VISIT-TS version 2: A multimedia tool for population studies on tic disorders [version 1; referees: 1 approved]

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Abstract

Population-based assessment of Tourette syndrome (TS) and other tic disorders produces a paradox. On one hand, ideally diagnosis of tic disorders requires expert observation. In fact, diagnostic criteria for TS explicitly require expert assessment of tics for a definite diagnosis. On the other hand, large-scale population surveys with expert assessment of every subject are impracticable. True, several published studies have successfully used expert assessment to find tic prevalence in a representative population (e.g. all students in a school district). However, extending these studies to larger populations is daunting.

We created a multimedia tool to demonstrate tics to a lay audience, discuss their defining and common attributes, and address features that differentiate tics from other movements and vocalizations. A first version was modified to improve clarity and to include a more diverse group in terms of age and ethnicity. The result is a tool intended for epidemiological research. It may also provide additional benefits, such as more representative minority recruitment for other TS studies and increased community awareness of TS.

This article is included in the Tic collection.
Introduction

Some important questions in Tourette Syndrome (TS) require large-scale epidemiological studies. To give one example, studies have not yet had the power to definitively establish whether TS is equally common in people of African versus European descent. In the U.S., although diagnosis and treatment are about twice as common in European Americans (CDC, 2009), three prior studies in the U.S., though limited in various ways, all found tics to be more common in minorities (Costello et al., 1996; Lapou & Monk, 1964; [Table 4]; personal communication Costello EJ to KJB, 1999; personal communication Peterson BS to KJB, 2008; Peterson et al., 2001). The results may differ so dramatically because of true genetic or epigenetic differences between racial groups (Robertson et al., 2009), or because social determinants of health care create barriers to diagnosis or treatment that create an artifactual difference in apparent prevalence (American Psychiatric Association, 2013, under Tic Disorders/Culture-Related Diagnostic Issues; CDC, 2009; Olsson et al., 2011). Setting this question will require large-scale prevalence studies that recruit an adequate, representative sample of minority populations.

Cubo (2012) reviews several factors that complicate epidemiological research on TS. One is that such studies generally must rely on assessments by lay interviewers. Although that approach has been very useful for psychiatric epidemiology in general, the validity may reasonably be questioned in the case of TS. There can be difficulties in conveying adequate descriptions of movements by words alone; probable miscategorization or failure to recognize some abnormal movements by both subjects and lay interviewers; the broad differential diagnosis of tics, including other movement disorders and normal movements; and misinterpretation of typical tics due to their intermittent nature, suppressibility and fluctuating severity over time or in response to the environment.

We were especially concerned that some respondents with tics, or whose children had tics, might not correctly interpret written descriptions of tics but would recognize the tics if they saw them. Supporting the potential importance of this concern, epidemiological studies that included expert examination (Comings et al., 1990; Cubo et al., 2011; Hornsey et al., 2001; Jin et al., 2005; Khalifa & von Knorring, 2003; Khalifa & von Knorring, 2005; Kurlan et al., 1994; Lanzì et al., 2004; Mason et al., 1998; Wang & Kuo, 2003) generally report a several-fold higher prevalence of tic disorders than do other epidemiological studies (CDC, 2009; Scahill et al., 2014).

To address these issues, we developed a multimedia screening interview to enhance population-based ascertainment of tic disorders by lay interviewers (“VISIT-TS”, Gordon et al., 2010). A video presented and discussed typical tics, and addressed a few difficulties in differential diagnosis, and then presented questions to gather the information required for diagnosis by DSM-IV-TR (American Psychiatric Association, 2000) and DSM-5 (American Psychiatric Association, 2013). After initial testing and application (unpublished report, Striley CW, Black KJ, Kelso N, and Vagelakos L), we revised the instrument. Here we describe the approach we took and the result: VISIT-TS v. 2.

Methods

We first reviewed previous methods including the Yale Child Study Center questionnaire (Findley et al., 1999; Jagger et al., 1982), the Kiddie SADS semi-structured interview (K-SADS-PL) (Kauffman et al., 1997) and the interviews used by Apter et al. (1993; Gillberg & Rasmussen (1982, Appendix); Hornsey et al. (2001) and Mason et al. (1998) who used the Apter questions and the National Hospital Interview Schedule for GTS (Rickards & Robertson, 2003; Robertson & Eapen, 1996); Appendix I in Khalifa & von Knorring (2003); and Table I in Linazasoro et al. (2006). We also reviewed the Diagnostic Confidence Index (Robertson et al., 1999), the YGTSS (Storch et al., 2005), and the parent and child self-report forms used by the Tourette Syndrome Association International Consortium for Genetics (1999). An expert in psychiatric epidemiology (CWS) developed the questions that would be posed, in consultation with a movement-disorders-trained neuropsychiatrist (KJB). The interview was designed to address both current (past month) and lifetime symptoms and included information needed for TSSG, DSM-IV-TR and DSM-5 criteria for TS.

We wrote a script addressing the following aims: demonstrate tics, discuss their defining and common attributes, and address features that differentiate tics from other movements and vocalizations. We then selected video clips from patients and research volunteers who gave written permission to re-use their video separately from patient care or the research study they had participated in. We also obtained permission from people with tics to re-use selected video clips that they had already made publicly available on YouTube. The final video was produced by Ty Travis (San Tan Valley, Arizona, USA). We dubbed the final product VISIT-TS, for “Video-Integrated Screening Instrument for Tics and Tourette Syndrome” (Gordon et al., 2010).

The first version of VISIT-TS was used in an initial reliability and validity study that provided experience and initial feedback from interviewees and staff (unpublished report, Striley CW, Black KJ, Kelso N, and Vagelakos L). It was also shared with about a dozen other movement disorders experts and we reviewed their feedback. In response to this initial experience and feedback, we made many changes, including new video clips, thereby reducing the amount of time the narrator is shown and showing more diversity in ethnicity and age. We added and improved graphics, including written text while examples of tics appear in the background. To avoid confusion, we removed videos showing examples of non-tic movements, and we eliminated medical terms unfamiliar to the general public, such as chorea. Finally, we restored an unintentionally omitted question on lower facial tics. We call the revised product VISIT-TS v. 2.

Results

The revised VISIT-TS multimedia tool includes almost 100 video clips defining and demonstrating tics including simple and complex motor tics as well as simple and complex vocal tics, edited to a length of 5 minutes. Following the clips, 16 questions are presented in written and spoken form, one at a time, accompanied in most cases by brief video of the phenomenon being ascertained (see Appendix 1; question 4 of the video, at about 6:09, is a good
brief demonstration). The questions take another 5 minutes. The video clips demonstrate adults and children in similar numbers, both sexes (male:female ≈ 5:4), and include some ethnic diversity (about 1 in 8 clips are Hispanic or non-white). VISIT-TS v. 2 is freely available for noncommercial use at https://zenodo.org/collection/user-kjb or at http://dx.doi.org/10.5281/zenodo.55604.

Discussion

This approach is based on the premise that survey respondents will respond more accurately about tics in themselves (or their children) after the interviewer shows them a brief video about tics than if they would if only asked about history by questionnaire or by cross-sectional lay observation. Because tics can come and go, can be suppressed, and often resemble intentional movement or vocalizations, diagnosis of tic disorders can be challenging (Black et al., 2016; Cubo, 2012).

Some data are available to judge the sensitivity of lay diagnostic instruments for tic disorders. In two studies, about half of the children who had previously been diagnosed with TS were missed by research screening: 1 of 2 in Landgren et al., 1996 and 8 of 15 in Snider et al., 2002. Conversely, routine clinical assessment for tics is also insensitive; Kadesjo & Gillberg (2000) report that a tic diagnosis had been considered during child psychiatric treatment in only 1 of 18 children with TS. Khalifa & von Knorring (2003) examined the sensitivity of their questionnaire but only by comparing questionnaire responses from parents to those from teachers. It appears Wang & Kuo (2003) collected physician examination data on questionnaire-negative children, but those data were not reported. Linazasoro et al. (2006) do not specify whether any of the tics diagnosed by a physician observing a classroom of students for 20 minutes were missed by parent or teacher questionnaires. Stefanoff et al. (2008) diagnosed a tic disorder in 6% of children whose parents and teachers noticed no tics; this is more remarkable given that the diagnosis rate in screen-positive children was only 18%. Cubo et al. (2011) found sensitivities of 36%–73% for questionnaires completed by teachers, observers or parents. In a recent study, a semstandardized diagnostic interview (the DISC) captured only about half the cases of TS, and there was little agreement between DISC results and expert clinician diagnosis (Lewin et al., 2014).

The most detailed data on the sensitivity of questionnaires for tic diagnosis come from the study of Mason et al. (1998). They gave questionnaires containing the 4 tic screening questions of Apter et al. (1993) to students, parents and teachers. They also screened for tics with direct classroom observation by Dr. Mason, a psychologist trained in tic detection at the Queen Square, London, TS center; she watched each classroom for an hour, 2 minutes per student. To confirm the diagnosis, Mason then directly examined all 16 consenting screen-positive students in a traditional clinical setting. Importantly, Mason also examined 8 students randomly chosen from screen negatives, i.e. those who had no tics reported by themselves, parents, or teachers, and no tics observed in the classroom. Remarkably, 3 of the 8 had at least one tic when examined directly, counted only if it had been present for at least a year by history! This very high rate of missed chronic tic disorders (37.5%) suggests that traditional questionnaires and interviews are insufficiently sensitive. VISIT-TS was designed to improve sensitivity by making sure subjects and parents have seen typical tics on video before answering questions about them.

Linazasoro & colleagues (2006) used a method somewhat similar to the VISIT-TS approach, i.e., they showed a videotape of tics as part of an initial lecture to parents and teachers, followed by a survey that included a short written description of tics. Independently, “all children were directly observed in the classroom by an expert clinician in the field of tics who diagnosed tics based exclusively on the characteristics of the movements”, with a limit of 20 minutes’ observation per classroom (p. 2107). However, the authors note limitations of their work including the fact that children were observed collectively, for a relatively brief period of time, and while engaged in school work, when tics may have been suppressed. The questionnaires actually identified more children (98) than the expert (57), suggesting either that parents and teachers overdiagnosed some movements as tics, or that they were describing tics present in the past but no longer present, or that the classroom observation was not an adequately sensitive clinical comparison. A videotape demonstrating tics was released (Tourette Syndrome Association, 1990), but it was intended for a professional audience rather than for epidemiological studies.

The VISIT-TS also has limitations. The DSM and TSSG criteria explicitly require application by properly trained experts, so VISIT-TS is primarily intended as a screening tool rather than as a substitute for clinical expertise. The sensitivity of VISIT-TS has not been reported. Nevertheless, a 5- to 10-minute video-illustrated questionnaire is probably a reasonable compromise for epidemiological or other tic studies that require screening large population samples for tic disorders.

Data availability

The video can be found here: http://dx.doi.org/10.5281/zenodo.55604 (Vachon et al., 2016).

Author contributions

MJV assembled video clips and wrote the first draft and KJB edited it. CWS and KJB drafted the questions. MJV, MRG and ECB identified video clips. CWS, MRG, JMK and KJB created the first version of the video. MJV, CWS, MRG, MLS and KJB contributed to the spoken script for the video. MJV and JMK edited video clips. All authors approved the final draft.

Competing interests

VISIT-TS v. 2 is published under a CC BY-NC-ND license, i.e. freely available for noncommercial uses. Funds from any future commercial application of VISIT-TS are intended to benefit research on or clinical care for tic disorders.
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Supplementary material
Appendix 1
Questions asked at the end of the VISIT-TS v. 2
1. Have you ever had eye movement tics?
2. Have you ever had repeated blinking or winking, like the tics in the video?
3. Have you ever had eyebrow tics?
4. Have you ever had mouth, tongue or jaw tics?
5. Does any part of your upper body, such as your head, shoulders, or arms jerk, turn or move unexpectedly or differently like in the video?
6. Has the middle or lower part of your body ever moved differently, like your stomach muscles contracting, your legs kicking or feet moving, over and over?
7. Have you sometimes repeated sounds over and over, like clicking, clucking, humming, grunting or smacking?
8. Have you sniffed or cleared your throat over and over, even when you didn’t have allergies or a cold and your nose wasn’t runny or itchy?
9. Have you had any repeated movements that are always done in the same way and involve more than one muscle group—like eyes and mouth, or shoulder plus arm? These tics may seem like they are being done on purpose or intentionally at times, but usually they are not.
10. Have you done things over and over like:
    a. Adjusting your clothes?
    b. Making obscene gestures?
    c. Patting yourself?
    d. Twisting your hair?
    e. Scratching your foot inside your shoe?
11. Have you sometimes repeated words over and over, or yelled out phrases over and over?
12. The video showed several behaviors, including making noises and movements, that people did over and over. Do you feel you have any tic-like behavior that has not already been mentioned? If yes, please describe it.

if Yes to any previous question, continue:
13. Did your “YES” answer, or answers, refer to something that started before your 18th birthday, that is when you were age 0 through 17 years old?
14. Did your “YES” answer, or answers, refer to something that happened many times a day most days for some period of time?
15. Did your “YES” answer, or answers, refer to something that kept happening over a period of a year or more?

if Yes to question 15, continue:
16. You said you had a year go by during which you had tics most of the time. During that year, did you ever have a time when all the tics were gone for 3 months straight?
References


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In the article “VISIT-TS version 2: A multimedia tool for population studies on tic disorders” by Vachon, MJ, Striley, CW, Gordon, MR, Schroeder, ML., Bihun, EC, Koller, JM, and Black, KJ, a new method of classifying tics by lay persons is proposed as an aid in larger scale population studies.

The authors provide a clear account for the need for improving existing methods of detecting and differentiating tics and present an interesting solution in the form of their new multimedia tool. It is a positive aspect that participants with tics include different age groups, races, and both sexes. However, some of the video clips present one separate tic, whereas several tics are present in the video.

The rationale for developing the screening tool is its integration in large-scale epidemiological studies. Even though a few epidemiological studies focusing on tics and Tourette have been carried through, future methods will probably offer possibilities for screening cohorts for several co-occurring conditions. The authors may thus consider the possibility of further pruning the video (almost 10 minutes) and comparing two versions (e.g. the long one and a shorter version focusing on the different types of tics, which are more or less presented in the first few minutes of the video) for their validity against expert ratings. The shorter the time used for a screening, the more it will be used in big cohort studies. This could be integrated into future perspectives at the end of the manuscript.

It would also be interesting to carry out similar tests of the sensitivity as mentioned in the article by comparing the VISTS-TS to clinician ratings.

Storch et al. (2005) is used as a reference for the YGTSS – the authors might consider adding the reference of Leckman’s (1989) original work.

Competing Interests: No competing interests were disclosed.

We have read this submission. We believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
Kevin J Black, Department of Psychiatry, Washington University in St Louis, USA

We appreciate the thoughtful suggestions from Drs. Plessen and Hagstrøm.

Our primary intent was to help lay people decide whether they (or their child) had any tics, not to distinguish which tics they had. I see from your response that we did not make that point clear. I think there are benefits to presenting and asking about different tics separately, even for our more limited goal, but your perspective also suggests an opportunity to shorten or otherwise improve on the video.

You wrote, “some of the video clips present one separate tic, whereas several tics are present in the video.” I believe you are saying that some video clips may show two tics while the narration is discussing only one of them. If I understood that correctly, it’s a good point that we had not considered. We may be able to address that in a future version of the video.

Competing Interests: No competing interests were disclosed.

Discuss this Article

Version 1

Reader Comment 10 Aug 2016

Irene Malaty, University of Florida, USA

The authors have attempted to address a longstanding problem in TS epidemiologic research, which is the challenge in capturing tic prevalence when recognition and trends in seeking healthcare are variable. Previous “real world” observational ascertainment techniques have often utilized limited face time in single environments. The authors have produced a video educational tool to help involve community members who have consistent time observing (or living with) potential tic behaviors across time and environments. This could increase the reach of tic screening and possibly increase sensitivity for detecting tic behaviors. The video has as its strengths simple language and short clips. I particularly like the inclusion of short clips that reinforce particular tics being discussed when cataloguing body regions and tic types in the second half of the video. No tool is perfect, and a novice evaluator may not obtain from this video the ability to distinguish actual causes of observed behaviors. For instance, one might misclassify idiopathic blepharospasm as a tic. A person with chorea would also be scored for having movement of the arms or legs, which would not be tics. Similarly, psychogenic movements could be classified as tics. On the other hand, even experts may need further questioning to distinguish such diagnoses from tics. In summary, specificity of using lay assessments may be questionable. This may or may not be acceptable in a screening tool. It would be interesting to learn how this tool may be implemented in practical terms. Process for consenting participants will need to be considered. Would schools be target sites for dissemination? How would it be distributed in a way that would sample a wider audience than prior tools have? This may indeed be a great educational tool for screening larger populations, and even for helping clinicians collect information from patients. Validating this tool against currently accepted means of diagnosing Tourette (involving expert assessments) in a subset of participants will help confirm its utility. I enjoyed watching the video and thank the authors for sharing this interesting project.
Competing Interests: I am on the medical advisory board of Tourette Association of America (TAA) and direct a TAA center of excellence. I have participated in Tourette research funded by Auspex and Neurocrine. None of these present conflicts with my comments.