Web-based Social Media Intervention to Increase Vaccine Acceptance: A Randomized Controlled Trial

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BACKGROUND: Interventions to address vaccine hesitancy and increase vaccine acceptance are needed. This study sought to determine if a Web-based, social media intervention increases early childhood immunization.

METHODS: A 3-arm, randomized controlled trial was conducted in Colorado from September 2013 to July 2016. Participants were pregnant women, randomly assigned (3:2:1) to a Web site with vaccine information and interactive social media components (VSM), a Web site with vaccine information (VI), or usual care (UC). Vaccination was assessed in infants of participants from birth to age 200 days. The primary outcome was days undervaccinated, measured as a continuous and dichotomous variable.

RESULTS: Infants of 888 participants were managed for 200 days. By using a nonparametric rank-based analysis, mean ranks for days undervaccinated were significantly lower in the VSM arm versus UC (P = .02) but not statistically different between the VI and UC (P = .08) or between VSM and VI arms (P = .63). The proportions of infants up-to-date at age 200 days were 92.5, 91.3, and 86.6 in the VSM, VI, and UC arms, respectively. Infants in the VSM arm were more likely to be up-to-date than infants in the UC arm (odds ratio [OR] = 1.92; 95% confidence interval [CI], 1.07–3.47). Up-to-date status was not statistically different between VI and UC arms (OR = 1.62; 95% CI, 0.87–3.00) or between the VSM and VI arms (OR = 1.19, 95% CI, 0.70–2.03).

CONCLUSIONS: Providing Web-based vaccine information with social media applications during pregnancy can positively influence parental vaccine behaviors.

WHAT’S KNOWN ON THIS SUBJECT: Many parents with concerns about childhood vaccines use the Internet and social media for vaccine information. The effectiveness of using Web-based vaccine information and social media to increase parental vaccine acceptance has not been evaluated.

WHAT THIS STUDY ADDS: By using a randomized control trial design, we found that a Web-based, social media intervention administered outside of the physician’s office can effectively improve childhood vaccine acceptance among pregnant women.
Between 10% to 15% of parents choose to delay or refuse 1 or more recommended vaccines for their children. This decision leaves children and their communities vulnerable to vaccine-preventable diseases. Parents who are hesitant to vaccinate their children also have complex information-seeking behaviors. They often start to weigh the risks and benefits of vaccination during pregnancy, seek information from many sources, and express interest in receiving vaccine information before routine well-child visits. Although physicians are trusted sources of health information for parents, vaccine-hesitant parents are inclined to distrust traditional sources of scientific authority and report using the Internet to gather information on vaccines.

Regardless of whether they are hesitant about childhood vaccination, parents who use the Internet to educate themselves must sift through vast amounts of vaccine information. Web sites range from government-sponsored, pro-vaccine resources that carefully present factual information to staunchly antivaccination Web sites that use anecdotes and social media to disseminate misinformation. Along this spectrum, numerous parenting message boards and blogs vigorously discuss vaccine-related topics. Exposure to antivaccine messages through social media appears to intensify parents’ worries and lower their intentions to vaccinate.

At the same time, social media may have the potential to allay parental vaccine concerns and improve immunization rates. An expert-moderated, interactive vaccine Web site could provide parents with a forum to voice their opinions, ask questions, and interact with other concerned parents and vaccine experts. This type of dynamic online environment could help build trust and combat misinformation. At present, the impact of using social media to improve vaccine acceptance is not known. In addition, we are not aware of any interventions targeting vaccine hesitancy that have effectively changed parental vaccination behavior in the United States. As part of the Colorado Vaccine Social Media study, we conducted a randomized controlled trial (RCT) to evaluate the effectiveness of Web-based vaccine information and social media interventions to increase vaccine acceptance.

METHODS

Study Overview
Between September 2013 and July 2016, we conducted a single-center RCT of vaccine information and social media interventions designed to reduce undervaccination among infants of women recruited during pregnancy. Our primary outcome was days undervaccinated from birth to age 200 days. We hypothesized that infants of women exposed to interventions during pregnancy will have less vaccine delay than infants receiving usual pediatric care.

Participants were randomly assigned to 1 of 3 groups: a Web site with vaccine information and social media components (VSM); a Web site with vaccine information (VI); or usual care only (UC). The Kaiser Permanente Colorado (KPCO) institutional review board approved this study.

Study Setting, Participants, and Randomization
All participants were members of the KPCO health plan, a nonprofit managed care organization serving ~628,000 individuals. Each year, ~5000 pregnant women and ~130,000 children receive healthcare at KPCO clinics.

Recruitment was conducted in 6-week waves between September 2013 and October 2015. At the beginning of each wave, we used electronic health records to identify pregnant women in the third trimester of pregnancy (13–6 weeks from delivery). Women had to be age 18 years or older, English speaking, have Internet access, and be enrolled in the KPCO health plan. Pregnant women were ineligible if they had a diagnosis for fetal death, miscarriage, or congenital anomaly. Eligible women received a combination of postcards, e-mails, and phone calls to elicit participation. Informed consent was obtained online by using a secure encryption program.

After consent, participants were administered a baseline survey to assess demographics and Internet use. Participants were also administered the Parent Attitudes and Childhood Vaccines (PACV) screening survey, which is a validated, 15-item instrument that assesses vaccine hesitancy on a scale of 0 to 100. Consistent with previous studies, participants scoring ≥50 were classified as “vaccine hesitant,” whereas participants with scores <50 were “nonhesitant.” To ensure balance across study arms, randomization was conducted independently within the 2 strata of hesitancy. Because only a small fraction of Web site visitors actively engage in social media activities, we used a randomization allocation ratio of 3:2:1 across the VSM/VI/UC arms to facilitate interaction. Randomization was done by an unblinded statistician using the SAS/STAT procedure Proc Plan. Although the participants and study team were not blinded to study arm assignment, the study team was blinded to participants’ hesitancy status.

To enhance security and prevent contamination, participants randomly assigned to the VSM and VI arms were required to create a login and password for the Web site. Infants of participants were managed for 200 days after birth to assess vaccination
status. To reflect how a Web-based resource would be used in practice, individuals in the VSM and VI arms were given access to the Web site but were not required to visit it.

Interventions
Separate interventions were developed for the VSM and VI arms. The theoretical basis for the VSM intervention was the multidirectional communication model, a social marketing strategy with 3 components. Component 1 is a standard, top-down process in which Web site developers create and present content to users. Component 2 is a bottom-up process that allows users to create content and interact with Web site developers. Component 3 is a side-to-side process in which users can interact with each other and share information. This model is intended to empower users by allowing them to become active participants in the communication process, thereby eliciting positive health behavior changes.

In contrast to the VSM intervention, the VI intervention only included the top-down component of the model.

The interventions were designed and pilot tested by using an adapted mental-models approach that included focus groups, individual interviews, surveys, and usability testing with parents and pregnant women. Details of this process have been described previously. In brief, our study team first developed the factual vaccine content, guided by the Health Belief Model and Theory of Planned Behavior.

We sought to present content that accurately represented the risks and benefits of vaccination, including information on vaccine-preventable diseases, vaccine safety, vaccine laws, the recommended immunization schedule, vaccine ingredients, vaccine development, and basic immunology. Information was labeled and arranged into short, easy-to-read sections, guided by best practices in risk communication and Web site design. Sources of information were carefully referenced and hyperlinked to help convey transparency and credibility. The information was focused on encouraging parents to receive recommended vaccines on time. Participants in the VSM and VI arms had access to the same base vaccine content.

In addition to vaccine content, participants in the VSM arm had access to social media technologies that included a blog discussion forum, chat room, and “Ask a Question” portal through which participants could directly ask our experts questions about vaccination. These technologies were designed to facilitate engagement and reinforce the factual content. Experts included a pediatrician, a vaccine safety researcher, and a risk communication specialist. Each month, the research team created 1 to 2 blog posts covering topics such as new vaccine safety research, vaccine-preventable disease outbreaks, changes in immunization policy, and the importance of adhering to the recommended immunization schedule. Posts were either text or audio (podcasts), and participants could contribute comments and ask questions. Each month, we hosted online chat sessions in which participants could engage in real-time conversations with experts. Participants were also encouraged to submit questions privately through e-mail; the team provided personalized responses within 2 business days. All participants in the VSM arm received monthly newsletters to encourage Web site participation and highlight new Web site content.

All interactive components were moderated to prevent bullying, disclosure of personal identifying information, and abusive language. Responses to comments and questions adhered to a consistent communication framework designed to convey dedication, expertise, and honesty. Intervention details (including the Hoffman’s template for intervention description and replication checklist and guide and screenshots of the intervention Web sites) are included in the Supplemental Information.

Outcome
Vaccination Status: Days Undervaccinated and Up-to-Date Status
Immunization data for infants were extracted from the electronic health record. We assessed vaccination status over the first 200 days of age to cover a majority of the recommended infant vaccines and minimize loss to follow-up. We assessed the following 6 vaccines recommended by the Advisory Committee on Immunization Practices: hepatitis B; rotavirus; diphtheria-tetanus-acellular pertussis; Haemophilus influenzae type b; pneumococcal conjugate vaccine; and polio. Our primary outcome was days undervaccinated, a continuous metric that measures differences between the time when vaccine doses were actually administered and when the doses should have been administered according to the Advisory Committee on Immunization Practices schedule. For example, the first dose of diphtheria-tetanus-acellular pertussis is due at age 2 months but is not considered late until age 92 days. Days undervaccinated for this dose would begin accruing on day 93. Of note, infants who did not receive the birth dose of hepatitis B vaccine were not considered delayed; days undervaccinated for the first dose of hepatitis B vaccine started accruing.
Days undervaccinated was analyzed both as a continuous measure and as a dichotomous variable (up-to-date with no delays, yes or no). The dichotomous variable of days undervaccinated was labeled as up-to-date vaccination status, representing a clinically meaningful measure for providers. Infants with 0 cumulative days undervaccinated at age 200 days were considered up-to-date. As a subanalysis, we assessed up-to-date status for measles-mumps-rubella (MMR) vaccine among infants with at least 489 days of follow-up, when days undervaccinated for the first dose of MMR begins to accrue.

**Statistical Methods**

The study was powered to detect a clinically meaningful odds ratio (OR) of 1.8–2.2 for up-to-date vaccination status between the study arms. For this effect size, we required 900 participants on the basis of an anticipated baseline vaccine hesitancy of 20%, a 3:2:1 allocation ratio, and a 2-sided α of .05. An a priori P value of <.05 was considered statistically significant. We conducted a modified intent-to-treat analysis by keeping the study arm assignment but excluding infants without outcome data from the analysis. Infants of participants were excluded if they disenrolled from KPCO after birth, enrolled after age 60 days, were not continuously enrolled during their follow-up period, or were not using KPCO for primary health services. These exclusions help ensure complete ascertainment of vaccination data. Participants were also excluded if they requested to be removed from the study or experienced a fetal demise or death of the child. Although we screened infants for documented contraindications to vaccines, premature infants were not excluded because they are to receive vaccines according to the recommended schedule.31

Days undervaccinated and up-to-date vaccination status were assessed from birth to age 200 days. Because of the skewed distribution of days undervaccinated, we used a nonparametric analysis and rank transformation approach.32 We ranked the days undervaccinated for all infants and then compared the mean ranks across study arms using 1-way analysis of variance. Up-to-date vaccination status was analyzed by using logistic regression to estimate ORs and associated 95% confidence intervals (CIs). Logistic regression was also used to assess MMR status among children with at least 489 days of follow-up. Data were analyzed with SAS 9.4 software (SAS Institute, Inc, Cary, NC).

**RESULTS**

**Study Participants and Baseline Characteristics**

A total of 1093 pregnant women were recruited into the study (Fig 1). By using a 3:2:1 randomization ratio, 542 participants were randomly assigned to VSM, 371 were assigned to VI, and 180 were assigned to UC. Baseline characteristics were evenly distributed across study arms (Table 1). Mean maternal age at enrollment was 31.6 years, and a majority of the population was white (86.9%) and college educated (82.8%). At enrollment, 14.1% of the population was classified as vaccine hesitant on the basis of the PACV screener, and >62% of participants reported using the Internet for health information at least weekly. Median vaccine hesitancy scores were 13, 17, and 15 for the VSM, VI, and UC arms, respectively (P = .44).

A lack of outcome data led to the exclusion of 205 infants (18.8%); infants were excluded because they were disenrolled from KPCO after birth (n = 16), enrolled after age 60 days (n = 21), had incomplete follow-up because of loss of insurance (n = 159), were not using KPCO for primary care services (n = 5), or had a fetal demise (n = 4) (Fig 1). There were no infants with documented contraindications to vaccines. Loss to follow-up ranged from 17.2% to 19.9% across study arms. Among participants lost to follow-up, median vaccine hesitancy scores were not significantly different across the arms (P = .97).

**Usage and Interaction**

Of 739 participants in the VSM and VI arms with 200 days of follow-up, 259 (35.0%) visited the Web sites at least once, with a mean of 1.8 (SD = 1.7) and range of 1 to 15 visits. Of 75 vaccine-hesitant participants, 33 (44.0%) visited the Web sites compared with 226 (34.0%) of the 664 nonhesitant participants. Over the study period, the VSM Web site offered 59 blog entries and 31 chat sessions. Participants in the VSM arm (n = 442) contributed 90 comments and questions. A majority of the interaction was between participants and the research team rather than between participants.

**Effectiveness**

Mean ranks for days undervaccinated were 438.5, 443.0, and 465.4 for the VSM, VI, and UC arms, respectively. Infants in the VSM arm had a lower mean rank for days undervaccinated than infants in the UC arm (difference = −26.9; P value = .02; Table 2). Mean ranks did not differ significantly between the VI and UC arms or the VSM and VI arms.

The proportion of infants up-to-date at the end of follow-up were 92.5, 91.3, and 86.6 for the VSM, VI, and UC arms, respectively. Infants in the VSM arm were more likely to be up-to-date at age 200 days than infants in the UC arm (OR = 1.92; 95% CI, 1.07–3.47; Table 3). Up-to-date status did not differ significantly between the VI and UC arms or the VSM and VI arms. The interaction between study arm and baseline vaccine hesitancy status was not statistically significant.
Among all infants enrolled from birth to age 200 days in KPCO (n = 8877) during the study period, the rate of up-to-date status was 86.3%, suggesting that the UC infant population was representative of the overall KPCO infant population.

For the MMR subanalysis, there were 776 (71%) infants with at least 489 days of continuous follow-up. The proportion of infants who received MMR by the end of follow-up were 95.6, 95.5, and 91.8 for the VSM, VI, and UC arms, respectively. Although none of the study arm comparisons were statistically significant, infants in the VSM and VI arms were ~2 times more likely to have received MMR than infants in the UC arm (Table 4).

**DISCUSSION**

This RCT of a Web-based vaccine information and social media intervention had a positive impact on early childhood immunization. Pregnant women exposed to the VSM arm were more likely to vaccinate their infants on time than participants receiving UC. These results suggest that interactive, informational interventions administered outside of the physician’s office can improve vaccine acceptance.

The authors of previous research have shown that the timing of vaccine information receipt is important to parents.5,7 Traditionally, vaccine information is provided to parents at well-child visits, although some parents make their vaccination decisions during pregnancy. In the
absence of accurate information during pregnancy, parents may tend to rely on the Internet, which may expose them to misinformation. We found that providing accurate online information with interactive technologies during pregnancy has a positive impact on infant-vaccine acceptance. Providing parents with information debunking vaccination falsehoods, such as the link between the MMR vaccination and autism, can cause vaccine-hesitant parents to become more entrenched in their antivaccination views and reduce their intentions to vaccinate. However, this backfire effect is likely modified by additional factors, such as the source, wording tone, and timing of information. Our intervention demonstrated that parental vaccine behaviors can be positively influenced with a carefully timed, interactive, informational online resource.
administered by their health care organization.

Although our VSM arm was designed to foster interaction between parents, almost all of the interaction was between parents and the research team. Parents who engaged in the social media applications were primarily interested in asking our experts questions to address their specific vaccine concerns. They did not appear to be interested in forming an ongoing vaccine-focused online community with other parents enrolled in the KPCO health plan. Given that only 1% of digital health social network members actively contributed to the interaction, it is possible that more participant-to-participant engagement would be observed if the intervention was scaled across the entire health plan or made publically available.

Web-based interventions are low-cost and broadly accessible approaches to deliver important public health messages. However, the VSM arm in our trial required significant resources to administer. A multidisciplinary, expert staff developed and reviewed new content, moderated chat room discussions, answered complicated questions related to the vaccination schedule, and addressed vaccine safety rumors as they surfaced. Therefore, it is unlikely that single clinician or clinic would have the means to manage their own social media interventions. This could be mitigated by creating a national, centralized social media vaccine resource, but additional research would need to determine if it would be trusted and used by parents. To help with these implementation decisions, a formal cost-effectiveness analysis of the VSM intervention is underway.

Social media technologies, Web site design preferences, and online information-gathering practices are constantly evolving. Such changes pose challenges to Web-based interventions. For example, our study period spanned more than 5 years from the development of the interventions through participant follow-up, data collection, and analysis. Over this time, newer social media platforms became increasingly popular among our target demographic population, including Twitter, Snapchat, and Instagram. Although it is not known if these platforms could be used to effectively address vaccine hesitancy, it is possible that our Web site appeared increasingly outdated and less appealing as the trial progressed. Therefore, future applications of our interventions would have to stay abreast of emerging technologies to continue to attract each new generation of parents.

This study had several limitations. The trial was conducted in a single, integrated health care system in Colorado, where the baseline vaccine hesitancy rate was 14.1%. Although this rate is similar to other investigations, there were only 99 hesitant participants in the analysis. As a result, we had limited statistical power to assess the interaction between study arm and vaccine hesitancy status.

Because this intervention was implemented as a pragmatic trial, we chose not to conduct a per-protocol analysis. Over the course of the trial, we gave participants in the VSM and VI arms unlimited access to the Web site, but they were not required to visit it. Of participants, ~35% visited the Web site at least once, and hesitant participants were more likely to access the Web site than nonhesitant participants. This implies that a per-protocol analysis in which researchers examine an association between Web site exposure and immunization outcomes would be biased by baseline hesitancy.

The overall loss to follow-up rate in the trial was 18.8%, which is largely attributable to parents who did not use KPCO health insurance for pediatric care after their children were born. Although this may have affected the trial’s generalizability, the rate of loss to follow-up did not differ significantly across the study arms. In addition, the rate of vaccine hesitancy among those lost to follow-up was similar across the arms, suggesting that excluding these individuals did not affect the internal validity of the results.

CONCLUSIONS

Despite these limitations, the results of this RCT demonstrate that Web-based vaccine information with social media technologies can positively influence parental vaccine decisions. As a complement to routine well-child care, the information appears to be effective when presented to parents before their children are born.

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ABBREVIATIONS

CI: confidence interval
KPCO: Kaiser Permanente Colorado
MMR: measles-mumps-rubella
OR: odds ratio
PACV: Parent Attitudes and Childhood Vaccines
RCT: randomized controlled trial
UC: usual care
VI: Web site with vaccine information only
VSM: Web site with vaccine information and interactive social media component
study design, provided input for the statistical analyses, and contributed to the first draft of the manuscript; and all authors reviewed and revised the manuscript and approved the final manuscript as submitted.

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REFERENCES


22. Montano DE, Kasprzyk D. Theory of reasoned action, theory of planned


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