Chemical Intolerance in Primary Care Settings: Prevalence, Comorbidity, and Outcomes

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ABSTRACT

PURPOSE This study extends previous community-based studies on the prevalence and clinical characteristics of chemical intolerance in a sample of primary care clinic patients. We evaluated comorbid medical and psychiatric disorders, functional status, and rates of health care use.

METHODS A total of 400 patients were recruited from 2 family medicine clinic waiting rooms in San Antonio, Texas. Patients completed the validated Quick Environmental Exposure and Sensitivity Inventory (QEESI) to assess chemical intolerance; the Primary Care Evaluation of Mental Disorders (PRIME-MD) screen for possible psychiatric disorders; the Dartmouth–Northern New England Primary Care Cooperative Information Project (Dartmouth COOP) charts for functional status; and the Healthcare Utilization Questionnaire.

RESULTS Overall, 20.3% of the sample met criteria for chemical intolerance. The chemically intolerant group reported significantly higher rates of comorbid allergies and more often met screening criteria for possible major depressive disorder, panic disorder, generalized anxiety disorder, and alcohol abuse disorder, as well as somatization disorder. The total number of possible mental disorders was correlated with chemical intolerance scores (P < .001). Controlling for demographics, patients with chemical intolerance were significantly more likely to have poorer functional status, with trends toward increased medical service use when compared with non–chemically intolerant patients. After controlling for comorbid psychiatric conditions, the groups differed significantly only regarding limitations of social activities.

CONCLUSIONS Chemical intolerance occurs in 1 of 5 primary care patients yet is rarely diagnosed by busy practitioners. Psychiatric comorbidities contribute to functional limitations and increased health care use. Chemical intolerance offers an etiologic explanation. Symptoms may resolve or improve with the avoidance of salient chemical, dietary (including caffeine and alcohol), and drug triggers. Given greater medication intolerances in chemical intolerance, primary care clinicians could use the QEESI to identify patients for appropriate triage to comprehensive nonpharmacologic care.

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INTRODUCTION

Between 13% and 33% of people in various populations report considering themselves to be “unusually” sensitive to common environmental chemicals, such as cleaning products, tobacco smoke, perfumes, pesticides, new carpet, and car exhaust.1-7 This condition, known as chemical intolerance,* is the hallmark of multiple chemical sensitivity (MCS), also known as multiple chemical intolerance or idiopathic environemental sensitivities.8 *Semantics in this area are a minefield. In this article, we use chemical intolerance to describe the loss of prior, natural tolerance to common foods and drugs that occurs in a subset of individuals, sometimes after an initial exposure, such as to pesticides or a “sick building.” Various authors have used toxicant-induced loss of tolerance.
mental intolerance. Chemical intolerance has a prevalence of 2% to 13% in population-based surveys. Symptoms are typically multisystem, that is, affecting cognitive, affective, musculoskeletal, gastrointestinal, genitourinary, and cardiovascular systems. Despite its relatively high prevalence in nonclinical samples, the diagnosis and etiology of chemical intolerance remain controversial and understudied. Skeptics and proponents often frame the debate in a dualistic manner, claiming that chemical intolerance is either completely psychogenic or completely toxicogenenic; however, accumulating data suggest that a more nuanced, multifactorial psychobiological process underlies the condition.

Apart from the debate over causality, chemical intolerance holds particular relevance for primary care clinicians. Patients who are chemically intolerant use health care services at increased rates (making an average of 23.3 visits to a medical professional per year). In addition, chemical intolerance is associated with poor quality of life and functional impairments leading to loss of employment and socioeconomic hardships. It is also associated with more medication prescriptions, greater use of physicians and hospitals after exposures, and more visits to environmental specialists. Nonetheless, coordinated multidisciplinary care substantially reduces health care costs in this population.

Previous studies show that persons meeting various criteria for MCS or the less severe, more common chemical intolerance also have increased rates of certain medical and psychiatric conditions. Comorbid diagnoses include heart problems, bronchitis, asthma, pneumonia, rhinitis, sinusitis, hypothyroidism and other autoimmune diseases, irritable bowel syndrome, fibromyalgia, and chronic fatigue syndrome. Family histories in persons having high scores for chemical intolerance show an increased prevalence of heart disease, diabetes mellitus, asthma, and rhinitis, as well as alcohol and drug abuse.

Other studies report associations with panic disorder, major depression, and childhood hyperactivity. A substantial subset of affected individuals also report multiple food intolerances, and several studies have demonstrated food triggers in a subset of children with attention-deficit/hyperactivity disorder (ADHD), migraine, and epilepsy, as well as adults with schizophrenia.

Although some investigators label these patients as having somatoform spectrum disorder, the mean age of onset is typically older than 30 years, a finding that is inconsistent with the diagnostic criterion requirement for the earlier age of onset in the Diagnostic and Statistical Manual of Mental Disorders (4th Edition) (DSM-IV) for classic somatization disorder. Apart from etiology, elevated levels of subjective mental distress (eg, somatization, anxiety, depression) are a major factor in increasing health care use across diagnoses in the general population. Persons with chemical intolerance score higher on scales measuring somatization, anxiety, and depression, although the scores on these psychiatric dimensions account for only a small portion of variance in total chemical intolerance scores outside clinical settings. Taken together, the evidence suggests that chemical intolerance may be an important yet unrecognized contributor to the clinical presentation and use patterns of patients in primary care.

The purposes of our study were to assess the prevalence of chemical intolerance and comorbid medical and mental disorders among primary care patients, and to use standardized instruments to evaluate these patients’ functional status and health care use.

METHODS
Participants
We recruited 400 adult patients (aged ≥18 years) from the waiting rooms of 2 primary care clinics in San Antonio, Texas, who were seeking care for nonacute conditions. The University Health Center–Downtown serves a low-income, predominantly Hispanic population, whereas the Leonard G. Paul Family Health Center serves a predominantly middle-class Hispanic and non-Hispanic white population. Patients were approached in the waiting room after signing in, by seeking the most recent patient to register, no eligible adult patient refused participation. On verbally agreeing to participate in a study about sensitivity to chemicals, patients gave written informed consent. Participants received monetary compensation ($5) for participation.

Instruments
All patients completed a questionnaire having 90 self-report items in either English or Spanish in 20 minutes before seeing the physician. The questions covered demographics, chemical intolerance, mental disorders, functional status, and health care use. The Hollingshead scale measured socioeconomic status (SES).
The Quick Environmental Exposure and Sensitivity Inventory (QEESI), a validated, 50-item, self-administered instrument with high sensitivity and specificity for differentiating individuals with chemical intolerance from the general population, gauged chemical intolerance.\textsuperscript{29} It has 4 scales—symptom severity, intolerance to inhaled chemicals (eg, diesel exhaust, fragrance, cleaning products), intolerance to other substances (eg, foods, medications, alcoholic beverages), and life impact—as well as a masking index, which is a measure of ongoing chemical exposures. Patients were defined as having chemical intolerance if they had a score of 40 or greater on both the chemical intolerance scale (range, 0-100) and the symptom severity scale (range, 0-100).

The Primary Care Evaluation of Mental Disorders (PRIME-MD) Patient Questionnaire screened for symptoms of mental disorders within the past month, including panic disorder, generalized anxiety disorder, major depressive disorder, alcohol abuse disorder, and somatization disorder. Its 25 items show both construct and concurrent validity.\textsuperscript{47}

The Dartmouth–Northern New England Primary Care Cooperative Information Project (Dartmouth COOP) charts’ 8 pictograph scales (scored 1 to 5, where 5 = greatest impairment) assessed functional status within the past month. The COOP charts provide more valid assessments for groups that are culturally diverse, have low literacy rates, or both. In addition to acceptable test-retest reliability (0.42-0.98), the COOP charts have good intraclass correlation coefficients (0.50-0.98) as well as concurrent and construct validity.\textsuperscript{48}

Finally, the Health Care Utilization Questionnaire evaluated use over the previous 2 months.\textsuperscript{49} This 8-item instrument asks individuals to report the number of visits they made to emergency settings (emergency department, minor emergency room, ambulance), physician offices, and mental health settings. Previous work has shown that patients can accurately recall their health care use over the past 2 months.\textsuperscript{50} Individuals also report how many times they were hospitalized in the preceding year.

Analysis

Associations between chemical intolerance and possible mental disorders were assessed using odds ratios. To adjust for the possible impact of other factors on functional status and health care use, we used multivariate analysis of variance (MANOVA) tests, first adjusting for demographic variables (Model 1) and then adjusting for both demographic variables and possible mental health disorders (Model 2). We used $P \leq 0.05$ as indicating significance, with $P \leq 0.10$ considered a trend.

RESULTS

Participants were middle aged on average, with a mean age of 47.4 years (SD, 14.7). Table 1 presents other demographics for the sample. Almost two-thirds of patients were women. One-third had less than a high school education. The majority of the sample fell largely into low-income and Hispanic groups.

Overall, 81 patients (20.3%) met criteria for chemical intolerance. The prevalence of chemical intolerance was related to social class: it was 9.1% among patients in Class I/II (highest SES status), 13.7% among those in Class III/IV, and 24.7% among those in Class V (lowest SES status) ($\chi^2 = 8.20, P = .04$). Patients who met criteria for chemical intolerance rated the severity of their intolerances or adverse reactions to medications, medical procedures, or medical materials significantly higher than did those without chemical intolerance (mean = 4.41 vs 1.87, $t = -5.42$, $P <.001$). Of the 81 patients who met criteria for chemical intolerance, only 19 (24%) reported having previously received this diagnosis.

Table 2 gives odds ratios for personal and family histories of self-reported medical and psychiatric conditions. In general, patients with chemical intolerance reported increased rates of most of the conditions

| Table 1. Patient Characteristics (N = 400) |
|------------------------------------------|
| **Characteristic**                        |
| **Clinic**                                |
| University Health Center–Downtown         | 270 (68) |
| Leonard G. Paul Family Health Center      | 130 (33) |
| **Sex**                                   |
| Male                                      | 148 (37) |
| Female                                    | 252 (63) |
| **Ethnicity**                             |
| Hispanic                                  | 343 (86) |
| Non-Hispanic white                        | 57 (14) |
| **Education**                             |
| Less than high school                     | 125 (32) |
| High school graduate                      | 102 (26) |
| Some college                              | 168 (43) |
| **Income**                                |
| <$20,000                                  | 205 (68) |
| $20,000-$40,000                           | 68 (23) |
| >$40,000                                 | 29 (7) |
| **Social class**                          |
| I                                         | 3 (1)   |
| II                                        | 19 (5)  |
| III                                       | 87 (22) |
| IV                                        | 51 (13) |
| V                                         | 235 (60) |

\* Missing excluded.
\* Based on Hollingshead and Redlich.\textsuperscript{44} Class I is highest socioeconomic status.
Also, they were more likely to report family histories of the same disorders, but only the family histories of gastrointestinal disorders, mood disorders, chemical intolerance, and systemic lupus erythematosus were significantly more common than in the group without chemical intolerance.

The 4 mental disorders previously linked with chemical intolerance (major depressive disorder, panic disorder, generalized anxiety disorder, alcohol abuse disorder) were significantly associated with chemical intolerance status (Table 3). In fact, Figure 1 shows that there was a linear relationship between the number of possible mental disorders and the prevalence of chemical intolerance. Similar to what has been found in earlier studies, PRIME-MD screening symptoms of possible somatization disorder were also significantly related to chemical intolerance.

Before controlling for demographics and comorbid psychiatric conditions, chemical intolerance was significantly associated with poorer functional status in all 8 areas of functioning, with trends toward significance for use of emergency services, medical specialists, and total ambulatory services, as well as hospitalization. In MANOVA controlling for demographic variables (Model 1),

| Table 2. Odds of a Personal History and Family History of Conditions by Chemical Intolerance Status |
|-----------------------------------------------|
| Condition | Personal History | Family History | Odds Ratio (95% CI) | Odds Ratio (95% CI) |
|-----------|-----------------|----------------|---------------------|---------------------|
| Asthma    | 14 (17)         | 34 (11)        | 1.75 (0.89-3.45)    | 22 (27)             | 59 (19)         | 1.64 (0.93-2.89) |
| Allergies | 43 (53)         | 129 (40)       | 1.67 (1.02-2.72)    | 36 (44)             | 132 (41)        | 1.13 (0.69-1.85) |
| Autism    | 1 (1.2)         | 1 (0.3)        | 3.98 (0.25-64.24)   | 2 (3)               | 8 (3)           | 0.98 (0.21-4.73) |
| Multiple sclerosis | 0 (0) | 2 (0.6) | – | 1 (1) | 7 (2) | 0.56 (0.07-4.59) |
| Arthritis | 27 (33)         | 75 (24)        | 1.63 (0.96-2.76)    | 31 (38)             | 97 (30)         | 1.42 (0.85-2.36) |
| Diabetes  | 22 (27)         | 63 (20)        | 1.52 (0.86-2.66)    | 39 (48)             | 122 (38)        | 1.50 (0.92-2.45) |
| Gastrointestinal disorder | 14 (17) | 54 (17) | 1.03 (0.54-1.96) | 21 (26) | 31 (10) | 3.25 (1.75-6.04) |
| Mood disorder | 44 (54) | 64 (20) | 2.83 (0.88-2.66) | 31 (38) | 56 (18) | 2.92 (1.71-4.96) |
| Chemical intolerance | 19 (24) | 24 (8) | 3.77 (1.94-7.30) | 16 (20) | 24 (8) | 3.03 (1.52-6.02) |
| Systemic lupus erythematosus | 3 (4) | 5 (2) | 2.42 (0.57-10.33) | 7 (9) | 9 (3) | 3.26 (1.19-9.03) |

a Values are number (%) unless otherwise indicated.
b Significant odds ratio.

| Table 3. Odds of Symptoms of Mental Disorders by Chemical Intolerance Status |
|-----------------------------------------------|
| Disorder | Chemically Intolerant, No. (%) (n = 81) | Chemically Tolerant, No. (%) (n = 319) | Odds Ratio (95% CI) |
|----------|------------------------------------------|----------------------------------------|---------------------|
| Major depressive disorder | 69 (85) | 106 (33) | 11.55 (6.00-22.26) |
| Generalized anxiety disorder | 63 (78) | 67 (21) | 13.16 (7.30-23.73) |
| Panic disorder | 44 (54) | 53 (17) | 5.97 (3.52-10.11) |
| Alcohol abuse disorder | 30 (37) | 63 (20) | 2.39 (1.41-4.06) |
| Somatization disorder | 74 (91) | 218 (68) | 4.90 (2.18-11.01) |

a Symptoms of the disorder were assessed with the Primary Care Evaluation of Mental Disorders Patient Questionnaire.
b Significant odds ratio.

Figure 1. Prevalence of Chemical Intolerance vs Number of Possible Mental Disorders

Note: Possible disorders were major depressive disorder, generalized anxiety disorder, panic disorder, and alcohol abuse disorder.
results were significant for the model overall \( (P < .001) \), with posthoc tests still revealing significantly greater impairments in all areas of functional status as well as increased use of emergency services, total ambulatory care, and hospitalizations for patients with chemical intolerance (Table 4). In MANOVA controlling for both demographics and PRIME-MD possible psychiatric disorders (Model 2), the differences in functional status and health care use were no longer significant with one exception: chemical intolerance remained significantly associated with more limitations in social activities.

### DISCUSSION

#### Key Findings

This study found that middle-aged patients from relatively lower SES groups who attend primary care clinics have a high prevalence of chemical intolerance. Compared with healthy controls from an earlier study in the same community, San Antonio, by one of the authors (C.S.M.), the current patients had significantly higher scores on the QEESI subscales for chemical intolerances, other intolerances (eg, drugs, alcohol, foods), and symptom severity. The present patients, however, scored lower for masking factors that might otherwise obscure awareness of an association between chemical exposures and symptoms. In other words, the current chemically intolerant patients may be more aware of their difficulties in tolerating everyday environmental exposures than are members of the healthy population.

Moreover, comparison of the current subset of clinic patients who met criteria for elevated chemical intolerance with the non–chemically intolerant subset revealed significant odds ratios only for self-reported allergies, mood disorders, and chemical intolerance histories. From the PRIME-MD screening data, persons with high chemical intolerance had markedly elevated odds ratios for possible major depression, generalized anxiety disorder, panic disorder, and alcohol abuse disorder, as well as somatization disorder. From the self-report questionnaire, the chemically intolerant also had significantly elevated odds of family histories of gastrointestinal disorders, mood disorders, systemic lupus erythematosus, and chemical intolerance.

Although patients with chemical intolerance reported general functional impairment and increased use of emergency, ambulatory, and hospital settings, these associations largely became nonsignificant when controlling for coexisting mental disorders, suggesting that comorbid psychiatric problems may substantially

### Table 4. Functional Status and Health Care Use by Chemical Intolerance Status

| Variable | Model 1\(^a\) | | Model 2\(^b\) | |
|----------|---------------|---|---------------|---|
|          | Chemically Intolerant \((n = 81)\) | Chemically Tolerant \((n = 319)\) | F \((P\ Value)\) | Chemically Intolerant \((n = 81)\) | Chemically Tolerant \((n = 319)\) | F \((P\ Value)\) |
| Limitations in functional status in past month\(^c\) | | | | | | |
| Daily activities | 3.17 | 2.27 | 39.31 \((< .001)\) | 2.69 | 2.39 | 3.61 \((0.058)\) |
| Social activities | 3.02 | 1.95 | 54.60 \((< .001)\) | 2.46 | 2.09 | 5.42 \((0.02)\) |
| Social support | 2.71 | 2.20 | 10.45 \((0.001)\) | 2.32 | 2.30 | 0.02 \((0.88)\) |
| Feelings | 3.11 | 2.07 | 51.15 \((< .001)\) | 2.26 | 2.28 | 0.02 \((0.90)\) |
| Physical fitness | 3.32 | 2.92 | 6.20 \((0.01)\) | 3.11 | 2.97 | 0.50 \((0.48)\) |
| Pain | 3.51 | 2.75 | 24.51 \((< .001)\) | 3.05 | 2.87 | 1.06 \((0.30)\) |
| Overall health | 3.66 | 2.84 | 44.39 \((< .001)\) | 3.11 | 2.98 | 1.07 \((0.30)\) |
| Quality of life | 3.06 | 2.51 | 22.28 \((< .001)\) | 2.59 | 2.63 | 0.13 \((0.72)\) |
| Health care use, number in past 2 months | | | | | | |
| Emergency services | 1.22 | 0.43 | 7.57 \((0.006)\) | 1.04 | 0.47 | 2.78 \((0.09)\) |
| Primary care offices | 2.06 | 2.04 | 0.01 \((0.94)\) | 1.90 | 2.08 | 0.25 \((0.62)\) |
| Specialty offices | 0.74 | 0.42 | 3.80 \((0.052)\) | 0.72 | 0.42 | 2.47 \((0.12)\) |
| Mental health services | 0.61 | 0.34 | 2.44 \((1.2)\) | 0.17 | 0.45 | 2.10 \((0.15)\) |
| Total ambulatory use | 4.63 | 3.22 | 5.00 \((0.03)\) | 3.83 | 3.42 | 0.32 \((0.57)\) |
| Hospitalizations, number in past year | 1.14 | 0.51 | 5.90 \((0.02)\) | 0.88 | 0.57 | 1.03 \((0.31)\) |

Note: Values are multivariate analysis of variance (MANOVA) means adjusted for demographics and symptoms of mental disorders, unless otherwise noted.

\(^a\) Adjusted for age, sex, and socioeconomic status; Wilk \(\Lambda = .797, F = 5.98 \((P < .001)\)\).

\(^b\) Adjusted for age, sex, socioeconomic status, and symptoms of depression, generalized anxiety disorder, panic disorder, and alcohol abuse disorder; Wilk \(\Lambda = .936, F = 1.58 \((P = .07)\)\).

\(^c\) Assessed with the Dartmouth–Northern New England Primary Care Cooperative Information Project (Dartmouth COOP) charts. Each scale has a possible score of 1 to 5, where 5 = greatest impairment.
 contribute to perceived impairments and elevated health care use in persons with chemical intolerance, similar to findings in previous studies of other primary care populations in which chemical intolerance was not assessed. Rates of alcohol abuse disorder were higher in the present chemically intolerant patients than in their unaffected counterparts, a finding that is not necessarily consistent across other samples. Although family histories for alcohol and drug problems are increased in MCS/chemical intolerance, many earlier studies observed increased levels of alcohol intolerance and lower levels of alcohol use in the index cases of chemical intolerance.

At the same time, there is no evidence that treating psychiatric symptoms alone will resolve chemical intolerance. Twin studies of chronic fatigue, a common overlapping diagnosis in chemical intolerance, indicate that psychiatric illness does not fully explain the clinical picture. Previous nonclinical studies showed that psychological distress per se accounts for only a proportion of the variance in chemical intolerance scores. In one study, patients with MCS pointed to chemical avoidance as the single most helpful intervention. These same patients rated psychotropic medications as least helpful for their condition. Our study and numerous earlier studies have found increased rates of medication intolerances in the chemical intolerance population.

Previous research has shown an association between chemical intolerance and work disability, problems with shopping, and difficulties with travel. Perhaps most troubling is the impact that chemical intolerance has on social functioning and relationships. In the present data, impairment in daily and social activities was independent of any associated mental disorders. Gibson et al found that people with chemical intolerance derive social support from involvement in chemical intolerance–related support groups and certain close relationships.

It is uncommon for primary care clinicians to recognize chemical intolerance in a given patient. Patients are not likely to report chemical intolerance as their chief complaint. The present data show that only 24% of patients who fulfilled criteria for the condition had previously received a related diagnosis. Nonetheless, earlier research found that 54% of community-dwelling people with high chemical intolerance reported at least 1 comorbid medical problem. In addition, patients with chemical intolerance were significantly more likely to report a family history of the condition as well. These data replicate previous findings on the familial nature of MCS and chemical intolerance.

Although a detailed discussion of possible biologic mechanisms is beyond the scope of this article, laboratory studies of persons with MCS or chemical intolerance indicate that both central nervous system sensitization and inflammatory processes contribute to the multisystem clinical picture. Both sensitization and chronic inflammation also play a role in the medical, psychiatric, and substance abuse conditions found at elevated rates in the personal histories, family histories, or both of patients with MCS described in the Introduction and identified in Table 2.

**Implications**

Our findings suggest that chemical intolerance is a common problem in primary care settings among low-SES primary care patients, but also that it often goes unrecognized and requires active investigation by the primary care physician. The presence of chemical intolerance among relatives, multiple medication intolerances, complex multisystem conditions, or a prior diagnosis of somatization disorder should raise the index of suspicion for chemical intolerance. Use of a standardized instrument such as the QEESI may be helpful in detecting chemical intolerance. If it is found, the physician should screen the patient for major depressive disorder, panic disorder, generalized anxiety disorder, and substance abuse, recognizing that these may be associated conditions, not necessarily causal.

The occupational medicine literature contains multiple reports of a higher prevalence of mood and anxiety disorders in chemically exposed workers. Chemical intolerance with multisystem symptoms, fatigue, anxiety, depression, irritability, and cognitive difficulties can also develop, after nonindustrial chemical exposures associated with home and office remodeling or extermination.

Caress et al found that some people with MCS or chemical intolerance had changed various products they used, purchased air filters, moved to a different residence, or used some combination of these measures. Although avoidance of known chemical triggers is recommended, evaluation of potential psychosocial factors is also important. In addition, because those with chemical intolerance are more intolerant of many medications and medical materials, the use of nonpharmacologic approaches to treatment where feasible may be particularly important for these patients. Fox et al reported on the benefits of a comprehensive, holistic, multidisciplinary treatment program for chemical intolerance, which ranged from lifestyle changes and nutritional supplementation to mindfulness-based stress reduction, physical therapy, group learning, guided imagery, and psychotherapy. Such programs have been shown to reduce impairment as well as health care use and costs. Referring chemically intolerant patients for a comprehensive specialized program of care may thus...
be the most pragmatic and cost-effective approach for selected patients.15,75

Study Limitations
This study has several limitations. First, we used self-report data in lieu of comprehensive but more costly and time-consuming clinical examinations and medical record reviews. Although prior research suggests that patients can accurately recall health care use over the past 2 months, the use data reported here were not verified through chart audit. Second, the PRIME-MD screen for mental disorders used in this study is not a diagnostic instrument; the actual rates of psychiatric disorders meeting DSM-IV criteria may differ. Our goal was to estimate the prevalence of chemical intolerance among primary care patients with chronic medical conditions. To reduce the risk of false-positives on the QEESI, we screened out patients who had acute problems. Finally, because the sites used in this study predominantly served low-income, middle-aged Hispanic patients, the findings may not be generalizable to other primary care populations; in fact, the prevalence of chemical intolerance was inversely related to social class in this study. Because prior studies have often focused on high-SES, white populations, however, this study contributes to our understanding of chemical intolerance in low-SES, minority primary care populations. Given their occupations and where they live, such individuals (86% Hispanic in our sample) are far more likely than higher-SES groups to have been exposed to solvents, cleaning agents, pesticides, and other substances now clearly linked with the development of chemical intolerance and associated neuropsychiatric symptoms, via a process referred to as toxicant-induced loss of tolerance.76

In conclusion, often unrecognized, chemical intolerance is prevalent and commonly comorbid with a range of medical and psychiatric conditions in primary care settings. The QEESI is a self-report tool for busy clinicians to use in recognizing chemical intolerance symptoms and developing a more effective treatment plan. This instrument may help pinpoint potential environmental chemical, food, and drug contributors to the clinical picture and lead to improved nonpharmacologic intervention strategies. The implications for the primary care physician are that chemical intolerance is prevalent in low-income, primary care settings but must be actively sought. Patients’ occupational and environmental exposures may have contributed to their condition. The presence of chemical intolerance among relatives, a history of medication intolerances/ adverse drug reactions, and multisystem symptoms including psychiatric symptoms (depression, cognitive difficulties, anxiety, panic attacks) should raise the index of suspicion for further chemical intolerance screening using the QEESI or a similar tool.

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Key words: chemical intolerance; idiopathic environmental intolerance; multiple chemical sensitivity; psychiatric comorbidity; functional impairments; health care utilization; practice-based research

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CORRECTIONS

Ann Fam Med 2012;10:365. doi:10.1370/afm.1429.

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When referring to the article by Margolius and colleagues in the text and reference, Margolius was spelled incorrectly as Margolis.

The Folsom Group. Communities of solution: the Folsom Report revisited. Ann Fam Med. 2012;10(3):250-260.

The footnote for Figure 2 shows a source of data for allied health as http://bhpr.hrsa.gov/healthworkforce/. This URL deviates from that published in the PDF of the article (http://bhpr.hrsa.gov/healthworkforce/reports/factbook02/FB101.htm), which was not available at the time of publication.