

Preconsult interactive computer-assisted client assessment survey for common mental disorders in a community health centre: a randomized controlled trial

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Abstract

Background: Access disparities for mental health care exist for vulnerable ethnocultural and immigrant groups. Community health centres that serve these groups could be supported further by interactive, computer-based, self-assessments.

Methods: An interactive computer-assisted client assessment survey (iCCAS) tool was developed for preconsult assessment of common mental disorders (using the Patient Health Questionnaire [PHQ-9], Generalized Anxiety Disorder 7-item [GAD-7] scale, Primary Care Post-traumatic Stress Disorder [PTSD-PC] screen and CAGE [concern/cut-down, anger, guilt and eye-opener] questionnaire), with point-of-care reports. The pilot randomized controlled trial recruited adult patients, fluent in English or Spanish, who were seeing a physician or nurse practitioner at the partnering community health centre in Toronto. Randomization into iCCAS or usual care was computer generated, and allocation was concealed in sequentially numbered, opaque envelopes that were opened after consent. The objectives were to examine the interventions' efficacy in improving mental health discussion (primary) and symptom detection (secondary). Data were collected by exit survey and chart review.

Results: Of the 1248 patients assessed, 190 were eligible for participation. Of these, 148 were randomly assigned (response rate 78%). The iCCAS ($n = 75$) and usual care ($n = 72$) groups were similar in sociodemographics; 98% were immigrants, and 68% were women. Mental health discussion occurred for 58.7% of patients in the iCCAS group and 40.3% in the usual care group ($p \leq 0.05$). The effect remained significant while controlling for potential covariates (language, sex, education, employment) in generalized linear mixed model (GLMM; adjusted odds ratio [OR] 2.2; 95% confidence interval [CI] 1.1–4.5). Mental health symptom detection occurred for 38.7% of patients in the iCCAS group and 27.8% in the usual care group ($p > 0.05$). The effect was not significant beyond potential covariates in GLMM (adjusted OR 1.9; 95% CI 0.9–4.1).

Interpretation: The studied intervention holds potential for community health centres to improve mental health discussion. Further research with larger samples should examine the impact on detection and enhance generalizability. **Trial registration:** ClinicalTrials.gov, no: NCT02023957, registered on Dec. 12, 2013.

The World Health Organization predicts that common mental disorders such as depression will rank first in the burden of diseases for high-income countries by 2030.¹ Depression can usually be effectively managed in primary care; however, most people with this illness in Canada do not get treatment.² The rate of depression in patients visiting family practices is high (14%–45%);^{3–6} for this reason, maximizing its identification in primary care has been considered as a possible way of increasing the number of people receiving appropriate treatment.^{7,8}

Ethnocultural and immigrant groups from non-European and low-income backgrounds in Canada are less likely to have

their common mental disorders (such as depression, anxiety and post-traumatic stress disorder [PTSD]) treated than their peers^{9,10} because they often encounter multifold barriers to accessing timely care.^{10–13} The development of community

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health centres has been effective in decreasing barriers to care for physical health for vulnerable communities such as immigrant and ethnocultural groups in Ontario.¹⁴ Studies report that these patients visit emergency departments less frequently and receive higher preventive care and chronic disease management compared with patients enrolled in other primary care models.^{15–18} Despite these successes, community health centre resources are stretched owing to the complex and growing needs of the vulnerable communities they aim to serve. In such situations, mental health problems may be less likely to be identified than physical health issues.

Increasing the determination of common mental disorders in community health centres could be one way of decreasing disparities in mental health treatment for vulnerable ethnocultural and immigrant groups. A recent innovation in this regard is the development of interactive and user-friendly pre-consult eHealth tools. These tools can be available in the waiting room and can be used to alleviate some patient barriers (e.g., knowledge gaps, stigma and communication difficulties) and provider constraints such as time.^{19,20}

We developed an interactive computer-assisted client assessment survey (iCCAS) tool in English and Spanish for common mental disorders.^{21–23} This study reports a randomized controlled trial (RCT) of the efficacy of the tool for improving discussion about mental health issues and detection of mental illness in an urban community health centre in Toronto.

Methods

The study protocol for the 2-arm pilot RCT was developed in collaboration with the partnering community health centre. We followed CONSORT guidelines for nonpharmacological interventions.²⁴

Study site

The partnering community health centre, Access Alliance, is located in Toronto, where more than 50% of the residents are immigrants and most identify themselves as members of racialized communities.²⁵ The 3 clinics of Access Alliance serve primarily immigrant, refugee and racialized ethnoracial communities who report English and Spanish as their top preferred languages. These clinics employed 4 family physicians and 5 nurse practitioners along with a multidisciplinary team at the time of study.

Intervention

The study intervention was a touch-screen iCCAS with validated screening scales for common mental disorders and was administered to patients in the waiting room. The tool produced individualized reports for patients and clinicians before the patient was seen by a family physician or nurse practitioner. Completion time ranged from 10 to 15 minutes. Data were encrypted and collected on a secure remote server. The research assistant printed the reports using a password protected web portal. The Patient Recommendation Sheet summarized the risks, provided information about community

resources and made suggestions to seek the clinician's advice when a risk was reported. The clinician risk report was a 1-page summary of the health risks and included scores and symptoms of the assessed common mental disorders in addition to possible referral pathways.

The development of the intervention included several steps. First, a comprehensive literature review of tools for identifying common mental disorders was undertaken. Twenty questionnaires used to identify major depression, generalized anxiety, PTSD and alcohol abuse in community or primary care samples were assessed on length, reliability, validity and use in diverse groups. The final selection of tools aimed for brevity while maintaining coverage of different common mental disorders important to diverse populations and cross-cultural validity. The final 52-item survey included the Patient Health Questionnaire 9 (PHQ-9);²⁶ Generalized Anxiety Disorder 7-item (GAD-7) scale;²⁷ Primary Care Post-traumatic Stress Disorder (PTSD-PC) screen;²⁸ and CAGE²⁹ (concern/cut-down, anger, guilt and eye-opener) scale, in addition to items pertaining to the social determinants of health (e.g., education, English language ability, housing, financial means to meet daily needs, immigration status and social support). Further details are described elsewhere.^{30,31}

Final survey and report structures were determined with input from researchers and community partners of Access Alliance; the survey was translated to and back-translated from Spanish, and a usability pilot was conducted with 9 clinicians and patients.

Participants and procedures

All family physicians and nurse practitioners at the study sites were eligible to participate, and all provided informed written consent permitting recruitment of their patients. Before patient recruitment, the physicians and nurse practitioners received a workshop with field updates by a mental health expert (K.M.) on the 4 included common mental disorders. Blinding of clinicians was not possible owing to nature of the intervention.

Patient inclusion criteria were age at least 18 years, proficiency in English or Spanish and visiting a consenting clinician. Exclusion criteria were being accompanied by a family member for interpretation, being a new patient, feeling unwell (self-report) or research assistant's inability to offer details (e.g., privacy or comprehension issues).

Patient recruitment occurred November 2013 to May 2014. Three trained research assistants (2 bilingual) approached patients in the waiting room and applied eligibility criteria. Eligible and willing patients received the study details in a separate room and provided informed written consent. The consenting patients were then randomized into the intervention (iCCAS) or control (usual care) group using 1:1 allocation. The randomized allocation sequence was computer generated by an off-site biostatistician before recruitment and concealed using sequentially numbered, opaque envelopes.³² The envelopes were opened after consent, keeping patients and recruiters blind to the assignment.

Before seeing the clinician, patients assigned to the iCCAS group completed the interactive survey using an iPad and received a printout of their tailored recommendation sheet. The Risk Report for clinicians was attached to each patient file before the consultation. Patients assigned to the control group continued to receive usual care with no health-risk assessments before the consultation. After the visit, patients in both groups completed an exit survey and received a resource list. At this time, all patients were asked whether they would like to see a counselor for issues covered in the study surveys.

Outcomes and data collection

The main outcomes were patient discussion (yes or no) on mental health and clinician detection (yes or no) of mental health symptoms. The former was measured by the exit survey and the latter by review of the medical charts (clinical notes section) for both groups. The mental health discussion could have been initiated by either patient or clinician. Patient sociodemographic and health care-related information were collected for both groups by the exit survey. The tool's acceptance and technological quality were assessed in the intervention group by including the Computerized Lifestyle Assessment scale^{33,34} for assessing perceived benefits, privacy barriers and interaction barriers on 5-point (disagree, not sure or agree) subscales, and 4 quality questions (easy or difficult)^{35,36} in the exit survey. The review of electronic medical charts was completed using a data extraction form that was developed with the assistance of the clinic's information technology team and piloted before its use.

Statistical analysis

A total recruitment of 150 patients was planned for this pilot study, with mental health discussion as the primary outcome of interest. A sample size of 75 patients per group was based on an anticipated improvement of a 20%–25% increase in the patient discussion rate as a result of the intervention.

We examined internal consistency of PHQ-9, GAD-7, PC-PTSD and CAGE scales using Cronbach α .

The sociodemographic characteristics of patients randomized to each of the 2 groups (iCCAS and control) were first compared using bivariate analyses, and subsequent analyses for health-related characteristics were performed; correlations within patients seen by the same medical staff (family physician or nurse practitioner) were accounted for in these analyses using the generalized linear mixed model (GLMM) approach. The associations between the 2 mental health outcomes (discussion and detection) and patient characteristics (sociodemographic and health-related) were examined in a similar manner. To evaluate the effectiveness of the intervention (iCCAS), while taking into account patient characteristics and patient correlations, multivariable analyses using GLMM were employed. The covariates of language, sex, education and employment were included in the multivariable analyses based on clinician investigators' input, literature review and statistical evidence; none of these variables had missing data. Statistical analyses were performed using SAS version 9.4 (SAS Institute) and SPSS version 22.

Ethics approval

Research ethics approval was obtained from York University (certificate no. e2013-291).

Results

The numbers of patients randomized and included in the final analyses are detailed in Figure 1. Of 1248 patients assessed, 190 were eligible to participate, and 148 of these were randomly assigned (response rate 77.9%). Six patients were found ineligible during chart review and excluded subsequently as recommended by Fergusson and colleagues.³⁷ One patient who felt unwell immediately after consent and provided no data was not included in the analysis. There were 76 patients in the intervention (iCCAS) group and 72 in the control (usual care) group. Patients in the iCCAS group reported "agreement" with the perceived benefits of the tool (mean \pm standard deviation [SD] 4.1 ± 0.6) and were "not sure" for the barriers to privacy (2.6 ± 0.8) and interaction (2.8 ± 0.9). More than 95% of participants found the technology easy to use.

Internal consistency of PHQ-9, GAD-7, PC-PTSD and CAGE scales were 0.86, 0.89, 0.75 and 0.63, respectively.

Sociodemographics

Overall, patient age ranged from 18 to 71 years (mean 37 ± 1.0), 61.2% identified as female and 78.2% participated in the study in English. A total of 97.9% were immigrants, and the top 3 regions of birth were Latin America (33.3%), South Asia (27.9%) and Africa or the Middle East (16.3%). There were no significant differences in the sociodemographic characteristics between the iCCAS and the usual care groups (Table 1 and Table 2).

Outcomes by sociodemographics

Mental health discussion occurred for 49.7% (73/147) of patients. The discussion frequency was higher for English-speaking (86.3%) than for Spanish-speaking (13.7%) patients, ($p \leq 0.05$). Patients who had a mental health discussion received mental health referral more often (30.1%) than patients who did not have a discussion (5.4%) ($p \leq 0.001$). None of the other sociodemographic or health-related variables were significantly associated with the mental health discussion. The secondary outcome of mental health detection occurred for 33.3% (49/147) of patients. The frequency of mental health detection was higher for English-speaking (89.8%) than for Spanish-speaking (10.2%) patients, ($p < 0.05$). Mental health detection was more common for unemployed (79.6%) than part- or full-time employed (20.4%) patients ($p \leq 0.05$). Patients who had a mental health detection received referral more often (42.9%) than patients without mental health detection (5.1%) ($p \leq 0.001$). Patients with mental health detection were also more likely to self-report poor or fair health (59.2%) than patients without detection (22.4%), ($p \leq 0.001$). Other sociodemographic or health-related variables were not significantly associated with mental health detection.

Outcomes by intervention

Analysis of the intervention and control groups shows that 58.7% of patients in the iCCAS group had a mental health discussion compared with 40.3% in the usual care group ($p \leq 0.05$) (Table 3). The intervention effect was significant on

mental health discussion in the unadjusted analyses (odds ratio [OR] 2.11, $p < 0.05$). In the final GLMM, the effect of the iCCAS on mental health discussion remained significant (adjusted OR 2.2, $p = 0.02$) while controlling for language, sex, education and employment (Table 3).

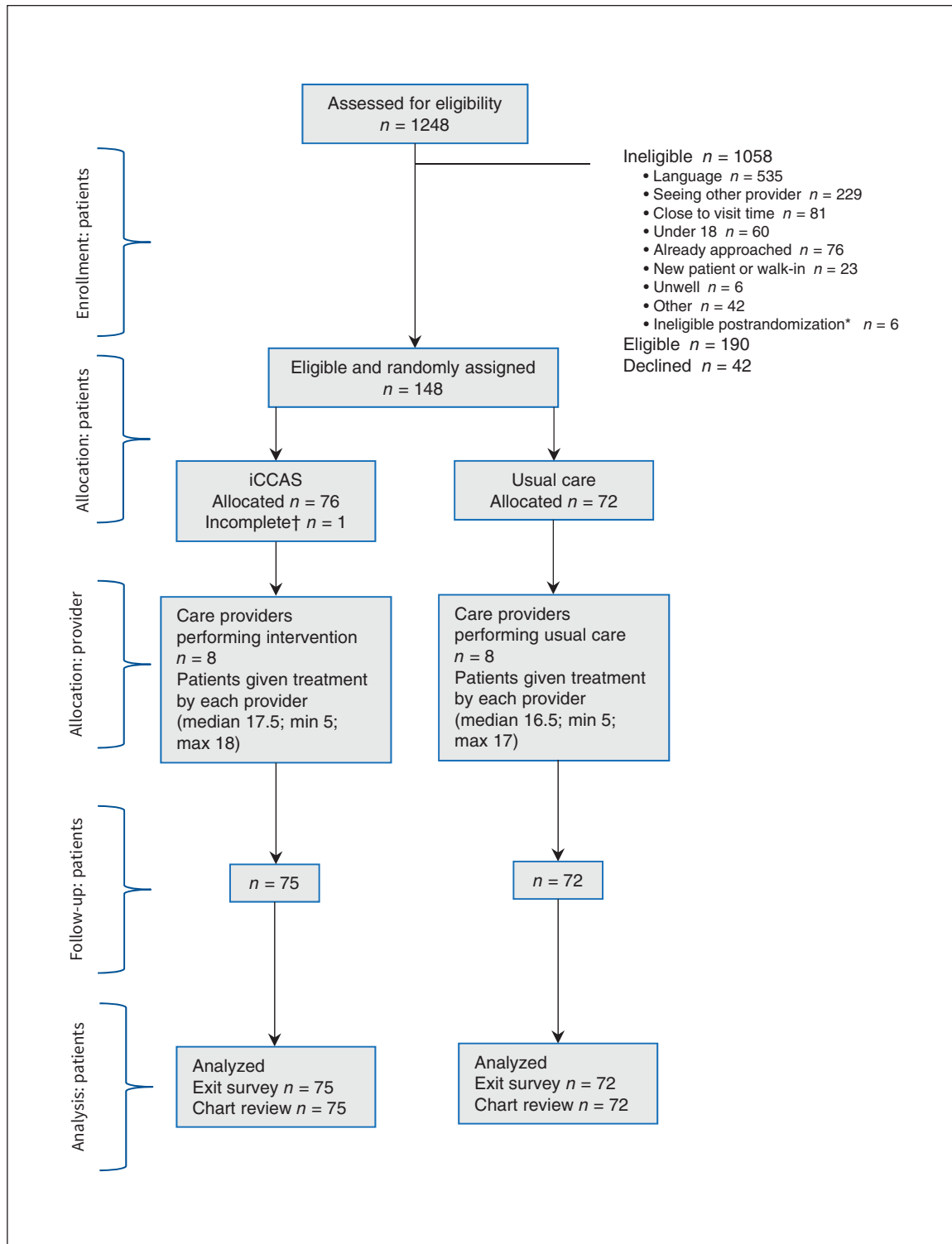


Figure 1: Study flow diagram. *Three patients already approached, 1 under age, 2 seeing other provider. †Patient felt unwell soon after consent and discontinued.

In terms of mental health detection, 38.7% of patients in the iCCAS group had a clinical detection of mental health symptoms compared with 27.8% in the usual care group ($p > 0.05$). The intervention effect was not significant on mental health detection in unadjusted analyses (OR 1.65, $p > 0.05$). In the final GLMM, the effect of iCCAS remained not signifi-

cant (adjusted OR 1.93, $p = 0.08$), whereas the effect of language, sex, education and employment were significant on mental health detection.

Interpretation

Our study, with a multicultural sample of patients, shows that an interactive, user-friendly, health risk assessment tool for common mental disorders administered in English and Spanish by touch-screen tablet was effective in significantly increasing the frequency of patient discussions about mental health with their attending clinicians at the collaborating community health centre. In addition, the study clinicians detected mental health symptoms more frequently in the iCCAS intervention group than in the usual care group, although the difference was not quite significant in multivariate analysis. At the same time, the outcomes of mental health discussion and detection were significantly correlated, and patients with mental health symptom detection had a significantly higher frequency of mental health referral compared with those without detection.

Patients in the iCCAS group reported minimal technological difficulties, agreed with the tool's benefits and neither agree nor disagree with the barriers in relation to privacy or provider interaction. Overall, the results of this pilot trial sug-

Table 1: Patient sociodemographic characteristics

Variable	iCCAS, no. (%) <i>n</i> = 75	Usual care, no. (%) <i>n</i> = 72
Age, yr, mean ± SD	36.5 ± 12.7	37.5 ± 12.2
Sex		
Male	26 (34.7)	27 (37.5)
Female	49 (65.3)	41 (56.9)
Transgender†	0 (0.0)	4 (5.6)
Language		
English	59 (78.7)	56 (77.8)
Spanish	16 (21.3)	16 (22.2)
Relationship status‡		
Married/common law	37 (49.3)	38 (52.8)
Separated/divorced/ widowed	11 (14.7)	10 (13.9)
Single, not in relationship	18 (24.0)	16 (22.2)
Single, in relationship	10 (13.3)	9 (12.5)
Have children	47 (62.7)	45 (62.5)
Immigrant	74 (98.7)	70 (97.2)
Citizenship status		
Canadian citizen/ permanent resident	54 (72.0)	50 (69.4)
Other	21 (28.0)	22 (30.6)
Years lived in Canada		
Up to 5 yr	41 (54.7)	36 (50.0)
5 or more yr	34 (45.3)	36 (50.0)
Highest level of education		
Up to grade 12	38 (50.7)	34 (47.2)
College or more	37 (49.3)	38 (52.8)
English reading/writing		
Poor or fair	19 (25.3)	18 (25.0)
Good or better	56 (74.7)	54 (75.0)
Employment status		
Full-time	15 (20.0)	9 (12.5)
Part time	11 (14.7)	16 (22.2)
Not employed	49 (65.3)	47 (65.3)
Annual household income before taxes		
< 20 000	48 (64.0)	46 (63.9)
20 000–30 000	12 (16.0)	18 (25.0)
30 000–40 000	8 (10.7)	2 (2.8)
40 000–60 000	2 (2.7)	4 (5.6)
> 60 000	5 (6.7)	2 (2.8)

Note: iCCAS = interactive computer-assisted client assessment survey.
 *Unless otherwise specified.
 †Subsequent analyses combined transgender with female (results were same on combining with male).
 ‡Total does not sum up because patients were allowed to select more than 1 response.

Table 2: Patient health-related characteristics

Variable	iCCAS, no. (%) <i>n</i> = 75	Usual care, no. (%) <i>n</i> = 72
Self-rated health		
Poor or fair	27 (36.0)	24 (33.3)
Good, very good or excellent	48 (64.0)	48 (66.7)
Purpose of visit		
New problem	21 (28.0)	12 (16.7)
Routine physical exam or follow-up	54 (72.0)	60 (83.3)
Follow-up suggested in index visit)	53 (70.7)	50 (69.4)
Internal referral provided		
Mental health	13 (44.8)	10 (40.0)
Other	16 (55.2)	15 (60.0)
External referral provided		
Maternal health	2 (9.1)	1 (4.5)
Other	20 (90.9)	21 (95.5)
Visit satisfaction		
Satisfied or very satisfied	59 (78.7)	61 (84.7)
Neutral	7 (9.3)	6 (8.3)
Dissatisfied or very dissatisfied	9 (12.0)	5 (6.9)

Note: iCCAS = interactive computer-assisted client assessment survey.

gest that iCCAS positively facilitated the continuum of discussion, detection and referral for concerns about common mental disorders. A wider adoption of such preconsult tools in the community clinics serving vulnerable communities could be useful in reducing patient barriers to disclosing mental health concerns and clinician challenges in conducting symptom

assessments,⁹⁻¹⁴ thereby facilitating diagnoses of common mental disorders in a timely manner. Future research with a larger sample and longitudinal arm is needed to enhance the generalizability of the reported results and to allow assessment of changes in mental health outcomes over time.

We were able to identify some systemic inequities in the

Table 3: Effect of intervention				
Variable	No. (%) of patients		Unadjusted OR (95% CI)	Adjusted OR (95% CI)
	Yes <i>n</i> = 73	No <i>n</i> = 74		
Mental health discussion				
Language				
English	63 (86.3)	52 (70.3)	2.67 (1.15–6.18)	2.80 (1.15–6.85)*
Spanish†	10 (13.7)	22 (29.7)		
Sex				
Female	49 (67.1)	45 (60.8)	1.32 (0.67–2.60)	1.54 (0.73–3.24)
Male†	23 (32.9)	29 (39.2)		
Education				
College or more	38 (52.1)	37 (50.0)	1.09 (0.57–2.09)	1.32 (0.65–2.66)
Less than college†	35 (47.9)	37 (50.0)		
Employed				
Full-time	9 (12.3)	15 (20.3)	0.49 (0.19–1.23)	0.51 (0.19–1.37)
Part-time	11 (15.1)	16 (21.6)	0.56 (0.23–1.13)	0.63 (0.25–1.59)
Not employed†	53 (72.6)	43 (58.1)		
Group				
iCCAS	44 (60.3)	31 (49.9)	2.11 (1.08–4.09)	2.23 (1.11–4.49)*
Usual care†	29 (39.7)	43 (58.1)		
Mental health detection				
Language				
English	44 (89.8)	71 (72.4)	3.35 (1.19–9.42)	3.02 (1.01–9.03)*
Spanish†	5 (10.2)	27 (27.6)		
Sex				
Female	29 (59.2)	65 (66.3)	0.74 (0.36–1.50)	0.79 (0.36–1.72)
Male†	20 (40.8)	33 (33.7)		
Education				
College or more	30 (61.2)	45 (45.9)	1.86 (0.92–3.76)	2.31 (1.08–4.96)*
Less than college†	19 (38.8)	53 (54.1)		
Employed				
Full-time	3 (6.1)	21 (21.4)	0.21 (0.06–0.76)	0.17 (0.04–0.66)*
Part-time	7 (14.3)	20 (20.4)	0.51 (0.19–1.34)	0.56 (0.21–1.56)
Not employed†	39 (79.6)	57 (58.2)		
Group				
iCCAS	29 (59.2)	46 (46.9)	1.64 (0.81–3.30)	1.93 (0.91–4.11)
Usual care†	20 (40.8)	52 (53.1)		

Note: CI = confidence interval, iCCAS = interactive computer-assisted client assessment survey, OR = odds ratio.
 **p* < 0.05.
 †Reference category.

discussion of mental health problems and identification. Compared with Spanish-speaking patients (whose visits were mediated by a professional interpreter), patients who spoke English had more discussion and detection of mental health symptoms. This could reflect problems related to the presence of an interpreter or language barriers in spite of an interpreter. In addition, patients with college or higher education had higher rates of mental health symptom detection. This could represent differences in health literacy linked to educational attainment.^{38,39} Patients who were unemployed had a higher rate of clinician detection of mental health symptoms than their employed counterparts. This could be related to clinicians' vigilance to low income as a risk for common mental disorders, or decreased functionality of patients due to undetected and untreated mental illnesses. Collectively, these findings point toward a complex interaction between social conditions and access to mental health care for vulnerable ethnoracialized groups.⁴⁰ There is an undeniable need for intersectoral approaches to address mental health inequities with attention to these social determinants.

Our study contributes to the area of computer-assisted health risk assessments for mental health that started in 1960s^{41,42} and grew with the emergence of digital technologies. Our study is not only specific to common mental disorders in primary care, but the interactive tool employs a "dual engagement" conceptual model by engaging patients and clinicians simultaneously and generating 2 types of individualized reports. This is a step forward in the field of computer-assisted health risk assessments wherein the results of assessment reports are often generated automatically for clinicians only. Our search for such primary care, dual-engagement tools with a focus on depression identified only a handful of studies in the last decade. Most of these studies have used small convenience samples,^{24,43-45} with the exceptions of eCHAT, Promote Health and My Own Health Report (MOHR).^{22,46,47} Each of these studies, although valuable, either does not include a control group (eCHAT), is limited to academic settings with an above average income population (Promote Health) or is not widely available for use (MOHR). In this emerging area of patient-centred clinical informatics, the iCCAS study advances the field by recruiting an immigrant, refugee and ethnoracially diverse patient population from a community health centre and using rigorous randomized controlled design. Further developments could also draw from emerging work on patient portals (e.g., PatientSite) linked to electronic medical record systems.^{48,49}

Limitations

We examined the intervention by collaborating with a single community health centre, which may limit generalizability. Further, availability of the tool only in English and Spanish allowed recruitment of a select group of patients at the clinic. The small sample size limited our ability to examine the intervention effect by sex, ethnicity or comorbidity. We could not collect data concerning nonresponders. The response rate of 78%, however, enhances confidence in the reported results. Participating providers were potentially more likely to ask

about mental health in the usual care group because of training for the study and non-masking of the intervention. However, because these biases diminish the group difference in outcomes, our estimates of the intervention effect are likely to be underestimates. Possible reasons for the lack of intervention effect on detection or case-finding warrants some discussion. On one side, our sample size was small; on the other, we collected data on the detection rates through chart reviews, which are a weak source of information because clinicians often take limited notes in charts. In addition, our review of charts was limited to the index visit, but case-finding for mental illnesses often requires multiple assessments, especially for patients from diverse cultural backgrounds and in need of interpreters to communicate with their clinicians. Further qualitative work (forthcoming) could assist in unpacking the underlying mechanisms through which the discussion rates improved.

Conclusion

Computer-assisted interactive and user-friendly health risk assessment tools in patients' languages of choice hold potential to improve timely discussion on common mental disorders among vulnerable immigrant communities served by the community health centre model. Such discussions are in turn anticipated to improve detection rates. Future research with larger samples and multiple primary care sites is needed to enhance the generalizability of the reported results.

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