# Prenatal and Postpartum Depression in Fathers and Its Association With Maternal Depression A Meta-analysis

James F. Paulson, PhD

Sharnail D. Bazemore, MS

HE PREVALENCE, RISK FACtors, and effects of depression among new fathers are poorly understood. Although a large body of research on maternal depression documents incidence rates between 10% and 30% and negative family and child developmental outcomes,1-3 paternal prenatal and postpartum depression has received little attention from researchers and clinicians.4 The emerging literature on paternal depression suggests that, like their maternal counterparts, fathers are at increased risk of depression in the postpartum<sup>5</sup> and gestational periods.6-8 Moreover, several studies have now documented negative child outcomes associated with paternal prenatal and postpartum depression.9,10

Although recent literature has addressed this phenomenon, studies in paternal prenatal and postpartum depression are troubled by inconsistent methods, clinical heterogeneity, and prevalence estimates that vary considerably.<sup>5,7,11-20</sup> To date, only 2 reviews on prenatal and postpartum depression in fathers have been published, but neither sought to quantitatively synthesize or resolve the discrepancies across studies, methods, or other issues.<sup>5,11</sup> We conducted the present meta-analysis of

CME available online at www.jamaarchivescme.com and questions on p 1987. **Context** It is well established that maternal prenatal and postpartum depression is prevalent and has negative personal, family, and child developmental outcomes. Paternal depression during this period may have similar characteristics, but data are based on an emerging and currently inconsistent literature.

**Objective** To describe point estimates and variability in rates of paternal prenatal and postpartum depression over time and its association with maternal depression.

**Data Sources** Studies that documented depression in fathers between the first trimester and the first postpartum year were identified through MEDLINE, PsycINFO, EMBASE, Google Scholar, dissertation abstracts, and reference lists for the period between January 1980 and October 2009.

**Study Selection** Studies that reported identified cases within the selected time frame were included, yielding a total of 43 studies involving 28 004 participants after duplicate reports and data were excluded.

**Data Extraction** Information on rates of paternal and maternal depression, as well as reported paternal-maternal depressive correlations, was extracted independently by 2 raters. Effect sizes were calculated using logits, which were back-transformed and reported as proportions. Random-effects models of event rates were used because of significant heterogeneity. Moderator analyses included timing, measurement method, and study location. Study quality ratings were calculated and used for sensitivity analysis. Publication bias was evaluated with funnel plots and the Egger method.

**Data Synthesis** Substantial heterogeneity was observed among rates of paternal depression, with a meta-estimate of 10.4% (95% confidence interval [CI], 8.5%-12.7%). Higher rates of depression were reported during the 3- to 6-month postpartum period (25.6%; 95% CI, 17.3%-36.1%). The correlation between paternal and maternal depression was positive and moderate in size (r=0.308; 95% CI, 0.228-0.384). No evidence of significant publication bias was detected.

**Conclusions** Prenatal and postpartum depression was evident in about 10% of men in the reviewed studies and was relatively higher in the 3- to 6-month postpartum period. Paternal depression also showed a moderate positive correlation with maternal depression.

JAMA. 2010;303(19):1961-1969

www.jama.com

depression in expecting and new fathers to (1) estimate paternal depression between the first trimester and 1 year postpartum; (2) describe differences across time within this period; (3) examine the association between paternal and maternal depression; (4) estimate the prevalence of maternal prenatal and postpartum depression identified in paternal depression studies; and (5) identify how published rates

Author Affiliations: Department of Pediatrics, Eastern Virginia Medical School, Norfolk.

**Corresponding Author:** James F. Paulson, PhD, Department of Pediatrics, Eastern Virginia Medical School, E. V. Williams Hall, 855 W Brambleton Ave, Norfolk, VA 23510 (paulsojf@evms.edu).

of paternal depression were affected by methodological factors such as measurement method, study location, and sample risk status.

## METHODS Search Strategy

We used 3 methods to identify studies for this meta-analysis. First, we used the reference lists of the most relevant reviews.5,11 Next, we searched MEDLINE, PsycINFO, Dissertation Abstracts International, EMBASE, and Google Scholar using the search terms depression, paternal, father, postnatal, postpartum, prenatal, antenatal, and perinatal. Finally, we used the "ancestry approach,"21 which involves consulting the reference lists of retrieved articles to find earlier relevant studies. Because of the emergent nature of this body of observational research literature, an inclusive approach to study selection was used.<sup>22,23</sup> Therefore, we included all relevant and accessible journal articles, dissertations, and book chapters that were produced between January 1980 and October 2009 that assessed paternal depression during pregnancy, the first postpartum year, or both.

#### **Study Selection**

Studies that reported an estimated number of depression cases among identified fathers were included. This resulted in the exclusion of several studies that reported mean scores for symptom severity because the exact number of cases could not be clearly determined. Several articles9,10,24-27 used data from common databases and were excluded to avoid duplication of data. Several studies measured depression on multiple occasions. In these cases, 1 depression measure per time period was selected based on these priorities: (1) structured interviews: (2) measures with demonstrated generality in men (eg, Beck Depression Inventory<sup>28</sup>) vs adaptations of maternal measures (eg, Edinburgh Postnatal Depression Scale<sup>29</sup>); and (3) measures with greater specificity for depression (eg, Beck Depression Inventory<sup>28</sup> vs General Health Questionnaire<sup>30</sup>). We excluded studies that selected fathers based on established maternal mental health problems because this could bias meta-analytic estimates. Also, because the identified studies of teen fathers were characterized by significant economic and social stressors, only studies of fathers aged 18 years or older were included.

## Data Abstraction and Quality Assessment

The 2 authors used a standardized coding manual (available from the authors on request) to extract the following data from articles: author names, publication year, sample size, period of assessment, study sample risk (0 or 1; high risk coded when a study denoted this clearly, including medically assisted pregnancies and infants with feeding, sleeping, or crying problems), location, sample size, response rate, number of fathers identified as depressed, number of mothers identified as depressed (when assessed), and correlation between maternal and paternal depressive symptoms. The coding manual was developed a priori and modified after use in several studies. Coding was done independently then aggregated, with disagreements resolved through discussion and consensus. Although quality assessment can be reliably conducted in metaanalyses of experimental studies, its use in observational research is controversial, with no clear consensus on rating methods or their appropriate use in analysis. As such, we used a simple objective rating system (based on the meta-analysis of similar data by Bennett et al<sup>2</sup>) that coded studies on a scale of 0 to 10, assigning 2 points each for sampling method (systematic or probability vs convenience or not reported), presence of clearly stated inclusion criteria, racial/ethnic diversity  $(\geq 20\%$  minority), educational diversity ( $\leq$ 80% at 1 educational level), and response rate (reported at  $\geq$  60%). Studies that did not report these methodological issues received lower scores. Because evidence on the validity of

quality ratings in observational research is lacking, we adopted the approach of Stroup et al<sup>23</sup> of broadly including studies and using sensitivity analysis to determine incremental effects of lower-quality studies.

### **Effect Size and Statistical Analysis**

Primary Outcome. The primary outcome was the point prevalence rate of paternal depression, defined as the number of cases divided by the total number of study participants. We coded these into both simple proportional effect sizes (by dividing the number of cases by the sample size) and logit units, as a direct transformation of these proportions. In this context, the logit transformation was used to form an unbounded (in contrast to the 0-to-1 bounded nature of proportions) estimate to facilitate moderator analysis.<sup>31</sup> After analysis, logit units were back-transformed to proportions for the purposes of reporting.

Secondary Outcomes. Secondary outcomes included rates of depression in female partners, which we coded as raw proportions and logit units, and standardized zero-order correlations between paternal and maternal depressive symptoms (when measured with a continuous or ordinal scale).

All major analyses were conducted with Comprehensive Meta-Analysis, version 2.0.32 In general, randomeffects models are argued to better address heterogeneity between studies and study populations, allowing for greater flexibility in parsing effect size variability. Moreover, they are less influenced by extreme variations in sample size.<sup>22</sup> Because studies in this meta-analysis are characterized by heterogeneity and highly variable sample sizes, randomeffects models were used. Heterogeneity among study point estimates was assessed with the Q statistic, with magnitude of heterogeneity being evaluated with the I<sup>2</sup> index.<sup>31</sup> When reported, all confidence intervals (CIs) reflect a 95% criterion.

We examined the following determinants of primary and secondary outcomes: period of measurement, risk sta-

**1962** JAMA, May 19, 2010—Vol 303, No. 19 (Reprinted)



Publication bias was assessed by visually inspecting funnel plots and applying the regression intercept of Egger et al.<sup>34</sup> In addition, we used the fail-safe procedure of Orwin,<sup>35</sup> which is based on effect sizes that would be considered practically insignificant rather than the traditional null-effect reference. This generated a number of unpublished studies with effects at the estimated population base rate for adult male depression<sup>36</sup> that would be needed to move estimates to a nonsignificant difference from base rates.

To assess the robustness of the results, we performed sensitivity analyses by sequentially removing each study and rerunning the analysis. We also conducted a separate analysis excluding studies with quality ratings in the lowest third to determine if potential methodological weaknesses influenced meta-analytic estimates.

### RESULTS Included Studies

Of the initial 256 identified studies, most (n=163) were excluded because they were not applicable to the present meta-analysis (eg, articles on other topics, depression not assessed in fathers, reviews or summaries, beyond postpartum period, infant death, teen parents). Of those that were reviewed in full text, 30 were excluded because the proportion of depressive cases was not reported, 16 reported on a sample



already included in the present study, and 4 were not retrievable (FIGURE 1). After a thorough review, 43 studies met the inclusion criteria for this metaanalysis<sup>6-8,13-20,37-68</sup> (TABLE). Of these studies, 23 reported rates of paternal depression at 2 or more time points and 20 reported a single observation. Because the inclusion of multiple effect sizes from a single sample would compromise the independence assumption of meta-analysis,<sup>22</sup> primary analyses used the earliest reported estimate, as this generally reflects a larger preattrition sample size.

The Table provides details on the characteristics of the 43 studies. Studies originated in 16 countries, with the United States contributing the most (n=17 studies). Most studies (n=40) used a self-report rating scale as the primary case definition method, with the remainder (n=3) using a structured or semistructured interview. Three studies enrolled men from higher-risk samples. Two studies used population-

based sampling procedures embedded within larger birth cohort studies, but most studies (n=30) recruited from maternity or postpartum units, the remainder coming from parenting/ prenatal classes and other health services. Thirty studies reported response rates greater than 70%. In addition to reporting paternal depression, 35 studies reported rates of partners' maternal depression and 14 reported the correlation between maternal and paternal depressive symptoms. Sample sizes varied widely across studies (N=23-10975), with a median of 130 participants (first quartile=80; third quartile=307). In all, using initial sample sizes across the 43 studies, a total of 28 004 participants are represented in this meta-analysis.

#### **Tests for Heterogeneity**

According to the criteria set by Higgins and Thompson,<sup>69</sup> the heterogeneity in published rates of paternal depression was statistically significant and

©2010 American Medical Association. All rights reserved.

(Reprinted) JAMA, May 19, 2010-Vol 303, No. 19 1963

Source (Study Location) and Time of Assessment	Depression Measure (Cutoff)	No. of Participants (Women) <sup>a</sup>	Depressed, No. (%)		Correlation
			Men	Women	Men and Women <sup>b</sup>
<b>Onset of paternal depression at</b> Areias et al, <sup>17</sup> 1996 (Portugal)	gestation <6 mo SADS				
6 mo gestation		42 (54)	2 (4.8)	9 (16.7)	
3 mo postpartum		12 (24)	2 (8.3)	17 (67)	
12 mo postpartum		42 (54)	10 (23.8)	20 (37)	
Condon et al, <sup>6</sup> 2004 (Australia)	EPDS (>12)	312	16 (5.2)		
en e me gectation	GHQ (>5)	0.12	57 (18.2)		
	MHI-5 (<17)		14 (4.6)		
3 mo postpartum	FPDS (>12)	276	5 (1.9)		
	GHQ (>5)	2.10	31 (11 3)		
	MHI-5 (<17)		4 (1.5)		
6 mo postpartum	EPDS (>12)	241	5 (2 1)		
	GHQ (>5)		27 (11.2)		
	MHI-5 (<17)		4 (1 7)		
12 mo postpartum	EPDS (>12)	222	5 (2.3)		
	GHQ (>5)		23 (10.4)		
	MHI-5 (<17)		7 (3.1)		
Fawcett and York <sup>14</sup> 1986 (US)	BDI (>9)		. (0.1.)		
3.5 mo gestation	88.(* 0)	23	1 (4 3)	6 (26 1)	
9 mo destation		24	2 (8 3)	8 (33 3)	
1.5 mo postpartum		23	3 (13)	6 (26.1)	
Field et al <sup>43</sup> 2006 (US)			- ()	0 (2011)	
5 mo gestation	CES-D (>15)	156	50 (32)	56 (36)	
Fletcher et al, <sup>44</sup> 2008 (Australia)					
Sometime during gestation	EPDS (>9)	307	16 (5.3)		
	EPDS (>6)	307	48 (15.5)		
Frost, <sup>45</sup> 1996 (US)	CES-D (>15)				
5 mo gestation		527	75 (14.2)	353 (67)	0.23
1 mo postpartum		476	67 (14)	100 (21)	0.16
4 mo postpartum		442	46 (10.6)	93 (21)	0.17
Matthey et al, <sup>20</sup> 2000 (Australia)	Multiple measures used to designate cases <sup>c</sup>				
5.5 mo gestation		152	8 (5.3)	19 (12.3)	0.18
1.5 mo postpartum		141	4 (2.8)	11 (7.7)	0.22
4 mo postpartum		125	4 (3.2)	12 (9.7)	0.18
12 mo postpartum		128	6 (4.7)	16 (12.4)	0.32
Ramchandani et al, <sup>55</sup> 2008 (UK)	EPDS (>12)	10975			0.26-0.31
4.5 mo gestation			426 (3.9)		
2 mo postpartum			399 (3.6)		
8 mo postpartum			378 (3.4)		
21 mo postpartum			425 (3.9)		
van den Berg et al, <sup>7</sup> 2009 (the Netherlands) 4 mo gestation	BSI (>15)	3083 (3822)	364 (11.8)	409 (10.7)	
Onset of paternal depression at	gestation 6-9 mo	. ,		. ,	
Atkinson and Rickel,37 1984 (US)	BDI (>9)	78			
8 mo gestation			10 (13)	23 (29)	
2 mo postpartum			10 (13)	20 (26)	
Bou <u>rne,<sup>60</sup> 2006 (US)</u>	CES-D (>8)				
8 mo gestation		120	17 (14)	48 (40)	0.17
12 mo postpartum		87	8 (9)	23 (27)	0.08
Escribè-Agüir et al, <sup>8</sup> 2008 (Spain) 8.25 mo gestation	EPDS (>10)	669 (687)	43 (6.5)	71 (10.3)	
Hall and Long,63 2007 (Canada)	CES-D (>16)	98 (91)			
8.75 mo gestation			11 (11.2)	30 (33)	0.27
2.5 mo postpartum			21 (21.4)	16 (17.6)	0.21
					(continued)

 Table.
 Characteristics of Studies Included in Meta-analysis

large in magnitude (Q=825.081; P < .001;  $I^2 = 94.910$ ;  $\tau^2 = 0.470$ ). Maternal depression also demonstrated significant heterogeneity across studies (Q=1394.968; P < .001;  $I^2 = 97.563$ ;  $\tau^2 = 0.792$ ), but the evidence for heterogeneity among correlations between maternal and paternal depressive symptoms was equivocal (Q=89.906; P < .001;  $I^2 = 85.540$ ;  $\tau^2 = 0.019$ ).

#### **Primary Outcomes**

The overall random-effects estimate of paternal depression was 10.4% (95% CI, 8.5%-12.7%) (FIGURE 2). Although no significant differences in depression rates were observed between higherand lower-risk samples (lower risk, 10.1%; 95% CI, 8.2%-12.4%; higher risk, 15.6%; 95% CI, 5.6%-36.5%; Q=0.721; P=.40, moderator analyses revealed 3 significant factors. First, there was considerable variability between different time periods vis-à-vis birth (Q=20.256; P < .001), with the 3- to 6-month postpartum period showing the highest rate (25.6%; 95% CI, 17.3%-36.1%) and the first 3 postpartum months showing the lowest rate (19 studies; 7.7%; 95% CI, 5.3%-11.1%). Second, national origin of the study accounted for variability in depression rates of fathers (Q=7.108; P=.008), with the US studies reporting an average rate of 14.1% (95% CI, 10.9%-18.0%) and international studies reporting an average rate of 8.2% (95% CI, 5.9%-11.1%). Finally, interviewbased case definition methods were associated with lower overall prevalence estimates (rating scale, 11.0%; 95% CI, 8.9%-13.5%; interview, 4.9%; 95% CI, 3.6%-6.7%; Q=18.236; P < .001). Because paternal age and family size were inconsistently reported, conclusions could not be drawn regarding the moderator effects of either.

Maternal depression had a metaanalytic point estimate of 23.8% (95% CI, 18.7%-29.7%). Time period was a significant determinant of maternal depression (Q=22.156; P<.001), with higher rates reported during the 3- to 6-month postpartum period (41.6%).

1964 JAMA, May 19, 2010-Vol 303, No. 19 (Reprinted)

Measurement method (rating scale,
25.5%; 95% CI, 20.0%-31.9%; inter-
view, 9.8%; 95% CI, 5.9%-15.8%) was
also a significant predictor of mater-
nal depression rate (Q=12.773;
P < .001). Study location (United States,
29.6; 95% CI, 19.3%-42.5%; interna-
tional, 19.7%; 95% CI, 15.0%-25.4%)
demonstrated a trend toward higher
rates in the United States (Q=2.599;
P = .107).

The overall random-effects estimate of maternal-paternal depressive symptom correlation was significantly larger than 0 and moderate in magnitude (r=0.308; 95% CI, 0.228-0.384).

#### **Tests for Publication Bias**

Visual inspection of funnel plots (available from the authors) revealed no obvious evidence of publication bias. Quantitative evaluation of publication bias, as measured by the Egger intercept, was nonsignificant (P=.15). Finally, the Orwin fail-safe procedure, using a base rate of 3%,<sup>36</sup> determined that 1444 unpublished studies at or below this level would be needed to bring the overall meta-analytic estimate of prenatal and postpartum depression to a nonsignificant difference from the base rate.

#### **Sensitivity Analyses**

Robustness of meta-analytic findings was examined by sequentially removing each study and reanalyzing the remaining data set (producing a new analysis for each study removed). No study affected the meta-analytic estimate more than 0.5%. Removing studies with quality ratings in the lowest 33% decreased the meta-analytic estimate of paternal depression by only 0.6% (from 10.4% to 9.8%). The pattern of differences across time periods, measurement methods, and study locations remained essentially unchanged in direction and magnitude.

### COMMENT

In this meta-analysis of paternal prenatal and postpartum depression and its correlation with maternal depres-

©2010 American Medical Association. All rights reserved.

Table.         Characteristics of Studies Included in Meta-analysis (continued)						
Source (Study Location)	Depression Measure	No. of Participants	Depressed, No. (%)		Correlation Between Men and	
and Time of Assessment	(Cutoff)	(Women) <sup>a</sup>	Men	Women	Women <sup>b</sup>	
Onset of paternal depression at Keeton et al 49 2008 (US)	CES-D (>15)	o (continued)				
9.04 mo destation	010 0 (> 10)	140	21 (15)	62 (44)		
1.3 mo postpartum			17 (12)	36 (26)		
4 57 mo postpartum			17 (12)	41 (29)		
6.68 mo postpartum			18 (13)	36 (26)		
12.81 mo postpartum			15 (11)	35 (25)		
Leathers and Kelley <sup>12</sup> 2000 (US)	CES-D (>16)	124	()	00 (20)		
6.5 mo gestation	020 2 (* 10)		9 (7.3)	38 (30.6)		
3.75 mo postpartum			8 (6.5)	14 (11.3)		
Morse et al <sup>52</sup> 2000 (Australia)	FPDS (>9)		- ()	(		
6.25 mo gestation		251	30 (12)	49 (19.5)		
9 mo gestation		204	18 (8 7)	45 (21.1)		
1 mo postpartum		166	10 (6)	38 (21.6)		
4 mo postpartum		151	9 (5.8)	23 (13.9)		
Baskin et al <sup>59</sup> 1990 (US)	CES-D (>15)	86	0 (0.0)	20 (10.0)		
8.5 mo destation	020 0 (2 10)	00	16 (18 6)	24 (28)	0.09	
2 mo postpartum			18 (21)	18 (21)	0.05	
Sandborg <sup>66</sup> 1086 (LIS)		50	10 (21)	10 (21)	0.00	
	DDI (~9)	50	8 (16)	24 (48)		
			۵ (۱۵) ۸ (۹)	17 (24)		
Onset of poternal depression at	nootnortum <2		4 (0)	17 (34)		
Ballard et al. <sup>19</sup> 1994 (UK)	postpartum <3	mo				
1.5 mo postpartum	EPDS (>12)	178	16 (9)	49 (27.5)		
6 mo postpartum	EPDS (>12)	148	8 (5.4)	38 (25.7)		
6 mo postpartum	PAS	148	6 (4.1)	23 (15.5)		
Carro et al, <sup>39</sup> 1993 (US)						
1 mo postpartum	BDI (>9)	70	7 (10)	20 (29)	0.25	
Davé et al, <sup>61</sup> 2005 (UK)		48				
1.25 mo postpartum	HADS (>7)		4 (8)			
	EPDS (>12)		4 (8)			
Edhborg et al, <sup>41</sup> 2005 (Sweden)	EPDS (>9)	106				
0.25 mo postpartum			3 (2.8)	22 (20.8)		
2 mo postpartum			1 (0.9)	10 (9.4)		
Edhborg, <sup>62</sup> 2008 (Sweden)	EPDS (>10)					
0.25 mo postpartum		132 (167)	4 (3)	40 (24) <sup>d</sup>		
2 mo postpartum		113 (155)	2 (1.8)	19 (12)		
Ferketich and Mercer, <sup>42</sup> 1995 (US)	CES-D (>15)	172				
0.5 mo postpartum			36 (20.9)			
1 mo postpartum			30 (17.4)			
4 mo postpartum			25 (14.5)			
8 mo postpartum			28 (16.3)			
Gao et al, <sup>46</sup> 2009 (China)						
1.5 mo postpartum	EPDS (>12)	130	14 (10.8)	18 (13.8)	0.37	
Goodman, <sup>47</sup> 2008 (US) 2.5 mo postpartum	EPDS (>9)	128	17 (13.3)	36 (28)	0.34	
Greenhalgh et al, <sup>48</sup> 2000 (UK)	EPDS (>12)					
0.25 mo postpartum		78	5 (6.4)			
1.5 mo postpartum		64	4 (6.3)			
Hjelmstedt and Collins, <sup>13</sup>						
2008 (Sweden)° 2 mo postpartum	EPDS (>9)	53	4 (7 5)			
Lane et al. <sup>50</sup> 1997 (Ireland)	EPDS (>12)		. (			
	- v · -/					

# 0.1 mo postpartum 175 (289) 6 (3) 33 (11.4) 1.5 mo postpartum 175 (224) 2 (1.2) 24 (10.7)

(continued)

(Reprinted) JAMA, May 19, 2010-Vol 303, No. 19 1965

Table. Characteristics of Studies Included in Meta-analysis (cor	ntinued)
--	----------

Source (Study Leastion)	Depression Measure (Cutoff)	No. of Participants (Women) <sup>a</sup>	Depressed, No. (%)		Correlation Between
and Time of Assessment			Men	Women	Women <sup>b</sup>
Onset of paternal depression at Madsen and Juhl, <sup>15</sup> 2007 (Denmark)	postpartum <3	mo (continued)			
1.5 mo postpartum	EPDS (>9)	542	27 (5)		
1.5 mo postpartum	GMDS (>12)	529	18 (3.4)		
Matthey et al, <sup>64</sup> 2001 (Australia) 1.6 mo postpartum	DIS	208 (230)	6 (2.9)	24 (10.4)	
Mezulis et al.65 2004 (US)	CES-D (>15)	350	. ,	. ,	0.12 <sup>f</sup>
1 mo postpartum	0101(10)		55 (15 6)	41 (11 6)	
4 mo postpartum			47 (13.3)	31 (8.8)	
12 mo postpartum			36 (10.2)	21 (5.0)	
Pinheiro et al, <sup>54</sup> 2006 (Brazil) 2.25 mo postpartum	BDI (>9)	386	46 (11.9)	91 (23.6)	0.529
Skari et al. <sup>56</sup> 2002 (Norway)	GHQ (>1) <sup>h</sup>		. ,	. ,	
0.25 mo postpartum		115 (126)	2 (1 7)	7 (5 6)	
1.5 mo postpartum		103 (109)	2 (1.9)	1 (0.9)	
6 mo postpartum		84 (91)	1 (1 2)	2 (2 2)	
0.25 mo postpartum		115 (124)	13 (11 3)	16 (37.1)	
		100 (124)	11 (10.9)	40 (07.1)	
		102 (106)	0 (10.7)	23 (21.3)	
		64 (91)	9(10.7)	17 (16.7)	
Soliday et al, <sup>37</sup> 1999 (US)	CES-D (>16)	51	13 (25 5)	20 (30 2)	0.20
Thorpo et al 68 1002 (LIK/Crasso)	010-0 (> 10)	51	10 (20.0)	20 (03.2)	0.23
1 mo postpartum	FPDS (>12)	267 (281)	2 (0 7)	35 (12 5)	
Wang and Chen, <sup>58</sup> 2006 (Taiwan)		201 (201)	2 (0.17)	00 (12.0)	
1.5 mo postpartum	BDI (>9)	83	26 (31.3)	33 (39.8)	
Onset of paternal depression at Bielawska-Batorowicz and Kossakowska- Petrycka, <sup>38</sup> 2006 (Poland) 4.5 mo postpartum	EPDS (>12)	<b>mo</b> 80	22 (27.5)	25 (31.2)	0.76
Dudley et al, <sup>40</sup> 2001 (Australia) <sup>e</sup>					
3.9 mo postpartum	EPDS (>10)	93 (158)	11 (11.8)	75 (47.5) <sup>a</sup>	0.33
	GHQ (>4)	93	43 (46.2)		0.27
	BDI (>9)	92	16 (17.4)		0.29
Smart and Hiscock, <sup>67</sup> 2007 (Australia) <sup>e</sup>		50 (71)	19 (20)	22 (45)İ	
	LI DO (23)	53 (71)	10 (00)	0 (15)	
4.5 mo posipartum		53 (59)	10 (19)	9(15)	
Bronte-Tinkew et al <sup>16</sup> 2007 (US)	postpartum >6	mo			
12 mo postpartum	CIDI-SF	2137	115 (5.4)	143 (6.7)	
Leathers et al, <sup>51</sup> 1997 (US)			. ,		
6 mo postpartum	CES-D (>15)	55	10 (18)	17 (31)	
Paulson et al, <sup>10</sup> 2006 (US) 9 mo postpartum	CES-D (>9)	5089	509 (10)	712 (14)	

Abbreviations: BDI, Beck Depression Inventory; BSI, Brief Symptom Inventory; CES-D, Center of Epidemiologic Studies Depression Scale; CIDI-SF, Composite International Diagnostic Interview Short Form; DIS, Diagnostic Interview Sched-ule; EPDS, Edinburgh Postnatal Depression Inventory; GHQ, General Health Questionnaire; GMDS, Gotland Male De-pression Scale; HADS, Hospital Anxiety Depression Scale; MHI-5, Mental Health Index of the Short Form–36 health survey; PAS, Psychiatric Assessment Schedule; SADS, Schedule for Affective Disorders and Schizophrenia.

<sup>a</sup>Numbers in parentheses represent the number of women who participated at each time point. If a number in parentheses does not appear in this column and a percentage of depressed women is reported in the table, the number of female participants is the same as the number of male participants. <sup>b</sup> All correlations are Pearson *r* correlation coefficients unless otherwise noted.

C Fathers were assessed for depression using the BDI (>16 from time 1 through time 4) and the GHQ (>7 at time 1 and time 4 only). Mothers were assessed using the BDI (>16 at time 1, time 3, and time 4), EPDS (>12 at time 2 only), and GHQ (>7 at time 1 and time 4 only).

<sup>d</sup>Percentage of depressed women based on EPDS greater than 9.

<sup>e</sup> The studies by Hjelmstedt and Collins<sup>13</sup> (child conceived through assisted reproductive technology), Dudley et al,<sup>40</sup> and Smart and Hiscock<sup>67</sup> (infant crying, sleeping, or eating problems in both) were considered to include high-risk individuals. <sup>f</sup>Point biserial.

<sup>g</sup>Spearman correlation

<sup>h</sup>A GHQ depression subscale case score greater than 1 indicates clinically important depression.

A GHQ total case score greater than 5 indicates clinically important psychological distress

Percentages of depressed women based on EPDS greater than 12.

1966 JAMA, May 19, 2010-Vol 303, No. 19 (Reprinted)

©2010 American Medical Association. All rights reserved.

sion, wide variation was observed in reported rates of depression for fathers and mothers. The overall metaanalytic rate of paternal depression between the first trimester and 1 year postpartum was 10.4%. Since recent national data on base rates of depression in men place the 12-month prevalence at 4.8%,<sup>70</sup> this suggests that paternal prenatal and postpartum depression represents a significant public health concern. It must be noted that considerable variability was observed in reported rates of paternal depression. Although timing of measurement, study location, and measurement method were significant predictors, they accounted for only a small amount of overall heterogeneity. In terms of timing, fathers experienced the highest rates of depression 3 to 6 months postpartum, although the small number of studies measuring paternal depression during this period suggests cautious interpretation. Differences were also observed across study locations, with higher rates of prenatal and postpartum depression reported in the United States (14.1% vs 8.2% internationally). Questionnaire methods of case identification produced somewhat higher rates than did interview methods, although this should be interpreted cautiously because of the small number of studies that used interviews. Surprisingly, sample risk status was not a determinant of depression rates.

Maternal depression demonstrated considerable heterogeneity. This varied by time period, with a peak rate of 41.6% in the 3- to 6-month postpartum period, and by measurement method (higher rates with rating scales). Our random-effects estimate is somewhat larger than that of some reports,<sup>1,2</sup> with the variability in rates being clearly observable.

In both men and women, the potential causes of unexplained heterogeneity are varied. Although interview vs self-report questionnaire methods were compared, there were too many different questionnaires and interviews to conduct an instrument-by-instru-

Figure 2. Prevalence of Paternal Birth-Related Depression From Gestation to 1 Year Postpartum

a clinical diagnosis typically more conservative in making tured interviews, a method that is cation observed in studies using struccases. This interpretation is also con-sistent with the lower rates of identifisive disorder were not used to identify diagnostic criteria for major depresscale cutoff scores suggests that strict nor depression, but the use of rating reported herein explicitly described mipressive disorder.71 Few of the studies duration of symptoms, for major decriteria, either by severity, number, or and impairment who do not meet strict dividuals with depressive symptoms pression, a category that includes incases that can be classified as minor deheterogeneity. One possible source of teristics may also contribute to ment moderator analysis. Sample loca-tion and undetermined sample characheterogeneity is the liberal inclusion of

yet available. tion, but strong evidence of this is not implications for screening and prevenare of great interest, particularly for their Studies that speak to direction of effect suggest direction of causal influence. studies included in this meta-analysis paternal depression,5,11,40 none of the pression may play some causal role in thors have suggested that maternal deby most standards. Although other autimate, 0.308), a moderate association tions were significant (meta-analytic esmeta-analysis, 12 of the 14 correlators of maternal depression.72 In our sion, is among the strongest predicsatisfaction, a close correlate of deprespression suggests that marital analysis of maternal postpartum dedren. Moreover, an extant metapaired with index mothers or chilexclusively in the context of fathers depression has been examined almost tum depression is important, as paternal text of paternal prenatal and postpartoms. Examining this effect in the conmaternal and paternal depressive sympanalysis was the correlation between Another area of focus for this meta-

First, because studies used variable methods of measuring and reporting de-

Study Postpartum: 3 to 6 mo Bielawska-Batrorwicz and Kossakowska-Petrycka,<sup>38</sup> 2006 Dudley et al.<sup>4,2</sup> 2001 Smart and Hiscock,<sup>67</sup> 2007 Postpartum: birth to 3 mo Ballard et al.<sup>19</sup> 1994 Carro et al.<sup>39</sup> 1993 Davé et al.<sup>61</sup> 2005 Edhborg et al.<sup>41</sup> 2005 Edhborg <sup>62</sup> 2008 Ferketich and Mercer,\*<sup>2</sup> 1995 Gao et al.<sup>46</sup> 2009 Goodman,<sup>47</sup> 2008 Greenhaldh et al.<sup>46</sup> 2000 Hjelmstedt and Collins,<sup>13</sup> 2008 Lane et al.<sup>50</sup> 1997 Matthey et al.<sup>52</sup> 2001 Mezulis et al.<sup>55</sup> 2004 Pinheiro et al.<sup>56</sup> 2006 Skari et al.<sup>56</sup> 2002 Soliday et al.<sup>57</sup> 1999 Thorpe et al.<sup>68</sup> 1992 Wang and Chen.<sup>58</sup> 2006 Leathers and Kelley,<sup>51</sup> 2000 Morse et al.<sup>52</sup> 2000 Raskin et al.<sup>59</sup> 1990 Sandberg.<sup>66</sup> 1996 Gestation: first and second trimester Areias et al,  $^{\rm 17}$  1996 Overall across all periods Test for heterogeneity:  $l^2 = 94.91$ ; P<.001 Postpartum: 6 to 12 mo Bronte-Tinkew et al, <sup>16</sup> 2007 Leathers et al, <sup>51</sup> 1997 Gestation: third trimester Atkinson and Rickel,<sup>37</sup> 1984 Bourne,<sup>60</sup> 2006 Condon et al.<sup>6</sup> 2004 Fawcett and York, <sup>14</sup> 1986 Field et al.<sup>43</sup> 2006 Fletcher et al.<sup>44</sup> 2008 Frost, <sup>45</sup> 1996 Escribè-Agüir et al,<sup>8</sup> 2008 Hall and Long,<sup>63</sup> 2007 Keeton et al,<sup>49</sup> 2008 Matthey et al,<sup>20</sup> 2000 Ramchandani et al,<sup>55</sup> 2008 van den Berg et al,<sup>7</sup> 2009 Overall Overall Overall Paulson et al,<sup>10</sup> 2006 Overall Overall 42 312 23 156 307 527 152 10 975 3083 Sample Size, No. 2137 55 5089 78 120 669 98 152 124 50 592 80 0.13 (0.07-0.22) 0.14 (0.09-0.22) 0.14 (0.09-0.22) 0.16 (0.06-0.19) 0.14 (0.09-0.2) 0.14 (0.09-0.2) 0.08 (0.04-0.14) 0.12 (0.08-0.17) 0.19 (0.12-0.28) 0.16 (0.08-0.29) 0.09 (0.09-0.14) 0.10 (0.65-0.20) 0.03 (0.01-0.08) 0.03 (0.01-0.08) 0.21 (0.15-0.28) 0.21 (0.16-0.28) 0.11 (0.06-0.17) 0.13 (0.08-0.02-0.14) 0.04 (0.02-0.11) 0.03 (0.01-0.07) 0.03 (0.01-0.07) 0.03 (0.02-0.05) 0.16 (0.12-0.20) 0.02 (0.00-0.07) 0.02 (0.00-0.07) 0.02 (0.00-0.07) 0.02 (0.00-0.07) 0.02 (0.00-0.07) 0.02 (0.00-0.07) 0.02 (0.00-0.07) 0.02 (0.00-0.07) 0.02 (0.00-0.07) 0.05 (0.01-0.17) 0.18 (0.14-0.23) 0.04 (0.01-0.25) 0.32 (0.25-0.40) 0.16 (0.12-0.17) 0.05 (0.03-0.10) 0.04 (0.04-0.04) 0.12 (0.11-0.13) 0.05 (0.05-0.06) 0.16 (0.09-0.29) 0.10 (0.09-0.11) 0.17 (0.11-0.27) 0.34 (0.23-0.48) 0.10 (0.08-0.13) 0.09 (0.05-0.15) 0.26 (0.17-0.36) 0.28 (0.19-0.38) 0.08 (0.05-0.11) 0.12 (0.09-0.15) 0.11 (0.06-0.18) Effect Size (95% CI) 0.00 • 🔳 + + ╷┿<sup>┿</sup> ŧ ł  $\Diamond$ ÷. • • 1 1 L 0.30 Event Rate (95% Cl) t.

Overall effects were calculated through random-effects model estimates, with separate calculations for overall effects within each period and across all periods. Effect sizes were calculated via a logit transformation of rates (number of reported cases divided by the sample size), which were back-transformed to proportions after estimates and standard errors were computed. Studies are stratified by period d assessment. For studies that assessed depression at multiple time points, only the earliest estimate is reported. Data marker size corresponds to study sample size.

0.60

(Reprinted) JAMA, May 19, 2010-Vol 303, No.

19 **1967** 

pression in different time periods, time frame-specific prevalence cannot be clearly established, limiting interpretation to the rate of depression observed at that point in time. Also, since point estimates are drawn from a pool of heterogeneous studies, many of which did not use strong populationbased sampling methods, there is a potential of bias in our results from studies' methodological weaknesses. These may not have been adequately accounted for by our simplified method of quality rating. The method of identifying depressed cases was highly variable across studies, thereby limiting the specificity of our primary outcome. However, this variability in case identification accurately reflects inconsistencies in both applied and basic research into prenatal and postpartum depression.<sup>5</sup> Removing relatively weaker studies in sensitivity analysis left effects essentially unchanged. We did not find substantial evidence of publication bias in this area, and fail-safe analysis suggested that our findings are robust to unpublished null findings.

With these limitations in mind, this meta-analysis allows us to draw several conclusions regarding paternal prenatal and postpartum depression. First, a significant number of expecting and new fathers experience depression during this period. Second, expecting and new fathers in the United States experience depression at marginally higher rates than do fathers internationally, a finding that bears further investigation vis-à-vis varying social norms and postpartum work practices cross-nationally. Third, there is a moderate correlation between depression in fathers and mothers. There are many implications of these findings. The observation that expecting and new fathers disproportionately experience depression suggests that more efforts should be made to improve screening and referral, particularly in light of the mounting evidence that early paternal depression may have substantial emotional, behavioral, and developmental effects on children.<sup>10,55</sup> The correlation between paternal and maternal depression also suggests a screening rubric<sup>73</sup>—depression in one parent should prompt clinical attention to the other. Likewise, prevention and intervention efforts for depression in parents might be focused on the couple and family rather than the individual.

Future research in this area should focus on parents together to examine the onset and joint course of depression in new parents. This may increase our capacity for early identification of parental depression, add leverage for prevention and treatment, and increase the understanding of how parental depression conveys risk to infants and young children.

Author Contributions: Dr Paulson had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Study concept and design: Paulson.

Acquisition of data: Paulson, Bazemore. Analysis and interpretation of data: Paulson, Bazemore. Drafting of the manuscript: Paulson, Bazemore. Critical revision of the manuscript for important in-

tellectual content: Paulson, Bazemore.

Statistical analysis: Paulson, Bazemore.

Administrative, technical, or material support: Paulson, Bazemore.

Study supervision: Paulson.

Financial Disclosures: None reported.

#### REFERENCES

**1.** Gotlib IH, Whiffen V, Mount J, Milne K, Cordy N. Prevalence rates and demographic characteristics associated with depression in pregnancy and postpartum. *J Consult Clin Psychol.* **1989**;57(2):269-274.

2. Bennett HA, Einarson A, Taddio A, Koren G, Einarson TR. Prevalence of depression during pregnancy: systematic review. *Obstet Gynecol*. 2004;103(4): 698-709.

**3.** Beardslee WR, Versage EM, Gladstone TR. Children of affectively ill parents: a review of the past 10 years. *J Am Acad Child Adolesc Psychiatry*. 1998; 37(11):1134-1141.

4. Cox J. Postnatal depression in fathers. *Lancet*. 2005; 366(9490):982.

5. Goodman JH. Paternal postpartum depression, its relationship to maternal postpartum depression, and implications for family health. J Adv Nurs. 2004; 45(1):26-35.

**6.** Condon JT, Boyce P, Corkindale CJ. The First-Time Fathers Study: a prospective study of the mental health and wellbeing of men during the transition to parenthood. *Aust N Z J Psychiatry*. 2004;38 (1-2):56-64.

 van den Berg MP, van der Ende J, Crijnen AA, et al. Paternal depressive symptoms during pregnancy are related to excessive infant crying. *Pediatrics*. 2009; 124(1):e96-e103.

 Escribè-Agüir V, Gonzalez-Galarzo MC, Barona-Vilar C, Artazcoz L. Factors related to depression during pregnancy: are there gender differences? J Epidemiol Community Health. 2008;62(5):410-414.

9. Ramchandani P, Stein A, Evans J, O'Connor TG;

ALSPAC Study Team. Paternal depression in the postnatal period and child development: a prospective population study. *Lancet*. 2005;365(9478):2201-2205.

**10.** Paulson JF, Keefe HA, Leiferman JA. Early parental depression and child language development. *J Child Psychol Psychiatry*. 2009;50(3):254-262.

**11.** Schumacher M, Zubaran C, White G. Bringing birth-related paternal depression to the fore. *Women Birth.* 2008;21(2):65-70.

**12.** Leathers SJ, Kelley MA. Unintended pregnancy and depressive symptoms among first-time mothers and fathers. *Am J Orthopsychiatry*. 2000;70(4): 523-531.

**13.** Hjelmstedt A, Collins A. Psychological functioning and predictors of father-infant relationship in IVF fathers and controls. *Scand J Caring Sci.* 2008; 22(1):72-78.

**14.** Fawcett J, York R. Spouses' physical and psychological symptoms during pregnancy and the postpartum. *Nurs Res.* **1986**;35(3):144-148.

**15.** Madsen SA, Juhl T. Paternal depression in the postnatal period assessed with traditional and male depression scales. *J Mens Health Gend*. 2007;4(1): 26-31.

**16.** Bronte-Tinkew J, Moore KA, Matthews G, Carrano J. Symptoms of major depression in a sample of fathers of infants: sociodemographic correlates and links to father involvement. *J Fam Issues*. 2007;28(1): 61-99.

**17.** Areias MEG, Kumar R, Barros H, Figueiredo E. Comparative incidence of depression in women and men, during pregnancy and after childbirth: validation of the Edinburgh Postnatal Depression Scale in Portuguese mothers. *Br J Psychiatry*. 1996;169(1):30-35.

**18.** Matthey S, Barnett B, Howie P, Kavanagh DJ. Diagnosing postpartum depression in mothers and fathers: whatever happened to anxiety? *J Affect Disord*. 2003;74(2):139-147.

**19.** Ballard CG, Davis R, Cullen PC, Mohan RN, Dean C. Prevalence of postnatal psychiatric morbidity in mothers and fathers. *Br J Psychiatry*. 1994;164 (6):782-788.

**20.** Matthey S, Barnett B, Ungerer J, Waters B. Paternal and maternal depressed mood during the transition to parenthood. *J Affect Disord*. 2000;60 (2):75-85.

**21.** Cooper HM. Synthesizing Research: A Guide for Literature Reviews. Vol 2. 3rd ed. Thousand Oaks, CA: Sage Publications; 1998.

**22.** Cooper H, Hedges L, Valentine JC. *Handbook of Research Synthesis and Meta-Analysis.* 2nd ed. New York, NY: Russell Sage Foundation; 2009.

23. Stroup DF, Berlin JA, Morton SC, et al; Metaanalysis Of Observational Studies in Epidemiology (MOOSE) Group. Meta-analysis of observational studies in epidemiology: a proposal for reporting. JAMA. 2000;283(15):2008-2012.

**24.** Buist A, Morse CA, Durkin S. Men's adjustment to fatherhood: implications for obstetric health care. *J Obstet Gynecol Neonatal Nurs*. 2003;32(2):172-180.

**25.** Boyce P, Condon J, Barton J, Corkindale C. Firsttime fathers' study: psychological distress in expectant fathers during pregnancy. *Aust N Z J Psychiatry*. 2007;41(9):718-725.

26. Hjelmstedt A, Widstrom AM, Wramsby H, Matthiesen AS, Collins A. Personality factors and emotional responses to pregnancy among IVF couples in early pregnancy: a comparative study. *Acta Obstet Gynecol Scand*. 2003;82(2):152-161.

**27.** Ballard CG, Davis R, Handy S, Mohan RNC. Postpartum anxiety in mothers and fathers. *Eur J Psychiatry*. 1993;7(2):117-121.

**28.** Beck AT, Steer RA. *Manual for the Beck Depression Inventory*. San Antonio, TX: Psychology Corp; Harcourt Brace; 1993.

1968 JAMA, May 19, 2010-Vol 303, No. 19 (Reprinted)

29. Cox JL, Holden JM, Sagovsky R. Detection of postnatal depression-development of the 10-item Edinburgh Postnatal Depression Scale. *Br J Psychiatry*. 1987; 150:782-786.

**30.** Berwick DM, Murphy JM, Goldman PA, Ware JE Jr, Barsky AJ, Weinstein MC. Performance of a fiveitem mental health screening test. *Med Care*. 1991; 29(2):169-176.

**31.** Lipsey MW, Wilson DB. *Practical Meta-analysis.* Thousand Oaks, CA: Sage Publications; 2000.

**32.** Comprehensive Meta-Analysis [computer program]. Version 2.0. Englewood, NJ: Biostat; 2009.

**33.** Affonso DD, De AK, Horowitz JA, Mayberry LJ. An international study exploring levels of postpartum depressive symptomatology. *J Psychosom Res.* 2000;49(3):207-216.

**34.** Egger M, Davey Smith G, Schneider M, Minder C. Bias in meta-analysis detected by a simple, graphical test. *BMJ*. 1997;315(7109):629-634.

**35.** Orwin RG. A fail-safe N for effect size in meta-analysis. *J Educ Stat.* 1983;8(2):157-159.

**36.** American Psychiatric Association. *Diagnostic and Statistical Manual of Mental Disorders*. 4th ed, text rev. Washington, DC: American Psychiatric Association; 2000.

**37.** Atkinson AK, Rickel AU. Postpartum depression in primiparous parents. *J Abnorm Psychol*. 1984; 93(1):115-119.

**38.** Bielawska-Batorowicz E, Kossakowska-Petrycka K. Depressive mood in men after the birth of their off-spring in relation to a partner's depression, social support, fathers' personality and prenatal expectations. *J Reprod Infant Psychol.* 2006;24(1):21-29.

**39.** Carro MG, Grant KE, Gotlib IH, Compas BE. Postpartum depression and child development: an investigation of mothers and fathers as sources of risk and resilience. *Dev Psychopathol*. 1993;5:567-579.

**40.** Dudley M, Roy K, Kelk N, Bernard D. Psychological correlates of depression in fathers and mothers in the first postnatal year. *J Reprod Infant Psychol*. 2001; 19(3):187-202.

**41.** Edhborg M, Matthiesen AS, Lundh W, Widstrom AM. Some early indicators for depressive symptoms and bonding 2 months postpartum: a study of new mothers and fathers. *Arch Womens Ment Health*. 2005;8(4):221-231.

**42.** Ferketich SL, Mercer RT. Predictors of role competence for experienced and inexperienced fathers. *Nurs Res.* 1995;44(2):89-95.

**43.** Field T, Diego M, Hernandez-Reif M, et al. Prenatal paternal depression. *Infant Behav Dev.* 2006; 29(4):579-583.

**44.** Fletcher R, Vimpani G, Russell G, Sibbritt D. Psychosocial assessment of expectant fathers. *Arch Womens Ment Health.* 2008;11(1):27-32.

**45.** Frost LA. Postpartum Distress in Fathers: Predicting Depressive Symptoms, Anxiety, and Anger at One

Month Postpartum [dissertation]. Madison: University of Wisconsin; 1996.

**46.** Gao LL, Chan S, Mao Q. Depression, perceived stress, and social support among first-time Chinese mothers and fathers in the postpartum period. *Res Nurs Health*. 2009;32(1):50-58.

**47.** Goodman JH. Influences of maternal postpartum depression on fathers and father-infant interaction. *Infant Ment Health J.* 2008;29(6):624-643.

**48.** Greenhalgh R, Slade P, Spiby H. Fathers' coping style, antenatal preparation, and experiences of labor and the postpartum. *Birth*. 2000;27(3):177-184.

49. Keeton CP, Perry-Jenkins M, Sayer AG. Sense of control predicts depressive and anxious symptoms across the transition to parenthood. J Fam Psychol. 2008;22(2):212-221.

**50.** Lane A, Keville R, Morris M, Kinsella A, Turner M, Barry S. Postnatal depression and elation among mothers and their partners: prevalence and predictors. *Br J Psychiatry*. 1997;171:550-555.

**51.** Leathers SJ, Kelley MA, Richman JA. Postpartum depressive symptomatology in new mothers and fathers: parenting, work, and support. *J Nerv Ment Dis.* 1997;185(3):129-139.

**52.** Morse CA, Buist A, Durkin S. First-time parenthood: influences on pre- and postnatal adjustment in fathers and mothers. *J Psychosom Obstet Gynaecol*. 2000;21(2):109-120.

**53.** Paulson JF, Dauber S, Leiferman JA. Individual and combined effects of postpartum depression in mothers and fathers on parenting behavior. *Pediatrics*. 2006; 118(2):659-668.

**54.** Pinheiro RT, Magalhaes PVS, Horta BL, Pinheiro KAT, da Silva RA, Pinto RH. Is paternal postpartum depression associated with maternal postpartum depression? population-based study in Brazil. *Acta Psychiatr Scand.* 2006;113(3):230-232.

**55.** Ramchandani PG, Stein A, O'Connor TG, Heron J, Murray L, Evans J. Depression in men in the postnatal period and later child psychopathology: a population cohort study. *J Am Acad Child Adolesc Psychiatry*. 2008;47(4):390-398.

**56.** Skari H, Skreden M, Fredrik U, et al. Comparative levels of psychological distress, stress symptoms, depression and anxiety after childbirth: a prospective population-based study of mothers and fathers. *Int J Gynaecol Obstet*. 2002;109:1154-1163.

**57**. Soliday E, McCluskey-Fawcett K, O'Brien M. Postpartum affect and depressive symptoms in mothers and fathers. *Am J Orthopsychiatry*. 1999;69(1): 30-38.

**58.** Wang SY, Chen CH. Psychosocial health of Taiwanese postnatal husbands and wives. *J Psychosom Res*, 2006:60(3):303-307.

**59.** Raskin VD, Richman JA, Gaines C. Patterns of depressive symptoms in expectant and new parents. *Am J Psychiatry*. 1990;147(5):658-660.

60. Bourne H. Gender Ideology, Depression, and Mari-

tal Quality in Working-Class, Dual-Earner Couples Across the Transition to Parenthood [dissertation]. Amherst: University of Massachusetts; 2006.

**61.** Davé S, Nazareth I, Lorraine S, Senior R. The association of paternal mood and infant temperament: a pilot study. *Br J Dev Psychol*. 2005;23(4):609-621.

**62.** Edhborg M. Comparisons of different instruments to measure blues and to predict depressive symptoms 2 months postpartum: a study of new mothers and fathers. *Scand J Caring Sci.* 2008;22(2): 186-195.

**63.** Hall WA, Long BC. Relations among prenatal role quality, life satisfaction, and dual-earner parents' postnatal depression. *J Prenat Perinat Psychol Health*. 2007; 21(3):231-248.

**64.** Matthey S, Barnett B, Kavanagh DJ, Howie P. Validation of the Edinburgh Postnatal Depression Scale for men, and comparison of item endorsement with their partners. *J Affect Disord*. 2001;64(2-3):175-184.

**65.** Mezulis AH, Hyde JS, Clark R. Father involvement moderates the effect of maternal depression during a child's infancy on child behavior problems in kindergarten. *J Fam Psychol*. 2004;18(4):575-588.

**66.** Sandberg J. The Learned Helplessness Model of Depression: Locus of Control, Experienced Control, and Transient Postpartum Depression in First-Time Parents [dissertation]. Provo, Utah: Brigham Young University; 1986.

**67.** Smart J, Hiscock H. Early infant crying and sleeping problems: a pilot study of impact on parental well-being and parent-endorsed strategies for management. *J Paediatr Child Health*. 2007;43 (4):284-290.

**68.** Thorpe KJ, Dragonas T, Golding J. The effects of psychosocial factors on the mother's emotional wellbeing during early parenthood: a cross-cultural study of Britian and Greece. *J Reprod Infant Psychol*. 1992; 10:205-217.

**69.** Higgins JP, Thompson SG. Quantifying heterogeneity in a meta-analysis. *Stat Med.* 2002;21 (11):1539-1558.

**70.** Kessler RC, Berglund P, Demler O, et al; National Comorbidity Survey Replication. The epidemiology of major depressive disorder: results from the National Comorbidity Survey Replication (NCS-R). *JAMA*. 2003;289(23):3095-3105.

**71.** Marchesi C, Bertoni S, Maggini C. Major and minor depression in pregnancy. *Obstet Gynecol*. 2009; 113(6):1292-1298.

**72.** Beck CT. A meta-analysis of predictors of postpartum depression. *Nurs Res.* 1996;45(5):297-303.

**73.** Freeman MP, Wright R, Watchman M, et al. Postpartum depression assessments at well-baby visits: screening feasibility, prevalence, and risk factors. *J Womens Health (Larchmt)*. 2005;14(10):929-935.