

The U.S. Preventive Services Task Force recommends based elsewhere in Pregruanc: Recommendations and from the AHRQ Web site and Vaginosis in Pregnancy; Recommendations and from the AHRQ Web site and cleantimehouse.

(1-800-358-9295).

Replicts are available from the AHRQ Web site at www.ahrq.gov/ (www.ahrq.gov), or in print through the National GuideLine Clearinghouse (1-800-358-9295).

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Medical Subject Headings (MeSH): bacterial vaginosis, vaginitis, pregnancy, prevention, health services, evidence-based medicine, MEDLINE, methods, mass screening (Am J Prev Med 2001;20(3S):62-72)

We found no benefit to routine BV screening and treatment. A subgroup of high-risk women may benefit from BV screening and treatment; however, there may be a subgroup for whom BV treatment could increase the occurrence of premature delivery.

Women randomized controlled trials reported results for premature delivery less than 34 weeks. Two trials of high-risk women found an increase in premature delivery less than 34 weeks in women who did not have BV but received BV treatment. Compared to patient populations, treatment regimens, and study designs did not pool different studies reporting results for premature delivery less than 34 weeks. The pooled estimate showed no benefit ($ARR = -0.04$; $90\% CI = -0.02$ to 0.09), but variation was noted among individual studies. Two trials of high-risk women found a reduction was pooled estimate showing heterogeneity ($ARR = 0.22$; $90\% CI = 0.13$ to 0.31) for premature delivery before 37 weeks. Four high-risk studies reported results for premature delivery less than 34 weeks. The heterogeneous trials, three homogeneous studies showed potential benefit of BV treatment ($ARR = 0.19$ to 0.04), whereas one showed no benefit ($ARR = -0.08$, 90% heterogeneity). They clustered into two groups; one with previous delivery, were statistically studies of high-risk populations, women with previous delivery, were statistically no benefit to BV treatment in average-risk women for any pregnancy outcome. Results of seven randomized controlled trials met inclusion criteria for the meta-analysis. We found no benefit to BV treatment in trials met inclusion criteria for the meta-analysis.

Seven randomized controlled trials measured intermediate intervals (CLs) for the effect of treatment. Mean and 90% confidence intervals (CLs) for the effect of treatment applied to assess heterogeneity, to pool studies when appropriate, and to calculate the absolute risk reduction [ARR]. A stepwise procedure based on the profile likelihood was rate of a given pregruanc outcome in the control group minus the treatment group (the For each study, we measured the effect of treatment by calculating the difference in the less than 250 grams, spontaneous abortion, postpartum endometritis, and neonatal sepsis. For each study, complications, rates of spontaneous abortion, and total premature delivery less than 34 weeks, premature rupture of membranes, low birth weight weeks and less than 34 weeks, prematurity, comorbidity, deliveries, and gestational age less than 37 weeks and less than 34 weeks, rates of spontaneous abortion and total premature delivery less than 37 weeks, complications, rates for previous deliveries such as previous comorbidities, demographic details, risk factors for premature delivery such as previous antibiotic interventions, timing of antibiotic treatment in pregruanc, criteria for treatment, and following information was abstracted: study design and blinding, diagnostic methods, no benefit to BV treatment in pregruanc outcome. All randomized controlled trials of BV treatment in pregruanc that specifically measured

pregruanc outcomes. All randomized controlled trials of BV treatment in pregruanc that measured intermediate intervals of treatment measured the effect of treatment on pregruanc outcomes. All randomized controlled trials of BV treatment in pregruanc that measured intermediate intervals of treatment measured the effect of treatment on pregruanc outcomes.

The following information was abstracted: study design and blinding, diagnostic methods, comorbidity, previous deliveries, gestational age, birth weight, and gestational age less than 37 weeks. To determine whether screening and treatment for the U.S. Preventive Services Task Force. Pregruanc outcomes, as part of an assessment for women for BV reduces adverse effects, pruritis, or malodor, but often women with BV are asymptomatic.

Bacterial vaginosis (BV) is a strong independent risk factor for adverse pregruanc outcomes. BV is found in 9% to 23% of pregruanc women. Symptoms include vaginal discharge, pruritis, or malodor, but often women with BV are asymptomatic.

Data Synthesis:

Effects:

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Screening for Bacterial Vaginosis in Pregnancy

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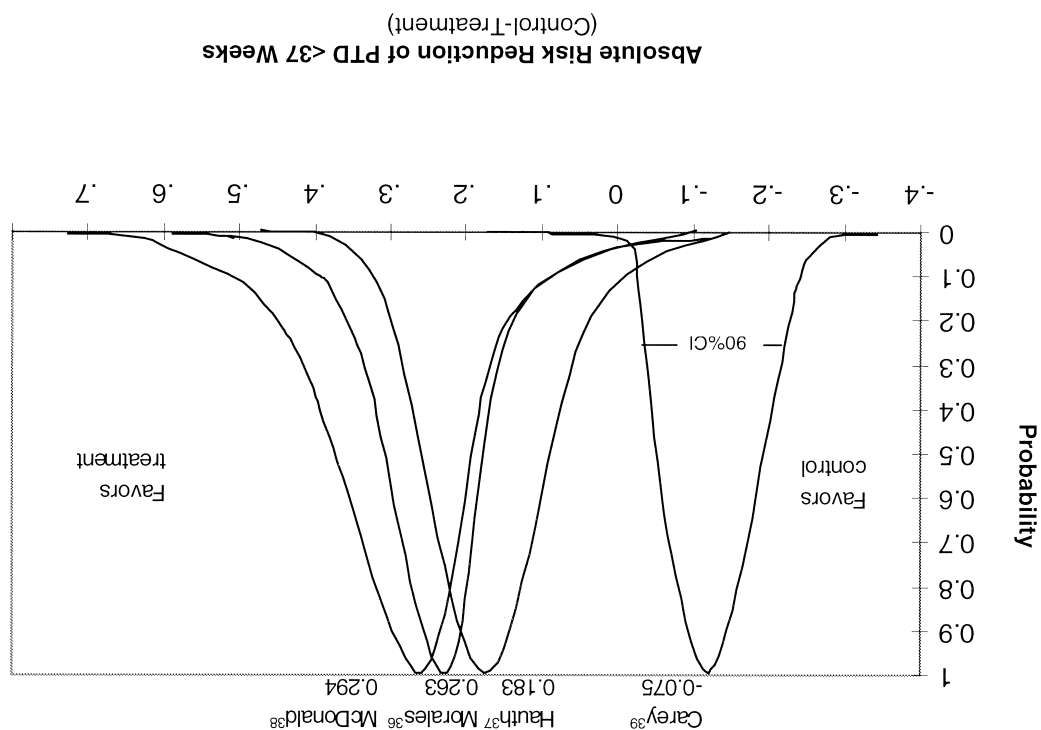
*Measures of quality of randomized controlled trials on a 3-point scale.

Morales et al.³⁶ reported benefit of treatment and reported a high proportion of women reporting two or high-risk women is not known.

However, the factor that defined this subgroup that was more likely to benefit from BV treatment, might have identified a "more selected" high-risk group Donald et al.³⁸ Morales et al.³⁶ and Hauth et al.³⁷ence for "general" high-risk patients, whereas McDonald et al.³⁹ may have portrayed the experts' results of Cary et al.³⁹ may have delivered the entire high-risk group (with and without BV) was 21%. The although the preterm delivery rate among the entire BV patients for the study by Vermullen and Brumse,³⁵ not possible to calculate the preterm delivery rate for the lowest rate, followed by McDonald et al.³⁸ at 35%, Morales et al.³⁶ at 39%, and Hauth et al.³⁷ at 57%. It was the placebo groups. The NICHD study,³⁹ at 23%, had the variation in preterm delivery rates before 37 weeks in ment (Table 2). The most striking difference was their of high-risk patients who may benefit from BV treatment to 0.057).

We examined all high-risk studies for factors that could explain variation and possibly define a subgroup of high-risk patients who may benefit from a subgroup of high-risk patients who may benefit from BV treatment. We were suffciently similar to pool with a pooled ARR of 0.259 (90% CI=0.149 to 0.369). A similar trend across were homogeneous and showed benefit (pooled ARR=0.22, 90% CI=0.13 to 0.31),³⁶⁻³⁸ indicating 22 weeks less than 37 weeks (Figure 2). Three of these studies were homogenous and showed benefit (pooled ARR=0.096 to 0.321). The smaller study, with a total sample size of 34, showed a stronger benefit.

Figure 2. Individual results of preterm delivery (PTD) before 37 weeks in high-risk patients. The measure of effect is a difference in probability of benefit from control minus treatment (absolute risk reduction [ARR]). The width of the curve indicates the precision of the estimate and is used to calculate 90% confidence intervals.



from studies by McDonald et al.³⁸ and Morales et al.³⁶ an ARR of -0.036 (90% CI=-0.101 to 0.030). Results in ARRs 0.288, 90% CI=0.149 to 0.427, McDonald et al.³⁸ reported an ARR of 0.176 (90% CI=0.024 to 0.329), whereas Cary et al.³⁹ showed no benefit, with 17 patients in each group.³⁸ When pooled, these studies had a slight trend toward benefit that was not significant statistically significant (pooled ARR=0.036; 90% CI=-0.021 to 0.092) (Figure 4).

Three studies reported results on preterm premature rupture of membranes.^{36,38,39} Results from the study by McDonald et al.³⁸ showed the greatest benefit among the entire sample of small sample size (17 patients in each group).³⁸ When pooled, these also had the least precision because of small sample size (the trend toward greatest benefit with an ARR of 0.118 the treatment group). The study that showed benefit in the treatment group, the study that showed four studies showed statistically significant improvements preterm delivery prior to 34 weeks. None of the four studies reported preterm delivery or spontaneous premature rupture of membranes. The total ARR for the entire study was 0.208 (90% CI=0.096 to 0.321). The smaller study, with a total sample size of 34, showed a stronger benefit.

These studies, the pooled ARR was 0.208 (90% CI=0.096 to 0.321). The NICHD trial³⁹ showed spontaneous preterm delivery before 37 weeks.^{37,38} In (Figure 3, curve at left). Two studies provided data for patients treated (ARR=-0.075, 90% CI=-0.189 to 0.039) no benefit (ARR=-0.075, 90% CI=-0.189 to 0.189) showed fewer deliveries at less than 37 weeks per 100 patients treated (Figure 3). The NICHD trial³⁹ showed fewer preterm deliveries at less than 37 weeks per 100 fewer patients treated (ARR=-0.075, 90% CI=-0.189 to 0.189) no benefit (ARR=-0.075, 90% CI=-0.189 to 0.189). The NICHD trial³⁹ showed fewer deliveries at less than 37 weeks per 100 patients treated (ARR=-0.075, 90% CI=-0.189 to 0.189) no benefit (ARR=-0.075, 90% CI=-0.189 to 0.189).

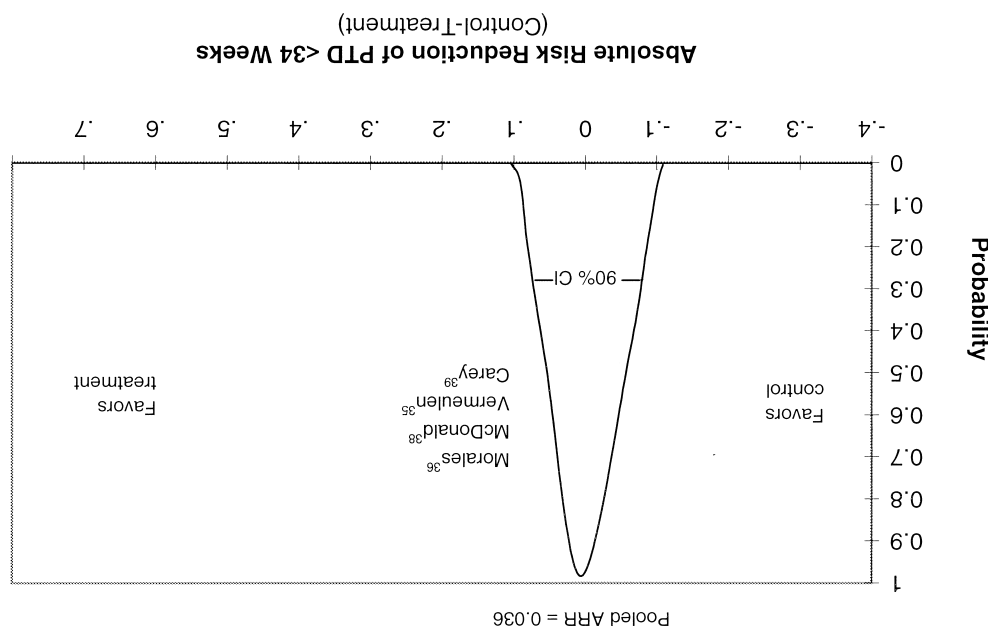
Four studies reported preterm delivery or spontaneous premature rupture of membranes. The total ARR for the entire study was 0.208 (90% CI=0.096 to 0.321). The smaller study, with a total sample size of 34, showed a stronger benefit.

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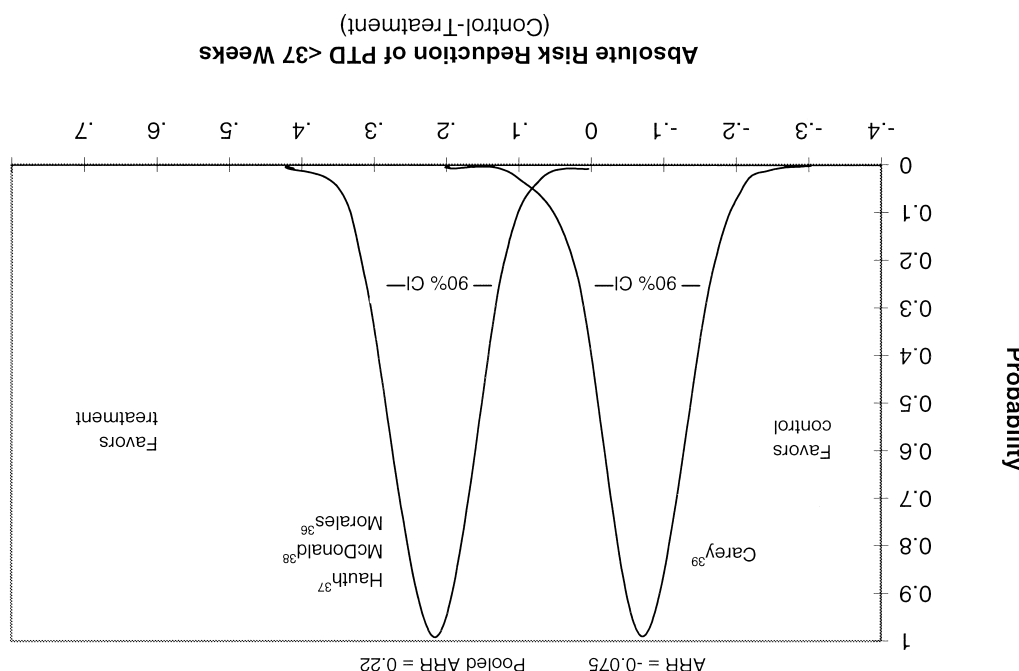
Figure 4. Pooled results of four studies reporting rates of preterm delivery (PTD) before 34 weeks in high-risk patients. None pooled effect was not statistically different from zero (0.036% ; 90% CI = -0.021 to 0.092%).
of the four studies reported a statistically significant decrease in preterm delivery before 34 weeks with treatment, and their



whereas Carey et al.,³⁹ Hauth et al.,³⁷ and McDonald et al.³⁸ asked if patients had ever experienced preterm delivery. The issue of timing and quantity of previous delivery and should be examined further.

provide data on the number of previous preterm deliveries before 37 weeks per 100 patients treated. Other studies did not provide previous deliveries. Other studies did not experience preterm delivery in their last pregnancy, this factor. Morales et al.,³⁶ also selected for patients who were in their populations, so we could not compare them on this factor. The ARR was -0.075% (90% CI = -0.189 to 0.039%), indicating seven additional preterm deliveries before 37 weeks per 100 patients treated.

Figure 3. Pooled results of four studies reporting rates of preterm delivery (PTD) before 37 weeks in high-risk patients. The pooled absolute risk reduction for Hauth et al.,³⁷ McDonald et al.,³⁸ and Morales et al.,³⁶ was 0.22% (90% CI = 0.13 to 0.31%), indicating 22 fewer preterm deliveries before 37 weeks per 100 patients treated. One study, Carey et al.,³⁹ was dissimilar from the others and did not pool. In that study, the ARR was -0.075% (90% CI = -0.189 to 0.039%), indicating seven additional preterm deliveries before 37 weeks per 100 patients treated.



BV, because the only trial of these four that examined delivery less than 34 weeks in high-risk patients without BV, might be associated with a higher rate of preterm deliveries than 34 weeks in high-risk patients without BV at the NICHD trial.³⁹ The second case scenario we also assumed that metronidazole therapy comes to treat BV-negative women (*i.e.*, the maximum plausible effect), as well as a worst-case scenario, using the lower 90% CI of the pooled estimate. In both scenarios to that of BV-negative women (i.e., the maximum plausible effect), all studies of BV reduces their risk of adverse pregnancy outcomes to that of other high-risk studies.³⁶⁻³⁸ Of note, the second scenario incorporates the pooled results of three other high-risk population studies that were not included in this base case, where assumed that treating these women used in this balance sheet used metronidazole therapy. The second scenario used a more conservative estimate of the NICHD study³⁹ for the latest outcomes. CIs from the NICHD study³⁹ for the general high-risk population incorporate the mean and 90% confidence intervals of screening for BV in 1000 patients from the general high-risk population and 1000 from a more selected population. The base case for the general population uses estimates of screening for BV in 1000 patients from the general high-risk population and 1000 from a more screened population. This base case for the general population that may benefit from screening for BV in women suggests that there may be a subgroup of high-risk pregnancies that benefit in some high-risk pregnancies, and preterm premature rupture of membranes. The findings of benefit in some high-risk pregnancies, and preterm premature rupture of membranes, grams, and preterm delivery to 34 weeks, low birth weight less than 2500 prior to 34 weeks, and preterm delivery of preterm delivery (women with a previous preterm delivery) for the clinical liability important outcomes of preterm delivery to screen all women at high risk for preterm benefit to screen all women at high risk found no treatment of BV in pregnancy.¹⁵ We similarly found no treatment with those of a recent Cochrane review of preterm delivery to screen all women. The findings for average-risk women are consistent with those of a recent Cochrane review of preterm delivery to screen all women.¹⁵ The findings for average-risk women in a summary, there appears to be no benefit to screening time, but we do not know which of the NICHD patients were BV negative at second treatment. These data emphasize the importance of the accuracy of screening tests to diagnose BV.

Summary of Benefits and Harms

In summary, but we do not know which of the NICHD patients were BV negative at second treatment. These data emphasize the importance of the accuracy of screening tests to diagnose BV.

Factor	Carey ³⁹	McDonald ³⁸	Hauth ³⁷	Morales ³⁶	Vermuelen ³⁵	PTD, preterm delivery; GA, gestational age.
PTD <37 weeks,	23%	35%	57%	39%		
placebo	16-23	24	22-24	13-20	26	
Treatment	Oral metronidazole	Oral metronidazole	Oral metronidazole	Oral metronidazole	Oral metronidazole	
GA (wk) at	2 g, repeat	400 bid, 2 d	250 mg tid, 7 days,	250 mg tid, 7 d	plus oral erythromycin	④ 48 hr
Second treatment	All, 24-29 wk	If positive test,	33 mg tid, 14 d	if positive test,	at 28 wk	if positive test, 4 wk post-treatment
						No second treatment All, 32 wk

symptoms. However, the details for identification of BV alone. Third, we defined patients as asymptomatic if they were most likely identified through routine clinical examination. Women who have been identified as having BV by this criterion have looked at the effectiveness of treatment in women who have been identified as having BV by this criterion have been identified as having BV by screening used in every day practice. One technique is screening by identification of clue cells alone. There are no studies of BV treatment that research studies may not reflect those methods used in based practices. Second, screening methods used in clinics and may not be generalizable to community clinics and may not be useful for public health conducted in tertiary referral centers or public health may be useful for patients. First, all U.S. studies were to determine whether screening and testing for BV there are several issues of generalizability to consider

Generalizability

Screening test for BV is below 80% (not shown). term delivery before 34 weeks if the specificity of the screening and treatment result in an increase in prevalence. In the more selected high-risk group, for example, test. In the changes in the accuracy of the screening sensitive to changes in the accuracy of the screening on preterm delivery less than 34 weeks is moderately harmful effect.³⁷ We also assumed that the screening 34 weeks in BV-negative patients, the effect of screening

Screen all women who have at least one previous PTD. Screen it there is more than one previous PTD or other risk factors. Males sign indicates a net increase, plus indicates a net decrease, in adverse effects.

Assumptions	Effect sizes, BV patients (probability: control group vs treated group) ^a	Effect sizes, patients without BV (probability: control group vs treated group) ^a	Results (n)	Outcomes ^c
Proportion of pregnant women who meet screening criteria	0.1	+0.22 (+0.13 to +0.31)	-0.075 (-0.19 to +0.04)	PTD <37 weeks
Relative risk of PTD in BV population	0.25	+0.29 (+0.15 to +0.43)	-0.04 (-0.101 to +0.03)	PPROM
Sensitivity of screening test	0.95	+0.075 (-0.06 to 0.15)	+0.012 (-0.06 to 0.08)	PTD <34 weeks
Specificity of screening test	0.95	+0.06 (+0.01 to +0.15)	-0.04 (-0.06 to 0.08)	PROM
Adherence to treatment	0.8	-0.06 (-0.13 to -0.01)	-0.02	PTD >34 weeks
Assumptions	0.03	-0.02 (-0.13 to 0.07)	0	Promotional
Beneft and relevant factors	"General" high-risk	"More selected"	0	Promotional

Table 3. Summary of benefits and harms of screening 1000 high-risk pregnant women for bacterial vaginosis

assumed a potential increase in preterm delivery before 24 (90% CI=17 to 40) patients screened. Because every at least than 37 weeks would be prevented for every patient screened; likewise, one case of preterm delivery than 34 weeks would be prevented for every 111 women screened. One case of preterm delivery at less than 34 weeks (90% CI=19 fewer to 2 additional cases) of preterm cases (90% CI=28 to 70), premature rupture of membranes (90% CI=25 to 59), 49 fewer cases of preterm weeks in 42 fewer preterm deliveries before 37 weeks (90% CI=28 to 70), 8 additional cases of preterm deliveries before 37 weeks (90% CI=7 fewer to 36 additional cases), 8 additional cases of preterm deliveries before 36 additional treatments here would be 14 additional preterm deliveries before 37 weeks (90% CI=4 fewer to 19 additional cases), 2 fewer cases of preterm deliveries before 34 weeks (90% CI=1 fewer to 11 additional cases).

In the second high-risk group, screening and treatment 190 of these complete diagnosis to have BV, and screened, 238 are correctly diagnosed to have BV, and adherence to treatment is 80%. In the general high-risk population, of 1000 women screened it has a sensitivity of 95% and specificity of 95%, the test has a sensitivity of 95% and specificity of 95%, the harmful effect.³⁷ We also assumed that the screening this effect found a clinically and statistically significant

34 weeks in BV-negative patients, the effect of screening 34 weeks in BV-negative patients, the effect of screening

Screen all women who have at least one previous PTD. Screen it there is more than one previous PTD or other risk factors. Males sign indicates a net increase, plus indicates a net decrease, in adverse effects.

Assumptions	Effect sizes, BV patients (probability: control group vs treated group) ^a	Effect sizes, patients without BV (probability: control group vs treated group) ^a	Results (n)	Outcomes ^c
Proportion of pregnant women who meet screening criteria	0.1	-0.02 (+0.01 to +0.15)	-0.075 (-0.13 to 0.04)	Promotional
Relative risk of PTD in BV population	0.25	+0.29 (+0.15 to +0.43)	-0.04 (-0.101 to +0.03)	PROM
Sensitivity of screening test	0.95	+0.075 (-0.06 to 0.15)	-0.04 (-0.06 to 0.08)	PTD <34 weeks
Specificity of screening test	0.95	+0.06 (+0.01 to +0.15)	-0.04 (-0.06 to 0.08)	PTD >34 weeks
Adherence to treatment	0.8	-0.06 (-0.13 to -0.01)	-0.02	Promotional
Assumptions	0.03	-0.02 (-0.13 to 0.07)	0	Promotional
Beneft and relevant factors	"General" high-risk	"More selected"	0	Promotional

assumed a potential increase in preterm delivery before 24 (90% CI=17 to 40) patients screened. Because every at least than 37 weeks would be prevented for every patient screened; likewise, one case of preterm delivery than 34 weeks would be prevented for every 111 women screened. One case of preterm delivery at less than 34 weeks (90% CI=19 fewer to 2 additional cases) of preterm cases (90% CI=28 to 70), 49 fewer cases of preterm weeks in 42 fewer preterm deliveries before 37 weeks (90% CI=28 to 70), 8 additional cases of preterm deliveries before 37 weeks (90% CI=7 fewer to 19 additional cases), 8 additional cases of preterm deliveries before 36 additional treatments here would be 14 additional preterm deliveries before 37 weeks (90% CI=4 fewer to 19 additional cases), 2 fewer cases of preterm deliveries before 34 weeks (90% CI=1 fewer to 11 additional cases).

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Recommendations for Future Research

Consequently, it is possible that women presenting for BV may represent a different group of patients with symptoms were also included. The evaluation of BV symptoms were also performed. The risk category for adverse pregnancy outcomes.

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University Evidence-based Practice Center is based on a more comprehensive Systematic Review, which is available online at www.ahrq.gov/. This article is based on a more comprehensive Systematic Review, which is available online at www.ahrq.gov/.

Evidence Review, which is available online at www.ahrq.gov/.

Chronic/pregnancy. That document was reviewed by committee members, including Sharon Hillier, PhD, University of Pittsburgh experts, including Mark Kleibaum, MD, MPH, National Institute of Child Health & Human Development, Sharon Hillier, PhD, University of Pittsburgh experts, including the American Academy of Family Physicians, the American College of Obstetricians and Gynecologists, the National Institute of Child Health & Human Development; the National Institutes of Health; the Centers for Disease Control and Prevention; and the U.S. Navy Bureau of Medicine and Surgery. Review by these individuals and groups does not necessarily imply endorsement of this article or of the accompanying recommendations of the U.S. Preventive Services Task Force.

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