Chronic inflammation and quality of life in older adults: a cross-sectional study using biomarkers to predict emotional and relational outcomes

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Abstract

Background: This study explores relationships between chronic inflammation and quality of life, making a case for biopsychosocial modeling of these associations. It builds on research from social and clinical disciplines connecting chronic conditions, and inflammatory conditions specifically, to reduced quality of life.

Methods: Data from Wave I of the National Social Life, Health, and Aging Project are modeled using ordinal logistic and ordinary least-squares regression techniques. Inflammation is measured using C-reactive protein; quality of life is conceptualized as happiness with life overall as well as intimate relationships specifically.

Results: For most NSHAP participants, chronic inflammation significantly predicts lower odds of reporting high QoL on both emotional and relational measures. Social structural factors do not confound these associations. Inconsistent results for participants with very high (over 6 mg/L) CRP measurements suggest additional social influences.

Conclusions: Findings echo strong theoretical justification for investigating relationships between CRP and QoL in greater detail. Further research should explore possible mediation of these associations by sociomedical sequelae of chronic disease as well as social relationship dynamics. Elaboration is also needed on the mechanisms by which social disadvantage may cause chronic inflammation.

Keywords: Inflammation, Quality of life, Health, NSHAP, Chronic conditions, Quantitative
Theoretical foundations
The inflammatory biomarker C-reactive protein (CRP) has a strong evidence basis for use in sociomedical research. Bodies produce CRP in response to inflammation [1-3]. CRP gets deposited wherever swelling occurs—including joints, mucous membranes, and muscles [3]. This substance is generated when inflammation of any type is present [1,2]. Consequently, it can capture inflammation from a wide variety of health conditions, even if the person has no diagnoses [3]. While CRP levels can reflect a certain amount of acute inflammation, they are particularly good for capturing chronicity.

Acute inflammation from injury or pathogenic illness is unlikely to introduce significant differential bias. Elevations in CRP levels from acute inflammation tend to be not only temporary, but also small [3]. By contrast, baseline CRP levels are generally stable over time [4]. Average CRP values increase with age, but the relative distribution of values according to health status remains largely stable over time [3,4]. Likewise, while baseline CRP levels cannot be used to diagnose any particular disease [5], high levels do provide excellent evidence of generalized inflammation. Elevated levels can also indicate progression of inflammation in people with chronic conditions [3,5].

High levels of CRP circulating freely in the blood generally indicate widespread inflammation affecting multiple tissues [4]. Indeed, researchers have discovered that levels of freely circulating CRP are determined exclusively by the body's rate of synthesizing this substance [6]. Likewise, acute conditions are unlikely to elevate the synthesis rate substantially or for very long [4]. In fact, CRP levels often drop rapidly when synthesis stops [4]. CRP thus constitutes a good predictor of chronic inflammation for a variety of different disease populations, despite its inherent limitations.

Only a few published studies have explicitly focused on the relationship between CRP and QoL [7-9]. While these studies do not delve deeply into potential psychosocial dynamics of the association between CRP and QoL, they do find that elevated levels of CRP significantly predict diminished levels of QoL [7]. An excellent foundation thus exists for further work on CRP and QoL using generalized linear modeling techniques. Researchers in a variety of disciplines can now extend this foundation to more diverse outcomes that capture different elements of overall QoL. Scholars can also explore synergistic effects of biological and social mechanisms on QoL, and potentially unique dynamics of these phenomena in older adults. Given robust linkage in sociological literature between health status and self-rated QoL [10], a strong theoretical connection exists between CRP and QoL outcomes.

Although research on CRP specifically as a determinant of QoL remains largely absent from the literature, research on chronic inflammatory disease and its relationship to QoL has expanded greatly within the past 20 years. Several researchers have focused on general associations between specific inflammatory chronic conditions and health outcomes, finding significant associations between symptom severity and reductions in QoL [11,12]. Recent research also suggests dynamic relationships between inflammation and patient affect as a potential mediator of associations between chronic inflammation and QoL for a variety of health conditions [13-15].

Literature on specific inflammatory diseases largely mirrors these general findings concerning inflammatory disease and QoL. For example, literature on arthritis documents robust linkage between severity of inflammation and reductions in QoL, both acutely and over time [16-24]. This literature also includes research on: asthma [25]; interstitial cystitis [26-36]; irritable bowel syndrome [37]; inflammatory bowel disease [37]; chronic ulcerative colitis [38]; Crohn's disease [39,40]; non-specific bowel inflammation [41-44]; chronic prostatitis [45]; vulvodynia [46]; and non-specific genital inflammation [46]. These scholars have focused on a wide range of QoL outcomes, spanning from self-rated enjoyment of life to sexual satisfaction to economic livelihood.

Psychosocial mechanisms implicated in the relationship between chronic inflammation and reduced QoL include: adverse interactional dynamics [47,48]; internal stress processes [49,50]; emotional strain of coping with negative social consequences of illness [13,51]; physical, emotional, and sexual abuse [47,48]; depression [48]; sexual dysfunction [36]; pursuit of "health-related normality" [52]; fear of having to deal with their symptoms in public places or in visible ways [43,51]; feelings of isolation and alienation [53,54]; lack of social support from others [55]; frustration in seeking a diagnosis [50,56]. Comorbidities are common among people with chronic inflammation [57], and often exacerbate the negative consequences of inflammation for QoL [35].

Both the literature on generalized inflammation and the literatures on specific inflammatory diseases as predictors of QoL indicate that inflammation can cause changes in QoL. Extant research also suggests that stress can cause and/or exacerbate inflammation itself [49,50]. Consequently, people whose QoL is already low for reasons not related to inflammatory pathology may experience higher levels of inflammation if their life circumstances induce psychological distress. Overall extant literature indicates that inflammation can cause reductions in QoL. These reductions may in turn be exacerbated by psychological distress [8] and resultant increases in allostatic load [58], creating a vicious cycle of cumulative inequality in health over time [59].

As suggested previously, diagnosis remains highly problematic because many people with these diseases go
untreated, never receive an accurate diagnosis, or find that no clinically recognized diagnosis yet exists for their condition [50,56]. Likewise, many patients who do receive one diagnosis do not ever learn the full extent of their condition due to the siloized structure of specialty medicine in the United States. These issues may affect women disproportionately, compounding an already significant array of disadvantages in health [60,61]. These studies thus present a strong case for studying associations between CRP and QoL in lieu of using diagnosis information.

In studying these associations, scholars should note that chronic inflammation itself may reflect the influence of multiple social factors. This is especially true for older adults [62], as cumulative processes may manifest strongly [59]. Link and Phelan’s [63] review of advancements in fundamental causation theories supports both this argument and the idea that social factors may condition the impacts of inflammation on QoL, once present [63]. The authors illuminate the potential for both material and psychosocial resources to impact people’s ability to cope with disease.

Research on resource multiplication illustrates the potential for relatively small disadvantages in psychosocial resiliency to compound into major consequences for health and QoL. This reasoning again introduces a cumulation perspective that mirrors the “weathering” phenomenon observed in Gerominus and colleagues’ [58,60,61] work on physical health. Consequently, data on older adults may offer particular insight into the effects of chronic inflammation on QoL across the life course. Multiple research teams [62,64] have conducted meaningful inquiry into chronic inflammation among older adults using the National Social Life, Health, and Aging Project (NSHAP) dataset collected by Linda Waite and colleagues.

Methods
Data and subject selection
This study uses data from Wave I of the NSHAP. This biopsychosocial dataset provides information on physical, mental, and social health among United States residents aged 57 to 85. Consequently, the NSHAP is an excellent resource for interdisciplinary scholars of health and QoL in older adults [65]. This dataset allows researchers to explore the synergistic relationships between biological, psychological, and social factors that may influence health status and perceived quality of life.

The NSHAP dataset was developed between 2005 and 2006 by researchers at the University of Chicago. As such, the NSHAP includes data on 3,005 individuals living in and around Chicago, Illinois. Oversampling of specific demographic groups (African Americans, Latinos, males, and persons 75 to 85 years of age) allows the dataset to be nationally representative [66]. Data for the NSHAP are collected via a combination of in-home interviews, supplemental questionnaires, blood spot blotters, and saliva cultures. As of Summer 2014 the NSHAP remains essentially cross-sectional, with only two waves of data available.

Life course sociology literature increasingly suggests that disadvantage in psychosocial outcomes accumulates over time from adverse social experience [67-73]. While the NSHAP data remain largely cross-sectional at present, the inclusion criteria guarantee at least 57 years elapsed between birth and data collection for a given subject. Analyzing an older population cross-sectionally allows researchers to capture long-term impacts from biosocial exposures that began in early life [67,68]. Consequently, the NSHAP can illuminate potential cumulative disadvantage in health and QoL from chronic inflammation due to inherited conditions.

QoL was examined in two different ways: emotionally (via self-reports of general happiness) and relationally (via self-reports of overall happiness with intimate relationships). Extant research suggests that relevant biological and social predictors may not influence all domains of QoL equally [74]. In addition, factor analysis for the two QoL outcomes yielded relatively high values for uniqueness, suggesting that discrete information about QoL was captured by each of the two measures. Assessing different types of QoL thus facilitated broader inference about the potential impact of chronic inflammation on these outcomes, and the social factors that may contribute to these patterns.

The main predictor for both QoL outcomes was CRP. This biomarker has a strong evidence basis supporting its use in biosocial research on inflammation. CRP also provides the best evidence of chronic inflammation among available variables in the NSHAP. In assessing the potential influence of CRP on QoL, the possible role of numerous social covariates was also examined as outlined in Table 1. These covariates were considered as both potential confounders of relationships between inflammation and QoL, and fundamental causes of inflammation itself.

Using correlational tables, all variables were checked for unique value as covariates—i.e., low collinearity. Certain covariates (e.g., feeling able to communicate openly with intimate partners) also correlated somewhat with one or both outcome variables. However, this did not present a methodological problem for two reasons. First, the NSHAP data documentation explains that the outcome variable measuring happiness with intimate relationships captures a variety of factors that may influence this perception, including emotional satisfaction, physical satisfaction, quality of communication, feelings of being able to rely on another person, and a variety of other elements [66]. Second, one of the goals of this study was to generate
hypotheses about potential social mediators of observed associations between CRP and QoL.

Correlational tables were also used to check for differential attrition owing to the lack of successful CRP measurements for about 35% of NSHAP participants. Challenges with inference related to the original measurement strategies in the NSHAP were also noted. For example, the dataset captures information on sex only as a binary construct. It also does not (despite calling its sex variable “gender”) collect any information on gender identity. Assessing possible effects from ethnoracial background presented similar challenges. Despite the NSHAP’s nationally representative sampling strategy, data could only be used for participants whose CRP serolevels had been successfully measured. Analyses related to race thus focused primarily on the potential impact of identifying as a member of any racial minority group.

As suggested above, some variables in the NSHAP may have problematic values due to measurement error. This concern is most pressing for the CRP variable. Although most values are clustered within a range of 10 mg/L, some values are closer to 50 and one person appears to have a value of 100. Comparing the CRP variable’s median value (1.531 mg/L) to its mean value (3.19 mg/L) suggests that outliers may be a problem for this measure. Since CRP levels usually remain in the single digits even for people with severe inflammation, these very high values are more likely attributable to

| Table 1 Descriptions and summary statistics for included variables |
|-----------------------------|-----------------|----------------|-----------------|-----------------|-----------------|
| Construct                  | Variable        | Units          | Cases | Mean  | St. Dev. | Min. | Max. |
| Quality of life            | Generally feeling happy | Points      | 2,995 | 3.58  | 0.88     | 1    | 5   |
|                            | Happiness with relationships |          | 2,890 | 5.89  | 1.61     | 1    | 7   |
| Chronic inflammation       | C-reactive protein | mg/L        | 1,939 | 3.20  | 6.03     | 0    | 100 |
| Sex                        | Female          | Yes/No        | 3,005 | 0.52  | 0.50     | 0    | 1   |
|                            | Male            |              |      | 0.48  | 0.50     |      |     |
| Age                        | Years of age    | Years         | 3,005 | 69.3  | 7.85     | 57   | 85  |
|                            | 57 to 64        | Yes/No        | 3,005 | 0.34  | 0.47     | 0    | 1   |
|                            | 65 to 74        |              |      | 0.36  | 0.48     |      |     |
|                            | 75 to 85        |              |      | 0.30  | 0.46     |      |     |
| Education                  | Years of education | Years      | 2,971 | 12.48 | 4.10     | 0    | 32  |
|                            | Less than high school | Yes/No   | 3,005 | 0.23  | 0.42     | 0    | 1   |
|                            | High school or equivalent |         |      | 0.26  | 0.44     |      |     |
|                            | Some college    |              |      | 0.28  | 0.45     |      |     |
|                            | Finished college |              |      | 0.22  | 0.41     |      |     |
| Race                       | White           | Yes/No        | 2,993 | 0.70  | 0.46     | 0    | 1   |
|                            | Any minority    |              |      | 0.30  | 0.46     |      |     |
|                            | Black           |              |      | 0.17  | 0.38     |      |     |
|                            | Hispanic        |              |      | 0.10  | 0.30     |      |     |
|                            | Other race      |              |      | 0.02  | 0.15     |      |     |
| Income                     | Household earnings | $/year      | 2,124 | 51,264 | 76,559  | 0    | 1,800,000 |
| Social support and communication | Open with partner | Points     | 2,012 | 2.71  | 0.54     | 1    | 3   |
|                            | Open with friends |              | 2,704 | 2.02  | 0.73     |      |     |
|                            | Relying on family | Points      | 2,793 | 2.57  | 0.65     | 1    | 3   |
|                            | Relying on friends |            | 2,680 | 2.30  | 0.71     |      |     |
| Relationship status        | Married         | Yes/No        | 3,005 | 0.60  | 0.49     | 0    | 1   |
|                            | Cohabitating    |              |      | 0.02  | 0.14     |      |     |
|                            | Separated       |              |      | 0.02  | 0.13     |      |     |
|                            | Divorced        |              |      | 0.11  | 0.31     |      |     |
|                            | Widowed         |              |      | 0.22  | 0.41     |      |     |
|                            | Never married   |              |      | 0.04  | 0.19     |      |     |
measurement issues. Indeed, data documentation for the NSHAP suggests that most implausible measurements on CRP owe to issues with data collection and processing.

This study was approved by the Florida State University Human Subjects Committee. All data management and analysis was conducted using Stata Version 12.

Analytic techniques
Net associations between chronic inflammation and QoL were examined. Hypotheses were tested concerning (1) potential unique effects of inflammatory pathology on QoL, and (2) the ability of CRP to mediate associations between fundamental social determinants and QoL. Recommendations were developed for further investigation of these hypotheses in future studies.

Generalized linear modeling techniques, specifically ordinal logistic regression, were used to regress QoL outcomes on CRP. While both QoL variables are numeric and additive in nature, they do not approximate continuity because of the relatively limited numbers of response categories (between 5 and 7) available for each item. Furthermore, QoL data in the NSHAP have significant left skew, consistent with evidence from the literature that life satisfaction is often relatively high among older adults [75].

Consequently, either ordinal logistic (ologit) regression or ordinal probabilistic (oprobit) regression was indicated [74]. Sensitivity analyses using both frameworks were performed to ensure that results did not differ substantially; ologit models are presented for ease of interpretation. Models with and without social structure and relationship covariates (enumerated in Table 1) were generated to check for potential confounding influences. Pursuant to the above discussion of parametric versus non-parametric methods, OLS regression was used to conduct additional sensitivity analyses, but did not meaningfully alter findings. Results from non-parametric methods are thus presented for optimal substantive validity.

In exploring the potential influence of social structure and relationship factors, OLS regressions were generated of CRP on social structure and relationship predictors respectively. These models built on findings from previous research [62,64] suggesting that social factors could fundamentally cause chronic inflammation. Both the OLS models and the more complex ologit models allowed for the development of mediation frameworks for formal analysis in future studies.

Model diagnostics were performed to check regression assumptions and assess the influence of possible outlying data. For ologit models, these included likelihood ratio and Wald tests of the proportional odds/parallel regression assumption. For OLS models, these included Breusch-Pagan tests for heteroskedasticity, and re-estimation of models using Huber-White standard errors when indicated. Potential outliers were investigated by generating standardized residuals with the “predict” command and plotting them by case identification number. When indicated, models were re-estimated excluding observations with standardized residual values greater than 2. While this process often strengthened findings slightly, unadjusted findings are presented to maintain conservatism in interpreting results.

Additional sensitivity analyses were conducted to address potential problems related to sampling and measurement. Alternate models were computed using Stata’s “survey” and/or “cluster” prefixes to incorporate sampling weights. The “robust” omnibus corrector option was also used to adjust for a variety of methodological issues. However, neither of these approaches significantly impacted substantive findings. This likely owed to the size and diversity of analytic samples, as well as the active incorporation of constructs addressed by NSHAP sampling weights. Unadjusted results are thus presented for ease of interpretation.

To compare models with and without statistical controls, fit statistics were computed, including pseudo R-squared values and information criteria. This helped to generate hypotheses about whether or not inflammatory pathology can confer additional effects on QoL in excess of the social factors that can fundamentally cause chronic inflammation. Findings from this study recommend exploring chronic inflammation in two distinct contexts: (1) as a fundamental biological cause of QoL, and (2) as a mediator in relationships between fundamental social determinants and QoL outcomes.

Results
Summary of findings
Ordinal logistic regression models of the two QoL outcomes on the CRP predictor were generated and examined. Simply regressing QoL scores on the raw CRP variable did not prove useful because this general level of analysis masked more nuanced trends in the data. Several sets of bivariate models were thus created, using different methods of coding the CRP variable, using different sets of inclusion criteria for CRP and income, and using models stratified by different levels of serum CRP. Rather than attempting to select a single series of final models, general trends were noted across different inclusion criteria and levels of inflammation. This manuscript takes a conservative approach to presenting findings. All values reported come from regressions using the full sample for each outcome variable, with no additional inclusion criteria.

Highly significant negative associations were identified between CRP levels and both QoL variables. Specifically, CRP significantly predicted NSHAP participants’ odds of reporting high levels of general happiness and relationship
happiness. Net associations with CRP for these QoL measurements mirrored one another in terms of magnitude. When exponentiated, both raw coefficients for the CRP variable yielded odds ratios of 0.90 (p < 0.001) for general happiness and 0.92 (p < 0.01) for happiness with relationships. Among people in the NSHIP sample, each additional milligram per liter of serum CRP is thus associated with 8 to 10 percent lower odds of experiencing high levels of QoL when measured either emotionally or relationally.

However, these findings only held true for levels of CRP below 6 mg/L. These distinctions were explored using regressions stratified by a 6 mg/L CRP threshold. Within each group, CRP was preserved as a continuous independent variable and used to predict QoL. People with more than 6 mg/L of serum CRP did not exhibit a significant association between their levels of the inflammatory biomarker and QoL on either outcome measure. Differences in sample sizes may partially account for this dichotomy: Samples for the CRP < 6 mg/L group were 1,683 and 1,626 for emotional and relational QoL outcomes respectively, versus only 255 and 246 for the CRP > 6 mg/L group. The discussion section of this manuscript explores potential (beyond simple sampling power issues) for which significant relationships were not observed between CRP and QoL above 6 mg/L.

Ensuing analyses thus focus only on results for people with less than 6 mg/L of serum CRP. Sample sizes for each stratum of inflammatory biomarker values are also displayed. For each outcome, the group of NSHIP participants with CRP levels below 6 mg/L represented approximately 87 percent of total variation in inflammatory biomarker values contained within the full dataset. Indeed, a growing body of clinical literature on CRP indicates that stratifying out values in the upper single digits and beyond can generally capture the vast majority of true variation within a given study population [62]. Even among older adults, pronounced inflammatory pathology often appears at very small numeric values of CRP [62]. Hypotheses were thus generated about how differences in inflammatory biomarker levels may become important sources of social inequality.

Figures 1a through 2b illustrate graphically how the probability of reporting different levels of QoL on each relevant measure varies according to CRP readings. Figures 1a and 2a show how scores for each outcome vary according to levels of CRP. By contrast, Figures 1b and 2b show variation within specific levels of the outcome variable across different amounts of CRP. Taken together, these figures illuminate both overall trends and specific dynamics of how chronic inflammation may impact emotional and relational QoL.

Figures 1a and 2a illustrate that while significant relationships between CRP serolevels and QoL can be observed for both the emotional and relational measures of QoL. Figures 1b and 2b add detail by illustrating two complementary patterns. Higher levels of CRP generally predict lower odds of reporting high levels of QoL on each outcome measure. By contrast, lower levels of CRP predict higher odds of reporting higher levels of QoL.

These graphics also reveal nuances of the observed relationships between inflammatory biomarker levels and QoL. Specifically, while clear overall correlations (shown with trendlines) present themselves between CRP and both QoL outcomes, the curves of best fit for these data do not closely approximate linearity in either case. Sociomedical phenomena that may contribute to these patterns are explored further in the discussion section.

Alternate model specifications

Bivariate models for both outcomes were also re-estimated incorporating social structural factors. These covariates represented formal statistical “controls” that could help to illuminate any potential confounding from structural disadvantage. Likewise, significant but non-confounding results for social structure covariates could illustrate potential fundamental causation of chronic inflammation by different types of disadvantage in social status and position.

Re-estimating models for both outcomes using social structure and relationship covariates did indeed return significant results for a variety of social factors. With respect to social structure, both female sex (β = 0.288, p < 0.001) and racial minority identity (β = 0.282, p < 0.01) predicted higher levels of CRP. By contrast, each additional 10,000 dollars of annual income (β = −0.001, p < 0.05) predicted slightly lower levels of CRP. Additional years of age did not significantly predict differences in CRP serolevels. With respect to social relationships, being married (β = −0.413, p < 0.01) and feeling able to communicate openly with intimate partners (β = −0.213, p < 0.05) both predicted lower levels of CRP. Other social relationship factors such as feeling able to communicate openly with friends, feeling able to rely on family members, and feeling able to rely on friends did not significantly predict differences in CRP serolevels.

Yet these results did not generally indicate a confounding influence by any of the structural factors assessed. In fact, some social factors (including race for the emotional outcome and income for the relational one) became non-significant when incorporated into more complex models for each QoL outcome. In general, results from computing more complex models suggested that fundamental causation of inflammation by structural disadvantage was occurring, in addition to potential independent influences on QoL from these same factors. Preliminary analyses of fundamental causation corroborated work by other scholars [62,64] suggesting that disadvantage in a variety of social domains can fundamentally cause chronic inflammation.
A notable exception to the overall pattern of CRP retaining at least some predictive significance was the relational QoL outcome when modeled incorporating social relationship factors. Upon incorporating these covariates, the CRP term became non-significant. However, the literature suggested that because social relationship dynamics can change dramatically in response to incipient or increasingly severe chronic disease [54,55] these factors were more likely to mediate associations between chronic inflammation and QoL rather than confounding them. Similar patterns appeared with the emotional QoL outcome, for which CRP remained a significant (albeit less so) predictor in the full model. Given that QoL outcomes directly addressing social relationships are likely to respond more strongly to changes in these dynamics, more sophisticated mediation analyses using longitudinal data should be conducted in the future.

Models were also re-estimated using interaction terms to check for moderating influences by social structure factors that showed independently significant relationships with one or both QoL outcomes. None of these interaction terms showed any statistical significance when incorporated into more complex models, and are thus not presented in depth using tables or graphs. These null results do not constitute unilateral evidence that structural disadvantage cannot moderate associations between chronic inflammation and QoL. Rather, they suggest that key aspects of social structure likely exert their main influences farther “upstream” [76] than stages of causation captured by moderation frameworks. Indeed, as with the analyses incorporating statistical controls as discussed above, preliminary moderation analyses merely provided more support for the idea that social factors can fundamentally cause chronic inflammation—and perhaps also QoL as a partial function of this inflammatory pathology.
Discussion

Implications of key findings

Findings from this study suggest that the progression of chronic inflammatory disease may exert a significant negative impact on QoL in both emotional and social domains. This study establishes a conceptual foundation that demonstrates the feasibility and importance of conducting more detailed and sophisticated research on the formative and summative sociomedical dynamics of chronic inflammation. Likewise, it can help to corroborate previous research on the fundamental social causes of inflammation itself, especially findings from other studies using NSHAP data [62].

While preserving a robust and representative sample of people in the NSHAP, stratifying my analyses by CRP levels below and above 6 mg/L also limited the extent to which outlying values attributable to measurement error could skew findings. Using evidence-based stratification techniques optimized internal validity without dismissing the potential contributions of very high CRP values to the overall study results. This facilitated consideration of why the much smaller samples of people with biomarker levels greater than 6 mg/L showed no significant associations between chronic inflammation and QoL, whereas their peers had markedly different results. As noted previously, among people with fewer than 6 mg/L of the CRP marker, chronic inflammation was significantly associated with both emotional and relational QoL.

Other sociological studies of QoL reflect that different QoL outcomes may be more or less salient to different people. They also suggest that depending on the particular elements of QoL they capture, these same measures may be more or less sensitive to influence from specific biological and social forces. When measured using CRP,
chronic inflammation appears to have the potential to
influence people’s overall happiness, as well as their sat-
isfaction in intimate relationships, to a small but consist-
ent degree. Other QoL outcomes may show different
patterns of association with CRP.

This same literature has important implications for the
remainder of my analyses of possible relationships be-
tween CRP and QoL. Any associations documented in
older populations are likely to be more pronounced
among younger groups, as older individuals may expect a
certain amount of decline in health and functionality as a
normal part of the experience of growing older. The social
role of “older adult” in contemporary American society
thus includes changes in physical health and functionality,
despite growing pressure to minimize these changes as
much as possible. Demonstrating evidence of impairment
in the course of performing their prescribed social role
thus affords older adults a certain amount of legitimacy
and credibility, even if they experience very large declines
in health and/or functionality.

Likewise, presenting evidence of diminished health and/or
functionality can lead older adults to begin re-
cieving assistance, either informally from family mem-
bors or formally from home health professionals. Extant
sociological literature suggests that while people often
experience negative feelings related to loss of independ-
ence when entering into a caregiving arrangement, re-
cieving care from other people can also exert significant
positive impacts on QoL. Researchers in a variety of dis-
ciplines have documented how living with any type of
persistent illness can produce feelings of isolation and
loneliness, both of which negatively affect QoL. These
feelings can become particularly pronounced for people
who experience a level of functional disability substantial
enough to make living alone difficult, but not severe
enough to warrant seeking outside assistance. Taken to-
gether, these trends may explain why a significant associ-
ation between CRP levels and QoL outcomes did not
emerge for NSHAP participants with CRP levels in ex-
cess of 6 mg/L.

As noted in the results section, visual analysis of
NSHAP data revealed nuances in the relationships be-
tween specific levels of CRP and the odds of reporting
high QoL across both outcome measures. While overall
correlations between CRP and QoL were consistently
negative and clear trendlines emerged, the curves of best
fit for these data did not entirely resemble straight lines.
This may owe in part to measurement considerations,
especially the fact that while higher levels of CRP do
predict chronic inflammation in an overall population,
specific individuals may have levels significantly higher
than average without experiencing significant pathology.

The higher average levels of CRP found among
female-bodied NSHAP participants may speak to this
phenomenon. As noted in the literature review, females
tend to have higher average levels of inflammatory bio-
markers regardless of disease status when compared to
their male peers. However, newer research suggests that
higher levels of CRP in females may owe strongly to
underlying social inequalities that impact females dispro-
portionately [61].

Differences in average levels of CRP among subgroups
within the full NSHAP sample may also stem from so-
 ciomedical phenomena related to the lived experience of
chronicity at particular levels of inflammatory pathology.
Across different levels of disease pathology, people’s ex-
pectations for their health and resultant QoL may differ.
Extant literature shows strong associations between cog-
nitive dissonance—the mental anguish people experience
when reality differs unfavorably from expectations—and
reduced QoL among people who develop chronic condi-
tions, including inflammatory diseases.

The fact that social structure factors did not confound
observed relationships between CRP levels and either QoL
outcome indicates that these forces may exert other types
of influences. Fundamental causation [77] of chronic in-
flammation by structural disadvantage is one of the most
likely ways in which these influences may operate. Indeed,
previous research using the NSHAP [62] supports concept-
ualizing chronic inflammation as a socially conditioned
outcome. Likewise, a variety of other studies have shown
strong associations between structural disadvantage and
the onset of different chronic inflammatory conditions
[59,78,79]. Future research should thus analyze inflamma-
tory biomarkers themselves as potential mediators of rela-
relationships between social disadvantage and QoL.

Strengths of this study include background and per-
spectives from multiple disciplines, parsimonious model
specifications, and use of data on older adults. The inter-
disciplinary approach used in this research helps to
bridge gaps in prior work on inflammation, health, social
life, and QoL. As such inquiry into chronic inflamma-
tion’s potential effects on QoL remains relatively new,
parsimonious models were used to guard against report-
ing of significant associations in cases where none were
present. The choice of NSHAP as a data source, and
specifically the advanced ages of its participants, allowed
for insight into potential cumulation of disadvantage in
QoL. This advantage persisted despite the limited num-
ber of waves currently available in the NSHAP.

Limitations of this study include lack of longitudinal data,
challenges with data capture in the NSHAP and narrow
generalizability. Two specific factors limiting generalizability
are the advanced ages of participants and relatively high
average levels of QoL among these individuals. Only
two waves of NSHAP data are currently available, so
the potential for using panel designs to study these
phenomena remains limited. NSHAP researchers also
experienced problems with certain measurements in the dataset, including the CRP assay. Although this study accounts for potentially problematic CRP values via stratification, it is nonetheless possible that some values in the focal range were also erroneous. Using data on older adults confers drawbacks in addition to the advantages noted above. Specifically, the relatively small range of ages captured by the NSHAP restricts the generalizability of these findings to late midlife and elderly adulthood. Similar processes may occur in children, adolescents, and younger adults without manifesting in the same ways.

Irrespective of the specific strengths and limitations of a particular study, it should also be noted that no research can unilaterally confirm or refute the potential influence of chronic inflammation on QoL. Rather, the statistical significance and reliability of results from this study lend support to the idea of inflammatory biomarkers as meaningful correlates of QoL. These findings likewise recommend further investigation of the topic in both community and clinical settings.

In addition to their consistent statistical significance, these findings may indeed possess some clinical significance. Specifically, the potential for a 1 mg/L difference in CRP levels to produce 8 to 10 percent lower odds of reporting high levels of QoL has important implications for people’s everyday lives. A person with severe inflammatory disease is likely to experience substantially elevated levels of CRP within a certain range [62]. Stratifying regression analyses using a cutoff value of 6 mg/L of CRP allowed for examination of variability across the vast majority of the analytic sample. For each computed model, the size of the population group with CRP serovalues between 0 and 6 mg/L exceeded that of its counterpart with levels in excess of 6 mg/L by approximately a factor of 8.

Conclusions

Taken together, findings from analyses of CRP’s potential net effects on measures of happiness and relationship satisfaction suggest that chronic inflammation may be able to exert a significant negative impact on QoL for older adults. The findings themselves do not constitute firm evidence of a causal association between CRP levels and QoL. However, the robust associations documented between CRP and QoL across a variety of measures indicates that the relationship may be causal, and deserving of further exploration.

Scholars of aging and health should note that the above patterns hold true only for community-dwelling older adults—i.e., the specific population sampled by the NSHAP. Elderly persons residing in long-term care (LTC) settings are often discredited and ignored [80]. This can occur even for residents with extremely strong cognitive and/or physical functioning [80]. Indeed, persons who would have been eligible for participation in NSHAP but moved to long-term care facilities before the study began were censored from the sample. The potential influence of chronic inflammation on the social lives and QoL of older adults in LTC settings thus represents a priority for future research using other data.

Future research should also contextualize chronic inflammation within broader biopsychosocial models of QoL. Extant research suggests that social factors can act as fundamental causes [77] of both chronic inflammation [62,64] and QoL [75]. Specifically, disadvantage in one or more social domains predicts both increased inflammatory biomarker levels [62,64] and reduced QoL [75]. However, research is needed on whether or not chronic inflammation can mediate associations between structural disadvantage and QoL.

Likewise, associations between inflammatory biomarker levels and QoL may themselves be mediated by additional social factors. These potential mediating influences should be explored in future studies. Extant literature provides ample support for social interaction dynamics as potential mediators of associations between health status characteristics and QoL outcomes. For example, trouble in communicating with intimate partners can intensify the impact of disadvantage in health on QoL [81,82]. Loss of social support from friends and family members who become emotionally exhausted may also lead to sharper reductions in QoL among those with persistent inflammation [83].

Similar interactional factors may also moderate associations between inflammation and QoL. If the dynamics of people’s social relationships remain relatively unchanged by severity of inflammation, then associations between inflammatory biomarkers and QoL outcomes may vary according to initial levels of interactional advantage. For example, people who communicate openly with their intimate partners may find that dealing with challenges in health actually brings them closer [16,17]. In turn, social support received from intimate partners may help to protect health [84] and preserve QoL [85]. Conversely, people in strained relationships may find that their partners become distant and unsupportive as their health declines [54,83].

Finally, further research can benefit from incorporating longitudinal data, as well as more sophisticated regression frameworks such as Structural Equation Modeling (SEM). Though only two waves of NSHAP data are available currently, future research can leverage panel analysis techniques to illuminate potential mediation, moderation, and fundamental causation influences in greater depth. Likewise, techniques such as SEM can more accurately capture the simultaneous, synergistic impacts of multiple social and clinical factors. As more waves of NSHAP data become available,
researchers can draw on these frameworks to achieve more nuanced and comprehensive understanding of how chronic inflammation may shape QoL in biopsychosocial context.

**Abbreviations**

CRP: C-reactive protein; NSHAP: National social life, health, and aging project; OLS: Ordinary least-squares; QoL: Quality of life; SEM: Structural equation modeling.

**Competing interests**

The author declares that she has no competing interests.

**Authors’ contributions**

The sole author of this manuscript (ACH Nowakowski) performed all background research, study planning, data analysis, and writing for the paper.

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