Validation of Non-Smoking Status by Spouse Following a Cessation Intervention

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Background: Following cessation interventions, self-reported smoking abstinence with biochemical verification is the “gold standard” for defining outcomes. Because obtaining biochemical verification is challenging in community studies, we compared self-reported cessation among smokers completing treatment to the smoking status reported by each participant’s spouse or proxy.

Method: Participants were smokers who had reported quitting 12 months after a cessation intervention. Participants had either attended a smoking cessation clinic or they were patients seen by physicians who had recently participated in a cessation-training program. Proxies living with these participants were interviewed by telephone to ask about their partner’s smoking status. We compared the participants’ responses to those from their spouses.

Results: At 12 months, 346 of 1423 baseline smokers had quit; 161/346 reported non-smokers were called and 140 proxies were interviewed. The participants averaged 51 years of age, 69% were women. At baseline, the mean number of cigarettes smoked per day was 20.1 (SD = 9.9) and the average number of quit attempts was 2.4 (SD = 1.2). Cessation methods used were medical advice (21%) and/or pharmacotherapy (79%). Of the 140 spouses interviewed, only 10 (7.1%) reported that their partners were currently smoking.

Conclusions: Proxy-reported data on smoking status could be used to validate self-report.

Background

The tobacco epidemic remains a major global public health threat and requires multiple strategies for tobacco control that includes not only the enactment of public policies, but also the promotion of smoking cessation services at the individual level (WHO, 2003). These strategies are intended to reduce the number of present and future smokers and related morbidity and mortality (Knofino et al., 2012).

Efficacy of a clinical smoking cessation intervention is evaluated 6 to 12 months after implementation by the Russell Standard criteria that were established to guide outcome assessments. These criteria require self-report of smoking status and biochemical verification of abstinence among others (West, Hajek, Stad & Stapleton, 2005). However, obtaining biochemical verification of abstinence is challenging in community studies and especially costly in low- and middle-income countries.

In public health surveys, self-reported status is widely used to report on population smoking prevalence (Wong, Leatherdale, Malaison & Hammond, 2012; Yeager & Krosnick, 2010). Self-report smoking status has been used as the principal population metric and misclassification was reported to be only 2% in a large US sample (Caraballo et al., 1998; Yeager & Krosnick, 2010) and 8.4% in a Canadian study (Wong et al., 2012). Self-report smoking status has also been used successfully in internet-based surveys (Ramo, Hall & Prochaska, 2011), among pregnant women (Kvalvik et al., 2012), and patients with chronic medical conditions (Ismail, Gill, Lawton, Houghton & MacFarlane, 2000; Wilson, Elborn, Fitzsimons & McCrum-Gardner, 2011). The use of self-report smoking status alone has been questioned as a definitive cessation outcome in a systematic review (Connor Gorber, Schofield-Hurwitz, Hardt, Levasseur & Tremblay, 2009), among patients with...
Participants and Recruitment

Potential participants for this study were selected from (i) 520 smokers who attended a smoking cessation clinic and (ii) 1,378 smoking patients seen by private-practice physicians who had participated in an educational program to help their patients quit from March 2009 to July 2011. Patients treated at the smoking cessation clinic were referred by their physicians and consented for follow-up telephone calls. The patients recruited from private practices were recruited from lists of patients seen by the physicians participating in the study, were called to ascertain smoking status and confirmed smokers were randomly selected and invited to participate in the study by responding to the surveys. Eligibility criteria for the current investigation included reporting continuous abstinence non-smoking status at 12 months after the cessation intervention and having a proxy (spouse or other household member) who could be phoned to answer questions about the participant’s quitting process and smoking status. All proxies of participants recruited from the tobacco cessation clinic and a 10% random sample of proxies for the private practice patients were selected for telephone interviews.

Methods

Setting

This study was based in the smoking cessation clinic at a university hospital primary care program and in selected internal medicine private practices located in Buenos Aires, Argentina. Participants were evaluated 12 months after they had completed the cessation intervention.

Participants and Recruitment

Interventions

The intervention at the clinic consisted of 8 to 12 weeks of individual treatment based on a cognitive behavioural approach, options for available pharmacotherapy, and support by clinicians. The patients recruited from private practices were seen by physicians who took part in a study aimed to test the effectiveness of an educational program to teach them cessation counselling techniques, referrals to services, and use of pharmacotherapy (University of California San Francisco, 2004).

Procedures

All patients received a telephone call 12 months after the date on which they had completed the cessation intervention or visited their physician. Continuous smoking abstinence was ascertained and only self-reported non-smokers were included in the current study. The spouses or persons who lived with the participant from the tobacco cessation clinic and a 10% random sample of patients from the private practices, who reported having quit smoking, were interviewed by telephone. Contacted proxies were asked about the smoking status and quitting process of their partner/spouse. The study protocols were approved by an NIH (National Institutes of Health) approved Institutional Review Board Centro de Investigación Clinica y Educación Medica (CEMIC).

Data Analysis

We compared the responses of the proxy respondents (mostly spouses) to the responses of the study participants. Data were analyzed using SAS and descriptive statistics reported means and standard deviations. Tobacco use was dichotomized as smoker or non-smoker with continuous abstinence. We reported response percentages of participants and their proxies.

Results

At 12 months, 1,423 of 1,898 participants were contacted (75% of those eligible) to assess their smoking status and 346 reported being non-smokers. Of those 346 participants, 172 were called again to confirm their smoking status and 161 reported being non-smokers. We asked to interview a proxy for each of these 161 non-smokers and 140 were interviewed. We were unable to reach the other 21. The 140 self-reported non-smoking participants averaged 51 years of age, 69% were women, and 49% had 12 years or more of education (see Table 1). The mean number of cigarettes per day at baseline survey was 20.1 (SD = 9.9) and the average number of quit attempts in the previous year was 2.4 (SD = 1.2). Reported cessation methods included physician advice or behavioural intervention only (21%), bupropion (56%), nicotine replacement therapy (20%), and varenicline (3%).

At 12 months, these 140 participants reported that they remained continuously abstinent of smoking but of the 140 proxies interviewed, 10 (7.1%) reported that their
Table 1
Characteristics of 140 smokers reporting abstinence 12 months after cessation intervention, 2009–2011, Buenos Aires, Argentina

|                          | Cessation Clinic at University | Private Practices 27 | Total Sample 140 |
|--------------------------|-------------------------------|----------------------|------------------|
| Age in years             |                               |                      |                  |
| 20–39                    | 28 (24.8)                     | 7 (25.9)             | 35 (25.0)        |
| 40–49                    | 16 (14.1)                     | 6 (22.2)             | 22 (15.7)        |
| 50–59                    | 37 (32.7)                     | 3 (11.1)             | 40 (28.6)        |
| ≥ 60                     | 32 (28.3)                     | 10 (37.0)            | 42 (30.0)        |
| Women                    | 77 (68.1)                     | 19 (70.3)            | 96 (68.5)        |
| Education in years       |                               |                      |                  |
| 6 or less                | 12 (10.7)                     | 4 (14.8)             | 16 (11.4)        |
| 7–12 years               | 45 (39.2)                     | 10 (37.0)            | 55 (39.2)        |
| ≥ 13 years               | 56 (50.0)                     | 13 (48.1)            | 69 (49.2)        |
| Cigarettes per day       |                               |                      |                  |
| 1–9                      | 3 (2.6)                       | 7 (25.9)             | 10 (7.1)         |
| 10–19                    | 29 (25.7)                     | 9 (33.3)             | 38 (27.1)        |
| More than 20             | 81 (71.7)                     | 11 (40.7)            | 84 (65.7)        |
| Cessation Therapies Used |                               |                      |                  |
| Nicotine Patch           | 10 (8.9)                      | 1 (3.7)              | 11 (7.9)         |
| Nicotine gum             | 17 (15.2)                     | 0                    | 17 (12.1)        |
| Bupropion                | 78 (68.7)                     | 1 (3.7)              | 79 (56.4)        |
| Varenicline              | 4 (4.6)                       | 0                    | 4 (2.7)          |
| Behavioural therapy      | 4 (4.6)                       | 25 (92.6)            | 29 (20.7)        |
| Self-reported Non-smoking status at 12 months | 113 (100) | 27 (100) | 140 (100) |
| Proxy report of non-smoking status | 103 (91.2) | 27 (100) | 130 (92.9) |

partner/household member was a current smoker. For all 10 discordant responses between participant and proxy, the participant was from the cessation program in the university hospital. These 10 patients averaged age of 61 years and smoked an average of 21 cigarettes per day at baseline; four of them had completed 7 years of education, six had 12 or more years of education, and six were women.

Discussion

Results from this report imply that proxy respondents report on smoking status could be used to validate self-reported results following a smoking cessation intervention in low- and middle-income countries in place of more costly biochemical validation. Although only 7% of proxies reported discordant smoking status from that of study participants, the added effort of an additional follow-up with a proxy can provide additional methodological strength to a study.

Use of biochemical validation has been considered the gold standard for the past 30 years in smoking cessation studies because it lends a methodological rigour in ascertaining outcomes. However, the operational challenge of collection and cost of testing samples inhibit formal evaluations of cessation programs. The strategy of asking the spouse or proxy, when available, offers a viable alternative to validate cessation status and may serve as a practical substitute for the more expensive biochemical validation that requires obtaining a biological sample. In our study, participants were selected after they reported having quit smoking in two previous telephone calls, so the strength of the strategy of asking proxies is that it is useful for identifying study participants who avoid saying they continue to smoke.

Our data were collected in the setting of a large urban centre and the results are only applicable to persons with a spouse who is willing to be contacted and respond to the study interviewer. However, one could possibly extend these findings by prospectively identifying a person who will know a participant’s smoking status and contact that person as the proxy. Furthermore, most of the participants were heavier smokers reporting more than 20 cigarettes per day prior to the intervention. It is interesting that all discrepant proxy reports were on patients in the more intensive cessation program.

One limitation of this study is that the conclusions are based exclusively on the participants and proxy’s self-reported answers without biochemical verification. However, the use of proxies has been used in public health studies and those results suggest that proxies can be used in smoking cessation studies. (Barnett et al., 1997; Gilpin et al., 1994; Kolonel et al., 1977; Mak et al., 2005).
utility of proxies for validation of their household members smoking status rests on the assumption that proxies provide more truthful responses. (Chen et al., 1995; McLaughlin, Dietz, Mehl & Blot, 1987). We were also unable to locate 21 proxies and if all had reported that the study participant was smoking, the rate of discordant results would be 19% (31 of 161).

In conclusion, our study suggests that proxy-reported data on smoking status could be used to confirm self-reported results in smoking cessation trials in low- and middle-income countries in place of biochemical verification but more research is needed, especially a study design that includes self-report, proxy-report, and biochemical validation.

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Conflict of Interest
None.

Ethical Standards
The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008.

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