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Laura Plantinga, Emory University
Ann Vandenberg, Emory University
Felicia Goldstein, Emory University
Brian Jones, Georgia Institute of Technology
Jeremy Johnson, Georgia Institute of Technology
Christopher Bowling, Emory University
Charmayne Dunlop-Thomas, Emory University
Sung Lim, Emory University
Cristina Drenkard, Emory University

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Patient and provider perceptions of a novel cognitive functioning report for patients with systemic lupus erythematosus: a qualitative study

Laura Plantinga 1,2, Ann Vandenberg,1 Felicia Goldstein,3 Brian Jones,4 Jeremy Johnson,4 Christopher Barrett Bowling,5,6 Charmayne Dunlop-Thomas,1 Sung Sam Lim 1,2, Cristina Drenkard 1,2

ABSTRACT

Objective To determine whether and how cognitive assessment data should be included in a report for patients with SLE and their providers.

Methods Leveraging experiences from prior studies, we created a report that included a hypothetical patient’s results on tests of multiple domains based on the NIH Toolbox Fluid Cognition Battery. In focus groups that comprised patients with SLE (two groups) and their providers (two groups), feedback was sought on the presentation of results as well as the potential value of the report in the clinical setting.

Results Feedback regarding the presentation of the report was generally positive. Both patients with SLE and their providers liked its simple graphics and use of a colour-graded scale to indicate performance. However, both groups stressed the importance of using non-stigmatising language in describing results. Several potential purposes of the report, including distinguishing cognitive versus other issues, explaining cognitive challenges, improving patient–provider interactions, guiding decision-making, improving functioning or preventing impairment and tracking cognitive function over time, were noted by the participants. Potential barriers, such as inadequate clinical staffing or time and lack of potential treatments for identified issues, were also discussed.

Conclusion In this exploratory study, we found that both patients with SLE and their providers were receptive to the idea of a patient-friendly report of cognitive test results. This study provides important information to guide future pragmatic research to optimise the delivery of cognitive information to patients with SLE.

INTRODUCTION

Cognitive impairment in patients with SLE, often described as a ‘lupus fog’, has been reported in up to 50% of patients with SLE.1–6 This impairment may limit independence and reduce quality of life for patients with SLE, regardless of age. Despite its importance, cognitive functioning is not well described in most medical charts7 and often not discussed in the clinical setting. Specific tools to improve the recognition of cognitive impairment and facilitate its discussion are currently lacking in SLE.

To address this gap, we leveraged our experiences in (1) creating a physical functioning report from a previous study in patients receiving dialysis8 and (2) completing detailed cognitive assessments as part of a pilot study of functioning in patients with SLE (Approaches to Positive, Patient-centered Experiences of Aging in Lupus; APPEAL)9 to create a prototype cognitive functioning report for patients with SLE and obtain feedback on the value and presentation of the report. Our objective was to determine whether and how cognitive assessment data should be included in a functioning report for patients with SLE and their providers.

METHODS

Study design and patient involvement

Four 90 min focus groups (two SLE patient (n = 18) groups and two SLE provider
groups (n=9)) were held in 2019. Patient participants were recruited from the APPEAL pilot, representing a subsample of the population-based Georgians Organized Against Lupus cohort; provider participants (physicians, nurses, medical assistants) were recruited from the local provider population based on their experience with patients with SLE. While the initial prototype report was created by the study team, without patient involvement, these focus groups were designed to identify patient and provider priorities for the report of cognitive functioning information in future pragmatic research and ultimately clinical care.

Data collection and analysis
Development of the cognitive functioning report
Based on a physical functioning report piloted in a population of patients receiving dialysis, our goal was to create a patient-friendly report for patients with SLE (figure 1). The domains of cognitive functioning included episodic memory, working memory, processing speed, inhibitory control and attention, and cognitive flexibility and were based on the NIH Toolbox assessments that were performed in the APPEAL pilot study. Everyday examples of tasks involving these cognitive domains were listed, discussed and selected by the entire team over the course of several meetings; the chosen examples were illustrated and provided along with a written description (figure 1). To gather feedback on preferred presentation of performance/scores, the hypothetical patient’s performance was displayed as both a colour-graded scale with categories of ‘impaired’, ‘borderline’, ‘low average’ and ‘average or better’ and as numeric scores, including age, sex, race, ethnicity and education-adjusted t-scores (mean=50, SD=10) with associated percentiles. Further explanations were provided on the back of the report (figure 1).

Results
Focus group participants
The patient participants had a mean age of 51 and were predominantly female (83%) and Black (83%), and the mean duration of SLE was 20 years. The provider participants had a mean age of 40; 89% were female and 33% were Black. The provider participants reported a mean of 6 years of experience treating SLE.
distinguish the effects of age-related or dementia-related problems from other issues. Even with cognitive test results, it might be impossible to do so. Providers also stated that, in some cases, the possibility of using the report to distinguish cognitive problems from other issues was not clear.

The proposed report was seen as a communications tool: ‘It actually gives them (providers) now a visual of what it is that you may have been trying to express.’ While patient participants focused on the way that the report would initiate the conversation around cognition, provider participants focused more on how it would inform their communication with patients. In terms of guiding decision-making, both patients and providers mentioned follow-up referrals, and providers discussed the possibility of adjusting medications as needed.

Patient and provider participants also had unique perspectives on the potential value and challenges of using the proposed report. First, patients less often mentioned the possibility of using the report to distinguish cognitive problems from other issues. Providers also stated that, even with cognitive test results, it might be impossible to distinguish the effects of age-related or dementia-related cognitive decline from the effects of SLE (including musculoskeletal problems that hinder testing, distracting pain, fatigue or depression). Second, several patients—no provider—mentioned the possibility of using the report to identify cognitive problems for prevention of further deterioration or improvement. A related anticipated problem was the potential lack of a solution to the cognitive problems identified in the report (eg, ‘I would feel comfortable telling them (the results), as long as I had something for them to do to improve it’ (provider)). Finally, providers voiced significant concern about not having the time and resources to add cognitive assessment into clinic visits: ‘I’m not sure of how easy it would be in a clinic where I’m just the sole provider…if it was more of a multidisciplinary clinic, where you had nurses and social workers involved in the patient care, I think it would really work.’

### Table 1: Patient and provider feedback on potential value of shared information about cognition for patients with SLE

| Purpose | Patients (n=18) | Providers (n=9) |
|---------|----------------|----------------|
| Distinguishing cognitive problems from other issues | ‘You wonder okay, now, if I’m having something else going on or if it’s a lupus fog.’ | ‘It would be really helpful to understand if it is a real problem, or if it is their mood, or just their perception.’ |
| Explaining cognitive problems | ‘I think it’s helpful information because it does break down the different types of memory loss, like, “cause I never thought about it. How it’s broke down.’ | ‘They might feel a little bit less frustrated if they can see that they do have some cognitive impairment, and maybe that’s why they’re having some trouble at work.’ |
| Improving patient–provider interactions | ‘I think it would break the ice. Sometimes when you go to the doctor, you’re scared or it just doesn’t come to you, certain things, you may just wanna ask. With all of these different segments…it will bring back some of the things that you may have wanted to ask.’ | ‘I think it would maybe help change maybe how you counsel the patient. If they’re more cognitively impaired, maybe give them much more simple instructions or involve the family a little bit more.’ |
| Guiding decision-making | ‘I want them to, you know, refer me to like a neurologist or something, that can see my brain and see what’s going on and how it’s functioning and, get me some type of help there.’ | ‘A lot of our patients are, surprisingly, professionals who are still doing things like teaching, or pharmacists, so some of their daily work is very important. If we found something like this, it might be something we would have to intervene on.’ |
| Improving cognitive functioning or preventing cognitive impairment | ‘[I]t would be nice to know what I can prevent now because luckily I’m still young. If there’s something I can do now to make it better when I’m 40, 50, 60…I don’t know, brain exercises.’ | – |
| Monitoring cognitive function over time | ‘[Y]ou want to know if you’ve improved or if you failed or if you slightly fell out, or whatever the case might be.’ | ‘This would be good in certain situations especially if you have a patient that always comes in and says they have the lupus fog, to sort of track them.’ |

### Potential value of the cognitive report

Table 1 describes several stated potential purposes of having information about SLE patient cognition, such as that provided in the cognitive report. General comments included ‘I think it’s going to be really helpful’ (provider) and ‘I would love to do this with my doctor’ (patient). Specific to the report, both patients and providers voiced its potential to improve patient–provider interactions; patient participants envisioned the report as a communications tool: ‘It actually gives them (providers) now a visual of what it is that you may have been trying to express.’ While patient participants focused on the way that the report would initiate the conversation around cognition, provider participants focused more on how it would inform their communication with patients. In terms of guiding decision-making, both patients and provider participants mentioned follow-up referrals, and providers discussed the possibility of adjusting medications as needed.

### Presentation of the cognitive report

The participants liked the overall appearance of the report, although a few patient participants suggested that the multiple results made it appear ‘complex’ at first glance. Feedback on graphic representations of cognitive domains was generally positive, with most participants understanding the tasks presented as relevant without the need for explanation. Table 2 presents feedback for
the various presentations of the cognitive test results. Participants liked the simplicity of the colour-gradated scale, although some suggested changing the word ‘impaired’ to something less judgemental such as ‘fair’. In addition, most patient and some provider participants viewed the colour-gradated scale as a more neutral way to present results than numeric scores. Across groups, many suggested that the colour-gradated scale alone should be the primary presentation of results, potentially with numeric scores on the back of the report for providers and patients who wish to track these results. For numeric scores, most participants had preferences for percentiles over t-scores, if numeric scores were presented (table 1). While both represent comparisons to the general population, with adjustment for age, sex, race/ethnicity and education, some patients expressed preferences to be compared with a lupus population or not to be compared with others at all (eg, ‘The description says, “Like you.” I don’t know what is like me…’).

**DISCUSSION**

In this exploratory study, we found that both patients with SLE and their providers were receptive to the idea of a patient-friendly report of cognitive test results. The report was seen by patients and providers as a tool to identify cognitive issues (or reassure patients of normal/stable functioning) and to facilitate better patient–provider interactions.

Importantly, this study, while limited by its qualitative nature and small sample size, identified several considerations for use of the report in the clinical or pragmatic research setting. First, expectations of patients with SLE regarding the potential for improvement in cognition should be managed. Few interventions to improve cognitive functioning have been studied in SLE to date; however, neither patients with SLE nor their providers mentioned the possibility of providing support (such as occupational therapy) to help manage day-to-day tasks in the setting of cognitive impairment; education in this area of both

| Performance reported by: | Representative quote(s) | |
|--------------------------|--------------------------|--------------------------|
|                          | Patients (n=18)           | Providers (n=9)           |
| Colour-gradated scale    | ‘…a quick glance through the color coding and the scale, it gives you a brief report as to where you stand within these particular categories.’ | ‘I like the color part too, and I think the patients understand red being bad versus green being better.’ |
|                          | ‘The word “impaired” makes you feel like you’re in dummy-mode.’ | ‘I think the way it’s laid out, actually being sort of skewed with just “average or better” at the top. Even with this patient that is 27% but almost in the middle, so it actually looks better than it really is.’ |
|                          | ‘The only thing I would worry about is the wording, like impaired, borderline, below average... If I saw this and I was below average...It might even make me more depressed.’ | |
| Numeric scores           | ‘I’m not sure that I want to see numbers. I want for my clinician to say to me, “You are doing well,” or, “You’re not doing so well,” or, “You could be doing a little better. Maybe we could do this.” I don’t need to see these numbers.’ | ‘I think that’s enough. I would take the numbers off. And maybe if the numbers could be on the back and more so a reference for the physician or whoever was seeing the patient.’ |
| t-scores                 | ‘I think the whole t-score...is sort of confusing.’ | ‘A t-score is going to be really confusing to patients. And then the doctor is going to have to spend a lot of time explaining what a t-score is. Because even really well-educated patients are not going to know what a t-score is.’ |
|                          | ‘Out of a 100 people they’re...39.’ (Representing misunderstanding of the t-score) | |
| Percentiles              | ‘Well the percentage’s okay but the t-score kind of threw me off.’ | ‘The percentile. It’s easier. Patients like percentages because they can remember it.’ |
|                          | ‘I think some of us are probably more worried about numbers because it’s kind of like, it’s kind of like, labeling us. That’s kind of a label, a percentage score.’ | ‘If you’re talking about which of these numbers is better, percentile I think is a lot easier to understand.’ |
|                          | ‘Dealing with someone whose processing information is 13th percentile and trying to explain to them...it’s really hard.’ | |
patients and providers might be needed. Second, presentation of results is important; some patients with SLE may just want reassurance that functioning remains ‘average or better’, but other patients, and providers, may wish to track numeric results to identify longitudinal patterns. Third, the context of the clinical setting would have to be considered. Having providers on staff who feel comfortable discussing cognitive test results and have adequate time for the discussion is critical. Additionally, rheumatologists and primary care providers may wish to be aware of these issues to help guide their clinical decision-making, but they should also have the ability to refer a patient to a neuropsychologist or other specialist specifically for further cognitive testing and/or to provide needed cognitive support. Partnership with geriatric providers could mitigate these issues by facilitating provider education and providing a cognitive support referral network.

In conclusion, these results provide important information to guide future pragmatic research to optimise the delivery of cognitive information to patients with SLE.

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ORCID iDs
Laura Plantinga http://orcid.org/0000-0003-0809-8981
Sung Sam Lim http://orcid.org/0000-0003-2361-0787
Cristina Drenkard http://orcid.org/0000-0002-6832-7291

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