasks are appraised is a factor affecting the experience of caregivers, such that positive appraisals of the caregiving role and its rewards leads to greater life satisfaction and lower levels of depression (Haley, et al., 2003). Greater number of positive social activities and expansive social networks were also linked to greater levels of life satisfaction in aging caregivers (Haley, et al., 2003). Having a
spouse to share financial burden and household responsibilities was also found to increase the life satisfaction experienced and alleviate caregiver burden among the elderly (Landry-Meyer, Gerard, & Guzell, 2005).

Loneliness plagues many caregivers when social relationships are hindered (Borg, et al., 2006). As previously mentioned, social support has positive effects on life satisfaction and is correlated with higher levels of contentment (Borg, et al., 2006; Grandon, et al., 2008; Hooker, et al., 1998; Morano, 2003; Waldron-Perrine, et al. 2009). In fact, Borg, et al., (2006) reported that one third of the caregivers they surveyed wished they could spend more time engaging in social interactions. A smaller group reported having no one to speak to in a time of need, which may bring about issues of distress, such as isolation (Borg, et al., 2006).

It has also been reported that caregivers receive support only infrequently (Borg, et al., 2006). Their most sought after form of support is economic assistance and speaking to someone who is also in a caregiving position about their problems (Borg, et al., 2006). Caregivers who are gainfully employed received the latter from their coworkers (Borg, et al., 2006), which may explain why people in this category report higher life satisfaction. In addition, there is very little agreement between what a caregiver would like to receive as a form of support and what they actually obtain (Borg, et al., 2006).

**Gender, caregiver burden, and life satisfaction.** Independent of personality characteristics and subjective appraisals, men and women’s life satisfaction has been reported to be affected by different objective factors (Waldron-Perrine, et al. 2009). In a study conducted by Waldron-Perrine, et al., (2009), the authors concluded that caregivers
who receive inadequate social support are more likely to be negatively affected by the care recipient's duration of illness and severity of symptoms, particularly among women (Waldron-Perrine, et al. 2009). Caregiver income and uncertainty about the care recipient's prognosis, however, had an equal effect among both genders (Waldron-Perrine, et al. 2009). Interestingly, the authors found that the relationship between the duration of the illness and life satisfaction was positively correlated for men and negatively correlated for women, which suggests that men begin to adapt to caregiving tasks over time, while women suffer from negative cumulative effects (Waldron-Perrine, et al. 2009).

Research shows that women are more negatively affected by caregiving responsibilities than men (Collins, et al., 1997; Hooker, et al., 2000; Kristensson Ekwall, et al., 2006; Waldron-Perrine, et al., 2009). Female caregivers have been shown to undergo more distress and strain when caring for others (Hooker, et al., 2000; Waldron-Perrine, et al., 2009) while men giving care to a loved one report being satisfied with their lives (Kristensson Ekwall, et al., 2006). Caregiving tasks have not been found to cause this discrepancy between men and women's burden since both sexes engage in the same amount of workload associated with caregiving (Collins, et al., 1997). Women were reported to feel unable to cope with their situation more often and felt obligated to continue providing care even though they would prefer to discontinue caregiving (Collins, et al., 1997). Nevertheless, there still remains the belief that women are better suited to taking over the caregiver role and that women are better equipped to handle it (Collins, et al., 1997).
Hooker, et al., (2000), found a clear difference in male and female caregivers. Women caring for their husbands suffering from dementia were found to have significantly worse mental health than husbands caring for their wives in the same group (Hooker, et al., 2000). In contrast, no differences in mental health were found between husband and wife caregivers in the group comprised of nondementia patients, even though the groups did not differ in their use of social support (Hooker, et al., 2000). The authors suggested that women fare worse than men in caregiving situations if cognitive deterioration is present.

Ethnicity and life satisfaction. As stated previously, the assessment and appraisal of caregiving and its responsibilities has an impact on the experience of life satisfaction and depression. For some ethnicities, this appraisal may be influenced by factors such as familism and filial piety (Kim, Knight, & Flynn Longmire, 2007; McCallum, et al., 2007; Pinquart, et al., 2005). Familism is conceptualized as the "subordination of individual interests to those of the family" (Rogers & Sebald, 1962, p. 26), and filial piety is the submission to one's elders which entails both emotional support as well as physical and financial assistance (Yeh & Bedford, 2003). Indeed appraisal of caregiver burden may be impacted by familism (Kim, et al., 2007; McCallum, et al., 2007). Strong familism beliefs accompanied by positive social support may improve health outcomes in caregivers (McCallum, et al., 2007). Conversely, the idea of being obligated to care for someone because of filial piety may lead to powerlessness and decreased life satisfaction (Anngela-Cole & Hilton, 2009). Ethnic groups in general reported the caregiving role to be more rewarding than Non-Hispanic White caregivers.
(Pinquart, et al., 2005). One explanation may be the influences of familism and filial piety.

Familism and positive filial beliefs are endorsed by African American caregivers, more so than Non-Hispanic White caregivers (McCallum, et al., 2007; Pinquart, et al., 2005). The endorsement of Familism and Filial Piety may have an impact on African American caregivers since they experience less depression and burden (Roth, Haley, Owen, Clay, & Goode, 2001), and greater life satisfaction than Non-Hispanic White caregivers (Haley, et al., 1995; Lawton, Rajagopal, Brody, & Kleban, 1992; Roth, et al., 2001). African Americans also reported greater reward and satisfaction obtained from the caregiver role (Lawton, et al., 1992) and experience less stress and feel more efficacious than Non-Hispanic White caregivers (Haley, Wadley, West, & Vetzel, 1994).

Familism is also seen among Hispanic caregivers (Pinquart, et al., 2005). Similarly to African Americans, Hispanic caregivers reported greater rewards and uplifts from caregiving (Pinquart, et al., 2005), and felt more competent in managing the caregiving role (Montoro-Rodriguez, et al., 2009) than Non-Hispanic White caregivers. However, they were found to experience more elevated levels of depression and distress than Non-Hispanic White caregivers (Pinquart, et al., 2005; Valle, Yamada, & Barrio, 2004). In fact, Hispanics fared worse than both Non-Hispanic White (Adams, et al., 2002; Cox, et al., 1990; Valle, et al., 2004) and African American caregivers (Adams, et al., 2002; Cox, et al., 1990; Morano, et al., 2005) evidenced by significantly higher levels of depression and psychological distress experienced (Adams, et al., 2002; Cox, et al., 1990). Although both African Americans and Hispanics reported the use of religion and spirituality as a powerful coping mechanism to handle the demands of caregiving, this
marked discrepancy may be because Hispanic caregivers felt they needed more social support (Adams, et al., 2002). Furthermore, the existence of familism beliefs without sufficient social support has been correlated with greater distress and caregiver burden (McCallum, et al., 2007).

Other ethnic groups may also experience caregiving differently from White Non-Hispanics due to their cultural beliefs. For example, Asian caregivers who engage in caregiving because of sense of obligation or filial piety were shown to experience less life satisfaction compared to those who did so because of a conscious choice to engage in caregiving (Anngela-Cole, et al., 2009). Specifically, Japanese caregivers were found more likely to care for their elders to avoid fear of social isolation or shame (Asai, 2002), which may affect their life satisfaction compared to other Asians in the caregiving role. Overall, Asian caregivers were found to report higher levels of depression and burden than their Non-Hispanic White counterparts (Pinquart, et al., 2005), which is problematic because Asian caregivers have at times lacked in services and outlets which reduce caregiver burden (Ho, Weitzman, Xingjia, & Levkoff, 2000).

Thus, on the basis of the research presented, it appears as though many factors interact to culminate in the experience of caregivers' life satisfaction. Indeed, factors such as familism and filial piety (Kim, et al., 2007; McCallum, et al., 2007; Pinquart, et al., 2005) as well as the age (Haley, et al., 2001) and gender (Collins, et al., 1997; Hooker, et al., 2000; Kristensson Ekwall, et al., 2006; Waldron-Perrine, et al., 2009) of the caregiver impact the perception of the caregiver role. Caregivers' view of the caregiving role has been shown to influence the levels of caregiver burden (Anngela-Cole & Hilton, 2009; Haley, et al., 2003; Kim, et al., 2007), which in turn affects the
experience of life satisfaction (Asai, 200; Haley, et al., 2003). The quality of the relationship between the caregiver and care receiver also affects the experience of caregiver burden (Lawrence, et al., 1998) and resulting life satisfaction. Furthermore, it is suggested in the literature that adaptive coping skills can reduce levels of distress and caregiver burden resulting higher levels life satisfaction (Haley, et al., 2003).

The purpose of the present study is to extend our current understanding of the effects of caregiver burden on life satisfaction by examining whether or not there are ethnic differences in coping strategies used to cope with caregiving. There is a lot of compelling evidence of effective strategies for coping with caregiving for a spouse or loved-one with a chronic condition, such that caregiver burden is lessened and life satisfaction may be maintained. There is evidence that both gender and ethnicity may impact caregiver burden, ways of coping with the stress of caregiving, and ultimately life satisfaction. It is also clear from the literature that there are ethnic and cultural differences in the methods and efficacy of coping with stress, and in particular coping with the stress associated with caregiving. Thus, older adults of different ethnic backgrounds may be at particular disadvantage in coping with caregiving because of the cultural beliefs that shape their family dynamics. Therefore, the current study will examine male and female older adult caregivers from different ethnic backgrounds in order to determine if there are differences in caregiver burden.

Moreover, on the basis of the literature it seems reasonable to surmise that effective coping can mediate any direct negative effects of the demographic factors (age, gender, ethnicity) on life satisfaction. That is, the established relationships that show lower life satisfaction in female caregivers of different ethnicities may be offset by
adaptive and effective coping. However, it is also likely that less adaptive coping may increase the negative effects of the demographic characteristics on life satisfaction. That is, the established relationships showing that female ethnic caregivers experience lower life satisfaction may be found to be the result of maladaptive coping strategies.

Understanding the cultural/ethnic differences in coping with caregiving in later life will help us to better understand how the stress of caregiving is assessed, what resources are particularly useful in particular ethnic groups, and ultimately, may highlight points of intervention to help caregivers of different ethnic groups cope better.

The current study will test several specific hypotheses in order to determine the linkages among age, gender, ethnicity (i.e. ethnicity, ethnic beliefs), caregiver burden, coping with caregiving, and life satisfaction. Specifically, it is hypothesized that:

1) There will be age, gender, and ethnic differences in the caregiver burden experienced. Based on extant literature it is expected that older, female, Hispanic caregivers will report greater caregiver burden.

2) There will be age, gender, and ethnic differences in the coping strategies used to manage caregiver burden. Consistent with the coping literature, it is expected that caregivers will use a varied array of different coping strategies, some of which have been shown to be effective and useful at decreasing stress and burden (e.g., acceptance, reframing, problem solving, seeking social support) and some that have been shown to be less effective at managing the distress associated with caregiving (e.g., escape, avoidance) and may even be associated with increased levels of distress and lower levels of life satisfaction. Thus, it is expected that there will be a wide range of coping strategies used across
caregivers and this study will determine how these patterns of coping vary across age, gender and ethnicity. Specifically it is expected, based on the extant literature that older, female, Hispanic caregivers will use more dysfunctional coping strategies, such as avoidance, and less active styles of coping aimed at changing, reducing, or negating the stress of caregiving.

3) There will be age, gender, and ethnic differences in mean levels of life satisfaction among the caregivers. Older, female, Hispanic caregivers will report less life satisfaction that their younger, male, non-Hispanic counterparts.

4) Coping strategies are hypothesized to mediate the relationship among demographic factors and life satisfaction. That is, part of the variability in life satisfaction will be explained by demographic factors such as age, gender, and ethnicity, however, coping strategies that are less effective will exacerbate the effects of caregiver burden on life satisfaction.

5) Lastly, on the basis of extant literature, it is hypothesized that there will be a significant relationship among burden and coping and life satisfaction, Specifically, higher rates of burden will be correlated with less effective coping and lower life satisfaction.
CHAPTER III
Methodology

Participants

Participants in this study were volunteers over the age of 60 years old who were giving care to a loved-one or spouse with a chronic medical condition. A total of 103 older adults were recruited, with 54 older Hispanic adults and 49 older Non-Hispanic White adults. There were 17 male caregivers and 86 female caregivers. The mean age of the sample was 67.42, with caregivers between the ages of 60 to 85. Because of the nature of the caregiving experience, older adults had to have been in the caregiving role for a minimum of one year. Additionally, all participants were screened to determine whether they suffered from memory problems or dementia using the Mini-mental status exam (Folstein, Folstein, & McHugh, 1975), and if so, they were excluded from participation. All older adult caregivers provided demographic information, self-reported health information, ethnocultural beliefs, completed the COPE inventory, the Burden Interview, The Satisfaction with Life scale, the Center for Epidemiological Studies-Depression Scale, and the State Trait Anxiety Scale.

Measures

Demographic data. A standard demographic questionnaire was given that gathers information on age, sex, education, occupation, socioeconomic status (as measured by the Hollingshead Index; Hollingshead, 1975). Additionally, ethnicity and language competencies were queried. Finally, as part of the demographic questionnaire a self-reported health index was given to assess the current health status of the caregiver.
and information will be gathered about the diagnosis, illness trajectory, severity and level
of functioning of the care recipient.

**Ethnocultural beliefs.** Literature suggests that there is an interaction among
ethnicity and cultural beliefs (Rozario, et al., 2008; Kim, et al., 2007). Therefore, to
accurately assess the distinction among ethnicity and cultural two measures of cultural
beliefs was administered. Familism was assessed using the Familism Scale (Losada,
Knight, Marquez-Gonzalez, Montorio, Etxeberria, & Peñacoba, 2008; Sabogal, Marin,
Otero-Sabogal, VanOss Marin, & Perez-Stable, 1987) adapted from the version created
by Bardis (1959). The nine item questionnaire assesses three components of familism:
familial obligations, perceived support from the family, and family as referents (Losada,
et al., 2008; Sabogal, et al., 1987). The participants rated the items based on a 5 point
likert scale ranging from (0) strongly disagree to (4) strongly agree. Higher scores
indicate higher levels of familism.

Filial piety was assessed using the Filial Piety Scale (Ho, 1994). The 22 item
questionnaire assesses aspects of filial piety, such as obedience, respect, ancestral
worship and providing for one's parent (Ho, 1994). There are 11 negative items on the
scale to measure diverging beliefs. Participants rated the items on a 6 point likert scale
ranging from (1) strongly disagree to (6) strongly agree. Higher ratings on the scale
indicate greater filial piety. Negative items were scored inversely (Ho, 1994).

**The COPE.** Coping strategies were assessed using the COPE scale (Carver, et
al., 1989). The questionnaire includes 60 questions, four assessing each of the following
coping behaviors: (1) active coping, (2) suppression of competing activities, (3) restraint
coping, (4) instrumental social support, (5) emotional social support, (6) positive
reinterpretation, (7) acceptance, (8) venting emotions, (9) turning to religion, (10) denial, (11) behavioral disengagement, (12) mental disengagement, and (13) planning. The instructions asked participants to answer each question about what they usually do to manage the stress associated with caregiving. Answers will range on a four point likert scale from (1) not very often to very often (4) (Carver, et al., 1989).

**The Burden Interview.** Levels of burden were measured using The Burden Interview (Zarit, et al., 1980). The 22 item questionnaire assesses subjective caregiver burden by using qualitative information (i.e., the affective response of the caregiver) to assess specific areas which are usually affected in a caregivers life (Zarit, et al., 1980). The instructions asked the participants to circle the response which best describes their feelings. Caregivers endorsed each item along a five point likert scale ranging from (0) never to (4) nearly always present (Zarit, et al., 1980). Higher ratings on the items indicated more elevated levels of burden.

**The Satisfaction with Life Scale.** Levels of life satisfaction were measured using The Satisfaction with Life Scale (Diener, et al., 1985). The narrowband five item questionnaire conceptualizes life satisfaction as a subjective appraisal of how close one’s living conditions are to their ideal situation (Diener, et al., 1985). Questions are worded so that participants evaluate their lives on the basis of their own individual beliefs as to what constitutes the ideal (e.g., "The conditions of my life are excellent"). Caregivers rated their agreement with each item based on a seven point likert scale ranging from (1) strongly disagree to (7) strongly agree. A higher score on the scale indicates elevated levels of life satisfaction.
**Center for Epidemiological Studies-Depression Scale.** Symptoms of Depression were measured using the Center for Epidemiological Studies-Depression Scale (CES-D; Radloff, 1977). The 20 item self-report scale measures depressive symptomology in the general population (Radloff, 1977). The instructions asked the participant to state how often they have felt symptoms of depression over the past week. Participants rate items on a four point likert scale ranging from (0) rarely or none of the time to (3) most or all of the time (Radloff, 1977). Higher scores indicate increased depressive symptoms. Four items on the scale are inversely scored.

**State Trait Anxiety Inventory.** The participants' levels of stress were assessed using the State Trait Anxiety Inventory (STAI; Speilberger, 1983). The measure consists of two separate 20-item self-report scales which evaluate both state anxiety (i.e., what they are experiencing at that very moment) and trait anxiety (i.e., how they generally feel; Speilberger, 1983). The instructions on the state anxiety scale asked the participant to indicate how they feel at the moment. Participants' rate the items on a four point likert scale ranging from (1) not at all to (4) very much so (Speilberger, 1983). Higher scores indicate elevated levels of stress. Ten items are inversely scored.

The trait anxiety scale is structured similarly to the state anxiety scale. The instructions for the trait anxiety scale asked the participant to indicate how they generally feel (Speilberger, 1983). Participants' rated the items on a four point likert scale ranging from (1) almost never, to (4) almost always (Speilberger, 1983). Higher scores indicate elevated levels of stress. Ten items are inversely scored.
Procedure

Participants were recruited from the community by reaching out to organizations that provide services to the elderly. Caregiver support groups and disease support groups were identified for recruitment, as well as day care centers for the elderly and senior recreational centers. At all locations information about the study was provided in flyers, leaflets, and announcements. Sign-up sheets with the principal investigator’s contact information were distributed. Interested participants who contacted the principal investigator were screened and if they met the inclusion criteria were scheduled an interview at the time and location of convenience for the participant. All interviews were be face-to-face format and conducted by the principal investigator and research assistants. Interviews were conducted in English or Spanish depending on the preference of the participant. Prior to beginning the interview proper, each participant was informed of the nature of the study and issues of confidentiality, all questions were answered and then, if the participant agreed, an Informed Consent was presented and signed. Then the interview began with the demographic data followed by the experimental measures.

Data Analytic Plan

Data analysis was done using SPSS. A power analysis was conducted using the statistical computer program G Power 3.1 and a sample size of 50 elderly participants per ethnic category was found to be sufficient. A total of 103 participants were assessed.

The data analytic approach to test each hypothesis is as follows:

1) In order to determine whether or not there are age, gender, and ethnic differences (i.e., ethnicity, ethnocultural beliefs) in the caregiver burden and mental health, t-tests and ANOVAs were performed to test this hypothesis, with
demographic variables as the independent variables and levels of burden and the levels of depression and anxiety as the dependent variable. In addition, Pearson Product Moment Correlational Analyses were conducted to further examine the hypothesis.

2) In order to determine if there are age, gender, and ethnic differences in the coping strategies correlational analyses were performed to determine if there are associations among the demographic factors and the dimensions of the COPE scale. For those variables that were statistically correlated, multiple regression analyses were performed in order to determine whether age, gender or ethnicity is the strongest predictor of coping styles employed. In these analyses, each demographic factor was treated as an independent variable and the outcome, coping, was the dependent variable. Thus, the relative weight of each demographic factor as a predictor of each coping strategy was assessed. A separate linear regression was performed for each major dimension of coping.

3) In order to determine whether there are demographic differences in life satisfaction a similar approach was taken. First, correlations were examined in order to determine which variables are related. Then, for those variables that were correlated, a hierarchical linear regression analyses were performed in order to determine whether age, gender or ethnicity is the strongest predictor of life satisfaction. In these analyses, each demographic factor was treated as an independent variable and the outcome, life satisfaction, was the dependent variable. Thus, the relative weight of each demographic factor as a predictor of each life satisfaction was assessed.
4) Next, coping strategies were hypothesized to mediate the relationship among demographic factors and life satisfaction. That is, part of the variability in life satisfaction was expected to be explained by demographic factors such as age, gender, and ethnicity, however, coping strategies that are less effective were expected to exacerbate the effects of caregiver burden on life satisfaction. In order to test the mediational role of coping a series of hierarchal linear regressions were used following Holmbeck (2006) and Kraemer, Kiernan, Essex and Kupfer, (2008). Specifically, in order to test for mediation four criteria must be met. First, there must be a significant relationship between the demographic factors and life satisfaction. Second, the relationship among the demographic factors and coping must be significant. Third, there must be a significant relationship between coping style and life satisfaction after controlling for demographic factors. And finally, the impact of demographic factors on life satisfaction should be significantly less after controlling for coping.

5) It is also possible that coping mediates the relationship between burden and life satisfaction. Therefore, in order to test the mediational role of coping a series of hierarchal linear regressions were used following Holmbeck (2006) and Kraemer, et al., (2008). Specifically, in order to test for mediation four criteria must be met. First, there must be a significant relationship between caregiver burden and life satisfaction. Second, the relationship among burden and coping must be significant. Third, there must be a significant relationship between coping style and life satisfaction after controlling for burden. And finally, the
impact of caregiver burden on life satisfaction should be significantly less after controlling for coping.
CHAPTER IV

RESULTS

Descriptive Data

Participants. A total of 103 participants were included in the present study, 17 (16.5%) males and 86 (83.5%) females. The mean age for the sample was 67.42 ($SD = 6.436$), with ages ranging from 60 to 85. There were 49 (47.6%) Non-Hispanic White participants and 54 (52.4%) Hispanic participants in this study.

Ethnocultural beliefs. Participants rated how they perceived their obligation to their family. The mean score for the Familism scale was 22.85 ($SD = 6.15$), with scores ranging from 3 to 36. The mean score for the Filial Piety scale was 62.89 ($SD = 10.87$), with score ranging from 34 to 96. Male ($M = 65.65, SD = 11.54$) caregivers scored higher on filial piety than female caregivers ($M = 62.35, SD = 10.72$). Male caregivers ($M = 23.12, SD = 4.94$) also scored higher on familism than female caregivers ($M = 22.77, SD = 6.39$). Differences were also present among scores for the Filial Piety and Familism scale for Hispanic and Non-Hispanic White caregivers. Hispanic caregivers ($M = 65.31, SD = 9.35$) scored higher on the Filial Piety scale than Non-Hispanic White caregivers ($M = 62.89, SD = 10.87$). Similarly, Hispanic caregivers ($M = 23.35, SD = 5.58$) scored higher on the Familism scale than Non-Hispanic White caregivers ($M = 20.04, SD = 5.55$).

Coping and Outcomes. Caregivers who participated in this study were asked to report the coping methods they used, their levels of depression, anxiety, life satisfaction, and caregiver burden.
Coping and Depression

The relationship between coping and depression was investigated using Pearson Product Moment Correlational Analyses. The results indicate that depression does not have a significant relationship with Positive Reinterpretation and Growth ($r = -.062, p = .536$), Use of Instrumental Social Support ($r = -.004, p = .971$), Religious Coping ($r = -.028, p = .781$), Humor ($r = .121, p = .224$), Restraint ($r = .184, p = .063$), Use of Emotional Support ($r = -.047, p = .640$), Substance Use ($r = .052, p = .605$), Suppression of Competing Activities ($r = .152, p = .126$), or Planning ($r = -.081, p = .413$). Thus, the use of these coping strategies does not relate to depression in these caregivers.

Depression was found to have a positive correlation with Mental Disengagement ($r = .382, p = .001$), Focus on Venting ($r = .346, p = .001$), Denial ($r = .411, p = .001$), and Behavioral disengagement ($r = .398, p = .001$). The results indicate that those caregivers who are more likely to both mentally and behaviorally disengage, more likely to be in denial, more likely to focus on venting negative emotions are also more depressed.

Conversely, depression was found to be negatively correlated with Active Coping ($r = -.227, p = .021$), and Acceptance ($r = -.301, p = .002$). Those caregivers who are more depressed are using less active coping and acceptance, two coping strategies found to be beneficial for reducing burden.

The relationship between coping and anxiety was also investigated using a Pearson Product Moment Correlations. Results show that state anxiety is not significantly correlated with Focus on Venting ($r = .178, p = .072$), Use of Instrumental
Social Support ($r = -0.115, p = 0.248$), Religious Coping ($r = 0.132, p = 0.184$), Humor ($r = -0.117, p = 0.238$), Restraint ($r = 0.104, p = 0.294$), Use of Emotional Support ($r = -0.127, p = 0.202$), Substance Use ($r = 0.077, p = 0.439$), Suppression of Competing Activities ($r = 0.017, p = 0.862$), or Planning ($r = -0.171, p = 0.085$).

Results also show that state anxiety is positively correlated with Mental Disengagement ($r = 0.279, p = 0.004$), Denial ($r = 0.285, p = 0.004$), and Behavioral Disengagement ($r = 0.341, p = 0.001$). Furthermore, results indicate that state anxiety is negatively correlated with Positive Reinterpretation and Growth ($r = -0.304, p = 0.002$), Active Coping ($r = -0.250, p = 0.011$), Acceptance ($r = -0.338, p = 0.0001$). Thus, those caregivers who are higher in state anxiety, anxiety related to the state of caregiving, are more likely to cope with caregiving through mental and behavioral disengagement and denial. These results suggest that anxiety associated with caregiving leads to decreased use of beneficial coping strategies and increased use of dysfunctional coping strategies.

The relationship between coping and trait anxiety was also investigated. Correlational analyses show that trait anxiety is not significantly correlated with Use of Instrumental Social Support ($r = -0.048, p = 0.632$), Active Coping ($r = -0.140, p = 0.159$), Religious Coping ($r = 0.026, p = 0.795$), Humor ($r = -0.049, p = 0.626$), Restraint ($r = 0.192, p = 0.052$), Use of Emotional Support ($r = -0.068, p = 0.493$), Substance Use ($r = 0.045, p = 0.653$), Suppression of Competing Activities ($r = 0.100, p = 0.315$), or Planning ($r = -0.035, p = 0.729$).

However, trait anxiety is positively correlated with Focus on Venting ($r = 0.418, p = 0.0001$), Mental Disengagement ($r = 0.348, p = 0.0001$), Denial ($r = 0.196, p = 0.047$), and Behavioral disengagement ($r = 0.321, p = 0.001$). Moreover, results indicate that trait
anxiety is negatively correlated with Positive Reinterpretation and Growth ($r = - .210, p = .033$), and Acceptance ($r = - .209, p = .034$). Again, caregivers who are more anxious are engaging in less adaptive coping strategies.

The relationship between coping and burden was also investigated. Results show that burden is not significantly correlated with Positive Reinterpretation and Growth ($r = - .056, p = .574$), Use of Instrumental Social Support ($r = .188, p = .057$), Denial ($r = .138, p = .165$), Religious Coping ($r = .163, p = .099$), Substance Use ($r = -.084, p = .401$), Suppression of Competing Activities ($r = .069, p = .487$), or Planning ($r = -.058, p = .561$).

Caregiver burden is significantly, positively correlated with Mental Disengagement ($r = .355, p = .0001$), Focus on Venting ($r = .438, p = .0001$), Behavioral disengagement ($r = .313, p = .001$), Restraint ($r = .305, p = .002$), and Use of Emotional Support ($r = .201, p = .041$). Moreover, results indicate that burden is negatively correlated with Active Coping ($r = -.198, p = .045$), Humor ($r = -.209, p = .034$), and Acceptance ($r = -.205, p = .038$). Thus, less adaptive coping strategies are highly correlated with greater caregiver burden while adaptive coping strategies correlate with less burden.

Finally, the relationship between coping and life satisfaction was investigated. Results show that burden is not significantly correlated with Positive Reinterpretation and Growth ($r = .155, p = .117$), Mental Disengagement ($r = -.138, p = .164$), Focus on Venting ($r = -.192, p = .052$), Use of Instrumental Social Support ($r = -.002, p = .983$), Active Coping ($r = .151, p = .128$), Denial ($r = .132, p = .183$), Religious Coping ($r = .033, p = .741$), Behavioral disengagement ($r = -.134, p = .177$), Restraint ($r = -.102, p = .431$).
Use of Emotional Support ($r = -.044, p = .660$), Substance Use ($r = .133, p = .180$), Acceptance ($r = .099, p = .318$), Suppression of Competing Activities ($r = -.073, p = .463$), or Planning ($r = -.058, p = .561$). However, one interesting positive correlation emerged. Results also show that life satisfaction is positively correlated with Humor ($r = .229, p = .020$). Thus, those caregivers that are able to find humor or cope through humor tend to report higher life satisfaction.

A series of correlational analyses were also conducted to examine the linkages among mental health outcomes: depression, state anxiety, trait anxiety, burden, and life satisfaction. As expected, depression was found to be positively correlated with state anxiety ($r = .509, p = .0001$), trait anxiety ($r = .698, p = .0001$) and burden ($r = .512, p = .0001$). Depression was also found to be negatively correlated with life satisfaction ($r = -.306, p = .002$). State and trait anxiety were found to be positively correlated ($r = .570, p = .0001$). State anxiety was positively correlated with burden ($r = .411, p = .0001$) and negatively correlated with life satisfaction ($r = -.421, p = .0001$). Similarly, trait anxiety was found to be positively correlated with burden ($r = .528, p = .0001$) and negative correlated with life satisfaction($r = -.603, p = .0001$). Lastly, there was a negative correlation between life satisfaction and caregiver burden ($r = -.504, p = .0001$). These results indicate perhaps a profile of negative mental health patterns in caregivers.

Consistent with Carver et al., (1989), the individual coping strategies were combined to create two different types of coping, maladaptive coping and adaptive coping. Further extending the results reported above, correlational analyses showed that maladaptive coping was significantly correlated with increased burden, ($r = .399, p = .0001$), increased state, ($r = .437, p = .0001$) and trait anxiety ($r = .375, p = .0001$), and
increased depression, ($r = 515, p = .001$). Thus, taken together results show that on the level of individual coping strategies and on the level of general coping styles dysfunctional coping leads to negative mental health outcomes for caregivers.

**Demographic Differences in Burden**

Hypothesis one examined demographic differences between burden and mental state among participants in the study. It was hypothesized that older caregivers would report greater levels of caregiver burden. Similarly, female and Hispanic caregivers were expected to experience greater caregiver burden than their male and Non-Hispanic White counterparts.

The relationship between age and burden was investigated using Pearson Product Moment Correlations. There was a small, marginally significant correlation between the two variables ($r = -.173, p = .080$). In other words, results suggest a trend that older caregivers are experiencing lower levels of caregiver burden, while younger caregivers experience higher levels of burden. But this result was not significant thus must be interpreted with caution.

An independent samples t-test was conducted to compare the burden levels for males and females. Results showed no significant difference $t(101) = .298, p = .78$, which indicates that while females may report greater burden ($M = 30.63, SD = 14.79$) than males ($M = 29.41, SD = 18.20$) this difference was not significant. Similarly, an independent samples t-test was conducted to compare the burden levels for Hispanic and Non-Hispanic White participants. Results showed no significant difference $t(101) = .424, p = .67$ between the two groups. Although Non-Hispanic White caregivers may report
higher levels of burden ($M = 31.10, SD = 15.48$) than Hispanic caregivers ($M = 29.81, SD = 15.26$), the results were not found to be significant.

An Analysis of Variance was also conducted to investigate the difference between levels of caregiver burden, depression, and anxiety among male and female caregivers. There was no statistical significance between gender and depression $F(1, 101) = .003, p = .956$, state anxiety $F(1, 101) = .010, p = .922$, trait anxiety $F(1, 101) = .623, p = .432$ or burden $F(1, 101) = .089, p = .766$.

Lastly, several Analyses of Variances were conducted to investigate the difference between levels of caregiver burden, depression, and anxiety among Hispanic and Non-Hispanic White caregivers. There was no statistical significance between ethnicity and depression $F(1, 101) = .440, p = .508$, state anxiety $F(1, 101) = .488, p = .486$, trait anxiety $F(1, 101) = 2.155, p = .145$ or burden $F(1, 101) = .180, p = .672$. In sum, contrary to initial hypotheses there were no differences in the demographic factors (age, gender, and ethnicity) and mental health outcomes.

**Demographics Differences in Coping**

The second hypothesis for this study examined the relationship between demographics and coping methods used. It was expected that older caregivers would use more dysfunctional methods of coping. Female and Hispanic caregivers were also expected to use less effective coping methods to manage their caregiving role.

The relationship between age and methods of coping used was investigated using the Pearson Product Moment Correlations. There was no significant correlation between age and Positive Reinterpretation and Growth ($r = -.059, p = .553$), Mental disengagement ($r = -.071, p = .474$), Focus on and venting of emotions ($r = -.056, p = .
Use of instrumental social support \((r = -0.100, p = .314)\), Active coping \((r = -0.044, p = .655)\), Denial \((r = 0.020, p = .839)\), Religious coping \((r = 0.136, p = .171)\), Humor \((r = -0.024, p = .813)\), Behavioral disengagement \((r = -0.055, p = .579)\), Restraint \((r = 0.058, p = .563)\), Use of emotional social support \((r = -0.116, p = .245)\), Substance use \((r = -0.163, p = .099)\), Acceptance \((r = 0.097, p = .331)\), Suppression of competing activities \((r = 0.142, p = .152)\), or Planning \((r = -0.004, p = .967)\). The results suggest that age does not have a significant relationship with these forms of coping.

An independent samples t-test was conducted to compare the coping methods between males and females. Results showed no significant difference between gender and Positive Reinterpretation and Growth \(t(101) = -1.286, p = .201\), with males \(M = 11.76, SD = 3.07\) reporting it less frequently than females \(M = 12.68, SD = 2.62\), Mental disengagement \(t(101) = -0.155, p = .877\), with males \(M = 8.12, SD = 2.71\) reporting it less frequently than females \(M = 8.22, SD = 2.46\), Focus on and venting of emotions \(t(101) = -0.991, p = .324\), with males \(M = 8.05, SD = 3.19\) reporting it less frequently than females \(M = 8.81, SD = 2.80\), Active coping \(t(101) = -1.327, p = .188\), with males \(M = 11.52, SD = 2.27\) reporting it less frequently than females \(M = 12.27, SD = 2.10\), Denial \(t(101) = -1.554, p = .123\), with males \(M = 4.76, SD = 1.56\) reporting it less frequently than females \(M = 5.79, SD = 2.62\), Humor \(t(101) = 0.197, p = .844\), with males \(M = 8.41, SD = 4.39\) reporting it more frequently than females \(M = 8.19, SD = 4.29\), Behavioral disengagement \(t(101) = -0.086, p = .931\) with males \(M = 5.88, SD = 2.09\) reporting it less frequently than females \(M = 5.93, SD = 2.85\), Restraint \(t(101) = -0.939, p = .350\), with males \(M = 9.12, SD = 2.39\) reporting it less frequently than females \(M = 9.79, SD = 2.75\), Acceptance \(t(101) = 0.567, p = .572\) with males \(M = 9.79, SD = 2.75\).
= 12.52, SD = 3.87) reporting it less frequently than females (M = 13.00, SD = 2.97), Suppression of competing activities t(101) = .291, p = .772, with males (M = 10.35, SD = 2.62) reporting it more frequently than females (M = 10.16, SD = 2.43), or Planning t(101) = .154, p = .878 with males (M = 12.18, SD = 3.30) reporting it more frequently than females (M = 12.05, SD = 2.80). Contrary to initial expectations, there are no gender differences in the use of these different coping strategies in this sample.

There was a significant difference, however, between men and women regarding Use of instrumental social support t(101) = -2.58, p = .011, with males (M = 7.94, SD = 3.54) reporting it less frequently than females (M = 10.37, SD = 3.53). Males (M = 8.59, SD = 4.62) were also significantly less likely to use Religious coping t(101) = -3.291, p = .001, than females (M = 12.26, SD = 4.11). Females (M = 10.02, SD = 3.38), also endorse a greater Use of emotional social support t(101) = -3.235, p = .002, than males (M = 7.18, SD = 2.96). There was a marginally significant difference between males and females regarding Substance use t(101) = 4.109, p = .071, with males (M = 5.82, SD = 3.57) reporting it more frequently than females (M = 4.14, SD = .65). Thus, gender differences did emerge in the use of instrumental social support, emotional support, and religious coping with females generally reporting more of these strategies than males.

An independent samples t-test was conducted to compare coping methods between Hispanic and Non-Hispanic White caregivers. Results showed no significant difference among the groups on Mental disengagement t(101) = -.080, p = .937 with, Focus on and venting of emotions t(101) = -1.327, p = .188, Use of instrumental social support t(101) = -1.154, p = .251, Active coping t(101) = .516, p = .607, Behavioral disengagement t(101) = .018, p = .985, Restraint t(101) = -.051, p = .960, Substance use
Suppression of competing activities $t(101) = -.776, p = .439$, or Planning $t(101) = .397, p = .692$. Contrary to initial hypotheses, there were no significant differences among Hispanics and Non-Hispanic Whites on these coping strategies.

Interestingly though, ethnic differences did emerge in the use of these coping strategies. Specifically, there was a significant difference, among Hispanic and Non-Hispanic White caregivers regarding Positive Reinterpretation and Growth $t(101) = 2.668, p = .009$, with Hispanic caregivers ($M = 13.20, SD = 2.10$) reporting it more frequently than Non-Hispanic White caregivers ($M = 11.79, SD = 3.10$). Thus, Hispanic caregivers seem to be able to derive more positive meaning from caregiving than their non-Hispanic White counterparts. Consistent with initial hypotheses, Hispanic caregivers ($M = 6.33, SD = 3.09$) report significantly greater use of Denial $t(101) = 3.270, p = .002$, than Non-Hispanic White caregivers ($M = 4.83, SD = 1.26$). Religious coping $t(101) = 3.224, p = .002$, was found to be lower in Non-Hispanic White caregivers ($M =10.22, SD = 4.95$) than Hispanic caregivers ($M = 12.94, SD = 3.38$). Hispanic caregivers ($M = 10.13, SD = 4.36$) are significantly more likely to use Humor $t(101) = 5.420, p = .0001$ than Non-Hispanic White caregivers ($M = 6.12, SD = 3.09$). And finally, a significant group difference was found for Acceptance $t(101) = 2.071, p = .041$ with Hispanic caregivers ($M = 13.52, SD = 2.85$) reporting it less frequently than Non-Hispanic White caregivers ($M = 12.27, SD = 3.29$). Although it did not reach significance, consistent with initial hypotheses and previous research, there was a marginally significant difference between Hispanic and Non-Hispanic White caregivers regarding Use of emotional social support $t(101) = -1.751, p = .083$, with Hispanic caregivers ($M = 8.98$, $SD = 3.38$) reporting it more frequently than Non-Hispanic White caregivers ($M = 7.33, SD = 2.10$).
-reporting it less frequently than Non-Hispanic White caregivers ($M = 10.18, SD = 3.29$).

Two Multivariate Analyses of Variance were performed to assess the relationship between gender, ethnicity, and coping methods used. The independent variables used were gender and ethnicity, and the dependent variable was coping. There was a marginally statistically significant difference in gender influencing coping $F(2, 100) = 2.943, p = .057$; Wilk's Lambda = .944; eta squared = .056. However, there was no significant relationship between ethnicity and coping $F(2, 100) = 2.424, p = .094$; Wilk's Lambda = .954; eta squared = .046. These findings suggest that gender accounts for more of the variance seen in coping skills used than ethnicity.

Multiple regression analysis was used to test if demographics significantly predicted coping methods used. A separate linear regression was done for each coping method that was highly correlated. The results of the regression indicated that demographics explained 9% of the variance, $R^2 = .090, F(3, 99) = 3.227, p = .024$ in the use of Positive Interpretation and Growth. It was found that ethnicity significantly predicted the use of Positive Interpretation and Growth (Beta = .071, $p = .007$). The results of the regression indicated that demographics explained 8% of the variance, $R^2 = .080, F(3, 99) = 2.869, p = .040$ in the Use of Instrumental Social Support. It was found that gender significantly predicted the Use of Instrumental Social Support (Beta = .061, $p = .012$). The results of the regression indicated that demographics explained 20% of the variance, $R^2 = .200, F(3, 99) = 8.251, p = .0001$ in the use of Religious Coping. It was found that gender significantly predicted the use of Religious Coping (Beta = .098, $p = .001$), as did ethnicity (Beta = .094, $p = .001$). The results of the
regression indicated that demographics explained 13% of the variance, $R^2 = .134$, $F(3, 99) = 5.120$, $p = .002$ in the Use of Emotional Support. It was found that gender significantly predicted the Use of Emotional Support ($\text{Beta} = .098, p = .002$). The results of the regression indicated that demographics explained 15% of the variance, $R^2 = .149$, $F(3, 99) = 5.798$, $p = .001$ in Substance Use. It was found that gender significantly predicted the Substance Use ($\text{Beta} = .144, p = .0001$). The results of the regression indicated that demographics explained 3% of the variance $R^2 = .034$, $F(3, 99) = 1.162, p = .328$ in the use of Suppression of Significant Activities, but it was not significant. The results of the regression indicated that demographics explained 12% of the variance, $R^2 = .124$, $F(3, 99) = 4.659$, $p = .004$ in Denial. It was found that ethnicity significantly predicted Denial ($\text{Beta} = .086, p = .002$). The results of the regression indicated that demographics explained 23% of the variance, $R^2 = .225$, $F(3, 99) = 9.596$, $p = .001$ in Humor. It was found that ethnicity significantly predicted Humor ($\text{Beta} = .225, p = .0001$). The results of the regression indicated that demographics explained 6% of the variance, $R^2 = .058$, $F(3, 99) = 2.031$, $p = .114$ in Acceptance, but it was not significant. Thus, the demographic factors show different patterns of predictive associations with different coping strategies.

**Demographics Differences in Life Satisfaction**

The third hypothesis sought out to determine if there existed a significant difference between demographics and life satisfaction. It was expected that older caregivers would report less life satisfaction. Similarly, female and Hispanic caregivers would also report less life satisfaction than their male Non-Hispanic White counterparts. The relationship between life satisfaction and age was investigated using a Pearson
Product Moment Correlations. Results indicate that there is no significant relationship between age and life satisfaction ($r = .0001, p = .996$).

An independent samples t-test was conducted to compare the life satisfaction between males and females. Results show that there is a marginally significant relationship between gender and life satisfaction $t(101) = -1.890, p = .062$, which indicates that although male caregivers ($M = 22.23, SD = 8.34$) reported lower life satisfaction than females ($M = 25.489, SD = 6.07$), the relationship is only marginally significant. An independent samples t-test was also conducted to compare life satisfaction between Hispanic and Non-Hispanic White caregivers. Results show that there is a significant difference in life satisfaction between Hispanic and Non-Hispanic White caregivers $t(101) = 2.000, p = .048$, signifying that Hispanic caregivers ($M = 26.17, SD = 6.59$) experience greater life satisfaction than Non-Hispanic White caregivers ($M = 23.61, SD = 6.34$) in this sample.

Multiple regression analysis was used to test if demographics significantly predicted life satisfaction. The results of the regression indicated that demographics explained 7% of the variance, $R^2 = .073, F(2, 100) = 3.913, p = .023$ in life satisfaction. It was found that ethnicity significantly predicted life satisfaction (Beta = .038, $p = .044$). Gender only marginally significantly predicted life satisfaction (Beta = .035, $p = .057$).

**The Role of Familism and Filial Piety**

Although it was not included in the hypotheses of the study, the role of familism and filial piety were examined as proxies of ethnicity. Results show that although familism was not significantly related to adaptive or maladaptive coping, it did have a significant influence on mental health outcomes. Specifically, familism was found to be
highly correlated with life satisfaction, \( r = .202, p = .04 \). Those caregivers who reported greater familistic beliefs also reported higher life satisfaction. Moreover, higher familism was associated with lower rates of burden, \( r = -.233, p = .02 \) and lower rates of trait anxiety, \( r = -.226, p = .02 \). Filial piety was found to be significantly correlated with maladaptive coping, \( r = .201, p = .04 \), indicating perhaps that some aspects of the concept of filial piety lead to less effective coping.

To further examine the role of familism a stepwise hierarchical linear regression analysis was performed with burden as the dependent measure and age, gender, ethnicity, familism and filial piety as the independent measures. The overall regression model was significant \( F(1, 102) = 5.81, p = .02 \). Of the carriers, only familism was a significant predictor of burden claiming twenty-three percent of the variance in burden (Beta =-.233, \( p = .02 \)). Similarly, when a stepwise hierarchical linear regression was computed with life satisfaction as the mental health outcome and age, gender, ethnicity, familism and filial piety as predictors, again the model was significant, \( F(1, 102) = 4.28, p = .04 \) and familism was the only carrier to enter the model and was found to predict twenty percent of the variance in life satisfaction (Beta = .202, \( p = .41 \)). The regression with depression as the outcome was not significant. Nor were the regressions predicting state or trait anxiety.

**Coping as a Mediator for Demographics and Life Satisfaction**

The fourth hypothesis stated that coping was expected to mediate the relationship among demographic factors and life satisfaction. In order to test the mediational role of coping methods a series of hierarchal linear regressions were used following Holmbeck (2006) and Kraemer, et al., (2008). Carver, et al., (1989), discussed a
theoretically driven and empirically supported conceptualization of coping methods being either adaptive or maladaptive. The author's distinction was used as a guide to consolidate all subscales from the COPE inventory (Carver, et al., 1989) to either adaptive coping methods or maladaptive coping methods. The following regressions were based on these two variables.

First, a hierarchical linear regression was conducted to determine if there is a significant relationship between the demographic factors and life satisfaction. Demographics are predictor variables expected to be significantly associated with life satisfaction. In order for coping to serve as a mediating factor, this relationship must exist. In step one of the hierarchical linear regressions used to test this linkage, age and gender were entered into the model ($R^2 = .034$), and in step two, ethnicity was added ($R^2 = .073$). The total models $R^2$ change was .039 and was significant ($p = .045$). The model as a whole was found to be marginally significant; $F(3, 99) = 2.587, p = .057$. In sum, the data suggests that there is a significant relationship between demographic factors and life satisfaction, thus meeting the first criteria necessary to show coping as a mediator.

Second, a hierarchical linear regression was conducted to determine if there is a significant relationship between the demographic factors and adaptive coping. In order
for coping to be a mediating variable, a significant linkage must exist between demographics factors and adaptive coping. In step one of the hierarchical linear regression conducted to investigate this relationship, age and gender were entered into the model ($R^2 = .055$), and in step two, ethnicity was added ($R^2 = .101$). The total models $R^2$ change was .46 and was significant ($p = .026$). The model as a whole was found to be significant, $F(3, 99) = 3.717, p = .014$. In sum, the data suggest that there is a significant relationship between demographic factors and adaptive coping, thus meeting one part of the second criteria. The linear regression shows that demographics account for 4.6% of the variance in the use of active coping. In order for coping to serve as a mediating variable between demographics and life satisfaction, this relationship had to exist to account for the impact that age, gender, and ethnicity have on life satisfaction as an outcome.

Third, a hierarchical linear regression was conducted to determine if there is a significant relationship between the demographic factors and maladaptive coping. A separate linear regression was necessary to ensure that coping, as a whole, is significantly related to demographics. In step one, age and gender were entered into the model ($R^2 = .000$), and in step two, ethnicity was added ($R^2 = .001$). The total models $R^2$ change was .001 but was not significant ($p = .765$). The model as a whole was not found to be significant; $F(3, 99) = .039, p = .990$. In sum, the data suggests that there is no significant relationship between demographic factors and maladaptive coping, thus failing to meet another part of the second criteria. The results indicate that age, gender, and coping, do not account for variance seen in the use of maladaptive coping. Since this
relationship does not exist, coping cannot be identified as a mediator between
demographics and life satisfaction.

**Coping as a Mediator for Burden and Life Satisfaction**

For the final hypothesis of the study, a series of hierarchal linear regressions were
used following Holmbeck (2006) and Kraemer, et al., (2008) in order to test the
mediational role of coping between burden and life satisfaction. Caregiver burden was
expected to be a predicting variable for life satisfaction, however, coping was thought to
mediate this relationship. First, a hierarchical linear regression was conducted to
determine if there was a significant relationship between caregiver burden and
life satisfaction. Caregiver burden was entered in to the model \( R^2 = .256 \) and was
significant \( (p = .0001) \). The model as a whole was found to be significant; \( F(2, 101) = 34.718, p = .0001 \). In sum, the data suggest that there is a significant relationship
between burden and life satisfaction, thus meeting the first criteria and signifying that
caregiver burden accounts for variance seen in levels of life satisfaction.

Second, a hierarchical linear regression was conducted to determine if there is a
significant relationship among burden and coping, which must exist to consider coping a
mediating variable between burden and life satisfaction. In step one, adaptive coping was
entered into the model ($R^2 = .011$), and in step two, maladaptive coping was added ($R^2 = .022$). The total models $R^2$ change was .11 and was significant ($p = .0001$). The model as a whole was found to be significant; $F(2, 100) = 9.525, p = .0001$. In sum, the data suggest that there is a significant relationship between burden and coping, thus meeting the second criteria. The data also show that burden is significantly related to coping as a whole, and accounts for 11% of the variance seen in the use of adaptive and maladaptive coping.

Third, a hierarchical linear regression was conducted to determine if there is a significant relationship between coping style and life satisfaction after controlling for burden. Further analyses were necessary to see if this model, which presents coping as a mediating variable, is acceptable. In step one, burden were entered into the model ($R^2 = .256$), and in step two, coping was added ($R^2 = .284$). The total models $R^2$ change was .28 but was not significant ($p = .206$). The model as a whole was found to be significant; $F(3, 99) = 13.085, p = .0001$. In sum, the data suggest that there is no significant relationship between coping and life satisfaction after controlling for burden, thus failing to meet the third criteria. The results indicate that caregiver burden and coping style do not significantly account for the variance seen in life satisfaction. Thus, coping cannot be seen as a mediator for burden and life satisfaction.
CHAPTER V  
DISCUSSION

The focus of the present research study was to extend the current understanding of how life satisfaction is impacted by caregiver burden by investigating whether or not there are demographic differences in coping strategies used to handle caregiving. The results indicate that there is indeed a relationship between burden and life satisfaction. More specifically, as caregiver burden increases, life satisfaction decreases. Coping strategies were not found to mediate this relationship in this particular sample. Caregiver burden was also found to relate to depression and anxiety. Results show that as caregiver burden increases, so does state anxiety, trait anxiety, and levels of depression experienced by caregivers. Furthermore, as these mental health variables increase, caregivers tend to experience lower levels of life satisfaction. Results show that women tend to employ more adaptive methods of coping than men, such as to improving their support systems and seeking advice from others. Ethnicity also influenced caregivers' choice of coping style. Hispanic caregivers were more likely to use religion, as well as humor, as a coping mechanism, which may explain why female and Hispanic caregivers were more likely to experience greater levels of life satisfaction. The results of the current study replicate earlier research and emphasize the negative mental health impact of caregiver burden and dysfunctional coping in the context of caregiving, and point to potential pathways for effective interventions to encourage more adaptive coping to reduce caregiver burden.
Outcomes of Caregiving

The main goal of the present research study was to investigate the relationship between caregiver burden and life satisfaction and examine the role that demographics play in caregivers’ choice of coping skills. Analyses conducted on the demographic differences in burden experience did not support the study's hypothesized relationship. Older caregivers were expected to experience greater levels of caregiver burden on the basis of empirical evidence suggesting age increases burden (Choo, et al., 2003; Minnes, et al., 2007; Roberto, 1995). However, results from the present study did not support that finding. Conversely, results indicate that as caregivers age, they experience lower levels of caregiver burden, while younger caregivers are experiencing higher levels of burden. Furthermore, there was no evidence that age is associated with a worsened mental state while in a caregiving role. One possible explanation for this may be that the caregiving elders in the present study were healthier and more physically able than counterparts in previous studies. Similarly, the resources available to caregivers in this sample may have been greater than in other previous samples. Given the large network of family and community resources available to Hispanic caregivers, perhaps burden is lessened due to the social support gleaned from these resources. However, these stipulations remain conjecture, as they were not empirically tested in the present study.

On the basis of the accumulated literature it was expected that female caregivers would experience greater burden (Choo, et al., 2003; Fitting, et al., 1986; Waldron-Perrine, et al., 2009). However, no significant differences were found between the levels of burden experienced between male and female caregivers. The lack of a significant relationship between gender and caregiver burden in the present study may be attributed
to the uneven distribution of male and female participants in the study. Male caregivers accounted for only a small percentage (16.5%) of all the caregivers interviewed. While historically caregiving has always fallen to females, and research suggests that females are, in fact, better suited for caregiving (Collins, et al., 1997), such a notion may be especially true in Hispanic families in which traditional gender roles are prevalent (Galanti, 2003). Thus, in recruiting caregivers for the present study, it was challenging to find male caregivers. The male caregivers who did participate may have been less burdened and still able to manage the care of their loved ones well, and thus they did not differ from significantly from the females in the sample. Again, caution must be used in extrapolating from the results, as this suggestion was not tested empirically.

Contrary to initial hypotheses, the present results also failed to support the relationship between ethnicity and caregiver burden. Hispanic caregivers were expected to experience higher levels of burden consistent with previous research showing that Hispanic caregivers experience higher levels of depression and distress than Non-Hispanic White caregivers (Pinquart, et al., 2005; Valle, et al., 2004). Other studies report that Hispanic caregivers stated that they are in need of more social support (Adams, et al., 2002), both of which contribute to increased burden (McCallum, et al., 2007). However, in the present study Hispanic and Non-Hispanic caregivers did not differ in terms of self-reported caregiver burden, depression, or anxiety, which may be the result of Hispanic caregivers endorsing ethnocultural beliefs, such as familism and filial piety (Kim, et al., 2007; McCallum, et al., 2007; Pinquart, et al., 2005). Research has shown that familism may affect levels of caregiver burden and, when accompanied with social support, may improve outcomes (Kim, et al., 2007; McCallum, et al., 2007).
The fact that Hispanics in the present sample had higher levels of life satisfaction could be seen as support for previous research that shows that Hispanic caregivers find their role to be more rewarding than Non-Hispanic White caregivers (Pinquart, et al., 2005). The results of the present study do show overwhelming evidence that familism increases life satisfaction, and reduces burden and anxiety. Thus, it is a beneficial belief when it comes to the experience of caregiving.

In the present study, it was anticipated that demographic factors such as age, gender, and ethnicity would influence the differential use of coping strategies. Specifically, in line with previous studies, it was expected that with advancing age, caregivers would employ more dysfunctional methods of coping as older caregivers tend to use more passive methods than their younger counterparts (Birkeland & Natvig, 2009; Folkman et al., 1987). However, in the present study there were no significant linkages among age and individual coping strategies. Moreover, when individual coping strategies were consolidated into adaptive coping and maladaptive coping, age still did not have a significant relationship with coping. The results found in the present study may be due to the fact that this caregiving sample was relatively young ($M = 67.42$ years old) compared to samples in previous research. Thus, the caregivers in the present sample may still be relatively healthy and capable of managing the daily requirements of caregiving and thus may not appraise their coping (Carver, et al., 1989). The literature suggests that middle-aged women who are caregivers are often more stressed than spousal caregivers giving care in later life, in part because the middle aged caregivers must give care to both the younger generation (their own children) as well as the older generation (their parents; See Brody, 1981; Stone, Cafferata, & Sangl, 1987; Treas,
However, in the present study the research sample was developmentally beyond middle-age but not yet old enough that their own health and physical limitations created added burden within the caregiving context.

Gender was also expected to influence the coping methods chosen by caregivers. It was hypothesized that female caregivers would employ more maladaptive methods of coping, derived from the literature that states that women tend to experience greater distress and difficulties with caregiving than men (Adams, et al., 2002; Navaie-Waliser, et al., 2002). Previous research also showed that females typically rely on emotion-focused coping strategies rather than problem focused coping for dealing with stress (Collins, et al., 1997; Hooker, et al., 2000; Kristensson Ekwall, et al., 2006; Waldron-Perrine, et al., 2009). However, the results from the present study indicate that women were more likely to choose Use of Instrumental Social Support, Use of Emotional Support, and Religious coping, which is contrary to the hypothesized pattern, since these three coping strategies have been categorized as adaptive coping methods (Carver, et al., 1989). Once again, the lack of significant gender differences may simply be the result of the largely discrepant sample sizes among genders in the present study.

In the present study, it was expected that Hispanic caregivers would employ more dysfunctional methods of coping than Non-Hispanic White caregivers. While there is little previous research suggesting differential use of coping among different ethnic caregivers, this relationship was anticipated due to research findings showing elevated levels of depression and distress among Hispanic caregivers compared with caregivers of other ethnic groups (Adams, et al., 2002; Pinquart, et al., 2005; Valle, et al., 2004). The present study, which explicitly examined different coping strategies, was supportive of
the notion that Hispanic caregivers employed maladaptive coping styles. Hispanic caregivers were more likely to use Denial, which is seen as a maladaptive coping skill (Carver, et al., 1989). Hispanic caregivers were also less likely to choose Acceptance as a coping skill and reported a lower Use of Emotional Support than Non-Hispanic White caregivers. Conversely, the results show that Hispanic caregivers chose Positive Reinterpretation and Growth, and Humor more frequently than Non-Hispanic White caregivers. Thus, while it was found that Hispanic caregivers are using some maladaptive coping strategies they were also found to be using some adaptive coping strategies. Taken together these individual strategies may balance out, and that may explain why Hispanic caregivers did not experience greater burden and why they still show higher life satisfaction from their caregiving experiences. Findings from the present study are important because a novel approach of investigating micro-level coping strategies as well as macro-level coping strategies was used. Whereas no ethnic differences emerged at the macro-level (i.e., adaptive versus maladaptive coping), ethnic differences did emerge at the micro-level (i.e., the level of specific strategies) and this highlights the importance of individual actions both in terms of managing coping effectively and in terms of pointing to specific techniques that could be learned to manage the stress and burden of caregiving.

Differences in the coping strategies employed are also of interest because of the impact these strategies have on levels of depression (Gallagher-Thompson, Gray, Dupart, Jimenez & Thompson, 2008; McQueeny, Stanton & Sigmon, 1997; Nolen-Hoeksema, 1987; Nolen-Hoeksema, 1991). Certain coping styles, such as Focusing on and Venting Emotions, may result in longer and more severe periods of depression (Nolen-Hoeksema,
Researchers have found that creating psychoeducational interventions where resources are provided to those struggling with symptoms of anxiety and depression can greatly improve the participants' outcomes (Gallagher-Thompson, et al., 2008; McQueeny, et al., 1997). Efforts to modify the use of ineffective coping skills by providing more adaptive alternatives have provided promising results. Caregivers of dementia patients have been found to have reduced overall life stress and depressive symptoms after such a treatment (Gallagher-Thompson, et al., 2008). Most importantly, this effect can be seen across ethnicities. Gallagher-Thompson, et al., (2008), provided psychoeducational interventions to both Hispanic and Non-Hispanic White caregivers in both English and Spanish in a treatment group focusing on modify coping skills used. Their intervention resulted in lower levels of depression, perceived psychological stress, and conditional bother (burden experienced by certain behaviors care recipients employed).

In line with a vast amount of previous research demonstrating the linkages among caregiver burden and life satisfaction, in the present study it was expected that demographic factors such as age, gender, and ethnicity would differentially influence these outcomes. Our results do not show any age effects and the explanation may be the age of the sample as mentioned above. Gender only marginally influenced life satisfaction in this sample of caregivers, such that female caregivers report slightly higher levels of life satisfaction than male caregivers. Once again, it must be stated that the unequal groups of male and female participants may account for the lack of a stronger relationship between the two variables.
Lastly, Hispanic caregivers were expected to report lower levels of life satisfaction than Non-Hispanic White caregivers because of research that states that Hispanic caregivers experience higher levels of depression than Non-Hispanic White caregivers (Adams, et al., 2002; Cox, et al., 1990; Valle, et al., 2004). Contrary to initial hypotheses, Hispanic caregivers experience greater levels of life satisfaction than Non-Hispanic White caregivers, possibly as a result of ethnocultural beliefs and how Hispanic caregivers perceive their caregiving role. Hispanic caregivers reported more elevated levels of familism and filial piety than Non-Hispanic White caregivers, which may be a factor influencing their life satisfaction.

Finally, the present study sought to examine the mediational role of coping to determine if more adaptive coping strategies may lessen the impact of burden. More specifically, on the basis of previous research demonstrating that male caregivers who use less adaptive methods of coping were found to experience higher levels of distress (Rose, et al. 1997), while women who use more adaptive methods of coping experienced lower levels of depression (Essex, et al., 1999). A model was created in which assumptions needed to be met in order to assume a mediational role of coping (Holmbeck, 2006; Kraemer, et al., 2008). While significant linkages among demographic factors and life satisfaction were found, and linkages among coping and life satisfaction were found, there were no significant linkages among demographic factors and coping therefore, it could not be concluded that coping mediates the relationship between demographic characteristics and life satisfaction. It is highly likely that the mediational role of coping would have emerged if the demographic factors had been more balanced across the sample. Not only did this study include small samples of caregivers within
each ethnic group, the imbalance among genders was a significant disadvantage. Moreover, it could be that the distinctions among adaptive and maladaptive coping strategies (Carver, et al., 1989) were not the proper approach at a constructual level. Evidence of this can be seen in the significant relations that emerged when individual coping strategies were examined that then were not found when examining coping at the level of adaptive/maladaptive. For several methodological reasons, it is possible that the analyses conducted here were not able to detect the mediational role of coping. In the present study, a significant relationship among burden and life satisfaction was found, yet burden and coping were not significantly related. The level of burden experienced in the current sample may be relatively low, in part because the caregivers were younger and perhaps more able to manage the stress of caregiving, which may explain the results obtained.

Limitations

The present study has several limitations that should be taken into account. First, it was a significant challenge to recruit caregivers into this study. Recruitment took nearly two years and it was especially hard to recruit the non-Hispanic White caregivers. Perhaps as a function of having to rely on a sample of convenience, the final sample consisted of low and unequal numbers of ethnic caregivers and a disproportionate number of female caregivers. The sample was also relatively young compared with previous caregiving studies. These factors taken together not only impacted the results derived from the study but signal the need to use caution in extrapolating or generalizing from the findings. It could also be that the recruitment of caregivers from support groups and referrals may lead to a sample of caregivers who are less burdened and coping more
effectively over all. This involvement in a supportive context may affect caregivers' outcomes by moderating their perceived levels of burden and life satisfaction. In addition to the ample resources available to caregivers in South Florida, there may be a significant difference in the socioeconomic status of Hispanic caregivers in the present study as opposed to caregivers in other parts of the country. More specifically, Hispanic caregivers in South Florida may have a higher socioeconomic status than Hispanic caregivers in other areas. Thus, the nature of the convenience sample suggests the need for caution in interpreting the results. Additionally, a limitation of the study is that it did not take into account the type of disease the care-recipient had, the level of dysfunction and type of caregiving required, or the amount of time the caregiver had been in that role. All these factors have been found to relate to the amount of distress and burden that caregivers experience (Hooker, et al., 1994; Hooker, et al., 2000; Hooker et al., 1998). Although all caregivers were required to have been caring for a loved one for over a year, the cumulative effects of caregiving after one year may not be similar to that of caregiving after five years. And thus, a final caveat to consider is the fact that this research was cross-sectional in nature and thus the ongoing and cumulative effects of the caregiving experience could not be examined. In essence, this research presents a snap-shot of the caregiving experience among a self-selecting sample of caregivers at one point in time. Given the dynamic and transitory nature of giving care to a loved one with a debilitating chronic illness, this snap-shot does not really reflect the totality of the caregiving experience.
Potential Implications

Nevertheless, the present study provides useful information on specific factors such as gender and ethnicity and how they relate to the use of different coping strategies within the context of caregiving in later life. The findings show that certain caregivers could benefit from learning more effective coping strategies to lessen their burden and improve their life satisfaction. The findings showed that certain methods of coping can lead to higher levels of depression, anxiety, and caregiver burden, as well as lower levels of life satisfaction. Knowledge of specific strategies that relate to better mental health outcomes can inform and shape psychosocial interventions aimed at caregivers by training and facilitating the use of adaptive coping to reduce negative outcomes and improve life satisfaction. Psychoeducational interventions focusing on the importance of employing adaptive coping methods and refraining from turning to maladaptive coping styles may be an asset to the interventions available for therapists working with this population.

In any research study the limitations may give rise to further investigation of important questions. From the current findings it would be useful for future research to investigate the cumulative effects of caregiving. Findings here suggest the need for longitudinal studies to better examine how demographics, coping, and caregiver outcomes interact over time. Moreover, a greater and more in depth analysis of the social and familial networks of ethnically diverse caregivers is very important. Clearly, ethnic difference exist in the experience of caregiving however we do not have a clear picture of what factors within different ethnic contexts may lead to better or more effective coping for caregivers. For example, is the higher life satisfaction that was found in Hispanics
due to decreased burden because of greater social support or greater involvement by others in the family system? Ultimately, this study has created several questions that future researchers may seek to answer.

In conclusion, the present study has highlighted points of interventions for therapists who seek to increase levels of life satisfaction and decrease levels of burden in caregivers. The effects of certain coping skills were reported in efforts to guide interventions aimed at improving the quality of coping methods employed by caregivers. Furthermore, this study has added to the literature by not only discussing the effects of demographics and coping on caregiver outcome, but also by expanding the opportunities for future research.
<table>
<thead>
<tr>
<th>Coping Methods</th>
<th>Depression</th>
<th>State Anxiety</th>
<th>Trait Anxiety</th>
<th>Life Satisfaction</th>
<th>Caregiver Burden</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive Reinterpretation and Growth</td>
<td>Pearson</td>
<td>-.062</td>
<td>-.273**</td>
<td>-.210*</td>
<td>.155</td>
</tr>
<tr>
<td></td>
<td>Correlation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sig.</td>
<td>.536</td>
<td>.005</td>
<td>.033</td>
<td>.117</td>
</tr>
<tr>
<td></td>
<td>(2-tailed)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mental Disengagement</td>
<td>Pearson</td>
<td>.382**</td>
<td>.298**</td>
<td>.348**</td>
<td>-.138</td>
</tr>
<tr>
<td></td>
<td>Correlation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sig.</td>
<td>.000</td>
<td>.002</td>
<td>.000</td>
<td>.164</td>
</tr>
<tr>
<td></td>
<td>(2-tailed)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Focus On Venting</td>
<td>Pearson</td>
<td>.346**</td>
<td>.166</td>
<td>.418**</td>
<td>-.192</td>
</tr>
<tr>
<td></td>
<td>Correlation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sig.</td>
<td>.000</td>
<td>.094</td>
<td>.000</td>
<td>.052</td>
</tr>
<tr>
<td></td>
<td>(2-tailed)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Use of Instructional Social Support</td>
<td>Pearson</td>
<td>-.004</td>
<td>-.120</td>
<td>-.048</td>
<td>-.002</td>
</tr>
<tr>
<td></td>
<td>Correlation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sig.</td>
<td>.971</td>
<td>.227</td>
<td>.632</td>
<td>.983</td>
</tr>
<tr>
<td></td>
<td>(2-tailed)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pearson Correlation</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------------</td>
<td>-----------</td>
<td>-----------</td>
<td>-----------</td>
<td>-----------</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sig. (2-tailed)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Active Coping</td>
<td>-.227**</td>
<td>.021</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Denial</td>
<td>.411**</td>
<td>.000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Religious Coping</td>
<td>.028</td>
<td>.781</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Humor</td>
<td>.121</td>
<td>.224</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Behavioral Disengagement</td>
<td>.398**</td>
<td>.000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Restraint</td>
<td>.184</td>
<td>.063</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Use of Emotional Support</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
</tr>
<tr>
<td>--------------------------</td>
<td>---------------------</td>
<td>----------------</td>
<td>---------------------</td>
<td>----------------</td>
<td>---------------------</td>
</tr>
<tr>
<td></td>
<td>-.047</td>
<td>.640</td>
<td>-.135</td>
<td>.172</td>
<td>-.068</td>
</tr>
<tr>
<td>Substance Use</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
</tr>
<tr>
<td></td>
<td>.052</td>
<td>.605</td>
<td>.073</td>
<td>.464</td>
<td>.045</td>
</tr>
<tr>
<td>Acceptance</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
</tr>
<tr>
<td></td>
<td>-.301**</td>
<td>.002</td>
<td>-.326**</td>
<td>.001</td>
<td>-.209*</td>
</tr>
<tr>
<td>Suppression of Competing Activities</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
</tr>
<tr>
<td></td>
<td>.152</td>
<td>.126</td>
<td>.007</td>
<td>.942</td>
<td>.100</td>
</tr>
<tr>
<td>Planning</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
<td>Sig. (2-tailed)</td>
<td>Pearson Correlation</td>
</tr>
<tr>
<td></td>
<td>-.081</td>
<td>.413</td>
<td>-.164</td>
<td>.098</td>
<td>-.035</td>
</tr>
</tbody>
</table>

**. Correlation is significant at the 0.01 level (2-tailed).
*. Correlation is significant at the 0.05 level (2-tailed).
Table 2

Results of Independent Sample t-Tests for Gender and Coping

<table>
<thead>
<tr>
<th>Outcome</th>
<th>Male Caregivers M (SD)</th>
<th>Female Caregivers M (SD)</th>
<th>t</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use of Instrumental Social Support</td>
<td>7.94 (3.54)</td>
<td>10.37 (3.54)</td>
<td>t(101) = -2.58</td>
<td>.011*</td>
</tr>
<tr>
<td>Religious Coping</td>
<td>8.59 (4.62)</td>
<td>12.26 (4.11)</td>
<td>t(101) = -3.291</td>
<td>.001**</td>
</tr>
<tr>
<td>Use of Emotional Support</td>
<td>7.12 (2.96)</td>
<td>10.02 (3.38)</td>
<td>t(101) = -3.235</td>
<td>.002**</td>
</tr>
<tr>
<td>Substance Use</td>
<td>5.82 (3.57)</td>
<td>4.12 (.65)</td>
<td>t(101) = 4.109</td>
<td>.071</td>
</tr>
</tbody>
</table>

**. Significant at the 0.01 level
*. Significant at the 0.05 level
<table>
<thead>
<tr>
<th>Outcome</th>
<th>Hispanic Caregivers $M (SD)$</th>
<th>Non-Hispanic White Caregivers $M (SD)$</th>
<th>$t$</th>
<th>$p$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive Reinterpretation and Growth</td>
<td>13.20 (2.10)</td>
<td>11.79 (3.10)</td>
<td>$t(101) = 2.668$</td>
<td>.009**</td>
</tr>
<tr>
<td>Denial</td>
<td>6.33 (3.09)</td>
<td>4.83 (1.26)</td>
<td>$t(101) = 3.270$</td>
<td>.002**</td>
</tr>
<tr>
<td>Religious Coping</td>
<td>12.94 (3.38)</td>
<td>10.22 (4.95)</td>
<td>$t(101) = 3.224$</td>
<td>.002**</td>
</tr>
<tr>
<td>Humor</td>
<td>10.13 (4.36)</td>
<td>6.12 (3.09)</td>
<td>$t(101) = 5.420$</td>
<td>.0001**</td>
</tr>
<tr>
<td>Acceptance</td>
<td>12.27 (3.29)</td>
<td>13.52 (2.85)</td>
<td>$t(101) = -2.071$</td>
<td>.041*</td>
</tr>
<tr>
<td>Use of Emotional Support</td>
<td>8.98 (2.91)</td>
<td>10.18 (3.29)</td>
<td>$t(101) = -1.751$</td>
<td>.083</td>
</tr>
</tbody>
</table>

**. Significant at the 0.01 level  
*. Significant at the 0.05 level
Table 4

*Correlations among Familism, Filial Piety and Outcomes*

<table>
<thead>
<tr>
<th>Variables</th>
<th>Familism</th>
<th>Filial Piety</th>
<th>Adaptive Coping</th>
<th>Maladaptive Coping</th>
<th>Depression</th>
<th>State Anxiety</th>
<th>Trait Anxiety</th>
<th>Life Satisfaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filial Piety</td>
<td></td>
<td>.417**</td>
<td>.105</td>
<td>.004</td>
<td>-.058</td>
<td>.088</td>
<td>-.027</td>
<td>.515*</td>
</tr>
<tr>
<td>Correlation</td>
<td>Sig.</td>
<td>.000</td>
<td>.291</td>
<td>.969</td>
<td>.559</td>
<td>.378</td>
<td>.784</td>
<td>.000</td>
</tr>
<tr>
<td>(2-tailed)</td>
<td>N</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
</tr>
<tr>
<td>Adaptive Coping</td>
<td>Pearson</td>
<td>.105</td>
<td>.012</td>
<td>.004</td>
<td>.088</td>
<td>.088</td>
<td>-.027</td>
<td>.515*</td>
</tr>
<tr>
<td>Correlation</td>
<td>Sig.</td>
<td>.291</td>
<td>.908</td>
<td>.041</td>
<td>.378</td>
<td>.784</td>
<td>.000</td>
<td>.000</td>
</tr>
<tr>
<td>(2-tailed)</td>
<td>N</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
</tr>
<tr>
<td>Maladaptive</td>
<td>Pearson</td>
<td>.004</td>
<td>.201*</td>
<td>.187</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coping</td>
<td>Correlation</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sig. (2-tailed)</td>
<td></td>
<td>.969</td>
<td>.041</td>
<td>.059</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N</td>
<td></td>
<td>103</td>
<td>103</td>
<td>103</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Depression</td>
<td>Pearson</td>
<td>-.058</td>
<td>.088</td>
<td>.027</td>
<td>.559</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Correlation</td>
<td>Sig.</td>
<td>.559</td>
<td>.378</td>
<td>.784</td>
<td>.000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(2-tailed)</td>
<td>N</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pearson Correlation</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------------</td>
<td>---------------------</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sig. (2-tailed)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>N</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>State Anxiety</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-.099</td>
<td>.125</td>
<td>-.177</td>
<td>.375**</td>
<td>.510**</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>.319</td>
<td>.207</td>
<td>.074</td>
<td>.000</td>
<td>.000</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Trait Anxiety</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-.226*</td>
<td>.149</td>
<td>-.076</td>
<td>.437**</td>
<td>.698**</td>
<td>.562**</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>.022</td>
<td>.133</td>
<td>.446</td>
<td>.000</td>
<td>.000</td>
<td>.000</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Life Satisfaction</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>.202*</td>
<td>.068</td>
<td>.106</td>
<td>-.082</td>
<td>-.306**</td>
<td>-.400**</td>
<td>-.603**</td>
<td></td>
</tr>
<tr>
<td></td>
<td>.041</td>
<td>.493</td>
<td>.285</td>
<td>.408</td>
<td>.002</td>
<td>.000</td>
<td>.000</td>
<td></td>
</tr>
<tr>
<td></td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td></td>
</tr>
<tr>
<td><strong>Caregiver Burden</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-.233*</td>
<td>-.001</td>
<td>.043</td>
<td>.399**</td>
<td>.518**</td>
<td>.419**</td>
<td>.533**</td>
<td>-.506**</td>
</tr>
<tr>
<td></td>
<td>.018</td>
<td>.995</td>
<td>.668</td>
<td>.000</td>
<td>.000</td>
<td>.000</td>
<td>.000</td>
<td>.000</td>
</tr>
<tr>
<td></td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
<td>103</td>
</tr>
</tbody>
</table>

* **Correlation is significant at the 0.01 level (2-tailed).**
* *. Correlation is significant at the 0.05 level (2-tailed).
References


Carver, C. S. (1997). You want to measure coping but your protocol’s too long: Consider the brief COPE. *Journal of Behavioral Medicine, 4*(1), 92-100.


APPENDICES
Appendix 1 - Informed Consent
ADULT CONSENT TO PARTICIPATE IN A RESEARCH STUDY
Correlations among Coping Skills and Life Satisfaction in Ethnic Older Caregivers

PURPOSE OF THE STUDY
You are being asked to participate in this research study. The purpose of this study is to look at how older caregivers from Hispanic and Non-Hispanic White ethnicities handle the stress of caring for a loved one. There is an interest in how this stress is managed because it may lead to improved or worsened life satisfaction among these caregivers. The goal is to see which methods of stress management lead to higher or lower levels of life satisfaction.

NUMBER OF STUDY PARTICIPANTS
If you decide to be in this study, you will be one of 100 people in this research study.

DURATION OF THE STUDY
Your participation will require approximately two and a half (2 1/2) hours.

PROCEDURES
If you agree to be in the study, you will be asked to do the following things:
1. You will be asked to participate in a face-to-face interview where your mental status will be assessed. Then you will be asked about your age, ethnicity, marital status, education level, occupation, religiosity, health, and the health of the person you care for. How you handle the stress of caring for your loved one and the level of burden you feel will also be measured. The life satisfaction you currently feel and your beliefs about obligation to your family will be assessed as well. You are encouraged to be as open and honest as possible.

RISKS AND/OR DISCOMFORTS
The following risks may be associated with your participation in this study: There are only minimal risks of emotional distress expected to be associated with your participation.
BENEFITS
The following benefits may be associated with your participation in this study: Your participation will lead to a better understanding of how older caregivers from Hispanic and Non-Hispanic White ethnicities handle the stress of caring for a loved one and how this impacts their life satisfaction. This information may lead to improved psychological treatment for caregivers who want to begin attending therapy sessions by allowing the therapist to adjust his or her services to the needs of the specific caregiver.

ALTERNATIVES
There are no known alternatives available to you other than not taking part in this study. However, any significant new findings developed during the course of the research which may relate to your willingness to continue participation will be provided to you.

CONFIDENTIALITY
The records of this study will be kept private and will be protected to the fullest extent provided by law. In any sort of report that might be published, no information that will make it possible to identify a subject will be included. Research records will be stored securely and only the researcher team will have access to the records. However, your records may be reviewed for audit purposes by authorized University or other agents who will be bound by the same provisions of confidentiality.

COMPENSATION & COSTS
You will not be receiving payment for your participation. You will not be responsible for any costs to participate in this study.

RIGHT TO DECLINE OR WITHDRAW
Your participation in this study is voluntary. You are free to participate in the study or withdraw your consent at any time. Your withdrawal or lack of participation will not affect any benefits to which you are otherwise entitled. The investigator reserves the right to remove you without your consent at such time that they feel it is in the best interest.

RESEARCHER CONTACT INFORMATION
If you have any questions about the purpose, procedures, or any other issues relating to this research study you may contact Didiana De La Osa at 305-878-9818, or by email at didiana.delaosa@yahoo.com.
IRB CONTACT INFORMATION
If you would like to talk with someone about your rights of being a subject in this research study or about ethical issues with this research study, you may contact the FIU Office of Research Integrity by phone at 305-348-2494 or by email at ori@fiu.edu.

PARTICIPANT AGREEMENT
I have read the information in this consent form and agree to participate in this study. I have had a chance to ask any questions I have about this study, and they have been answered for me. I understand that I am entitled to a copy of this form after it has been read and signed.

________________________________           __________________
Signature of Participant      Date

________________________________
Printed Name of Participant

________________________________    __________________
Signature of Person Obtaining Consent    Date
Appendix 2 - Informed Consent in Spanish
CONSENTIMIENTO DE UN ADULTO PARA PARTICIPAR EN UN ESTUDIO INVESTIGATIVO.
Correlación entre métodos de adaptación y satisfacción en la vida para personas de mayor edad y cierto grupo étnico que cuidan de otra persona.

PROPÓSITO
A usted se le pide su participación en un estudio investigativo. El propósito de este estudio es analizar cómo las personas mayores de descendencia hispana y blancos no hispanos, manejan el esfuerzo y la tensión por cuidar de un ser querido. Existe interés en cómo se maneja la tensión creada por los cuidados que se le dan a otra persona, y como estos afectan en mejorar o empeorar la satisfacción en la vida entre los responsables de cuidar a un ser querido. El objetivo es llegar a establecer los métodos para controlar la tensión que llevan a niveles altos o bajos de satisfacción en la vida.

CANTIDAD DE PARTICIPANTES
Si usted está de acuerdo en participar, será uno de 100 participantes.

DURACIÓN DEL ESTUDIO
Su participación requiere aproximadamente 2 horas y media.

PROCEDIMIENTO
Si usted decide participar, se le pedirá lo siguiente:

1. Tendrá que le pedirá participar en una entrevista cara a cara donde su estado mental será evaluado. Se le preguntará su edad, grupo étnico, estado civil, nivel de educación, ocupación, religión, salud y sobre la salud de la persona que recibe sus cuidados. Como usted maneja la tensión por ser proveedor de cuidados a un ser querido y el nivel de sobrecarga que siente también serán evaluados. La satisfacción en la vida que usted siente en este momento y lo que cree sobre su obligación a su familia serán evaluados en adición. Le pedimos su total y completa honestidad y sinceridad.

RIESGOS E INCOMODIDADES
Experimentará los siguientes riesgos, asociados con este estudio: Solo existen riesgos mínimos de angustia emocional asociadas con su participación.
**BENEFICIOS**
Los siguientes beneficios pueden ser obtenidos por su participación en este estudio:
Su participación puede ayudar para un mejor entendimiento de cómo las personas mayores de descendencia hispana y blancos no hispanos, manejan el esfuerzo y la tensión por cuidar de un ser querido y cómo impacta su satisfacción en la vida. Este estudio investigativo puede tener como consecuencia mejorías en el tratamiento psicológico de las personas que ofrecen cuidados a otros si deciden en algún momento asistir a sesiones terapéuticas, permitiendo al terapeuta ajustar sus servicios a las necesidades específicas de la persona que le da cuidados a un ser querido.

**ALTERNATIVAS**
No se conocen alternativas que no sea negarse a participar en este estudio. No obstante, cualquier conclusión de importancia obtenida durante el estudio, que pueda tener relación con su disposición a continuar su participación, le será informada.

**CONFIDENCIALIDAD**
Los registros de este estudio investigativo serán privados y tendrán la máxima protección ofrecida por la ley. Aún cuando se publiquen reportes o datos, no se incluirá ninguna información que haga posible la identificación de los participantes. Los registros serán guardados en un lugar seguro y solamente los miembros del grupo investigativo tendrán acceso a la información. En un momento determinado los registros pueden ser revisados mientras se conduce una auditoría por personal autorizado de la Universidad o sus agentes, todos ellos serán regidos por las condiciones de confidencialidad arriba descritas.

**COMPENSACIÓN Y COSTOS**
Usted no recibirá pago por su participación ni tendrá costo alguno.

**DERECHO A NEGARSE O RETIRARSE**
Su participación en este estudio es totalmente voluntaria. Usted es libre de participar en el estudio o retirar su consentimiento en cualquier momento cuando lo estime. Esta decisión no afectará su elegibilidad para ningún beneficio. El investigador se reserva el derecho de retirarlo del estudio investigativo en cualquier momento y sin su consentimiento según lo estime conveniente.

**INFORMACIÓN DEL INVESTIGADOR**
En caso de tener alguna pregunta sobre el propósito, los procedimientos o cualquier otro aspecto de este estudio investigativo, puede comunicarse con Didiana De la Osa por teléfono al 305 878 9818 o por correo electrónico a: didiana.delaosa@yahoo.com

**INFORMACIÓN DE IRB**
Si desea hablar con alguien sobre sus derechos al participar en este estudio o sobre la ética de este estudio puede comunicarse con la Oficina de Integridad Investigativa de FIU por teléfono al 305 348 2494 o por correo electrónico a: ori@fiu.edu.
ACUERDO DEL PARTICIPANTE
He leído y entendido toda la información brindada en este consentimiento y estoy de acuerdo en participar en este estudio investigativo. He tenido la oportunidad de hacer preguntas sobre el estudio y he recibido respuestas. Entiendo que tengo derecho a obtener una copia de esta forma después de firmarla.

----------------------------------------------------   --- -----------------------
Firma del participante      Fecha

----------------------------------------------------   --- ------------------------
Nombre del participante

----------------------------------------------------   --- ------------------------
Firma de la persona que obtiene el Consentimiento  Fecha
Appendix 3 - Mini Mental Status Exam
Mini-Mental Status Exam
(Mini-examen de status mental)

I would like to check your memory and concentration.

(Con su permiso, Voy a examinar su memoria y su concentracion)

Orientation

SCORE

( ) What is the: (day of the week) (month) (date) (year) (season)? 5 points

Cual es: (el dia de la semana) (mes) (fecha) (año) (epoca del año)?

( ) Where are we: (hospital) (floor) (town) (county) (state)? 5 points

Donde estamos? (hospital) (piso) (pueblo) (condado) (estado)

( ) Repeat the 3 words house, chair, table. Remember these 3 words. 3 points

Repita estas palabras: CASA, SILLA, MESA. Acuerdese de estas 3 palabras

( 1 point/correct response, only 1 attempt)

Attention and Calculation

( ) Serial 7: Subtract 7 from 100. Repeat for a total of 5 (93, 86, 79, 72, 65). 5 points

(1 point for each correct answer on the first attempt)

Por favor reste 7 de 100 y siga restando 7 hasta que yo le diga

Answer: noventa y tres, ochenta y seis, setenta y nueve, setenta y dos, sesenta y cinco 93 86 79 72 6 5

OR

( ) Spell HOME backwards. (Score 1 point for each letter in the correct place, D/C after first failure)

Deletrée HOGAR alrevés o sea de atras para adelante Answer: R A G 0 H
Recall

( ) Repeat the 3 words (1 point/correct response, only 1 try), 3 points
Por favor repita las 3 palabras que le dije que se acordara

Language

( ) Name a pencil (lapiz) and a watch (reloj). 2 points
Como se llama esto? Answer: lapiz y reloj

( ) Repeat the following: "No ifs, ands or buts."
Repita lo siguiente: "Ni porque ni porcuanto" 1 point

( ) 3 stage command: Take this paper with your right hand, fold it in half and give it back to me
Tome este papel con su mano derecha, doblelo por la mitad y regreseme el papel

Read and obey the following:

Lea y obedezca lo siguiente:

( ) Close your eyes (Cierre sus ojos) 1 point

( ) Write a sentence (Escriba cualquier frase u oracion). 1 point

( ) Copy this drawing (Copie este dibujo) 1 point

Total Score

DERIVING THE TOTAL SCORE

Add the number of correct responses. The maximum is 30.

Total Score

23-30 = Normal / 19-23 = Borderline / <19 = Impaired
CLOSE YOUR EYES
(CIERRE SUS OJOS)

WRITE A SENTENCE
(ESCRIBA CUALQUIER FRASE U ORACIÓN)

COPY THIS DRAWING
(COPIE ESTE DIBUJO)
Appendix 4 - General Information Survey
General Information Survey

Date: _______________     Interviewer:____________

Language preferred?  ENGLISH _____     SPANISH _____

I. DEMOGRAPHIC INFORMATION

1. How old are you? _______

2. Where were you born? ________________________________
   (town, state, country)

3. Are you? :        MALE _______     FEMALE _______

4. What is your ethnic background (check one)?
   ______ African American
   ______ Asian
   ______ Hispanic
   ______ West Indian
   ______ White Caucasian
   ______ Other

5. What is your religious affiliation (Check one)?
   ______ Christian
   ______ Catholic
   ______ Protestant
   ______ Orthodox
   ______ Jewish
6. What is your current marital status (check one)?

   _____ Married
   _____ Divorced
   _____ Separated
   _____ Widowed
   _____ Never Married

7. If married, how long have you been married? _______

8. Do you have any children?     YES _______ NO _______

9. If so, how many? _______

10. How many years of school have you completed (check the highest grade completed)?

     _____ Graduate or professional degree
     _____ College graduate
     _____ Partial college
     _____ High school graduate
     _____ Completed grade school

11. If you are married, how many years of school has your SPOUSE completed (check the highest grade completed)?

     _____ Graduate or professional degree
     _____ College graduate
______ Partial college
______ High school graduate
______ Completed grade school

12. What is your current employment status (check one)?
   ______ Employed full-time
   ______ Employed part-time
   ______ Student full-time
   ______ Student part-time
   ______ Housewife
   ______ Unemployed
   ______ Volunteer
   ______ Retired

13. What is/was your current/previous occupation (be very specific)?

__________________________________________________________________

14. If you are/were married, what is/was your SPOUSE’S current/previous occupation (be very specific)?

__________________________________________________________________

   Code SES: __________

II. HEALTH INFORMATION

15. In general, would you say your health is (check one)?
    ______ Excellent
    ______ Good
    ______ Fair
16. Would you say your health is better, about the same, or not as good as others your age (Circle one)?

- Better
- About the same
- Not as good

17. Please indicate how frequently you experience any of the following symptoms, using the scale below:

<table>
<thead>
<tr>
<th>Symptom</th>
<th>1 Never or almost never</th>
<th>2 Less than 3-4 times per year</th>
<th>3 Every month or so</th>
<th>4 Every week or so</th>
<th>5 More than once every week</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eyes water</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Itching/painful eyes</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ringing in ears</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Temporary deafness/hard of hearing</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lump in throat</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Choking Sensations</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sneezing spells</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Running nose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Congested nose</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Asthma or wheezing</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coughing</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Out of breath</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Swollen ankles</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chest pains</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Racing heart</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cold hands (even in hot temps)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leg cramps</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Insomnia</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Toothaches</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Upset Stomach</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Indigestion</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heartburn</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Severe pain/cramps in stomach</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diarrhea</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Constipation</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hemorrhoids</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- Swollen Joints
- Stiff muscles
- Back Pains
- Sensitive or tender skin
- Face flushes
- Severe itching
- Skin breaks out in rash
- Acne or pimpls on face
- Acne or pimpls other than face
- Boils
- Sweating (even in cold temps)
- Strong reactions to insect bites
- Headaches
- Sensations of pressure in head
- Hot flashes
- Chills
- Dizziness
- Feel faint
- Numbness or tingling (anywhere)
- Twitching of eyelid
- Twitching other than eyelid
- Hands tremble or shake
- Stiff joints
- Sore muscles
- Sore throat
- Nausea
18. Do you currently have health problems that worry or concern you? If yes, please describe:

________________________________________________________________________

III. CARE RECIPIENT DISEASE BACKGROUND

19. What illness has the care recipient been diagnosed with?

________________________________________________________________________

____

20. When was the illness diagnosed?

________________________________________________________________________

21. Who diagnosed the care recipient’s illness (General Practitioner, Specialist)?

________________________________________________________________________

22. How severe do you think the care recipient’s disease is (Check one)?

     _______ Not very severe
     _______ Mildly severe
     _______ Moderately severe
     _______ Quite severe
     _______ Very severe

23. Is the care recipient currently taking medication?

    YES _______    NO _______

24. If yes, what medication(s) do they take?

________________________________________________________________________
25. How effective are the medications in relieving their symptoms (check one)?

_____ Very effective
_____ Quite effective
_____ Moderately effective
_____ Mildly effective
_____ Not very effective

26. Does the care recipient have any problems with their memory? YES ___  NO ___

27. If the care recipient does have problems with their memory would you say those problems are (check one)?

_____ Mild
_____ Moderate
_____ Severe
Appendix 5 - General Information Survey in Spanish
Fecha: _____________________

Nombre de la persona que conduce la entrevista:

_____________________________________

Idioma preferido?    Inglés _______  Español ________

I. INFORMACION DEMOGRÁFICA

1. Que edad tiene?  __________

2. Dónde nació? (Especifique pueblo o ciudad, estado o provincia y país)

____________________________________

3. Cuál es su sexo?  Masculino _____  Femenino _____

4. Indique su ascendencia étnica, (por favor marque solo una de las siguientes opciones):

_________ Afro-americano
_________ Asiático
_________ Hispano
_________ Indio
_________ Blanco Caucásico
_________ Otro

5. Que religión profesa? (Por favor indique una)

_________ Cristianismo
_________ Catolicismo
_________ Protestante
_________ Ortodoxo
_________ Judío
_________ Musulmán
6. Cuál es su estado civil? (Por favor marque una):
    _______ Casado
    _______ Divorciado
    _______ Separado
    _______ Viudo
    _______ Soltero (nunca se ha casado)

7. Si es casado, por cuántos años? ____________________

8. Tiene hijos?  Si ___________  No ___________

9. Cuántos hijos tiene? ________________

10. Cuál es el último grado escolar terminado? (Por favor marque el más alto grado terminado)

    _______ Graduado o título profesional (Maestría o Doctorado)
    _______ Graduado de colegio (Técnico o especialista)
    _______ Universidad o Colegio especializado parcial
    _______ Bachillerato o Pre-Universitario
    _______ Escuela Secundaria o Primaria

11. Si es casado, cuál es el último grado escolar terminado de su ESPOSA(O)? (Por favor marque el más alto grado terminado)

    _______ Graduado o título profesional (Maestría o Doctorado)
    _______ Graduado de colegio (Técnico o especialista)
    _______ Universidad o Colegio especializado parcial
    _______ Bachillerato o Pre-Universitario
    _______ Escuela Secundaria o Primaria
12. Cuál es su estado laboral? (Por favor marque uno)

[________ Tiempo Completo
________ Tiempo Parcial
________ Estudiante tiempo Completo
________ Estudiante Tiempo Parcial
________ Ama de Casa
________ Desempleado
________ Voluntario
________ Retirado]

13. Cuál es su ocupación o empleo actual o pasado si es retirado o desempleado? (Por favor especifique.)

____________________________________________________________

14. Cuál es la ocupación o empleo, actual o pasado si retirada(o) o desempleada(o) de su ESPOSA(O) ? Por favor especifique.

___________________________________________

Código SES: _______________

II. INFORMACION DE SALUD

15. Como considera usted su salud? (Marque una opción)

[________ Excelente
________ Buena
________ Regular
________ Mala]

16. Piensa usted que su salud es mejor, igual o peor que otras personas de su edad? Marque una opción por favor.

[________ Mejor
________ Igual
________ Peor]
17. Por favor indique cuánto frecuentemente experimenta usted los siguientes síntomas, usando la escala ofrecida a continuación:

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nunca o casi nunca</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Menos de 3 ó 4 veces al año</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Todos los meses</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Todas las semanas o casi todas las semanas</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- Ojos Llorosos
- Picazón o dolor en los ojos
- Sonido o ruido en los oídos
- Sordera temporal o dificultad para oír
- Masa o sensación de nudo en la garganta
- Atoros o ahogamientos
- Estornudos
- Secreción nasal continuada (Agua por la nariz)
- Congestión nasal
- Asma o silbido en el pecho
- Tos
- Falta de aire
- Tobillos inflamados
- Dolor de pecho
- Rapidez en los sonidos del corazón
- Manos frías (aunque la temperatura sea alta)
- Calambres en las piernas
- Insomnia
- Dolores de muela o de dientes
- Molestias estomacales
- Acidez
- Dolores fuertes de estómago
- Diarrea
- Extreñimiento
- Hemorroides
- Articulaciones inflamadas
- Espasmos musculares
- Dolor de espalda
- Piel sensitiva
- Calores y enrojecimiento en la cara
- Picazones
- Urticarias o erupciones de la piel
- Acné, granitos o comedones en la cara
- Acné, granitos o comedones en otro lugar que no sea la cara
- Furúnculo
- Sudores aunque la temperatura no sea muy caliente
- Reacciones fuertes a las picaduras de insecto
- Dolores de cabeza
- Sensación de presión en la cabeza
- Calores repentininos
- Escalofríos
- Mareos
- Desmayos
- Entumecimientos
- Movimientos involuntarios de los párpados
- Movimientos involuntarios en otra parte del cuerpo
- Temblores en las manos
- Articulaciones rígidas
- Dolores musculares
- Dolor de garganta
- Nausea
III. ENFERMEDADES DE LA PERSONA QUE RECIBE LOS CUIDADOS

18. Qué enfermedades le han sido diagnosticadas a la persona que cuida?
________________________________________________________________________
________________________________________________________________________
________________________________________________________________________

19. Cuándo fue hecho el diagnóstico?
__________________________________________

20. Quién hizo el diagnóstico? (Especifique especialista, médico general)
________________________________________________

21. Indique según su criterio, la severidad de la(s) enfermedades de la persona que usted cuida?
   ___ No es severa
   ___ Poco severa
   ___ Severidad moderada
   ___ Severa
   ___ Muy severa

22. Está la persona que cuida tomando medicamentos? Si ______        No _____

23. Liste los medicamentos que toma
________________________________________________________________________
________________________________________________________________________

24. Cuán efectivos son los medicamentos en aliviar los síntomas? Marque una opción
   ______ Muy efectivo
   ______ Algo efectivo
   ______ Moderadamente efectivo
   ______ Poco efectivo
   ______ No muy efectivo

25. Tiene la persona que usted cuida algún problema con la memoria?
   Si ___   No___
25. Si la respuesta es afirmativa a la pregunta anterior, por favor especifique la magnitud de los problemas de memoria

_____ Ligeros  
_____ Moderados  
_____ Severos
Appendix 6 - Familism Scale
Familism Scale

Below is a list of issues concerning the family in general, not your own. Please read all statements very carefully and respond to all of them on the basis of your own true beliefs without consulting any other persons. Do this by reading each statement and then writing, in the space provided at its left, only one of the following numbers: 0, 1, 2, 3, 4. The meaning of each of these figures is:

0: Strongly disagree  1: Disagree  2: Undecided  3: Agree  4: Strongly agree

(For research purposes, you must consider all statements as they are, without modifying any of them in any way.)

1. ___ One should make great sacrifices in order to guarantee a good education for his/her children.

2. ___ One should help economically with the support of younger brothers and sisters.

3. ___ When someone has problems s/he can count on help from his/her relatives.

4. ___ When one has problems, one can count on the help of relatives.

5. ___ One can count on help from his/her relatives to solve most problems.

6. ___ Much of what a son or daughter does should be done to please the parents.

7. ___ One should be embarrassed about the bad things done by his/her brothers or sisters.

8. ___ Children should live in their parents’ house until they get married.

9. ___ One of the most important goals in life is to have children.
Appendix 7 - Familism Scale in Spanish
Escale de Familismo

A continuación se presenta una lista de declaraciones en cuanto a las familias en general, no sobre su propia familia. Lea todas las instrucciones cuidadosamente y responda a todas las declaraciones basado en sus propias creencias verdaderas sin consultar a las demás personas. Haga esto mediante la lectura de cada declaración y, a continuación, escriba, en el espacio proporcionado a su izquierda, sólo uno de los siguientes números: 0, 1, 2, 3, 4. El significado de cada una de estas cifras es:

0: Totalmente en desacuerdo 1: En desacuerdo 2: Indecisos
3: De acuerdo 4: Totalmente de acuerdo

(Con fines de investigación, debe tener en cuenta todas las declaraciones como son, sin modificar las de ninguna manera.)

1. ___ Uno debería hacer grandes sacrificios con el objetivo de garantizar una buena educación para sus hijos
2. ___ Uno debería ayudar económicamente en el sostenimiento de sus hermanos y hermanas pequeños
3. ___ Cuando alguien tiene problemas, puede contar con la ayuda de sus familiares
4. ___ Cuando uno mismo tiene problemas, puede contar con la ayuda de sus familiares
5. ___ Uno puede contar con la ayuda de sus familiares para solucionar la mayoría de los problemas
6. ___ La mayoría de lo que hace un hijo o una hija debería hacerse para agradar a sus padres
7. ___ Uno debería avergonzarse por las cosas malas hechas por sus hermanos o hermanas.

8. ___ Deben vivir los hijos en casa de sus padres hasta que se casen.

9. ___ Uno de los objetivos más importantes en la vida es tener hijos.
Appendix 8 - Filial Piety Scale in English
Filial Piety Scale

The following items are concerned with filial piety. We would like to know what your opinions are on these items. There are no "right" or "wrong" answers. So please respond according to your own personal opinions.

For each item, select only one of the following six alternatives:

1 = Strongly agree
2 = Disagree
3 = Mildly disagree
4 = Mildly agree
5 = Agree
6 = Strongly agree

Please do not skip any item.

1. ___ Sons and daughters may protest against being unreasonably scolded by their parents. (N)
2. ___ There is no place under the sun for both oneself and the enemy of one's father.
3. ___ If there is a reason for doing so, one may rely on an old people's home to provide for one's aged parents. (N)
4. ___ Any sacrifice is worthwhile for the sake of filial piety.
5. ___ Sons and daughters should not go to faraway while their parents are still living.
6. ___ In choosing a spouse, sons and need not follow "the parents' command". (N)
7. ___ The main reason for sons and daughters not to do dangerous things is to avoid getting their parents worried.

8. ___ Parents should not interfere with their children's freedom to choose a vocation. (N)

9. ___ The great debt that you have to repay your parents is as endless as the sky.

10. ___ "Rearing sons to provide for oneself in one's old age" should no longer be the main purpose of raising children. (N)

11. ___ No matter how their parents conduct themselves, sons and daughters must respect them.

12. ___ After the father has passed away, sons and daughters must conduct themselves according to the principles and attitudes he followed while he was still living.

13. ___ If there is a quarrel between one's wife and one's mother, the husband should advise his wife to listen to his mother.

14. ___ After their parents have passed away, sons and daughters do not necessarily have to finish the business left unfinished by their parents. (N)

15. ___ "Spreading one's fame to glorify one's parents" should not be the most important reason for getting ahead. (N)

16. ___ To worship their ancestors regularly on the proper occasions is the primary duty of sons and daughters.

17. ___ To continue the family line is not the primary purpose of marriage. (N)

18. ___ Sons and daughters do not necessarily have to seek parental advice and may make their own decisions. (N)
19. ___ Sons and daughters do not necessarily have to respect the people respected and loved by their parents. (N)

20. ___ After children have grown up, all the money they earn through their own labor belongs to themselves, even though their parents are still living. (N)

21. ___ "There is no crime worse than being unfilial".

22. ___ As a son or daughter, one must obey one's parents no matter what.
Appendix 9 - Filial Piety Scale in Spanish
Piedad Filial Escala (FP)

Los siguientes incisos están relacionados con la piedad filial. Quisiéramos saber su opinión acerca de cada uno de ellos. No existe una respuesta correcta o incorrecta, solo responda lo que usted cree usando las opciones que se ofrecen a continuación:

1 = Totalmente en desacuerdo
2 = Desacuerdo
3 = Desacuerdo parcial
4 = Acuerdo parcial
5 = Estoy de acuerdo
6 = Totalmente de acuerdo

Por favor exponga su opinión en todos y cada uno.

1. ___ Los hijos tienen derecho a protestar ante una reprimenda exagerada de sus padres.
2. ___ No existe lugar en el mundo donde puedan co-existir un hijo(a) y el enemigo de su padre.
3. ___ Si existe una razón válida para hacerlo, está bien ingresar a padre o madre en un asilo de ancianos.
4. ___ Cualquier sacrificio vale la pena po mantener la piedad filial
5. ___ Los hijos no deben alejarse mucho de sus padres mientras estos vivan.
6. ___ Cuando los hijos escogen esposa(o) no necesariamente deben seguir el consejo de los padres.
7. ___ La razón principal por la que los hijos no hacen cosas que impliquen peligro es para no preocupar a sus padres.

8. ___ Los padres no deben interferir con la libertad de sus hijos para decidir su vocación.

9. ___ La gran deuda que los hijos tienen con sus padres no tiene límites.

10. ___ Nunca los padres criarán hijos con el único propósito de tener quien los cuide en la vejez.

11. ___ No importa como se comporten los padres, los hijos siempre los respetarán.

12. ___ Después que los padres se mueran, los hijos deben conducir su vida de acuerdo a los principios y actitudes que sus padres observaron mientras en vida.

13. ___ Si existen diferencias entre una esposa(o) y una madre, siempre el esposo(a) hará hincapié en escuchar y obedecer a la madre.

14. ___ Después del fallecimiento de los padres, los hijos no son responsable de terminar u ocuparse de los asuntos que ellos dejaron pendientes.

15. ___ El afán de glorificar a un padre o madre no es la razón más importante para batallar por salir adelante en la vida.

16. ___ La principal responsabilidad de los hijos es venerar regularmente sus antecesores en cada ocasión que tengan.

17. ___ El principal propósito del matrimonio no es perpetuar un apellido

18. ___ Los hijos deben tomar decisiones propias y no necesariamente tienen que pedir consejo a sus padres.
19. ___ Los hijos no tienen que respetar y querer las mismas personas que sus padres querían y respetaban.

20. ___ Después de llegar a la edad adulta, los hijos pueden gastar o decidir que hacer con el dinero ganado por ellos mismos producto de su trabajo aunque sus padres estén vivos.

21. ___ No hay peor falta que no ser filial

22. ___ Los hijos siempre deben obedecer a los padres sin importar nada más.
Appendix 10 - The COPE Scale
The COPE

We are interested in how people respond when they confront difficult or stressful events in their lives. There are lots of ways to try to deal with stress. This questionnaire asks you to indicate what you generally do and feel, when you experience stressful events. Obviously, different events bring out somewhat different responses, but think about what you usually do when you are under a lot of stress.

Then respond to each of the following items by blackening one number on your answer sheet for each, using the response choices listed just below. Please try to respond to each item separately in your mind from each other item. Choose your answers thoughtfully, and make your answers as true FOR YOU as you can. Please answer every item. There are no "right" or "wrong" answers, so choose the most accurate answer for YOU--not what you think "most people" would say or do. Indicate what YOU usually do when YOU experience a stressful event.

1 = I usually don't do this at all
2 = I usually do this a little bit
3 = I usually do this a medium amount
4 = I usually do this a lot

1. I try to grow as a person as a result of the experience.
2. I turn to work or other substitute activities to take my mind off things.
3. I get upset and let my emotions out.
4. I try to get advice from someone about what to do.
5. I concentrate my efforts on doing something about it.

6. I say to myself "this isn't real."

7. I put my trust in God.

8. I laugh about the situation.

9. I admit to myself that I can't deal with it, and quit trying.

10. I restrain myself from doing anything too quickly.

11. I discuss my feelings with someone.

12. I use alcohol or drugs to make myself feel better.

13. I get used to the idea that it happened.

14. I talk to someone to find out more about the situation.

15. I keep myself from getting distracted by other thoughts or activities.

16. I daydream about things other than this.

17. I get upset, and am really aware of it.

18. I seek God's help.

19. I make a plan of action.

20. I make jokes about it.

21. I accept that this has happened and that it can't be changed.

22. I hold off doing anything about it until the situation permits.

23. I try to get emotional support from friends or relatives.

24. I just give up trying to reach my goal.

25. I take additional action to try to get rid of the problem.

26. I try to lose myself for a while by drinking alcohol or taking drugs.
27. I refuse to believe that it has happened.
28. I let my feelings out.
29. I try to see it in a different light, to make it seem more positive.
30. I talk to someone who could do something concrete about the problem.
31. I sleep more than usual.
32. I try to come up with a strategy about what to do.
33. I focus on dealing with this problem, and if necessary let other things slide a little.
34. I get sympathy and understanding from someone.
35. I drink alcohol or take drugs, in order to think about it less.
36. I kid around about it.
37. I give up the attempt to get what I want.
38. I look for something good in what is happening.
39. I think about how I might best handle the problem.
40. I pretend that it hasn't really happened.
41. I make sure not to make matters worse by acting too soon.
42. I try hard to prevent other things from interfering with my efforts at dealing with this.
43. I go to movies or watch TV, to think about it less.
44. I accept the reality of the fact that it happened.
45. I ask people who have had similar experiences what they did.
46. I feel a lot of emotional distress and I find myself expressing those feelings a lot.
47. I take direct action to get around the problem.
48. I try to find comfort in my religion.

49. I force myself to wait for the right time to do something.

50. I make fun of the situation.

51. I reduce the amount of effort I'm putting into solving the problem.

52. I talk to someone about how I feel.

53. I use alcohol or drugs to help me get through it.

54. I learn to live with it.

55. I put aside other activities in order to concentrate on this.

56. I think hard about what steps to take.

57. I act as though it hasn't even happened.

58. I do what has to be done, one step at a time.

59. I learn something from the experience.

60. I pray more than usual.
Appendix 11 - The COPE Scale in Spanish
COPE en Español

Estamos interesados en la forma en que las personas responden cuando tienen que hacer frente a acontecimientos difíciles o estresantes en sus vidas. Hay muchas formas de intentar manejar el estrés. Este cuestionario le pide que indique lo que generalmente hace y siente cuando experimenta acontecimientos estresantes. Obviamente, sucesos diferentes provocan respuestas algo diferentes, pero piense acerca de lo que hace habitualmente cuando está bajo un estrés intenso. Entonces responda a cada uno de los siguientes ítems marcando el número que corresponda, utilizando las opciones de respuesta que se presentan abajo. Por favor, intente responder a cada ítem de forma separada a los demás. Elija sus respuestas cuidadosamente, y responda de la forma más sincera que le sea posible. Por favor responda todos los ítems. No hay respuestas correctas o incorrectas, de modo que elija las respuestas que más se le ajuste a usted, no la que usted piense que la mayoría de la gente diría o haría. Indique lo que USTED habitualmente hace cuando experimenta un acontecimiento estresante.

1= No suelo hacer esto en absoluto
2= Suelo hacer esto un poco
3= Suelo hacer esto moderadamente
4= Suelo hacer esto mucho.

1. Intento desarrollarme como persona como resultado de la experiencia.
2. Me concentro en el trabajo u otras actividades sustitutivas para alejar el tema de mi mente.
3. Me altero y dejo aflorar mis emociones.
4. Intento conseguir consejo de alguien sobre qué hacer.
5. Concentro mis esfuerzos en hacer algo acerca de la situación.
6. Me digo a mí mismo: “Esto no es real”
7. Confío en Dios.
8. Me río acerca de la situación.
9. Admito que no puedo con ello y dejo de intentarlo.
10. Me disuado a mí mismo de hacer algo con demasiada rapidez.
11. Hablo de mis sentimientos con alguien.
12. Consumo alcohol o drogas para sentirme mejor.
13. Me acostumbro a la idea de lo que sucedió.
14. Hablo con alguien para saber más acerca de la situación.
15. Evito distraerme con otros pensamientos o actividades.
16. Sueño despierto con otras cosas diferentes.
17. Me altero y soy realmente consciente de la situación.
18. Pido la ayuda de Dios.
19. Hago un plan de acción.
20. Hago bromas sobre la situación.
21. Acepto que ha sucedido y que no puede cambiarse.
22. Demoro hacer algo sobre el tema hasta que la situación lo permita.
23. Intento conseguir apoyo emocional de amigos o familiares.
24. Simplemente abandono en el intento de lograr mi objetivo.
25. Tomo medidas adicionales para intentar librarme del problema.
26. Intento evadirme un rato bebiendo alcohol o tomando drogas.

27. Me niego a creer que haya sucedido.


29. Intento verlo de una forma diferente, para que parezca más positivo.

30. Hablo con alguien que pudiera hacer algo concreto acerca del problema.

31. Duermo más de lo habitual.

32. Intento encontrar una estrategia acerca de qué hacer.

33. Me concentro en el manejo del problema y si es necesario aparto otros temas un poco.

34. Consigo la compasión y comprensión de alguien.

35. Bebo alcohol o tomo drogas para pensar menos en ello.

36. Bromeo sobre ello.

37. Renuncio a intentar lograr lo que quiero.

38. Busco algo bueno en lo que está sucediendo.

39. Pienso acerca de cómo podría manejar mejor el problema.

40. Actúo como si realmente no hubiera sucedido.

41. Me aseguro de no empeorar las cosas por actuar demasiado pronto.

42. Intento evitar que otras cosas interfieran con mis esfuerzos de manejar la situación.

43. Voy al cine o veo la televisión para pensar menos en ello.

44. Acepto la realidad del hecho que ha sucedido.

45. Pregunto a personas que han tenido experiencias similares qué hicieron.

46. Siento un gran malestar emocional y me encuentro expresando estos sentimientos un montón.
47. Llevo a cabo una acción directa en torno al problema.

48 Intento encontrar consuelo en la religión.

49. Me obligo a mi mismo a esperar el momento oportuno para hacer algo.

50. Hago bromas de la situación.

51. Reduzco la cantidad de esfuerzo que dedico a resolver el problema.

52. Hablo con alguien acerca de cómo me siento

53. Utilizo alcohol o drogas para ayudarme a superarlo.

54. Aprendo a vivir con ello.

55. Dejo de lado otras actividades para concentrarme en el problema

56. Pienso profundamente acerca de qué pasos tomar.

57. Actúo como si nunca hubiera ocurrido.

58. Hago lo que hay que hacer, paso a paso.

59. Aprendo algo de la experiencia.

60. Rezo más de lo habitual.
Appendix 12 - The Burden Interview
ZARIT BURDEN INTERVIEW

Instructions: The following is a list of statements, which reflect how people sometimes feel when taking care of another person. After each statement, indicate how often you feel that way, never, rarely, sometimes, quite frequently, or nearly always. There are no right or wrong answers.

1) Do you feel that your relative asks for more help than s/he needs?
   Never Rarely Sometimes Quite Frequently Nearly Always
   0 ( ) 1 ( ) 2 ( ) 3 ( ) 4 ( )

2) Do you feel that because of the time you spend with your relative, you don’t have enough time for myself?
   Never Rarely Sometimes Quite Frequently Nearly Always
   0 ( ) 1 ( ) 2 ( ) 3 ( ) 4 ( )

3) Do you feel stressed between caring for your relative and trying to meet other responsibilities for your family or work?
   Never Rarely Sometimes Quite Frequently Nearly Always
   0 ( ) 1 ( ) 2 ( ) 3 ( ) 4 ( )

4) Do you feel embarrassed over your relative’s behavior?
   Never Rarely Sometimes Quite Frequently Nearly Always
   0 ( ) 1 ( ) 2 ( ) 3 ( ) 4 ( )

5) Do you feel angry when you are around your relative?
   Never Rarely Sometimes Quite Frequently Nearly Always
   0 ( ) 1 ( ) 2 ( ) 3 ( ) 4 ( )
6) Do you feel that your relative currently affects your relationship with other family members or friends in a negative way?

<table>
<thead>
<tr>
<th>Frequency Levels</th>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

7) Are you afraid what the future holds for you relative?

<table>
<thead>
<tr>
<th>Frequency Levels</th>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

8) Do you feel your relative is dependent upon you?

<table>
<thead>
<tr>
<th>Frequency Levels</th>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

9) Do you feel strained when you are around your relative?

<table>
<thead>
<tr>
<th>Frequency Levels</th>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

10) Do you feel your health has suffered because of you involvement with your relative?

<table>
<thead>
<tr>
<th>Frequency Levels</th>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

11) Do you feel that you don’t have as much privacy as you would like because of your relative?

<table>
<thead>
<tr>
<th>Frequency Levels</th>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>
12) Do you feel that your social life has suffered because you are caring for your relative?

Never  Rarely  Sometimes  Quite Frequently  Nearly Always
0 ( )  1 ( )  2 ( )  3 ( )  4 ( )

13) Do you feel uncomfortable about having friends over because of your relative?

Never  Rarely  Sometimes  Quite Frequently  Nearly Always
0 ( )  1 ( )  2 ( )  3 ( )  4 ( )

14) Do you feel that your relative seems to expect you to take care of him/her as if you were the only one he/she could depend on?

Never  Rarely  Sometimes  Quite Frequently  Nearly Always
0 ( )  1 ( )  2 ( )  3 ( )  4 ( )

15) Do you feel that you don’t have enough money to care for your relative, in addition to the rest of your expenses?

Never  Rarely  Sometimes  Quite Frequently  Nearly Always
0 ( )  1 ( )  2 ( )  3 ( )  4 ( )

16) Do you feel that you will be unable to take care of your relative much longer?

Never  Rarely  Sometimes  Quite Frequently  Nearly Always
0 ( )  1 ( )  2 ( )  3 ( )  4 ( )

17) Do you feel you have lost control of your life since your relative’s illness?

Never  Rarely  Sometimes  Quite Frequently  Nearly Always
0 ( )  1 ( )  2 ( )  3 ( )  4 ( )
18) Do you wish you could just leave the care of your relative to someone else?

<table>
<thead>
<tr>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

19) Do you feel uncertain about what to do about your relative?

<table>
<thead>
<tr>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

20) Do you feel you should be doing more for your relative?

<table>
<thead>
<tr>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

21) Do you feel you could do a better job in caring for your relative?

<table>
<thead>
<tr>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>

22) Overall, how burdened do you feel in caring for your relative?

<table>
<thead>
<tr>
<th>Never</th>
<th>Rarely</th>
<th>Sometimes</th>
<th>Quite Frequently</th>
<th>Nearly Always</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 ( )</td>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
</tr>
</tbody>
</table>
Appendix 13 - The Burden Interview in Spanish
CUESTIONARIO DE SOBRECARGA DEL CUIDADOR

(Escala de Zarit)

INSTRUCCIONES: A continuación se presentan una lista de frases que reflejan como se sienten algunas personas cuando cuidan a otra persona. Después de leer cada frase, indique con qué frecuencia se siente usted de esa manera, escogiendo entre Nunca, Casi Nunca, A Veces, Frecuentemente y Casi Siempre. No existen respuestas correctas o incorrectas.

CON QUE FRECUENCIA (rodee con un círculo la opción elegida)

1) ¿Con que frecuencia siente usted que su familiar/paciente solicita más ayuda de la que realmente necesita?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

2) ¿Con que frecuencia siente usted que, a causa del tiempo que gasta con su familiar/paciente, ya no tiene tiempo suficiente para usted mismo?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

3) ¿Con que frecuencia se siente estresada(o) al tener que cuidar a su familiar/paciente y tener además que atender otras responsabilidades? (Ej: con su familia o en el trabajo)

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>
4) ¿Con qué frecuencia se siente avergonzada(o) por el comportamiento de su familiar/paciente?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

5) ¿Con qué frecuencia se siente irritada(o) cuando está cerca de su familiar/paciente?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

6) ¿Con qué frecuencia cree que la situación actual afecta a su relación con amigos u otros miembros de su familia de una forma negativa?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

7) ¿Con qué frecuencia siente temor por el futuro que le espera a su familiar/paciente?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

8) ¿Con qué frecuencia siente que su familiar/paciente depende de usted?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>
9) ¿Con qué frecuencia se siente agotada(o) cuando tiene que estar junto a su familiar/paciente?

<table>
<thead>
<tr>
<th></th>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>()</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
</tr>
</tbody>
</table>

10) ¿Con qué frecuencia siente usted que su salud se ha visto afectada por tener que cuidar a su familiar/paciente?

<table>
<thead>
<tr>
<th></th>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>()</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
</tr>
</tbody>
</table>

11) ¿Con qué frecuencia siente que no tiene la vida privada que desearía a causa de su familiar/paciente?

<table>
<thead>
<tr>
<th></th>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>()</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
</tr>
</tbody>
</table>

12) ¿Con qué frecuencia Siente cree que sus relaciones sociales se han visto afectadas por tener que cuidar a su familiar/paciente?

<table>
<thead>
<tr>
<th></th>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>()</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
</tr>
</tbody>
</table>

13) (SÓLO SI EL ENTREVISTADO VIVE CON EL PACIENTE). ¿Con qué frecuencia siente se siente incómoda(o) para invitar amigos a casa, a causa de su familiar/paciente?

<table>
<thead>
<tr>
<th></th>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>()</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
<td>( )</td>
</tr>
</tbody>
</table>
14) ¿Con qué frecuencia cree que su familiar/paciente espera que usted le cuide, como si fuera la única persona con la que pudiera contar?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frequentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (    )</td>
<td>2 (    )</td>
<td>3 (    )</td>
<td>4 (    )</td>
<td>5 (    )</td>
</tr>
</tbody>
</table>

15) ¿Con qué frecuencia cree usted que no dispone de dinero suficiente para cuidar de su familiar/paciente, además de sus otros gastos?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frequentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (    )</td>
<td>2 (    )</td>
<td>3 (    )</td>
<td>4 (    )</td>
<td>5 (    )</td>
</tr>
</tbody>
</table>

16) ¿Con qué frecuencia siente que no va a ser capaz de cuidar de su familiar/paciente durante mucho más tiempo?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frequentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (    )</td>
<td>2 (    )</td>
<td>3 (    )</td>
<td>4 (    )</td>
<td>5 (    )</td>
</tr>
</tbody>
</table>

17) ¿Con qué frecuencia siente que ha perdido el control sobre su vida desde que la enfermedad de su familiar/paciente se manifestó?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frequentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (    )</td>
<td>2 (    )</td>
<td>3 (    )</td>
<td>4 (    )</td>
<td>5 (    )</td>
</tr>
</tbody>
</table>

18) ¿Con qué frecuencia desearía poder encargar el cuidado de su familiar/paciente a otra persona?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frequentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (    )</td>
<td>2 (    )</td>
<td>3 (    )</td>
<td>4 (    )</td>
<td>5 (    )</td>
</tr>
</tbody>
</table>
19) ¿Con qué frecuencia se siente insegura(o) acerca de lo que debe hacer con su familiar/paciente?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

20) ¿Con qué frecuencia siente que debería hacer más de lo que hace por su familiar/paciente?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

21) ¿Con qué frecuencia cree que podría cuidar a su familiar/paciente mejor de lo que lo hace?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>

22) En general, ¿con qué frecuencia se siente muy sobrecargada(o) al tener que cuidar de su familiar/paciente?

<table>
<thead>
<tr>
<th>Nunca</th>
<th>Casi Nunca</th>
<th>A Veces</th>
<th>Frecuentemente</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( )</td>
<td>2 ( )</td>
<td>3 ( )</td>
<td>4 ( )</td>
<td>5 ( )</td>
</tr>
</tbody>
</table>
Appendix 14 - The Satisfaction with Life Scale
Satisfaction with Life Scale

Instructions for administering the scale are: Below are five statements with which you may agree or disagree. Using the 1-7 scale below, indicate your agreement with each item by placing the appropriate number on the line preceding that item. Please be open and honest in your responding.

The 7-point scale is: 1 = strongly disagree, 2 = disagree, 3 = slightly disagree, 4 = neither agree nor disagree, 5 = slightly agree, 6 = agree, 7 = strongly agree.

_____  1. In most ways my life is close to my ideal.
_____  2. The conditions of my life are excellent.
_____  3. I am satisfied with my life.
_____  4. So far I have gotten the important things I want in life.
_____  5. If I could live my life over, I would change almost nothing.
Appendix 15 - The Satisfaction with Life Scale in Spanish
Escales sobre la Satisfaccion con La Vida

Instrucciones para la administración de la escala son: a continuación se presentan cinco declaraciones con las que usted puede estar en acuerdo o en desacuerdo. Utilizando la escala de 1-7 a continuación, indique su acuerdo con cada elemento, colocando el número apropiado en la línea anterior a ese tema. Por favor, sea abierto(a) y honesto(a) en su respuesta.

La escala de 7 puntos: 1 = totalmente en desacuerdo, 2 = desacuerdo, 3 = ligeramente en desacuerdo, 4 = ni de acuerdo ni en desacuerdo, 5 = ligeramente de acuerdo, 6 = acuerdo, 7 = muy de acuerdo.

1. El tipo de vida que llevo se parece al tipo de vida que siempre soñé llevar.
2. Las condiciones de mi vida son excelentes.
3. Estoy satisfecho con mi vida.
4. Hasta ahora he obtenido las cosas importantes que quier en la vida.
5. Si pudiera vivir mi vida de nuevo, me gustaría que todo volviese a ser igual.
Appendix 16 - Center for Epidemiological Studies - Depression
### Exhibit 6.3

*Note: Items 4, 8, 12, and 16 have their scores reversed before totaling.*

**Instructions for questions:** Below is a list of the ways you might have felt or behaved. Please tell me how often you felt this way during the past week.

<table>
<thead>
<tr>
<th>During the past week:</th>
<th>Rarely or none of the time (less than 1 day)</th>
<th>Some or a little of the time (1–2 days)</th>
<th>Occasionally or a moderate amount of the time (3–4 days)</th>
<th>Most or all of the time (5–7 days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. I was bothered by things that usually don't bother me</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>2. I did not feel like eating; my appetite was poor</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>3. I felt that I could not shake off the blues even with help from my family or friends</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>4. I felt that I was just as good as other people</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>5. I had trouble keeping my mind on what I was doing</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>6. I felt depressed</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>7. I felt that everything I did was an effort</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>8. I felt hopeful about the future</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>9. I thought my life had been a failure</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>10. I felt fearful</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>11. My sleep was restless</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>12. I was happy</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>13. I talked less than usual</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>14. I felt lonely</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>15. People were unfriendly</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>16. I enjoyed life</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>17. I had crying spells</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>18. I felt sad</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>19. I felt that people dislike me</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>20. I could not get &quot;going&quot;</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Durante la semana pasada</th>
<th>Raramente o ninguna de las veces (menos de 1 día)</th>
<th>Alguno o poco tiempo (1-2 días)</th>
<th>Ocasionalmente o una cantidad moderada de tiempo (3-4 días)</th>
<th>La mayoría o todo del tiempo (5-7 días)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Me molesté por cosas que usualmente no me molestan</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>2. No tenía ganas de comer, mi apetito fue pobre</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>3. Me sentí que no podía despejarme de mi mal estado de ánimo aún con la ayuda de mi familia o amigos/as</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>4. Sentí que era tan bueno/a como otras personas</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>5. Tuve problemas manteniendo mi mente en lo que estaba haciendo</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>6. Me sentí depresivo/a</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>7. Sentí que todo lo que hacía era un esfuerzo</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>8. Me sentí esperanzado hacia el futuro</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>9. Pensé que mi vida había sido un fracaso</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>10. Me sentí atemorizado</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>11. Mi sueño no tenía descanso</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>12. Estuve feliz</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>13. Hablé menos de lo usual</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>14. Me sentí solitario/solitaria</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>15. Las personas no eran amistosas</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>16. Disfruté la vida</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>17. Tuve períodos de llanto</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>18. Me sentí triste</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>19. Me sentí que no le caía bien a las personas</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>20. No podía “progresar/avanzar”</td>
<td>0</td>
<td>1</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>
Appendix 18 - State Trait Anxiety Form Y1
# SELF-EVALUATION QUESTIONNAIRE

Developed by Charles D. Spielberger
in collaboration with
R. L. Gorsuch, R. Luzzette, P. R. Vagg, and G. A. Jacobs

**STAI Form Y-1**

Name ___________________________________________ Date __________ S __
Age ________ Sex: M ______ F ______ T ______

DIRECTIONS: A number of statements which people have used to describe themselves are given below. Read each statement and then blacken in the appropriate circle to the right of the statement to indicate how you feel right now, that is, *at this moment*. There are no right or wrong answers. Do not spend too much time on any one statement but give the answer which seems to describe your present feelings best.

<table>
<thead>
<tr>
<th></th>
<th>NOT AT ALL</th>
<th>SEMI-MIDLY</th>
<th>MODERATELY</th>
<th>VERY MUCH</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. I feel calm</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>2. I feel secure</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>3. I am tense</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>4. I feel strained</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>5. I feel at ease</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>6. I feel upset</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>7. I am presently worrying over possible misfortunes</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>8. I feel satisfied</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>9. I feel frightened</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>10. I feel comfortable</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>11. I feel self-confident</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>12. I feel nervous</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>13. I am jittery</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>14. I feel indecisive</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>15. I am relaxed</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>16. I feel content</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>17. I am worried</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>18. I feel confused</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>19. I feel steady</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
<tr>
<td>20. I feel pleasant</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
<td>[ ] [ ] [ ] [ ]</td>
</tr>
</tbody>
</table>
Appendix 19 - State Trait Anxiety Form Y1 Spanish
# Questionario de Evaluación Propia

Creado por Charles D. Spielberger  

*en colaboración con*  

R. L. Gorsuch, R. Lushene, P. R. Vagg, and G. A. Jacobs  

**STAI Forma Y-1**

<table>
<thead>
<tr>
<th>Nombre</th>
<th>Fecha</th>
<th>S</th>
</tr>
</thead>
<tbody>
<tr>
<td>Edad</td>
<td>Sexo: H</td>
<td>M</td>
</tr>
</tbody>
</table>

**DIRECCIONES:** Un número de declaraciones que las personas han usado para describirse a sí mismo están abajo. Lea cada declaración y después llene el apropiado círculo a la derecha de la declaración para indicar como usted se siente *ahora*, es decir, *en este momento*.  

No hay respuestas correctas o incorrectas. No pierda mucho tiempo en ninguna de las declaraciones, pero dé la respuesta que parece describir mejor sus sentimientos en este momento.

<table>
<thead>
<tr>
<th>No</th>
<th>Del Todo</th>
<th>Algo</th>
<th>Moderadamente</th>
<th>Completamente Positivo</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>7</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>8</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>9</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>10</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>11</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>12</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>13</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>14</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>15</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>16</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>17</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>18</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>19</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td>20</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
</tbody>
</table>
Appendix 20 - State Trait Anxiety Form Y2
<table>
<thead>
<tr>
<th>Number</th>
<th>Statement</th>
<th>Scale</th>
</tr>
</thead>
<tbody>
<tr>
<td>21</td>
<td>I feel pleasant</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>22</td>
<td>I feel nervous and restless</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>23</td>
<td>I feel satisfied with myself</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>24</td>
<td>I wish I could be as happy as others seem to be</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>25</td>
<td>I feel like a failure</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>26</td>
<td>I feel rested</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>27</td>
<td>I am &quot;calm, cool, and collected&quot;</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>28</td>
<td>I feel that difficulties are piling up so that I cannot overcome them</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>29</td>
<td>I worry too much over something that really doesn't matter</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>30</td>
<td>I am happy</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>31</td>
<td>I have disturbing thoughts</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>32</td>
<td>I lack self-confidence</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>33</td>
<td>I feel secure</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>34</td>
<td>I make decisions easily</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>35</td>
<td>I feel inadequate</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>36</td>
<td>I am content</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>37</td>
<td>Some unimportant thought runs through my mind and bothers me</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>38</td>
<td>I take disappointments so keenly that I can't put them out of my mind</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>39</td>
<td>I am a steady person</td>
<td>1 2 3 4</td>
</tr>
<tr>
<td>40</td>
<td>I get in a state of tension or turmoil as I think over my recent concerns and interests</td>
<td>1 2 3 4</td>
</tr>
</tbody>
</table>
Appendix 21 - State Trait Anxiety Form Y2 Spanish
Questionario de Evaluación Propia
STAI Forma Y-2

Nombre ___________________________________________ Fecha __________________

DIRECCIONES: Un número de declaraciones que las personas han
usado para describirse a sí mismo están abajo. Lea cada declaración
y después llene el apropiado círculo a la derecha de la declaración
para indicar cómo usted se siente _generalmente_. No hay respuestas
correctas o incorrectas. No pierda mucho tiempo en ninguna de las
declaraciones, pero dé la respuesta que parece describir como usted
generalmente se siente.

<table>
<thead>
<tr>
<th></th>
<th>Casi Nunca</th>
<th>Algunas Veces</th>
<th>Amenudo</th>
<th>Casi Siempre</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>21. Me siento complacido/a…………………………………….1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>22. Me siento nervioso y sin descanso…………………………1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>23. Me siento satisfecho/a de mi mismo/a…………………………1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>24. Yo desearía ser tan feliz como otros parecen ser………………1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>25. Me siento como un fracaso……………………………………1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>26. Me siento cansado/a…………………………………………1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>27. Yo estoy “calmado, sereno y sereno/a de mi mismo/a”…1</td>
<td>2</td>
<td>3</td>
<td>4</td>
<td></td>
</tr>
</tbody>
</table>
| 28. Siento que las dificultades se van amontonando y yo
  no puedo superarlas…………………………………………1 | 2 | 3 | 4 |
| 29. Me preocupo demasiado sobre algo que realmente
  no tiene importancia………………………………………………1 | 2 | 3 | 4 |
| 30. Estoy feliz…………………………………………………………1 | 2 | 3 | 4 |
| 31. Tengo pensamientos que me perturban…………………………1 | 2 | 3 | 4 |
| 32. Me falta confianza en mí mismo/a…………………………..1 | 2 | 3 | 4 |
| 33. Me siento seguro/a……………………………………………1 | 2 | 3 | 4 |
| 34. Tomo decisiones fácilmente……………………………………1 | 2 | 3 | 4 |
| 35. Me siento inadecuado/a……………………………………..1 | 2 | 3 | 4 |
| 36. Estoy contento/a………………………………………………1 | 2 | 3 | 4 |
| 37. Algunos pensamientos sin importancia pasan por
  mi mente y me molestan………………………………………………1 | 2 | 3 | 4 |
| 38. Tomo los disgustos tan seriamente que no los
  puedo sacar de mi mente…………………………………………1 | 2 | 3 | 4 |
| 39. Soy una persona estable………………………………………………1 | 2 | 3 | 4 |
| 40. Me encuentro en un estado de tensión y desorden
  según pienso en mis recientes preocupaciones e
  intereses……………………………………………………………1 | 2 | 3 | 4 |

163
Appendix 22 - Debriefing Form
Debriefing Form

Thank you for participating in the present study which analyzes the interaction between demographics, caregiver burden, and coping styles in elderly caregivers. This study tests the linkages among demographic factors (such as age, gender, and ethnicity), perceived caregiver burden, coping with caregiving and their impact on life satisfaction.

Once again, you participation is greatly appreciated. If you know of anyone who is eligible for the present study, we ask that you do not discuss it with them until they have had a chance to participate as well. Any knowledge of the questions asked, prior to the interview, may change how a person responds. Thank you for your cooperation.

If you have any questions regarding this study, please feel free to ask the researcher at this time. You may also contact the researcher, Didiana De La Osa, by email at didiana.delaosa@yahoo.com or by phone at 305-878-9818. If you would like to talk with someone about your rights of being a subject in this research study or about ethical issues with this research study, you may contact the FIU Office of Research Integrity by phone at 305-348-2494 or by email at ori@fiu.edu.

If you would like to learn more about the topic of this research study, you may consult:


In the event that you feel psychologically distressed by participation in this study, we encourage you to call and schedule an appointment for psychological treatment at:

THE GOODMAN CENTER FOR PSYCHOLOGICAL SERVICES
2173 NW 99TH AVE
MIAMI, FL 33172
305-592-7860

NORTH MIAMI FOUNDATION FOR SENIOR CITIZENS
620 NE 127TH STREET
NORTH MIAMI, FL 33161
305-893-1450

JACKSON NORTH COMMUNITY MENTAL HEALTH CENTER
20201 N.W. 37TH AVENUE
MIAMI, FL 33056
786-466-2700

14701 N.W. 27TH AVENUE
OPA-LOCKA, FL 33054
786-466-2700

CRISIS STABILIZATION UNIT (24-HOUR)
(786) 466-2834

ADULT CASE MANAGEMENT AND PSYCHOSOCIAL REHABILITATION SERVICES
(786) 466-1340
Appendix 23 - Debriefing Form in Spanish
Gracias por participar en el estudio actual que analiza la relación entre factores demográficos, la sobrecarga del cuidador, y los estilos de manejar el esfuerzo y tensión en cuidadores de ancianos. Este estudio pone a prueba los vínculos entre los factores demográficos (tales como edad, género y etnia), carga percibida del cuidador, y formas de hacerle frente a los cuidados y su impacto en la satisfacción con la vida.

De nuevo se le agradece su participación. Si usted sabe de alguien que es elegible para este estudio, le pedimos que no lo hable con ellos sobre los questionarios hasta que hayan tenido la oportunidad de participar también. Conocimiento de las preguntas antes de la entrevista puede cambiar la forma en que una persona responde. Gracias por su cooperación.

Si usted tiene alguna pregunta relacionada con este estudio, por favor no dude en preguntar al investigador en este momento. También puede comunicarse con el investigador, Didiana De La Osa, por correo electrónico a didiana.delaosa@yahoo.com, o por teléfono al 305-878-9818. Si desea hablar con alguien acerca de sus derechos de ser un participante de este estudio de investigación o sobre los aspectos éticos con este estudio de investigación, puede comunicarse con la Oficina de Integridad de la Investigación de FIU por teléfono al 305-348-2494 o por correo electrónico a Ori @fiu.edu.

Si desea obtener más información sobre el tema de esta investigación, puede consultar:


En caso de que usted se siente psicológicamente angustiados por la participación en este estudio, le animamos a que llame y haga una cita para tratamiento psicológico en:

THE GOODMAN CENTER FOR PSYCHOLOGICAL SERVICES
2173 NW 99TH AVE 
MIAMI, FL 33172
305-592-7860

NORTH MIAMI FOUNDATION FOR SENIOR CITIZENS
620 NE 127TH STREET
NORTH MIAMI, FL 33161
305-893-1450

JACKSON NORTH COMMUNITY MENTAL HEALTH CENTER
20201 N.W. 37TH AVENUE  14701 N.W. 27TH AVENUE
MIAMI, FL 33056  OPA-LOCKA, FL 33054
786-466-2700  786-466-2700

UNIDAD DE ESTABILIZACIÓN DE CRISIS (24-HORAS)
(786) 466-2834

MANEJO DE CASOS DE ADULTOS Y SERVICIOS DE REHABILITACIÓN PSICOSOCIAL
(786) 466-1340