Symptoms of chronic rhinosinusitis differentially impact general health-related quality of life*

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Abstract

Background: The degree to which different sinonasal symptoms contribute to the overall quality of life (QOL) detriment in chronic rhinosinusitis (CRS) patients remains unknown. In this study we sought to characterize the effect of different CRS symptoms on the general health-related QOL in patients.

Methodology: We performed a prospective cross-sectional study of 131 adult patients with CRS. Sinonasal symptoms were evaluated using the 22-item Sinonasal Outcomes Test (SNOT-22) and general health-related QOL was evaluated using the EuroQol 5-Dimensional general health-related QOL survey (EQ5D) and visual analog scale (EQ5D-VAS). Health utility values (HUV) were determined using responses to the EQSD. SNOT-22 scores were broken down into subdomain scores for sleep, nasal, otologic/facial pain and emotional function symptoms.

Results: The otologic/facial pain subdomain score consistently had the largest impact on EQ5D-VAS and HUV. After otologic/facial pain, the sleep subdomain score had the second largest effect while the nasal subdomain score had the least impact on general health-related QOL.

Conclusions: Different types of CRS symptoms—most prominently otologic/facial pain and sleep-related symptoms—and their underlying pathophysiologic mechanisms may differentially affect the general health-related QOL detriment associated with CRS. These findings raise the possibility that treatment of the various symptoms associated with CRS may lead to differential improvement in general-health related QOL.

Key words: chronic rhinosinusitis, quality of life, sinonasal symptoms, SNOT-22, EQ5D

Introduction

Chronic rhinosinusitis (CRS), characterized by chronic inflammation of the sinonasal mucosa, leads to sinonasal symptomatology such as congestion, mucopurulent drainage, facial pain/pressure and hyposmia1,2. From the standpoint of pathophysiology, CRS is a multifactorial process without one specific underlying etiologic process3,4. Derangements in innate immunity, the sinonasal epithelial barrier, and response to bacterial flora as well as allergic inflammation and sinonasal obstruction have all been implicated in the pathogenesis of CRS5-9. The complexity of CRS is further highlighted by the frequent lack of correlation between sinonasal symptomatology and objective findings, such as radiographic disease severity10,11. Regardless of etiology, the goal of treatment of CRS is to reduce sinonasal symptomatology. CRS sinonasal symptomatology has previously been shown to have a significant negative impact on patient quality of life (QOL) that is similar to or worse than other chronic diseases, such as asthma or cardiac disease12,13. Care of patients with CRS results in billions of dollars in cost every year14. This expense includes direct healthcare costs that arise from physician visits, medical and surgical treatment, as well as indirect costs that arise from lost productivity. Various validated measures exist to quantify patient QOL; some metrics, such as the 22-item Sinonasal Outcomes Test (SNOT-22), are specific to CRS while others measure general QOL and are applicable to many different conditions14,15. Measures of
Materials and methods
Study participants
This study was approved by the Massachusetts Eye and Ear Infirmary Human Studies Committee. A total of 131 adult patients (with age 18 years or older) with CRS were recruited prospectively and provided informed consent for inclusion in this study. All participants met consensus guideline established criteria for CRS\(^\text{1,2}\). Exclusion criteria included comorbid diagnoses of: 1) vasculitis, 2) cystic fibrosis, 3) sarcoidosis, and 4) immunodeficiency. Any patient having undergone ESS within 6 months was also excluded.

Study design and data collection
This investigation is a prospective cross-sectional study of the impact of CRS symptomatology on general health-related QOL. All data were collected at enrollment. The age and gender of all participants were recorded and all participants completed the validated SNOT-22\(^\text{15,16}\), which reflects sinonasal symptomatology (and CRS-specific QOL), and the EuroQol 5-dimensional (EQSD) general health survey and visual analog scale (EQSD-VAS)\(^\text{17-19}\), which reflects general health-related QOL. Health utility values were derived, as previously described\(^\text{15}\); from the responses to the EQSD. At enrollment, participants were assessed by the evaluating rhinologist for a history of 1) aeroallergen hypersensitivity based on formal testing, 2) asthma, and 3) nasal polyps.

Classification and quantification of sinonasal symptoms into subdomain scores based on SNOT-22 responses
We classified sinonasal symptoms in four categories: sleep, nasal, otologic/facial pain and emotional function symptoms.

Statistical analysis
All analysis was performed with the statistical software package R (www.r-project.org). Correlations were assessed using Pearson’s method. A total of 131 participants were recruited to power this study to detect a statistically significant association of medium effect (\(r=0.15\))\(^\text{20}\) in a multivariable linear regression between EQSD-VAS and nine independent variables (any one of our four symptom subdomains and up to five additional covariates) at a significance level of 0.05 with power of 0.90 using the pwr package. PCA was performed on the SNOT-22 responses of all study participants. Determination of variable loading onto each PC was determined with a varimax rotation on the original PCA using the principal function of the psych package. The four PCs, which explained the greatest amount of data variability, were identified as the first four PCs. Univariate and multivariable associations between PC-derived composite scores and subdomain scores (as independent variables) and EQSD-VAS and HUV (as dependent variables) were calculated using linear regression models.

Results
Study participants
A total of 131 participants (53.4% male and 46.6% female) were enrolled with mean age of 51.4 (standard deviation [SD]: 15.4). Of these participants, 49.6% had a history of aeroallergen hypersensitivity, 29.8% had a diagnosis of asthma, 46.6% had nasal polyps and 33.6% had a history of prior endoscopic sinus surgery for CRS. The mean SNOT-22 score was 36.2 (SD: 23.1), the mean EQSD-VAS was 73.7 (SD: 19.2) and the mean HUV was 0.86 (SD: 0.14).

PCA performed on the SNOT-22 responses revealed that the first four principal components (PCs) from PCA performed on SNOT-22 responses of CRS patients are dominated by sleep, nasal, otologic/facial pain and emotional function symptoms, respectively\(^\text{21}\). Upon confirming this to be the case for our study participants by performing a PCA on their SNOT-22 responses, we calculated a PC-derived SNOT-22 subdomain score for each of these symptom categories by weighting the normalized score for each SNOT-22 item by the loadings on the corresponding PC, as previously described\(^\text{22}\). In the second method, we calculated an un-weighted SNOT-22 subdomain score for each symptom category. Based on the dominant symptoms that we found in each PC, un-weighted SNOT-22 subdomain scores were calculated by simply adding the responses to questions #11 through #18 for the un-weighted sleep symptoms subdomain score; questions #1 through #6, #21 and #22 for the un-weighted nasal symptoms subdomain score; questions #7 through #10 for the un-weighted otologic/facial pain subdomain score; and questions #19 and #20 for the un-weighted emotional function subdomain score.
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associated with HUV by univariate association (Table 1). After multivariable analysis controlling for all PC-derived SNOT-22 subdomain scores as well as age, gender, aeroallergen hypersensitivity, asthma and nasal polyps, all four PC-derived SNOT-22 subdomain scores exhibited a statistically significant association with both EQ5D-VAS and HUV (Table 1). The PC-derived otologic/facial pain subdomain score had the greatest impact on EQ5D-VAS ($b = -6.49$, 95%CI: -9.47 – -3.51, $p<0.001$) and on HUV ($b = -0.06$, 95%CI: -0.08 – -0.03, $p<0.001$). The PC-derived sleep subdomain score had the second largest effect on QOL and the PC-derived nasal subdomain score had the smallest effect on QOL (Table 1).

We similarly checked un-weighted sleep, nasal, otologic/facial pain and emotional function SNOT-22 subdomain scores for association with EQ5D-VAS (Figure 4) and HUV (Figure 5). All of these SNOT-22 subdomain scores were significantly associated with EQ5D-VAS while only the sleep, otologic/facial pain, and emotional function PC-derived subdomain scores were significantly associated with HUV by univariate association (Table 1). After multivariable analysis controlling for all PC-derived SNOT-22 subdomain scores as well as age, gender, aeroallergen hyper-sensitivity, asthma and nasal polyps, all four PC-derived SNOT-22 subdomain scores exhibited a statistically significant association with both EQSD-VAS and HUV (Table 1). The PC-derived otologic/facial pain subdomain score had the greatest impact on EQSD-VAS ($\beta = -6.49$, 95%CI: -9.47 – -3.51, $p<0.001$) and on HUV ($\beta = -0.06$, 95%CI: -0.08 – -0.03, $p<0.001$). The PC-derived sleep subdomain score had the second largest effect on QOL and the PC-derived nasal subdomain score had the smallest effect on QOL (Table 1).

The total SNOT-22 score was significantly correlated with the EQSD-VAS ($r = -0.53$, 95%CI: -0.64 – -0.39, $p<0.001$) and the EQSD-derived HUV ($r = -0.53$, 95%CI: -0.64 – -0.40, $p<0.001$). We next checked whether PC-derived sleep, nasal, otologic/facial pain and emotional function subdomain scores were associated with EQSD-VAS (Figure 2) and HUV (Figure 3). We found that all PC-derived subdomain scores were significantly associated with EQSD-VAS while only the sleep, otologic/facial pain, and emotional function PC-derived subdomain scores were significantly associated with HUV by univariate association (Table 1). After multivariable analysis controlling for all PC-derived SNOT-22 subdomain scores as well as age, gender, aeroallergen hypersensitivity, asthma and nasal polyps, all four PC-derived SNOT-22 subdomain scores exhibited a statistically significant association with both EQSD-VAS and HUV (Table 1). The PC-derived otologic/facial pain subdomain score had the greatest impact on EQSD-VAS ($\beta = -6.49$, 95%CI: -9.47 – -3.51, $p<0.001$) and on HUV ($\beta = -0.06$, 95%CI: -0.08 – -0.03, $p<0.001$). The PC-derived sleep subdomain score had the second largest effect on QOL and the PC-derived nasal subdomain score had the smallest effect on QOL (Table 1).

We similarly checked un-weighted sleep, nasal, otologic/facial pain and emotional function SNOT-22 subdomain scores for association with EQSD-VAS (Figure 4) and HUV (Figure 5). All of these SNOT-22 subdomain scores were significantly associated with EQSD-VAS and HUV by univariate analysis (Table 2). However, after multivariable analysis, only the un-weighted sleep subdomain score significantly associated with EQSD-VAS ($\beta = -0.55$, 95%CI: -0.96 – -0.13, $p=0.011$) and only the un-weighted otologic/facial pain subdomain score was significantly associated with HUV ($\beta = -0.01$, 95%CI: -0.02 – -0.00, $p=0.004$).
Discussion

CRS causes a significant impairment in QOL that leads to lost productivity at home and at work, which translates to billions of dollars in cost every year\textsuperscript{2,12}. Many studies have shown that the severity of CRS-specific symptomatology, taken as a whole, is associated with diminished general health-related QOL\textsuperscript{20}. However, no study has examined whether specific CRS symptoms may differentially contribute to this general health-related QOL impairment. In this study, we now show that the pathophysiology of CRS relating to otologic/facial pain and sleep symptomatology most contribute to general health-related QOL impairment while nasal symptoms contribute the least to general health-related QOL impairment. Investigating the impact of specific symptomatology may be particularly important because previous work has shown that the different CRS symptom types are associated with specific
effects on CRS severity. Sleep quality has been shown to be negatively impacted by CRS and this association is even greater in the setting of comorbid obstructive sleep apnea\(^26\). Cognitive dysfunction that arises as a result of distractibility or reduced concentration in the setting of CRS is associated with more severe CRS symptomatology\(^27\). More severe CRS symptomatology is also associated with higher reports of pain and increased risk of depression\(^28\) as well as decreased olfactory-specific QOL\(^29\). The severity of CRS-specific symptomatology may also be modified by patient characteristics. Factors such as aeroallergen hypersensitivity and comorbid asthma are associated with more severe CRS symptomatology\(^30,31\). These previous disease-specific QOL studies have been critical to our understanding of how patients are affected by CRS-spe-
cific symptom severity. However, while assessment of disease-specific QOL is meant to maximize the sensitivity of detecting intra-individual disease-specific changes\(^{(21,22)}\), assessment of general health-related QOL allows for detection of differences between individuals at the general population level\(^{(15)}\). Additionally, general health-related QOL can be used to calculate health utility values associated with disease states, which may then be used to perform cost utility analyses of different treatment interventions. In fact, treatment of CRS leads to improvement in general health-related QOL. Recent economic analyses using general health-related QOL assessments also have shown the cost-effectiveness of CRS treatments such as ESS\(^{(19)}\). As the cost effectiveness of various treatments is better defined, treatment algorithms will ideally identify those CRS patients who may stand to gain the greatest QOL improvement with different treatment options.

By identifying how certain symptoms relate to general health-related QOL, we may be able to maximize QOL gains with different CRS treatments that focus on alleviating specific symptoms. We and others have previously shown that CRS symptomatology may broadly be categorized as sleep-, nasal-, otologic/facial pain-, and emotional function-related symptoms\(^{(21,22,33)}\). Due to the intuitive grouping of CRS symptomatology into these four categories, as well as previous work suggesting that these four symptom categories may represent different underlying pathophysiologic mechanisms of CRS, in this current study we sought to determine how these four symptom patterns impacted general health-related QOL in CRS patients. We specifically hypothesized that these categories of CRS symptoms may be differentially associated with the general health-related QOL impairment seen in CRS patients.

There are multiple methods for collectively quantifying the severity of similar CRS symptoms. Using the SNOT-22, calculation of CRS subdomain scores for specific categories of symptoms has been previously performed in two different ways. Because the notion of CRS symptom subdomains was derived using structural equation modeling methods such as factor analysis or PCA of the SNOT-22\(^{(21,22,33)}\), PC-derived subdomain scores may be calculated as the sum of SNOT-22 item responses that are weighted by PC-derived weights (also referred to as loadings\(^{(22)}\). For example, the sleep subdomain score could be calculated by summing the score of each SNOT-22 item that is weighted by the sleep PC loadings. Alternatively, a simpler approach to calculating subdomain scores is to assign specific SNOT-22 items to each subdomain and simply add the scores for those SNOT-22 items in an un-weighted manner\(^{(34)}\). The PCA-based method for deriving SNOT-22 subdomain scores benefits from accounting for the exact relationship of dominant symptoms with each other as well as by accounting for small contributions from all SNOT-22 items, rather than accounting for just the most dominant symptoms for each subdomain score. However, the PCA-based method of calculating subdomain scores is analytically intensive and not amenable to convenient use in routine clinical practice. The alternative approach to accounting for only the dominant symptoms in each SNOT-22 subdomain and adding the scores of the corresponding SNOT-22 items in an un-weighted manner is much simpler and convenient to use. This approach, however, does not account for the subtle mathematical relationships and correlations between the SNOT-22 item responses/scores.

In our study, we confirmed sleep, nasal, otologic/facial pain and emotional function symptomatology as distinct CRS disease patterns using a PCA. We then assessed these four symptom categories using both of the aforementioned methods for calculating SNOT-22 subdomain scores. We found that PC-derived SNOT-22 subdomain scores were significantly associated with EQ5D-based measures of general health-related QOL. Un-weighted SNOT-22 subdomain scores, however, were less sensitive for detecting association with EQ5D-based measures of general health-related QOL after accounting for other subdomain scores and other confounding variables. For both methods of calculating SNOT-22 subdomain scores, the otologic/facial pain subdomain demonstrated the greatest magnitude of association with our EQ5D-based measures of general health-related QOL. The sleep subdomain had the second largest effect on general health-related QOL and nasal symptoms had the smallest effect. Our findings with respect to PC-derived SNOT-22 subdomain scores further demonstrate that there likely exist four distinct pathophysiologic processes in CRS that result in distinct symptom patterns and that each of these pathophysiologic processes has a significant, but different, impact on CRS patients’ general health-related QOL. However, further study is needed to confirm whether our findings are due to different pathophysiologic mechanisms of CRS.

Our findings may also have important implications for the consideration of different treatment strategies for patients with CRS. Targeting CRS symptoms that are most associated with a general health-related QOL impairment may lead to the greatest cost utility in the treatment of CRS. Further longitudinal study is needed to determine whether focusing interventions on the subdomains most dominant for patients will optimize gains in QOL. In the meantime, knowledge of the differential impact of CRS symptoms on patient QOL may allow a greater understanding of the overall effect of CRS on patients’ lives.

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**Authorship contribution**
LPH, KMP, RWB, DSC, STG performed the study and wrote/revised the manuscript. ARS designed, performed the study and
Conflict of interest
The authors declare no conflicts of interest.

References

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