

**ORIGINAL RESEARCH**

# Use of Mobile Health (mHealth) Tools by Primary Care Patients in the WWAMI Region Practice and Research Network (WPRN)

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**Purpose:** The purpose of this study was to determine the prevalence of mobile health (mHealth) use among primary care patients and examine demographic and clinical correlates.

**Methods:** Adult patients who presented to 1 of 6 primary care clinics in a practice-based research network in the northwest United States during a 2-week period received a survey that assessed smartphone ownership; mHealth use; sociodemographic characteristics (age, sex, race/ethnicity, health literacy); chronic conditions; and depressive symptoms (2-item Patient Health Questionnaire). Data analysis used descriptive statistics and mixed logistic regression.

**Results:** Of 918 respondents (estimated response rate, 67.4%), 55% owned a smartphone, among whom 70% were mHealth users. In multivariate analyses, smartphone ownership and mHealth use were not associated with health literacy, chronic conditions, or depression but were less common among adults >45 years old (adjusted odds ratio, 0.07–0.39;  $P < .001$ ). Only 10% of patients learned about mHealth tools from their physician, and few (31%) prioritized their provider's involvement.

**Conclusions:** Use of mHealth technologies is lower among older adults but otherwise is common among primary care patients, including those with limited health literacy and those with chronic conditions. Findings support the potential role of mHealth in improving disease management among certain groups in need; however, greater involvement of health care providers may be important for realizing this potential. (J Am Board Fam Med 2014;27:780–788.)

**Keywords:** Chronic Disease, Mobile Health, Practice-based Research, Primary Health Care

Mobile health (mHealth), which refers to the application of mobile or wireless communication technologies to health and health care,<sup>1</sup> barely ex-

isted 5 years ago; it has since emerged as a multi-billion-dollar industry.<sup>2</sup> Recent data suggest that >90,000 consumer smartphone health applications (“apps”) are now available for download.<sup>3</sup> Few of these have been the subject of scientific study, including their potential risks,<sup>4</sup> or to US Food and Drug Administration review or approval.<sup>1</sup>

According to estimates from the Pew Internet & American Life Project, in 2012 nearly one third of mobile phone owners, and more than half of smartphone owners, have used their phone to look up health information—a rate that has nearly doubled since 2010.<sup>5</sup> In Pew's general population survey, mHealth use was more common among individuals <50 years old, Latinos, African Americans, and those with higher socioeconomic status.

The potential for mHealth tools to represent an important advancement in health care, in particular for chronic disease care, has been recognized.<sup>1</sup>

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Whereas an estimated 69% of the US adult population track at least 1 health indicator, such as activity, weight, or symptoms, only about 20% of those use technology to facilitate their tracking.<sup>6</sup> Although health tracking is significantly more common among people living with chronic conditions, mHealth use has been reported as significantly less common among these individuals.<sup>5,6</sup> Primary care is the setting in which most chronic disease care occurs, and although there some literature describes mHealth tools available to facilitate chronic disease management, the actual patterns of mHealth use among patients in primary care settings have not been documented.<sup>7-9</sup> To address this gap in knowledge, this study aimed to determine the prevalence of mobile phone ownership and mHealth use among primary care patients in community-based clinics in the WWAMI region Practice and Research Network (WPRN) and to examine demographic and clinical correlates of use.

## Methods

### Setting

The WPRN is a primary care practice-based research network in the 5-state WWAMI region: Washington, Wyoming, Alaska, Montana, and Idaho. The 6 sites that participated in the study came from 4 of these states (Washington, Wyoming, Alaska, and Idaho) and included 4 hospital-associated outpatient practices, 1 office practice, and 1 federally qualified health center. The clinics serve many low-income patients; the proportion of patients who are uninsured or receiving Medicaid ranges from 22% to 62%. A champion at each clinic site, who had the opportunity to provide feedback on the survey design, assisted in study coordination, facilitated implementation of the study procedures, and returned completed surveys to the investigators, was engaged throughout the project. Participating sites received an administrative stipend of \$500.

### Participants and Procedures

This study used a card study method, an established means of collecting data in primary care settings, described by Westfall and colleagues<sup>10</sup> as a methodology used to conduct a prevalence study by collecting data from patients who consecutively present to a clinic. All adult patients ( $\geq 18$  years old) seen for a visit in one of the participating sites

during a 2-week period in June 2013 were eligible to participate. When checking in for their appointments at the front desk, patients were given a 1-page questionnaire designed to be completed in  $< 5$  minutes. The questionnaire was anonymous, and patients were informed that participation was voluntary and would not affect their health care. Participants returned the surveys to a collection box in the waiting area to ensure anonymity. Each questionnaire had a unique tracking number that was used to facilitate estimation of the response rate. Each day, front desk staff recorded the tracking number of the first and last questionnaire distributed for the day. If a participant declined to complete the questionnaire, the questionnaire form was discarded or placed in the collection box blank to avoid reuse. Questionnaires that were not distributed to patients were returned to the investigators after the completion of data collection. This study was considered minimal risk and was granted an exempt determination by the institutional review board at the University of Washington.

### Measures

#### Dependent Variables

Two questions were adapted from the Pew Internet & American Life project to assess mobile phone and smartphone ownership.<sup>5</sup> Mobile health use was assessed by asking participants if they have ever used their phone to “find health or medical information,” “download or use a health ‘app,’” or “track or manage a health issue (your diet or weight, activity, mood, blood pressure, etc.)” Participants were coded as mHealth users if they reported at least one of these uses and endorsed both mobile and smartphone ownership. Mobile health users indicated the frequency of use and completed 5 additional items about their mHealth use. The questionnaire included skip logic so that participants who did not endorse mobile phone ownership were not asked about smartphone ownership or mHealth use and were coded as “no” for these items. Participants who did not endorse smartphone ownership were not asked about mHealth use and were coded as not using mHealth tools. Regardless of phone ownership or mHealth use, all participants were asked if they use an Internet patient portal, how useful they believe receiving health materials from their provider on their phone would be, and their comfort with the security of

health information that is shared with their doctor or stored by a third party.

### Independent Variables

Demographic information included participants' age, sex, and self-reported race/ethnicity. Patients reported whether they had any of the following common chronic medical conditions: high blood pressure, diabetes, heart disease, asthma, chronic pain, depression, or any other chronic condition not listed. The 2-item Patient Health Questionnaire was used to assess the level of current depressive symptoms. Scores on the Patient Health Questionnaire range from 0 to 6, and a cutpoint of 3 has been used to identify patients with probable major depression, with high sensitivity and specificity.<sup>11,12</sup> Health literacy was assessed with 3-item instrument that has been validated against established measures of health literacy in a variety of settings.<sup>13–15</sup> Each item was scored from 1 to 5, and the scores were summed to yield a total score ranging from 3 (poor health literacy) to 15 (no health literacy limitations). Consistent with prior research, scores were coded dichotomously as having no health literacy limitations (score of 15) versus any limitation (scores of 3–14).<sup>16,17</sup>

### Data Analysis

Descriptive statistics, including frequencies, means, and standard deviations, were used to characterize patients' demographic and clinical features, mobile and smartphone ownership, and mHealth use. Associations between patient characteristics and technology ownership and use were examined with both unadjusted and adjusted mixed logistic regression models, clustered by clinic site. Specifically, separate mixed effects logistic regression models estimated the unadjusted associations between each patient characteristic (age, sex, race/ethnicity, health literacy, chronic conditions, and depression) and each dependent variable (smartphone ownership and mHealth use). Subsequently, for each dependent variable, a multivariate mixed logistic regression model was specified; all patient variables were entered simultaneously to estimate the independent associations between patient characteristics and smartphone ownership and use after adjusting for the effect of the remaining patient covariates. The primary analyses were conducted on the full cohort of patients in the sample. In sensitivity analyses, the regression models included patient cohorts that

**Table 1. Demographic and Clinical Characteristics of Patients Sampled from 6 Primary Care Practices in Washington, Wyoming, Alaska, and Idaho (June 2013)**

Characteristics	Patients, n (%)
Age, years (n = 858)	
18–24	130 (15.2)
25–34	183 (21.3)
35–44	164 (19.1)
45–54	163 (19.0)
55–64	140 (16.3)
≥65	78 (9.1)
Female sex (n = 859)	643 (74.9)
Race/ethnicity (n = 867)	
White	694 (80.1)
Native American	18 (2.1)
Asian/Pacific Islander	31 (3.6)
African American	21 (2.4)
Latino	42 (4.8)
Other/multiracial	61 (7.0)
Any health literacy limitation (n = 844)	526 (62.3)
Any chronic medical condition (n = 909)	575 (63.3)
Depression (n = 808)*	334 (41.3)

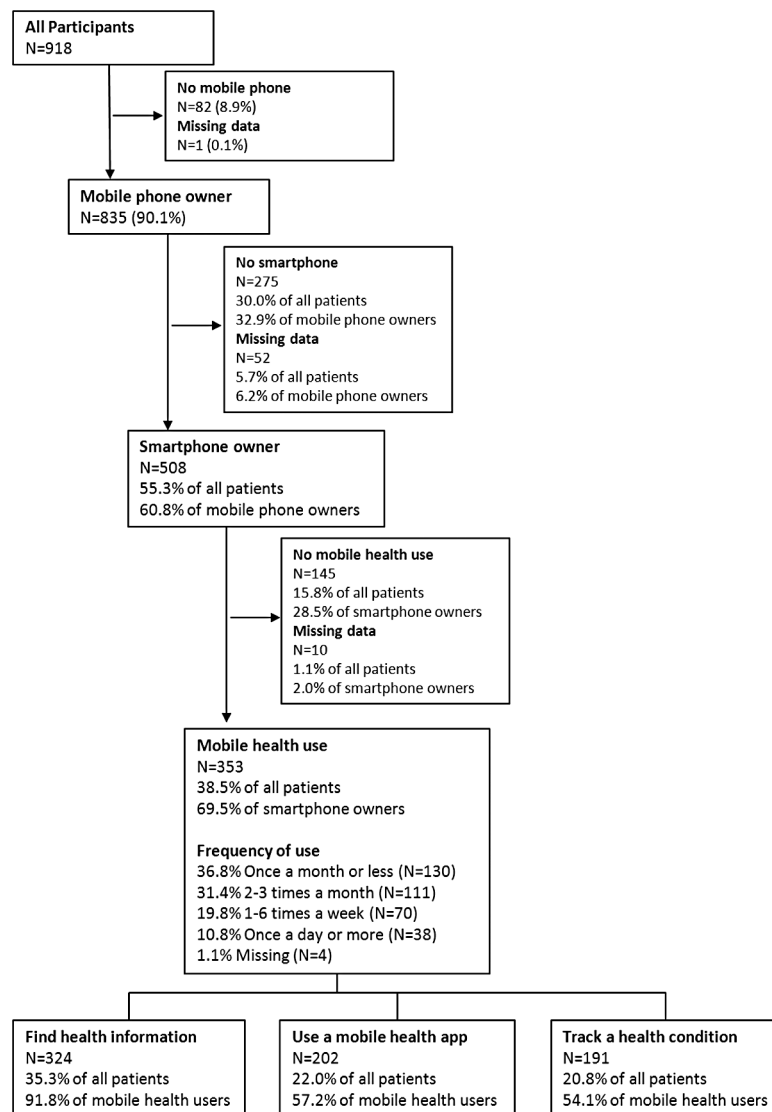
\*Depression is defined as a 2-item Patient Health Questionnaire score  $\geq 3$  or a history of depression endorsed as a medical condition.

were constructed based on the questionnaire skip logic (ie, regression models predicting smartphone ownership were restricted to mobile phone owners and models predicting mHealth use were restricted to smartphone owners). Multiple imputation (m = 40) was used to impute missing independent variables in multivariate models using “mi” commands in Stata software (StataCorp, College Station, TX). All variables were included in the imputation and assumed to have a multivariate normal distribution. Multivariate regressions were performed on each imputed data set, and results were combined using the rules defined by Rubin.<sup>18</sup>

### Results

Based on the tracking procedures, an estimated 1363 questionnaires were distributed and 918 were completed, for an estimated response rate of 67.4% (range, 31.8–84.0% across sites). Participants varied considerably in age; most (62%) had some health literacy limitations, most (63%) reported at least 1 chronic condition, and many (41%) reported current or prior depression (Table 1). As depicted in Figure 1, nearly all patients (91.0%, n = 835)

**Figure 1. Mobile phone ownership and mHealth use among primary care patients.**



owned a mobile phone and a majority owned a smartphone (55.3% of all patients [n = 508] or 60.8% of mobile phone owners). Almost 70% (n = 353) of smartphone owners reported mHealth use, which corresponds to 38.5% of all patients surveyed. Nearly all mHealth users reported using their phones to find health or medical information (91.8%, n = 324). A majority also reported downloading or using a health app (57.2%, n = 202) or using their phone to track or manage a health condition (54.1%, n = 191). As shown in Table 2, most patients who used mHealth tools did so infrequently: 69.1% (n = 241) reported use  $\leq 3$  times per month, whereas 10.8% (n = 38) reported use on a daily basis.

### **Types of Mobile Health Use**

Among 353 mHealth users, over one-third listed a general health app such as WebMD, iTriage, or Mayo Clinic as their favorite app (Table 2). The next most popular types were activity or fitness apps and weight loss or diet apps, which together were listed as the favorite by one quarter of the sample. Apps for chronic disease management, including those that monitor glucose or mood, were cited as a favorite by only 6 patients (2.6% of responses). Many respondents (30.4%) reported using an app for a short period of time then stopping, most frequently (48.0%) because it was too time-consuming. Patients rated appointment reminders as the most useful potential feature, fol-



**Table 2. Description of mHealth Use Among Primary Care Patients Who Report Such Use (n = 353)**

mHealth Use	Primary Care Patients Reporting Use
Mobile health use and type	
Find health information	324 (91.8)
Use health apps	202 (57.2)
Track or manage health condition	191 (54.1)
Frequency of mHealth use (n = 349)	
Once a month or less	130 (37.3)
2–3 times a month	111 (31.8)
1–6 times a week	70 (20.1)
Once a day or more	38 (10.9)
Favorite app (n = 235)	
General health app (eg, WedMD, iTriage, Mayo Clinic)	85 (36.2)
Activity/fitness (eg, My Fitness Pal)	34 (14.5)
Weight/diet (eg, Weight Watchers, LoseIt)	24 (10.2)
Reproductive (menstrual, pregnancy, or infant trackers)	19 (8.1)
Web search (eg, Google, Bing)	18 (7.7)
More than one favorite app	18 (7.7)
Other (eg, goal trackers, smoking logs)	14 (6.0)
Patient portal (eg, e-care, MyChart)	10 (4.3)
Medication (eg, pharmacy app, pill trackers)	7 (3.0)
Disease specific (eg, glucose monitoring, mood monitoring)	6 (2.6)
Ever stopped using an app after a short time (n = 310)	122 (39.4)
Reasons cited for stopping use of an app after a short time (n = 98)	
Took too much time	47 (48.0)
Didn't do what you wanted	32 (32.7)
Problem with login/password	6 (6.1)
Other	13 (13.3)
How mHealth users learned about apps (n = 277)	
Doctor/clinic	27 (9.8)
Family/friend	37 (13.4)
Website/Internet	150 (54.2)
Flier/mail/other ad	7 (2.5)
Other (eg, "app store")	56 (20.2)
Importance for PCPs to know about mHealth use (n = 311)	
Very important	37 (11.9)
Important	59 (19.0)
A little important	93 (29.9)
Not at all important	122 (39.2)
My doctor has recommended a health app (n = 322)	22 (6.8)

*Continued*

**Table 2. Continued**

mHealth Use	Primary Care Patients Reporting Use
How useful this feature would be on your phone*	
Appointment reminders (n = 318)	4.0 ± 1.4
Medication reminders (n = 312)	3.6 ± 1.7
General health information (n = 314)	3.5 ± 1.4
Track progress (eg, mood, weight) (n = 310)	3.5 ± 1.5
Help changing a habit (n = 308)	3.2 ± 1.7
Feedback on how I'm doing (n = 312)	3.2 ± 1.6
Tell doctor how I'm doing (n = 311)	3.2 ± 1.7
Stress management/coping (n = 310)	3.1 ± 1.7
Support group/social network (n = 308)	2.3 ± 1.8
Tell friend/family how I'm doing (n = 305)	1.9 ± 1.8

Data are number (%) or mean ± standard deviation.

\*Usefulness of the features was rated on a scale of 0 (least useful) to 5 (most useful).

PCPs, primary care providers.

lowed closely by medication reminders, general health information, and health tracking, whereas social features such as support groups and updating friends or family were rated as less useful (Table 2).

Much of the patients' mHealth use occurred outside of the context of health care (Table 2). Fewer than 10% of respondents learned about mHealth apps from their doctor or clinic; only 6.8% of respondents reported that their physician had recommended a health app to them. Respondents did not feel it was important for their physicians to know about their use of health apps: Nearly 70% reported that this was "not at all" or "a little bit" important.

#### **Demographic and Clinical Correlates of Smartphone Ownership and Mobile Health Use**

In unadjusted models, older patients and those with a chronic medical condition were significantly less likely to own a smartphone than younger patients and those without chronic conditions. Latino patients were significantly more likely to own a smartphone than whites (Table 3). In multivariate analyses, age was significantly independently associated with smartphone ownership, whereas Latino ethnicity and chronic conditions were no longer significantly independently correlated. Similar findings emerged from analyses restricted to mobile phone owners (Online Appendix Table 1).

**Table 3. Correlates of Smartphone Ownership Among All Patients**

	Unadjusted Models			Multivariate Model* (n = 866)	
	Patients (n)	OR (95% CI)	P Value	Adjusted OR (95% CI)	P Value
Age (years)	809				
18–24		Reference		Reference	
25–34		1.13 (0.65–1.95)	.68	1.13 (0.65–1.98)	.66
35–44		0.72 (0.41–1.25)	.24	0.77 (0.44–1.35)	.36
45–54		0.36 (0.21–0.60)	<.001	0.39 (0.22–0.68)	.001
55–64		0.18 (0.10–0.30)	<.001	0.21 (0.12–0.37)	<.001
≥65		0.06 (0.03–0.13)	<.001	0.08 (0.04–0.17)	<.001
Sex					
Male		Reference		Reference	
Female	811	0.87 (0.63–1.21)	.41	0.75 (0.51–1.09)	.13
Race/ethnicity	818				
White		Reference		Reference	
Native American		0.89 (0.33–2.45)	.83	0.78 (0.26–2.39)	.66
Asian/Pacific Islander		1.25 (0.56–2.79)	.58	0.74 (0.3–1.83)	.51
African American		2.61 (0.85–8.07)	.10	1.67 (0.51–5.47)	.39
Latino		2.04 (1.00–4.16)	.05	1.18 (0.55–2.53)	.67
Other/multiracial		1.05 (0.60–1.84)	.86	0.74 (0.41–1.37)	.34
Any health literacy limitation	797	0.87 (0.65–1.17)	.37	0.86 (0.62–1.2)	.38
Any chronic medical condition	857	0.5 (0.37–0.67)	<.001	0.74 (0.53–1.04)	.08
Depression	762	0.82 (0.61–1.10)	.18	0.81 (0.58–1.14)	.23

All models were adjusted for clustering within clinics.

\*The multivariate model reports results from a single model that includes all of the independent variables in the table. Multiple imputation was used for missing variables in the multivariate model.

CI, confidence interval; OR, odds ratio.

Like smartphone ownership, mHealth use was significantly associated with younger age in both unadjusted and multivariate models (Table 4), a pattern that was evident among analyses restricted to smartphone owners (Online Appendix Table 2). In unadjusted analyses, the presence of a chronic medical condition was associated with a significantly lower likelihood of mHealth use and Latino ethnicity was associated with a significantly higher likelihood of mHealth use, but neither pattern persisted in multivariate analyses or among analyses restricted to smartphone owners. Among smartphone owners, women were somewhat more likely than men to be mHealth users. Depression and health literacy were not associated with mHealth use.

### Discussion

This study found comparable or greater rates of mHealth use among primary care patients than previously reported in the general population. Smartphone ownership among primary care patients (55%) is approaching the penetration of In-

ternet access among primary care patients a decade ago (65%),<sup>19</sup> and our findings indicate that most patients with smartphones use mHealth tools. The extremely rapid adoption of mobile technologies among the general population is contributing to significant shifts in how people interact and how they seek information. This digital revolution is beginning to transform health care, presenting opportunities for improved service delivery as well as challenges for patients and providers who grapple with incorporating novel technologies into their daily lives and clinical workflows.

Our results, which suggest that patients may use mHealth to augment their use of formal clinical services, have important implications for health care providers. Of potential concern are the findings that few patients believe it is important for health care providers to know about their mHealth use. We did not assess reasons for this limited communication, and further research is warranted. Most mHealth users sought health information, and few used tools that were disease specific or otherwise connected to their health care providers.

**Table 4. Correlates of mHealth Use Among All Patients**

	Unadjusted Models			Multivariate Model* (n = 879)	
	Patients (n)	OR (95% CI)	P Value	Adjusted OR (95% CI)	P Value
Age (years)	822				
18–24		Reference		Reference	
25–34		1.00 (0.62–1.62)	.99	1.00 (0.61–1.62)	.99
35–44		0.64 (0.39–1.05)	.08	0.66 (0.4–1.1)	.11
45–54		0.31 (0.18–0.51)	<.001	0.32 (0.19–0.55)	<.001
55–64		0.14 (0.08–0.25)	<.001	0.16 (0.09–0.3)	<.001
≥65		0.05 (0.02–0.12)	<.001	0.07 (0.03–0.16)	<.001
Sex					
Male		Reference		Reference	
Female	825	1.28 (0.92–1.79)	.14	1.15 (0.79–1.66)	.47
Race/ethnicity	831				
White		Reference		Reference	
Native American		1.33 (0.50–3.52)	.57	1.16 (0.4–3.34)	.78
Asian/Pacific Islander		1.11 (0.52–2.37)	.79	0.72 (0.31–1.66)	.44
African American		0.93 (0.36–2.42)	.89	0.63 (0.23–1.74)	.37
Latino		2.29 (1.20–4.37)	.01	1.47 (0.73–2.93)	.28
Other/multiracial		1.30 (0.76–2.25)	.34	0.91 (0.5–1.64)	.75
Any health literacy limitation	810	1.00 (0.74–1.33)	.98	0.98 (0.7–1.36)	.89
Any chronic medical condition	870	0.60 (0.45–0.79)	<.001	0.91 (0.66–1.26)	.58
Depression	774	0.96 (0.71–1.29)	.78	0.95 (0.68–1.32)	.75

All models are adjusted for clustering within clinics.

\*The multivariate model reports results from a single model that includes all of the independent variables in the table. Multiple imputation was used for missing variables in the multivariate model.

CI, confidence interval; OR, odds ratio.

Given the type of tools now available, patients may not view physicians as a useful resource in advising them about mHealth tools and may prefer recommendations from their social network, peers, or other patients. Communication about mHealth tools may evolve as the tools themselves advance, with more tools allowing data-sharing with providers, or when health care systems offer mobile portals and apps directly to patients. However, other potential explanations warrant consideration. A recent report found that providers are suspicious of mHealth tools for patients; more than half of younger physicians have concerns that mHealth would foster too much patient independence, and nearly one quarter of such physicians discouraging use of mHealth tools.<sup>20</sup> Lack of communication between health care providers and patients about mHealth use represents a missed opportunity for providers to support patients' self-management activities.<sup>9</sup>

Several findings have implications for the design and development of the next generation of mHealth tools and for their integration into health

care systems. Consistent with research demonstrating that women use more health services and have greater health care help-seeking than men,<sup>21,22</sup> we found that women reported somewhat greater mHealth use. If this pattern is replicated, considering how technology design may be able to reduce gender-based disparities in health care engagement would be important. Among our sample, mHealth use was not associated with race/ethnicity, health literacy, or depression, suggesting that mHealth tools may reach populations traditionally underserved in health care systems and thus may have a role in reducing pervasive disparities in care. As expected, smartphone ownership and mHealth use lag among older adults, although this gap will narrow as the younger generations of digital natives age and contribute their skills with technology and their expectations of its role to their interactions with health care providers.

Similar to the Pew mobile health survey,<sup>5</sup> our unadjusted analyses revealed that mHealth use was less common among individuals with a chronic medical condition; however, this pattern did not

hold in multivariate analyses. Thus, the apparent lower use of mHealth among individuals with chronic diseases may be related to other patient characteristics (such as older age) rather than the presence of a chronic disease per se, a distinction that is relevant in considering the potential reach of mHealth tools. For example, patients who develop chronic diseases at younger ages may experience particular benefit from technologies that support improved disease self-management to prevent future disability and complications; however, most of the tools that patients commonly use now do not target disease management, implying that future development should prioritize disease management tools that meet patients' needs. In contrast, tools intended for older adults should account for age in their design, for example, by offering large print interfaces or similar design elements, regardless of whether they are intended for patients with or without chronic diseases.

Most patients using mHealth tools were infrequent users, and many reported stopping use after a short time, most often because it took too much time to use, suggesting that existing tools may have limited appeal for patients managing chronic diseases that require self-management activities to be sustained over time. The application of patient-centered design methods would help ensure that tools are effective at meeting needs identified by patients. Concurrently, research examining how best to incorporate mHealth tools into clinical practice settings, including how to integrate them into clinical workflows, will be important for successful dissemination of tools in health care settings.

Several study limitations deserve mention. Data were obtained from a cross-sectional survey and, therefore, all information was based on self-report. The study was conducted in the northwest region of the United States and included predominantly white patients; therefore it is unclear to what extent the findings generalize to other regions or populations of different racial/ethnic composition. Potential selection bias could have occurred if respondents differed systematically from the general clinic population, for example, in their general or technology literacy. No data about patients who were offered the survey but declined were available. It is possible that some eligible patients were not offered the survey; however, we have no reason to suspect any systematic omissions that would con-

tribute to selection bias. Although we cannot rule out the possibility of some selection bias, the overall response rate was high (67.4%); the sample included patients across a wide range of ages, many of whom had health literacy limitations; and our findings share many patterns similar to those from the national Pew survey of the general population.

Our results support the notion that mHealth tools are rapidly becoming an acceptable and scalable means of empowering patients, yet health care providers play a limited role in assessing or advising patients on their use of mHealth tools. This finding is particularly notable because the 2 features that patients rated as most likely to be useful—appointment reminders and medication reminders—are directly related to their use of medical care, suggesting that patients may be open to using mHealth tools to enhance their relationships with health care providers. Although data are sorely needed to establish the evidence base for mHealth tools and interventions, health care providers who routinely ask their patients about their mHealth use may be especially effective at eliciting important information about patients' self-management activities, which may help these providers to be more adept supporting chronic disease care. To fulfill the promise for transforming health care service delivery that mobile health offers, however, the next generation of mHealth tools will need to align the capabilities of mHealth technologies with effective models of health care delivery by providing well-integrated tools that address the needs of both patients and providers.

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**Appendix Table 1. Correlates of Smartphone Ownership Among Mobile Phone Owners**

	Unadjusted Models			Multivariate Model* (n = 783)	
	Patients (n)	OR (95% CI)	P Value	Adjusted OR (95% CI)	P Value
Age (years)	738				
18–24		Reference		Reference	
25–34		1.15 (0.60–2.20)	.67	1.14 (0.6–2.17)	.69
35–44		0.64 (0.34–1.21)	.17	0.69 (0.36–1.31)	.25
45–54		0.27 (0.15–0.49)	<.001	0.29 (0.16–0.54)	<.001
55–64		0.14 (0.08–0.27)	<.001	0.16 (0.09–0.31)	<.001
≥65		0.05 (0.02–0.11)	<.001	0.06 (0.03–0.14)	<.001
Sex					
Male		Reference		Reference	
Female	736	0.81 (0.56–1.17)	.26	0.69 (0.46–1.05)	.09
Race/ethnicity	743				
White		Reference		Reference	
Native American		0.93 (0.30–2.83)	.89	0.71 (0.21–2.46)	.59
Asian/Pacific Islander		1.45 (0.58–3.58)	.42	0.87 (0.3–2.51)	.80
African American		2.07 (0.67–6.41)	.21	1.27 (0.38–4.26)	.70
Latino		1.95 (0.90–4.20)	.09	1.05 (0.46–2.43)	.91
Other/multiracial		1.26 (0.66–2.37)	.48	0.86 (0.43–1.73)	.67
Any health literacy limitation	725	0.92 (0.67–1.27)	.62	0.92 (0.63–1.32)	.64
Any chronic medical condition	776	0.56 (0.41–0.77)	<.001	0.86 (0.6–1.25)	.43
Depression	690	0.84 (0.61–1.15)	.28	0.79 (0.55–1.16)	.23

All models are adjusted for clustering within clinics.

\*The multivariate model reports results from a single model that includes all of the independent variables in the table. Multiple imputation was used for missing variables in the multivariate model.

CI, confidence interval; OR, odds ratio.

**Appendix Table 2. Correlates of mHealth Use Among Smartphone Owners**

	Unadjusted Models			Multivariate Model* (n = 498)	
	Patients (n)	OR (95% CI)	P Value	Adjusted OR (95% CI)	P Value
Age (years)	473				
18–24		Reference		Reference	
25–34		0.83 (0.43–1.59)	.58	0.83 (0.42–1.61)	.58
35–44		0.72 (0.37–1.42)	.35	0.71 (0.35–1.43)	.33
45–54		0.41 (0.21–0.83)	.01	0.43 (0.21–0.89)	.02
55–64		0.26 (0.12–0.56)	.001	0.26 (0.12–0.6)	.001
≥65		0.17 (0.05–0.57)	.004	0.19 (0.05–0.68)	.011
Sex					
Male		Reference		Reference	
Female	471	1.77 (1.14–2.74)	.01	1.59 (0.99–2.55)	.06
Race/ethnicity	476				
White		Reference		Reference	
Native American		3.51 (0.43–28.75)	.24	3.02 (0.38–24.18)	.30
Asian/Pacific Islander		0.95 (0.35–2.59)	.93	0.88 (0.3–2.57)	.81
African American		0.44 (0.15–1.29)	.13	0.36 (0.11–1.1)	.07
Latino		1.77 (0.70–4.44)	.23	1.43 (0.55–3.73)	.47
Other/multiracial		1.49 (0.65–3.38)	.35	1.21 (0.51–2.85)	.67
Any health literacy limitation	464	1.2 (0.79–1.81)	.39	1.2 (0.77–1.88)	.43
Any chronic medical condition	493	0.95 (0.64–1.41)	.81	1.2 (0.77–1.88)	.41
Depression	436	1.28 (0.83–1.99)	.27	1.2 (0.75–1.94)	.44

All models are adjusted for clustering within clinics.

\*The multivariate model reports results from a single model that includes all of the independent variables in the table. Multiple imputation was used for missing variables in the multivariate model.

CI, confidence interval; OR, odds ratio.