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1 **Postpartum Psychiatric Disorders**

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17

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22

23

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32

33

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35 Introduction (S.M-B. and all co-authors); Epidemiology (T.M-O.); Mechanisms/pathophysiology (I.J.  
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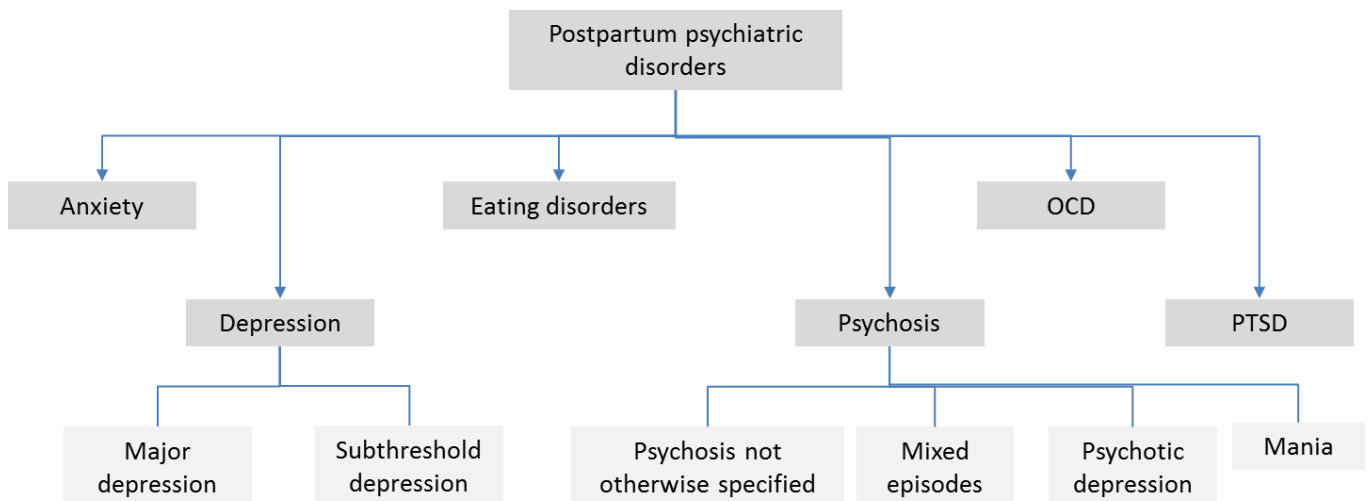
38 **Abstract**

39 Pregnancy is a complex and vulnerable period that presents a number of challenges to women,  
40 including the development of postpartum psychiatric disorders (PPDs). These disorders can include  
41 postpartum depression and anxiety, which are relatively common, and the rare but more severe  
42 postpartum psychosis. In addition, other PPDs can include obsessive–compulsive disorder, post-  
43 traumatic stress disorder and eating disorders. The aetiology of PPDs is a complex interaction of  
44 psychological, social and biological factors, in addition to genetic and environmental factors. The  
45 goals of treating postpartum mental illness are reducing maternal symptoms and supporting  
46 maternal-child and family functioning. Women and their families should receive psychoeducation  
47 about the illness, including evidence-based discussions on risks and benefits of each treatment  
48 option. Developing effective strategies in global settings that allow the delivery of targeted therapies  
49 to women with different clinical phenotypes and severities of PPD is essential.

50  
51

## 52 [H1] Introduction

53 Pregnancy and the first year after childbirth (which collectively can be referred to as the perinatal  
54 period) is arguably one of the most transformative times in a woman's life. This timeframe is also a  
55 complex and vulnerable period that presents a number of challenges for women. In particular, an  
56 increased risk for onset or worsening of psychiatric illness including mood disorders, anxiety  
57 disorders and psychosis exists during the first three months postpartum. All types of psychiatric  
58 disorders can occur during the postpartum period, with many chronic disorders starting before  
59 pregnancy and persisting throughout pregnancy into the postpartum period<sup>1</sup>. In this Primer, we  
60 focus specifically on the postpartum psychiatric disorders (PPDs). Collectively, the postpartum  
61 psychiatric disorders (PPDs) can include postpartum depression, which is relatively common;  
62 common anxiety disorders such as generalized anxiety disorder (GAD; which can include anxieties  
63 about the health of the baby); post-traumatic stress disorder (PTSD; which can occur due to a  
64 traumatic childbirth experience or can reflect pre-existing symptoms due to previous traumas  
65 before conception or during pregnancy); and the rarer, but usually severe presentation of  
66 postpartum psychosis. Other PPDs include eating disorders (which can worsen or recur  
67 postpartum, particularly when the infant is undergoing weaning), and obsessive–compulsive  
68 disorder (OCD)



70

71 In pregnancy, depressive and anxiety disorders are common with recent population estimates of  
72 11% for depressive disorders and 15% for anxiety disorders.<sup>2</sup> Further, antenatal anxiety and  
73 depression are one of the greatest risk factors for postpartum psychiatric disorders (PPD).<sup>3</sup>  
74 Inadequate social support and a history of adverse life events increases the risk for PPDs in all  
75 countries and levels of society<sup>4, 5</sup>. However, this risk is increased in poorer socioeconomic  
76 populations and lower income countries, due to poverty and limited access to health care.<sup>6</sup>

77 In recent years, awareness of the potentially serious adverse consequences in both the mother  
78 and the baby associated with untreated perinatal psychiatric illness has increased. Maternal  
79 suicide due to postpartum mood disorders (including unipolar and bipolar depressive disorders) is  
80 a leading cause of maternal mortality.<sup>7, 8, 9</sup> In addition, perinatal mood disorders are associated with  
81 an increased risk for low birth weight and premature birth, impaired mother-infant attachment, and  
82 infant malnutrition during the first year of life<sup>10, 11</sup>.

83 In this Primer, we focus on maternal PPDs, as they are common, morbid, and have a growing  
84 literature on the underlying pathophysiology. These disorders should not be confused with the so-  
85 called 'baby blues', which are usually described as transient mild mood and anxiety symptoms that  
86 often persist  $\leq 2$  weeks and usually resolve spontaneously with no sequelae.<sup>12</sup> If the symptoms of  
87 the 'baby blues' worsen and/or persist, they are considered PPDs. Herein we discuss the  
88 epidemiology of PPDs, and their underlying mechanisms and pathophysiology. We mainly focus on  
89 maternal PPDs, although paternal disorders are mentioned in some instances (Box 1). Importantly,  
90 we review the latest evidence on diagnosis, screening and prevention as well as management of  
91 PPDs. Lastly, we hope to put in context the public health impact of these disorders on mothers,  
92 their children and families to encourage wide scale adoption of strategies that make maternal  
93 mental health a global priority<sup>13</sup>.

94

## 95 **[H1] Epidemiology**

96 Data on the incidence of postpartum depression are from studies conducted in countries across  
97 the world, and variable incidence and prevalence are reported between countries<sup>14</sup>. By  
98 comparison, studies estimating the incidence and prevalence of postpartum psychosis are  
99 primarily carried out in Europe<sup>15</sup>, and demonstrate less variability in reported incidence and  
100 prevalence<sup>16, 17</sup>. Several methodological factors have influenced these differences. <sup>18</sup> For example,  
101 particularly for postpartum depression, the definition of the postpartum period is variable between  
102 studies, and has been defined as up to 4 weeks, 3 months, 6 months or 12 months postpartum<sup>19</sup>.  
103 Differences in study designs, such as using different tools to define case-groups and phenotypes  
104 can lead to variability in the reported number of cases. Data sources for postpartum depression  
105 and postpartum psychosis include self-reports and interviews, in addition to some population-  
106 based register data<sup>16</sup>. Moreover, the incomplete availability of longitudinal data that is needed to  
107 distinguish between first-time and recurrent psychiatric episodes might impede calculations of the  
108 true incidence and prevalence of PPDs. Consequently, a variation in reported incidence and  
109 prevalence could be explained by differences in methodologies between studies, which make  
110 direct comparisons difficult. In addition, the diverse symptoms of PPDs pose specific challenges to  
111 the estimation of prevalence and incidence of these disorders<sup>20</sup>.

112 As the literature surrounding the epidemiology of PPDs continues to grow with well-designed  
113 studies, we will have a better understanding of if differences in the incidence and prevalence of  
114 postpartum depression and postpartum psychosis are due to local/regional and national  
115 differences, or if the differences are due to variable study designs and data sources. This  
116 knowledge will assist hypothesis generating that might provide clues for the aetiology of these  
117 disorders.

118

119 **[H2] Mood disorders and anxiety**

120 Postpartum depression, comprising major depressive disorder and subthreshold depression, has  
121 an estimated point prevalence of 13% in high-income countries<sup>11</sup>, and ~20% in low-income and  
122 middle-income countries, 3 months postpartum (Box 2)<sup>21</sup>. In women with a history of any eating  
123 disorder, the prevalence of postpartum depression has been estimated at 35%<sup>22</sup>. Studies of  
124 postpartum depression often rely on self-reported questionnaires, including the commonly used  
125 Edinburgh Postnatal Depression Scale (EPDS)<sup>14, 23</sup>.

126 Although the prevalence estimates for postpartum mood disorders ranges between studies,  
127 guidelines are available that state that these disorders pose substantial public health risks and  
128 consequently, must be identified and treated<sup>24,25</sup>. Moreover, there is consensus that childbirth is a  
129 strong and potent risk factor for bipolar disorder. Indeed, the risk of underlying bipolarity in first-  
130 onset depression that occurs in the postpartum period is higher than in first-onset depression that  
131 occurs outside the perinatal period. In addition, women with bipolar disorder have a high risk of  
132 postpartum episodes, including depression, anxiety, mania and psychosis<sup>26, 27</sup>.

133 The estimated prevalence of postpartum anxiety disorders is ~10%, with a prevalence of 6% for  
134 GAD<sup>28</sup>. Anxiety disorders have substantial comorbidity with postpartum depression and other  
135 disorders, including postpartum PTSD, eating disorders and the exacerbation of personality  
136 disorders<sup>17</sup>.

137

## 138 [H2] Postpartum psychosis

139 The onset of a severe mental disorder requiring acute inpatient psychiatric treatment in the first  
140 postpartum months is ~1 per 1,000 births<sup>29-32</sup>, and are considered some of the most severe forms  
141 of illness in psychiatry<sup>18</sup>. These severe psychiatric disorders that have an onset in the immediate  
142 postpartum period are often called postpartum psychosis, which is an umbrella term for disorders  
143 recorded as, for example, mania, mixed episodes, psychotic depression, or psychosis not  
144 otherwise specified<sup>33</sup>

145 [http://journals.sagepub.com/doi/abs/10.1177/0004867414564698?url\\_ver=Z39.88-](http://journals.sagepub.com/doi/abs/10.1177/0004867414564698?url_ver=Z39.88-)  
146  [. Women with bipolar  
147 disorder have the highest risk for postpartum psychosis than women with other psychiatric  
148 diagnosis, as the risk for postpartum relapse in women with bipolar disorder is on average 37%<sup>18</sup>.  
149 However, variations also occur within bipolar disorder, as the risk of a severe episode \(ie  
150 postpartum psychosis\) is greater for women with bipolar I disorder than women with bipolar II  
151 disorder<sup>34</sup>. Additionally, the risk of symptom recurrence is particularly high for women with bipolar  
152 disorder who are not receiving medication during pregnancy<sup>18</sup> .](http://journals.sagepub.com/doi/abs/10.1177/0004867414564698?url_ver=Z39.88-2003&rfr_id=ori%3Arid%3Acrossref.org&rfr_dat=cr_pub%3Dpubmed&)

153 Relapse of psychosis can also occur in women with other psychiatric disorders, such as women  
154 with schizophrenia, although this is less common<sup>35</sup> (~16% within the first 12 months postpartum)  
155 Yes—it has been added now. <https://www.ncbi.nlm.nih.gov/pubmed/19188541> , and manifests  
156 differently from what is observed in women with bipolar disorder<sup>16</sup>.

157 Despite the widespread use of the term postpartum psychosis, this diagnosis is not recognized in  
158 current classification systems, including both ICD-10 and DSM-5<sup>36</sup>. However, it is clear that  
159 psychotic episodes are more prevalent during the postpartum period than in other periods in a  
160 woman's life, and evidence clearly suggests a particular vulnerability in women with bipolar  
161 disorder<sup>29</sup>.

162  
163 In women with severe postpartum psychiatric illness, maternal suicide is often a predominate  
164 concern. Although maternal suicide is a leading cause of maternal mortality<sup>7</sup>, the rates of  
165 completed suicide in postpartum women are lower than those in age-matched women without  
166 children<sup>37</sup>. Nonetheless, the prevention of maternal suicide is paramount and requires careful  
167 monitoring during the postpartum period and possibly extending beyond the first year. For  
168 example, one study demonstrated that most postpartum suicides occurred between 9 and 12



169 months postpartum and that the perinatal suicides were by highly lethal means (such as via  
170 firearm), suggesting that limiting follow-up to 1, 3 or 6 months postpartum might be insufficient.<sup>38</sup>

171

## 172 **[H1] Mechanisms/pathophysiology**

173 As previously mentioned, childbirth is one of the most potent triggers of psychiatric illness. Given  
174 that postpartum mental health disorders are one of the few occurrences in psychiatry whereby a  
175 biological trigger occurs at a known time point, elucidating the pathophysiology of these disorders  
176 may shed light on the mechanisms of mood and psychiatric disorders more broadly, and is vital for  
177 developing new treatment approaches.

178 The aetiology of all psychiatric disorders, including PPDs, is a complex interaction of psychological,  
179 social and biological factors, including the effect of genetic and environmental influences on risk  
180 (Figure 1)<sup>12</sup>. The involvement of particular combinations of aetiological factors differs between  
181 specific PPDs<sup>34</sup>; for example, biological factors might have a greater role in the triggering of  
182 postpartum psychosis, whereas psychosocial factors might have an important contribution in  
183 postpartum depression.<sup>34</sup> These are areas of intense investigation and future research is needed to  
184 extend our understanding of the many ways that psychological and biological processes interface.  
185 Stopping or changing medications in women with a prior history of psychiatric disorders due to  
186 concerns about the safety of medication during pregnancy could be considered a simple  
187 explanation for the triggering of PPDs. However, continuing medication in pregnancy is protective  
188 against mood disorders in a subset of patients<sup>39</sup>. Similarly, discontinuation of medication does not  
189 guarantee that a woman will relapse<sup>40</sup>. However, there is much that is still not understood about  
190 PPDs and the onset of PPDs reflects the outcome of many different pathways that manifest in  
191 vulnerable women. Future research will need to disentangle the mechanisms of depression in  
192 women before, during and after pregnancy to increase our understanding of the similarities and  
193 differences between perinatal depression and depression occurring at other times in life. We will

194 next discuss current theories on psychosocial and biological contributions that increase risk for  
195 PPDs.

196

## 197 ***[H2] Psychosocial factors and comorbidities***

223 Psychological and social stressors contribute to the development of maternal PPDs and are  
224 associated with poorer outcomes in the infants or children<sup>41, 42</sup>. In particular, adverse life events  
225 and a history of trauma have a greater prevalence in women that develop postpartum mood  
226 disorders, compared with mood disorders outside of the perinatal period<sup>4, 43, 44</sup>. A history of  
227 adverse early life experiences can substantially affect a mother's ability to have a strong  
228 attachment with her infant<sup>45</sup>, and adverse parent–infant interactions and worse attachment are  
229 associated with development of PPDs<sup>46-48</sup>.

230 Social support has a vital role in either contributing to or mitigating the impact of postpartum mood  
231 disorders on both the mother and child<sup>49</sup>. Indeed, social support, or the degree of tangible support  
232 provided from the social network of the mother and from the partner (such as financial support or  
233 assisting with infant care), have the greatest effects on postpartum depression<sup>50</sup>.

234

235 Other psychosocial risk factors include a past history of a mood disorder, which is consistently  
236 associated with an increased risk of postpartum depression and postpartum psychosis<sup>34</sup>. In  
237 addition, although the strength of the association between risk factors and PPDs is variable  
238 between high-income countries and low-income and middle-income countries, one of the strongest  
239 psychosocial risk factors in both settings is domestic violence and previous abuse, including abuse  
240 during childhood.<sup>4, 51</sup> Other risk factors with a medium to strong association with PPDs include  
241 marital difficulties, migration status and antenatal depression or anxiety<sup>17, 52</sup>. In addition, poverty,  
242 young age (between 14 and 21 years of age), substance misuse, increased parity, multiple births,  
243 an unwanted pregnancy, neuroticism, pregnancy complications including obesity and comorbidities

244 (for example diabetes, hypertension and pre-eclampsia) and neonatal problems are associated with  
245 PPDs<sup>17</sup>.

246

## 247 **[H2] Genetic factors**

248 Data from twin and adoption studies have implicated genetic factors in the aetiology of psychiatric  
249 disorders outside of the perinatal period, including schizophrenia and mood disorders<sup>53, 54</sup>.

250 However, only recently have genetic investigations, primarily of postpartum depression and  
251 postpartum psychosis, been conducted.

252 Genetic epidemiological and linkage studies for postpartum depression have demonstrated the  
253 involvement of genetic factors<sup>55, 56, 57</sup> and two studies have demonstrated the increased heritability  
254 of postpartum depression compared with depression outside of the perinatal period<sup>58, 59</sup>. To date,  
255 studies have suggested that episodes of postpartum psychosis are a marker for a more-familial  
256 form of bipolar disorder and that the puerperal triggering of bipolar illness is familial.<sup>60, 61</sup> However,  
257 a genome-wide association study (GWAS) for either postpartum depression or postpartum  
258 psychosis using modern genomics methods has not yet been carried out. Future genetic studies of  
259 postpartum mood disorders using modern genomics methods will require international  
260 collaborations and consortia to include large number of patients; these studies are currently  
261 underway<sup>62, 63</sup>.

262 Psychological stressors and early life adverse events have a lasting negative impact and can result  
263 in pathophysiological changes and altered gene expression due to increased allostatic load (the  
264 cumulative stress on the body that is a sum of lifetime exposure to stress)<sup>64</sup>. Potential mechanisms  
265 underlying how stressful life events change gene expression include epigenetic modification (such  
266 as DNA methylation and histone modification that change DNA accessibility and chromatin  
267 structure, subsequently regulating gene expression)<sup>65</sup>, changes in transcriptional control of stress-  
268 responsive pathways<sup>66</sup>, and shortened telomere length<sup>67, 68</sup>. Epigenetic alterations have been

269 reported in two genes, *HP1BP3* and *TTC9B*, which have different methylation patterns in women  
270 with postpartum depression, depending on whether the mood symptoms begin during pregnancy  
271 and continue into the postpartum period, compared with symptoms that develop postpartum only<sup>69</sup>.  
272<sup>70</sup>. These data indicate that different gene patterns might arise based on the timing of symptom  
273 onset. However, given the history of non-replication in many genetic studies, these findings require  
274 replication and overall, the mechanism of action in postpartum depression remains to be  
275 established.

276

## 277 **[H2] Sleep Disruption**

278 An almost universal feature of pregnancy and childbirth is disruption to sleep. In addition, sleep  
279 and circadian rhythm disruption can trigger the onset of psychiatric disorders, particularly episodes  
280 of mania in the postpartum period<sup>71, 72</sup>. Thus, that circadian rhythm disruption has not received  
281 more attention as a potential mechanism in PPDs is surprising.

282 Numerous studies have demonstrated profound changes in maternal sleep patterns during the  
283 perinatal period. Pregnant women experience poorer subjective sleep quality, increased waking,  
284 and more sleep-wake transitions than women who are not pregnant<sup>73</sup>. In the postpartum period,  
285 new mothers have frequent night waking, decreased night-time sleep, increased daytime napping,  
286 and a more irregular sleep-wake schedule, which is speculated to increase the risk of PPDs<sup>74</sup>. The  
287 mechanisms underlying the reported disrupted maternal sleep patterns in the perinatal period have  
288 been reported in two cross-sectional studies. The first study demonstrated a blunted melatonin  
289 amplitude in postpartum women, compared with non-pregnant women,<sup>75</sup> and the second study  
290 demonstrated differences in circadian rhythms between perinatal women with depression and  
291 perinatal women without depression; indeed, in the second study, women with depression had  
292 clinically-significant circadian rhythm phase shifts.<sup>74</sup> Further research is needed to better

293 understand the mechanisms of sleep disruption that might trigger PPDs and potential interventions  
294 that target the circadian rhythm disruptions during the perinatal period<sup>76</sup>.

295

## 296 ***[H2] Reproductive Hormones***

297 One important hypothesis for the aetiology of PPDs is based on the temporal onset of these  
298 disorders immediately after childbirth, which is a time of major physiological change for women,  
299 including alterations in hormonal systems. Multiple lines of evidence have demonstrated that  
300 fluctuations in reproductive hormones (such as oestrogen and progesterone) during the perinatal  
301 period are substantial contributors to the development of postpartum mood disorders in vulnerable  
302 women. Gonadal steroid hormones (such as oestrogen and progesterone) are produced at very  
303 high levels during pregnancy, but rapidly decrease to pre-pregnancy levels after childbirth. One  
304 study simulated this pattern of hormone expression and demonstrated substantial mood symptoms  
305 (such as sadness, anhedonia and anxiety) during the withdrawal period in five of eight women with  
306 a history of postpartum depression, but in none of the eight women with no history of postpartum  
307 depression<sup>77</sup>. Thus, women who are vulnerable to postpartum psychiatric episodes might not have  
308 gross abnormalities in endocrine physiology (such as no differences in the absolute levels of  
309 hormones), but might have an abnormal response to the hormonal fluctuations of pregnancy and  
310 childbirth.

311 Reproductive hormones have important functions in the central nervous system. Oestrogen and  
312 progesterone receptors are expressed throughout the brain and can modulate neurotransmission  
313 and neuroplasticity via both genomic and non-genomic mechanisms. For example, rodent studies  
314 have shown that ovariectomy reduces and estradiol administration increases brain-derived  
315 neurotrophic factor (BDNF) levels in the hippocampus and the forebrain<sup>78</sup>; BDNF levels are  
316 decreased by stress and depressive symptoms and are increased following treatment with  
317 antidepressants<sup>79</sup>. The rapid fall in oestrogen levels in the postpartum period might, therefore,

318 reduce BDNF levels and increase susceptibility to PPDs in women who are vulnerable. Similarly,  
319 progesterone has an important role in regulating neurotransmitter synthesis, release and transport  
320 <sup>80</sup> and, has been shown to up-regulate BDNF expression in the hippocampus and cerebral cortex  
321 in rodent models<sup>81</sup>.

322 The neurosteroid, allopregnanolone, which is a major metabolite of progesterone, might also have  
323 an important role in the aetiology and, potentially, in the treatment of postpartum depression<sup>82, 83</sup>.  
324 Allopregnanolone is a positive allosteric modulator of synaptic and extrasynaptic GABA<sub>A</sub> receptors<sup>84</sup>,  
325 <sup>85</sup> and animal models have demonstrated that intravenous allopregnanolone administration  
326 significantly reduces anxiety and depressive symptoms<sup>86</sup>. Allopregnanolone concentrations reach  
327 peak physiological levels during the third trimester of pregnancy and rapidly decrease following  
328 childbirth<sup>87, 88</sup>. The failure of GABA<sub>A</sub> receptors to adapt to the rapid fluctuations in allopregnanolone  
329 levels at childbirth is hypothesized to be a trigger for postpartum depression<sup>89, 90</sup>. This line of inquiry  
330 is being further explored by the development of brexanolone (a proprietary formulation of  
331 allopregnanolone) as a treatment for postpartum depression<sup>91,92</sup>.

332 Oxytocin is a neuroactive hormone that supports childbirth, lactation, maternal behavior, and social  
333 bonding<sup>93</sup>. Some studies have demonstrated an inverse association between circulating oxytocin  
334 levels and postpartum depression<sup>94</sup> although other studies have not found this<sup>95</sup>. The alterations in  
335 the oxytocin system that occur during pregnancy and childbirth do not occur in isolation, and the  
336 role of oxytocin in postpartum depression is likely to be complex and not accounted for by absolute  
337 levels, similar to the roles of other neuroactive hormones <sup>83</sup>.

338 Further, recent neuroimaging data increases our understanding of the neurobiological basis  
339 underlying perinatal mood disorders and the development of maternal behavior. Indeed, one study  
340 demonstrated that the effects of a polymorphism in *BDNF* (*BDNF* Val<sup>66</sup>Met) on hippocampal  
341 function are selectively modulated by estradiol<sup>96</sup>. This work lends further data to the importance of  
342 the role of sex steroids on the regulation of behavioural functions associated with psychiatric

343 disorders, such as emotional processing, arousal, cognition, and motivation. Thus, it follows the  
344 involvement of sex steroids of brain function could be revealed using neuroimaging. Indeed,  
345 multiple cortical and subcortical brain regions have altered activity observed using functional MRI  
346 or PET (such as, measurement of brain MAO-A<sup>97</sup>) in mothers with depression, in response to infant  
347 and non-infant emotional cues<sup>98,99</sup>. These alterations might impact important neuronal networks  
348 that are associated with learned reward, reaction to stimuli, stress, motivation and executive  
349 functioning. In addition, recent research from functional MRI studies shows distinct neurobiological  
350 patterns that distinguish anxiety and depression occurring in the perinatal period, compared with  
351 other times of a woman's life, and these patterns may have significant impact on the mother-infant  
352 relationship<sup>100</sup>.

353

## 354 **[H2] Other factors**

355 **[H3] Stress axis.** The postpartum period is a time of great flux for the HPA stress axis<sup>101</sup>. Indeed,  
356 alterations in the hypothalamic–pituitary–adrenal (HPA) axis occur during pregnancy, such as  
357 corticotropin-releasing hormone (CRH) production by the placenta, resulting in significantly  
358 increased levels during pregnancy, which abruptly decline postpartum<sup>102</sup>, and rising levels of  
359 gonadal steroids that contribute to puerperal hypertrophy of the pituitary and adrenal glands,  
360 leading to increases in ACTH and cortisol levels<sup>103</sup>. CRH fluctuations during the perinatal period  
361 might trigger HPA axis dysregulation and contribute to the onset of depressive and anxiety  
362 symptoms in a subset of vulnerable women<sup>104</sup>; however, inconsistent findings have been  
363 reported<sup>105, 106</sup>.

364

365 **[H3] Thyroid hormones.** Thyroid hormones have been implicated in the development of  
366 postpartum mood disorders. Thyroid binding globulin (TBG, which transports thyroid hormones in  
367 the blood) concentrations increase during pregnancy and might be an index of sensitivity to

368 elevated oestrogen levels. Some data also suggest that decreased [Au:OK? YES] TBG levels are  
369 a predictor of perinatal depression<sup>107</sup>. In addition, first-onset postpartum autoimmune thyroid  
370 disorders often co-occur with postpartum mood disorders<sup>108</sup>. The occurrence of both disorders  
371 coincides with the postpartum rebound phenomena of the maternal immune system, suggesting an  
372 overlap in aetiology<sup>109</sup>. Supporting this hypothesis, women with increased thyroid peroxidase  
373 antibodies during pregnancy have an increased risk for postpartum psychiatric episodes<sup>109, 110</sup>.  
374 Accordingly, the assessment of thyroid function (such as measurement of thyroid-stimulating  
375 hormone levels, tri-iodothyronine (T3) and tetraiodothyronine (T4)) is an essential part of diagnostic  
376 evaluations in women with postpartum psychiatric episodes.

377

378 **[H3] Neuroimmune pathways.** Neuroimmune pathways might also have a role in the  
379 pathophysiology of postpartum mood disorders<sup>101, 111-113</sup>. The transition from pregnancy into the  
380 postpartum period is characterized by an accelerated immune response (mediated through both  
381 pro-inflammatory and anti-inflammatory mediators for healing and involution) during labor that  
382 continues into the early postpartum period<sup>114</sup>. Consequently, immune changes at the end of  
383 pregnancy might predict postpartum depression. IL-6 levels are increased in women with  
384 postpartum depression compared with postpartum women who do not have depression in some,  
385 but not all studies.<sup>112,115</sup> However, leptin (a protein hormone made by adipose cells that regulates  
386 energy and has inflammatory functions) might also be associated with postpartum depression.  
387 Indeed, decreased maternal serum leptin levels during delivery are associated with a higher risk for  
388 postpartum depression, and might potentially serve as a biomarker for postpartum depression<sup>115</sup>.  
389 Lower levels of Clara cell protein (CC16, an endogenous anti-inflammatory compound) are  
390 associated with PPD a few weeks later<sup>116, 117</sup>. Furthermore, decreased levels of  $\omega$ 3  
391 polyunsaturated fatty acids (PUFAs) at the end of the third trimester are suggested to associate  
392 with increased risk of PPD in the early postpartum period<sup>118</sup>. The underlying mechanism is  
393 hypothesized to be increased peripheral inflammation<sup>118</sup> owing to the anti-inflammatory effects of



394  $\omega$ 3 PUFAs. In summary, dysregulation of the crosstalk between the immune system and the HPA-  
395 stress axis is hypothesized to be associated with the onset of postpartum depression<sup>101, 119</sup>

396 Interestingly, first pregnancies are more often followed by psychiatric episodes than subsequent  
397 pregnancies, which may illustrate the dysregulation of psychoneuroimmune systems. This effect is  
398 hypothesized to be due to the biological differences between first and subsequent pregnancies,  
399 and has raised the possibility of an aetiological link with other medical conditions that have a  
400 similar increase in prevalence in first pregnancies such as pre-eclampsia<sup>120</sup>. Intriguingly, pre-  
401 eclampsia and postpartum psychosis are both associated with immune dysregulation, for example  
402 the increased rates of postpartum autoimmune thyroiditis<sup>108, 109</sup> and alterations in immune  
403 biomarkers (such as, CNS autoantibodies) in women with postpartum psychosis<sup>121</sup>. In addition,  
404 abnormalities in monocyte and T cell function and tryptophan breakdown has been demonstrated  
405 in patients with postpartum psychosis or mania, compared with postpartum women without any  
406 psychiatric symptoms.<sup>122</sup> Patients with postpartum psychosis had monocytosis and failed to  
407 demonstrate the physiological T- cell increase that is normally observed during the postpartum  
408 period. These findings support the notion that immune system dysregulation contributes to  
409 affective instability and severe postpartum episodes<sup>122</sup>.

410 Future studies are needed to extend our understanding of the ways in which psychological and  
411 biological processes interact in PPDs. For example, social support might exert a stress-buffering  
412 effect via the downregulation of stress responses, including inflammatory reactivity to stressors and  
413 dampened sympathetic and hypothalamic-pituitary-adrenal (HPA) axis activity<sup>123, 124</sup>.

### 414 **[H1] Diagnosis, Screening and Prevention**

415 As with all psychiatric disorders, the diagnosis of postpartum depression is reached by a  
416 comprehensive clinical interview and diagnostic criteria that provide an operationalised definition of  
417 the disorder, using classification systems such as the Diagnostic and Statistical Manual of Mental  
418 Disorders, Fifth Edition (DSM-5)<sup>125</sup> or International Classification of Diseases, Tenth Edition

419 (ICD10)<sup>126</sup>. Diagnostic criteria are similar across the DSM-5 and ICD-10, which are the two most  
420 common classification systems; however, the DSM-5 uses the term 'depression with peripartum  
421 onset' to refer to the onset of depression during pregnancy and into the first month postpartum,  
422 whereas ICD10 does not use a primary code referring to the perinatal period, although a second  
423 code denoting postpartum onset is available (which is not used in practice) . However, the  
424 diagnosis of psychiatric disorders is more than a list of symptoms and the impact on symptoms of  
425 functioning; diagnosis should also include an understanding of predisposing aetiological factors,  
426 triggers and maintenance factors, which are elicited by a comprehensive biopsychosocial  
427 assessment<sup>24</sup>.

428 Postpartum depression is one of the most common postpartum psychiatric disorders, and can be  
429 mild and relatively self-limiting lasting only a few weeks, or can be more severe, with severe  
430 episodes potentially including psychotic symptoms<sup>17</sup>. Some symptoms of depression such as  
431 fatigue, sleep disturbance and appetite disturbance need careful enquiry, as a woman with a baby  
432 will be more tired than usual and have disrupted sleep (due to the baby needing a feed), although  
433 appetite might not be affected as breastfeeding will stimulate appetite despite a low mood.

434 Checking whether the mother is able to sleep when the baby is asleep, whether the fatigue persists  
435 after rest and the interest in food will help establish whether the symptoms are pathological and if  
436 they are indicative of postpartum depression. Notably, anxiety might be a prominent symptom in  
437 postpartum depression<sup>63</sup>, or can be a symptom of a comorbid anxiety disorder<sup>127</sup>. Diagnostic  
438 assessment should evaluate a history of manic or hypomanic symptoms, as first-onset postpartum  
439 depression can indicate underlying bipolar disorder<sup>26</sup>. Diagnostic challenges include barriers to  
440 disclosing symptoms due to stigma<sup>128</sup> and variations in the manifestation of symptoms, which might  
441 reflect cultural or educational differences<sup>129</sup>. In addition, it is important to ensure symptoms are not  
442 due to an underlying medical condition such as thyroid disease or an early presentation of first  
443 episode psychosis.

444 Although postpartum psychosis is not included as a primary diagnosis in the DSM-5 or ICD-10, this  
445 disorder is still recognised clinically and is usually considered to be a severe mood disorder.<sup>16</sup>  
446 Women with postpartum psychosis often have a history of bipolar disorder.<sup>16</sup> Most women have  
447 prodromal symptoms before the overt onset of postpartum psychosis; however, some women have  
448 acute onset of severe symptoms<sup>130, 131</sup>. Evaluation of women with postpartum psychosis includes  
449 assessment of manic, depressed, anxious and psychotic symptoms, and the assessment of the  
450 risk of causing harm to herself or her baby. Women with postpartum psychosis can present with  
451 either low or high mood (both elation and irritability), or frequently can present with a mixed state,  
452 including symptoms of both mania and ; a minority of women have an atypical symptom profile with  
453 disorientation and/or disturbance of consciousness<sup>132</sup>. Symptoms can also manifest as delusions,  
454 hallucinations, and particularly confusion and perplexity and patients can also have severe mood  
455 swings, insomnia, agitation and rapid deterioration. Postpartum psychosis is usually a rapid onset  
456 severe psychosis, typically starting within the first 2-4 weeks after birth, and is considered a  
457 psychiatric emergency as a lack of self-care and an inability to care for the infant can lead to  
458 suicide and/or, in rare cases, infanticide<sup>16</sup>. Accordingly, assessment should be carried out quickly  
459 (for example, The National Institute for Health and Care Excellence (NICE) recommend  
460 assessment within 4 hours of clinical presentation; in clinical practice this means within 24 hours of  
461 the acute onset of severe psychiatric symptoms) to ensure the woman can be cared for safely and  
462 appropriately. Excluding other medical disorders, such as cerebral space occupying lesions,  
463 thyroid disorders or infections is important as part of the diagnostic work-up. In many cases the  
464 mother's partner or family ask for psychiatric evaluation when the mother is irritated or agitated and  
465 not aware that she is seriously ill.

466  
467 Anxiety disorders (such as GAD and panic disorder), OCD and PTSD can all manifest in the  
468 postpartum period. OCD is characterised obsessive thoughts. Obsessions are intrusive, repetitive  
469 thoughts, images or impulses that are unacceptable and/or unwanted and give rise to subjective

470 resistance. By contrast, delusions that occur in psychotic disorders are fixed, false beliefs<sup>133</sup> that  
471 need appropriate psychological intervention<sup>17</sup>. Postpartum OCD poses a particular diagnostic  
472 challenge, as intrusive thoughts about harm befalling the infant (such as, what if I drop my infant or  
473 I accidentally cut the infant with a knife when I'm cooking) might be perceived as delusional and  
474 could lead to concerns about the safety of the infant. However, these thoughts are not associated  
475 with actual direct harm and the obsessions remain very ego-dystonic and highly distressing to the  
476 patient. Traumatic childbirth experiences can trigger PTSD, particularly in women with prior  
477 histories of trauma<sup>134</sup>. Differentiating the exacerbation of PTSD symptoms in women with past  
478 trauma and new onset PTSD owing to traumatic childbirth is important<sup>135</sup>. Past trauma history  
479 should include assessment of prior childhood abuse, adult interpersonal or other violence, among  
480 other forms of trauma. In addition, many women with PTSD or OCD present with symptoms of  
481 anxiety and mood symptoms, making the diagnosis of any one particular disorder a challenge<sup>63, 136</sup>.

482

483 Women with a previous history of psychiatric disorders will often experience a worsening of  
484 symptoms during the postpartum period, although few studies have examined strategies to  
485 mitigate this exacerbation of symptoms<sup>39</sup>. For women without a prior history of psychiatric  
486 disorders, the acute onset of psychiatric symptoms in the postpartum period is often highly  
487 distressing. However, whether the first onset of psychiatric symptoms in the postpartum period  
488 indicates the beginning of a more persistent and chronic mood disorder, or is a condition that will  
489 be restricted only to the postpartum period is unclear. This is an important area for future study.

490

## 491 **[H2] Screening**

492 Screening for postpartum depression has attracted widespread interest from researchers, clinicians  
493 and policy makers due to the high prevalence and associated sequelae in terms of maternal  
494 morbidity and adverse child outcomes. In many countries, screening for postpartum depression

495 during routine obstetrical care, including care by health visitors, is inconsistent, and this strategy  
496 has become an area of focus in many countries. In addition, up to 60% of perinatal women with  
497 depression do not seek help<sup>137</sup>. However, given the availability of screening instruments and  
498 effective treatments<sup>138</sup>, Clinical Practice Guidelines and recommendations are increasingly  
499 supportive of routine screening<sup>139, 140</sup>. More generally, international guidelines reflect a consensus  
500 that improved identification of PPDs is vitally important<sup>141,142</sup>. Accordingly, several national  
501 campaign to increase awareness of PPDs are underway, such as the Maternal Mental Health  
502 Alliance<sup>143</sup> in the UK. This alliance is a coalition of organizations that are dedicated to achieving  
503 consistent, accessible and quality mental health care in the first year after giving birth. In addition,  
504 state mandates for perinatal depression screening are increasing in the United States, including  
505 the US Preventive Services Task Force recommendation<sup>140</sup>. However, although this task force has  
506 concluded that the evidence base that is sufficient to recommend screening for perinatal  
507 depression when combined with adequate support systems<sup>25, 140</sup> this conclusion has also been  
508 criticised by others<sup>144</sup>.

509 The most widely researched and used screening tool for postpartum depression is the brief 10-item  
510 Edinburgh Postnatal Depression Scale (EPDS<sup>145</sup>), which was designed to exclude symptoms that  
511 can be normal features of the perinatal period, such as poor sleep, but that are often included in  
512 other self-report measures. A cut-off score of 13 is most commonly used to recommend further  
513 diagnostic assessment<sup>146, 147</sup>. In addition, the EPDS includes question about thoughts of self-harm,  
514 which can help to mobilise risk assessment and can predict suicidal intent<sup>148</sup>. The EPDS has been  
515 used prenatally and validated in a number of languages, its properties are relatively well-  
516 understood<sup>147</sup> and it appears to be highly acceptable in the target population<sup>149,150</sup>. High EPDS  
517 scores can reflect several psychiatric diagnoses. For example, among the 826 screen-positive  
518 women out of a sample of 10,000 women, the most common primary diagnosis was unipolar  
519 depressive disorder (found in 68.5% of women), but almost two-thirds of women had co-morbid  
520 anxiety disorder and 22.6% had a bipolar disorder<sup>151</sup>. These data highlight another potential

521 benefit of the EPDS: that most women with a false-negative result for unipolar depression have  
522 another diagnosable, and potentially treatable, psychiatric condition.

523 Other generic or perinatal-specific depression measures have been used to identify perinatal  
524 depression, but are not as well validated in perinatal women as the EPDS. Other measures include  
525 the Postpartum Depression Screening Scale<sup>152</sup>, the Beck Depression Inventory-II<sup>153</sup> and the  
526 Patient Health Questionnaire-9<sup>154,155</sup>. Alternatively, two case finding questions (the so-called  
527 Whooley questions <sup>141, 156, 157</sup>) can be asked to women to determine whether further mental health  
528 assessment should be carried out, and the use of these questions is recommended by NICE  
529 guidelines in the United Kingdom. The Whooley questions can also be used to detect any  
530 psychiatric disorder, and are not limited to depression<sup>2</sup>.

531 As previously mentioned, postpartum depression are frequently co-morbid with anxiety (in 4.3% of  
532 women). As anxiety substantially impacts maternal functioning and fetal and infant development<sup>28,</sup>  
533 <sup>158 159</sup>, this has spurred efforts to screen for postpartum anxiety. Three sub-items of the EPDS (the  
534 so-called EPDS-3A) can be used to identify perinatal anxiety disorders and sub-syndromal  
535 anxiety<sup>160</sup>. Other screening instruments for anxiety disorders include the Perinatal Anxiety  
536 Screening Scale (PASS)<sup>161</sup> and the generalized anxiety disorder scale (GAD-7)<sup>162</sup>. Screening tools  
537 for perinatal OCD and PTSD are also available, such as the specific perinatal OCD screening scale  
538 (The POCS),<sup>163</sup> and a short screening scale for PTSD (SPAN), respectively.<sup>164</sup>.

539 The utility of routine screening for postpartum psychosis, hypomanic and manic symptoms and  
540 bipolar disorder faces several barriers including a lack of evidence base of effectiveness and the  
541 reduced predictive value of screening for a relatively rare condition. Despite steady progress in this  
542 area<sup>165, 166</sup> a consensus test with well-known precision and an agreed cut-off has not been  
543 identified<sup>167,168</sup>. However, the Mood Disorder Questionnaire (MDQ) has shown solid psychometric  
544 properties for assessing bipolar disorder and is increasingly used <sup>169</sup>. Taking a full personal and  
545 family history might help to identify vulnerability to bipolar disorders which could trigger further

546 diagnostic assessment, given the strong association between bipolar disorder and increased risk  
547 for PPDs<sup>141, 142</sup>.

548 In general, screening programs in the postpartum period should include a clear pathway from  
549 screening, to diagnostic assessment and treatment<sup>170</sup>. Best practice guidelines agree that all  
550 women who have a positive screen need subsequent assessment, during which, co-morbidities  
551 and the woman's wider psychosocial context can be explored. Currently, only such well-resourced,  
552 integrated management programs have provided evidence that perinatal mental health is improved  
553 by depression screening<sup>171, 172</sup>. In this regard, e-screening and e-treatments to facilitate integrated,  
554 cost-effective care might be useful<sup>173</sup>. Few well-understood, validated screening approaches for  
555 PPDs that can ultimately improve morbidity and mortality are available. Indeed, further building of  
556 the evidence-base for screening, including the cost-effectiveness of perinatal depression screening  
557 as a policy direction is required<sup>142, 174</sup>.

558

## 559 **[H2] Prevention**

560 Interventions for the prevention of postpartum depression or postpartum anxiety are intended to  
561 prevent the onset, duration, or recurrence of these disorders. Prevention can reduce the mental  
562 health, physical health and socio-economic burden associated with postpartum depression for  
563 mothers, their offspring and families, as well as for health systems. The effectiveness of prevention  
564 of postpartum depression is facilitated by the fact that pregnant women are motivated to address  
565 factors that will affect their baby<sup>175</sup>. The assessment of risk factors for PPDs helps with diagnosis  
566 and formulation, but is also important for identifying potentially modifiable targets for prevention  
567 and treatment (**Box 3**)<sup>176</sup>. Thus, it is a requirement for both symptom screening and risk  
568 assessment that systems exist for adequate follow-up and support. Furthermore, women and  
569 clinicians should be informed that the established risk factors might have limited predictive value

570 for individual patients and, therefore, do not guarantee which women will develop or not develop  
571 postpartum depression.

572 Some psychosocial and psychological interventions have reduced the risk of women developing  
573 postpartum depression, although no single intervention type or modality appears superior to  
574 others. Data from trials included in a Cochrane review<sup>176</sup> as well as randomized controlled trials  
575 included in a qualitative review,<sup>175</sup> point towards particularly positive impacts when interventions  
576 target at-risk groups (such as women with a previous episode of depression or a recent life  
577 stressor), or include interpersonal therapy (IPT). As relationship challenges and lack of social  
578 support constitute strong risk factor for PPD, the interpersonal focus of the IPT intervention,  
579 therefore aims to address this causative or aggravating factor. Interventions with the most promise  
580 include interventions targeting at-risk groups (such as women with a previous episode of  
581 depression or a recent life stressor).

582 Trials included in these reviews were conducted among high risk women, based on various factors,  
583 as well as women enrolled from the general perinatal population. Trials assessing the use of  
584 interpersonal therapy, cognitive behavioural therapy, peer support, parental preparedness, and  
585 person-centred approaches for prevention of postpartum depression have demonstrated  
586 significantly positive results, whereas trials assessing the use of cognitive behavioural therapy for  
587 postpartum depression have demonstrated mixed results. These results, disaggregated for  
588 universal, selective or indicated prevention strategies are summarized in a more recent systematic  
589 review and meta-analysis<sup>177</sup>. The interventions were delivered using several modalities, including  
590 home visits and telephone support, provision by professional and lay practitioners, individual and  
591 group-based sessions, through multiple contact sessions and at postpartum initiation<sup>176</sup>.

592 There is conflicting evidence for the treatment of vulnerable women with antidepressants for the  
593 prevention of depressive episodes or anxiety symptoms during the perinatal period as well as  
594 anxiety symptoms has conflicting evidence<sup>178</sup>. One of the earliest studies demonstrated a



595 reduction in recurrence of postpartum major depression with prophylactic antidepressant treatment  
596 <sup>179</sup>. Small but emerging literature has suggested hormonal therapies, light therapy and other forms  
597 of circadian manipulation might be promising therapies for prevention of depression<sup>180</sup>. There is no  
598 strong evidence for the use of hormonal therapies, acupuncture, supplementation with omega-3  
599 polyunsaturated fatty acids, light therapy and other forms of circadian manipulation for prevention  
600 of postpartum depression. <sup>177,180</sup>

601 Interventions for the prevention of postpartum psychosis include careful monitoring for symptom  
602 development in women at high risk and adjustments of prophylactic medication, especially in  
603 women with bipolar disorder<sup>18, 24</sup>. Prophylactic treatment during pregnancy might reduce the rate of  
604 postpartum relapse in women with bipolar disorder, although no evidence from randomized  
605 controlled trials for this is available. For women with previous postpartum psychosis, prophylactic  
606 treatment with lithium or antipsychotics immediately postpartum might reduce relapse<sup>18</sup>.

607

## 608 **[H1] Management**

609 The goals of treating mental illness in the postpartum period are to reduce maternal psychiatric  
610 symptoms and to support maternal-child and family functioning. All women and their families  
611 should receive education about the illness and the potential treatment options, including the  
612 potential benefits and harms of each treatment option. Social support should be optimized and  
613 physical and psychiatric comorbidities should be addressed. In addition, strategies to assist women  
614 in obtaining sleep and a stable circadian rhythm are helpful, given that sleep deprivation is  
615 common during the postpartum period. In many cases, the symptoms of PPDs influence maternal-  
616 child interactions, which should be observed and discussed in a non-judgmental way.

617 Although specific recommended treatments depend on the underlying diagnosis, in general, a  
618 stepped care approach is advocated, in which the intensity of the intervention matches the severity  
619 and acuity of the clinical presentation. For example, women with mild symptoms of depressive,

620 anxiety, obsessive–compulsive and/or trauma or stressor-related disorders should first be offered  
621 the lowest-intensity interventions such as peer support and guided self-help, whereas women who  
622 do not respond to these treatments might require formal psychotherapeutic interventions, such as  
623 psychological therapies. For women with severe symptoms, who do not respond to non-  
624 pharmacological treatment, or who have bipolar disorder or psychosis, pharmacological  
625 interventions are likely to be introduced as a first-line treatment, used alone or in combination with  
626 a lower-intensity intervention. In such cases, the well-established benefits of breastfeeding on the  
627 infant must be considered in the context of maternal mental wellbeing, the passage of psychotropic  
628 medication into breast-milk and the infant, and the potential effects of medications on the neonate.  
629 Indeed, when breast-feeding is challenging, and/or when frequent nighttime feedings leads to  
630 sleep disruption, symptoms of depression or anxiety might be precipitated or exacerbated. In these  
631 cases, the benefits of breastfeeding must be weighed against the risk of ongoing maternal mental  
632 illness, and formula feeding is a viable and often recommended alternative. Other somatic  
633 treatments, such as electroconvulsive therapy (ECT), can be considered in women with treatment-  
634 refractory disorders. Throughout, monitoring progress to determine when or if to move to a higher-  
635 intensity intervention, and to ensure safety for mother and child is important. In the initial  
636 assessment and during treatment the patient and her family should be asked if thoughts of suicide  
637 or infanticide have occurred. Safety concerns and/or evidence of active psychosis are medical  
638 emergencies that require specialist consultation, emergency hospitalization and treatment.

639

## 640 **[H2] Mood disorders and anxiety**

641 Treatment of postpartum depression and other non-psychotic mental disorders (such as anxiety,  
642 OCD and trauma and stressor-related disorders; **Box 4**) depends on the severity of the initial  
643 presentation and the level of functional impairment, including the effect on the maternal–child  
644 interaction.<sup>128</sup> For women with a past history of mental illness, the previous treatment response and  
645 the time to response of previous episodes should be considered. The patient’s treatment

646 preference, in addition to as access to care and utilization of care should also be considered in all  
647 women, as patients who receive their preferred treatment are most likely to benefit from this  
648 treatment than other treatments.<sup>181</sup> Most women with non-psychotic mental disorders often prefer  
649 psychotherapy over pharmacological treatments, although the uptake and effectiveness of this  
650 therapy can be limited due to barriers in attending appointments, such as unpredictable infant  
651 schedules and competing childcare responsibilities.<sup>182</sup> Similarly, fathers also prefer psychological  
652 treatments to pharmacological therapy.<sup>183</sup> However, some women prefer pharmacological  
653 treatment alone, so individualizing treatments based on patient preferences is important. Treating  
654 maternal postpartum depression might not always improve the maternal–infant relationship, and  
655 additional interventions aimed at the mother–infant dyad or the family as a whole might be  
656 required.<sup>184</sup>

657

658 **[H3] Psychological Interventions.** Most trials for postpartum depression have focused on non-  
659 pharmacological treatments. For women with mild postpartum depression, psychosocial treatments  
660 including peer support, guided self-help, and supportive counseling by trained professions such as  
661 public health nurses (at home, or in support groups) can improve symptoms. For example, one  
662 systematic review of 5 trials demonstrated a 1 year remission rate of 68% in women with  
663 postpartum depression who received psychosocial treatments compared with a remission rate of  
664 54% in women treated with standard primary care.<sup>185</sup> For women with moderate symptoms of  
665 depression, or women who do not responded to psychosocial strategies, psychotherapies such as  
666 cognitive-behaviour therapy (CBT) and interpersonal therapy (IPT) that specifically address the  
667 psychological and related challenges of transitioning to parenthood are effective when delivered in  
668 individual, group, and partner-assisted formats, and either in-person, by telephone, or online.<sup>186</sup> A  
669 systematic review of 4 CBT and 1 IPT trials demonstrated a pooled remission rate of 60.3% for  
670 these interventions, compared with a rate of 48.1% for usual care.<sup>185</sup>

671

672 In addition, a CBT-based program was demonstrated to reduce worry and depressive symptoms in  
673 women with postpartum anxiety disorders, including GAD, social phobia and OCD, compared with  
674 symptoms at baseline.<sup>187</sup> The effectiveness of CBT for postnatal OCD symptoms was confirmed in  
675 a small RCT.<sup>188</sup> Although additional research is required, CBT-based interventions for postpartum  
676 anxiety disorders, and specifically interventions such as eye movement desensitization and  
677 reprocessing (EMDR) and trauma-focused CBT for trauma and stressor related disorders, can be  
678 used, although the latter two interventions have not been specifically evaluated in postpartum  
679 women [<sup>189</sup>.

680 The increasing use of internet-based CBT and the development of mobile apps that use this  
681 treatment modality demonstrates the power of digital health, which is often more accessible than  
682 traditional psychotherapy, and extends to individuals who can't participate in psychotherapy. A  
683 good example of this is MumMoodBooster, which was developed in Australia<sup>173</sup>.

684

685 **[H3] Drug therapies.** Antidepressants, such as selective serotonin reuptake inhibitors (SSRIs) and  
686 serotonin norepinephrine reuptake inhibitors (SNRIs) are the mainstay of pharmacological  
687 treatment for postpartum anxiety and depressive disorders. These therapies can be used alone or  
688 in combination with psychosocial or psychological treatments. In a systematic review, the pooled  
689 remission rate was significantly higher in patients receiving SSRIs (46.0%) compared with those  
690 receiving placebo (25.7%).<sup>190</sup> Although pharmacological therapies can be used alone or in  
691 combination with psychosocial or psychological treatments, whether combinatorial therapy is  
692 superior to either one alone has not been evaluated in postpartum women. However, combinatorial  
693 treatment does not lead to further improvements in functional outcomes compared with medication  
694 alone, in non-perinatal populations.<sup>191, 192</sup> To our knowledge, there are no drug treatment trials in  
695 maternal anxiety disorders, or in paternal postpartum depression or anxiety. However, SSRI and  
696 SNRI medications are first line pharmacological treatments for anxiety and depression outside the  
697 postpartum period. The duration of antidepressant therapy required for new-onset postpartum

698 depression, anxiety or a related disorder has not been studied, but clinicians are recommended to  
699 follow guidelines for these disorders in the general population. For depression, the initial treatment  
700 should be continued for 6 months to 1 year after remission; longer durations are required for  
701 severe and/or recurrent illness.<sup>193</sup>

702

703 Many women and their partners are concerned about the safety of using antidepressant drugs  
704 during breastfeeding. However, the use of antidepressants during the postpartum period is not a  
705 contraindication to breastfeeding, and indeed, the avoidance of medication when needed for  
706 severe illness is associated with maternal suicides<sup>194</sup>. The passage of SSRIs and SNRIs into  
707 breast-milk is variable between drugs, but most pass into the breast-milk at <10% of the maternal  
708 dose, which is compatible with breast-feeding.<sup>195</sup> As such, changing the antidepressant drug or the  
709 dose during the perinatal period to switch to a drug with lower breast-milk passage is generally not  
710 recommended. For new-onset postpartum depression, sertraline is often recommended as a first-  
711 line pharmacological treatment due to very minimal passage into breast-milk (Figure 2). However,  
712 in patients with a prior history of psychiatric disorders, therapies that have previously demonstrated  
713 efficacy should be considered, even those that have less data regarding safety during  
714 breastfeeding. Other SSRIs, SNRIs or mirtazapine (an atypical antidepressant) also have minimal  
715 passage into breast-milk, so these drugs are unlikely to be a cause for concern. Bupropion is  
716 generally not given to lactating women due to case reports of infant seizures associated with  
717 exposure to this drug.<sup>196</sup> In cases of severe depression and/or anxiety (with or without psychotic  
718 features), older antidepressants, other therapies such as benzodiazepines or antipsychotics might  
719 be indicated.

720

721 **[H3] Other treatments.** Given the likely role of hormonal fluctuations in the aetiology of postpartum  
722 depression, hormonal treatments have been evaluated. Transdermal oestrogen therapy reduced  
723 the symptoms of postpartum depression in one small study, but further trial are required. Progestin

724 therapy worsens postpartum depressive symptoms.<sup>197</sup> One trial of demonstrated the superiority of  
725 allopregnenalone to placebo in improving depressive symptoms in 21 women with severe  
726 postpartum depression,<sup>92</sup> although this requires further investigation. Complementary and  
727 alternative medicine treatments (for example, folate, s-adenosylmethionine, massage and  
728 acupuncture) are not well-supported by evidence.<sup>198</sup> However, using aerobic exercise for  
729 postpartum depression was recently examined in a systematic review and has good supporting  
730 evidence for mild symptomatology <sup>199</sup>. ECT can be considered in severe or treatment-refractory  
731 cases of depression.

732

## 733 **[H2] Postpartum psychosis**

734 Fluctuations in symptoms are common in women with postpartum psychosis, and thoughts of  
735 infanticide or suicide are often well hidden. Thus, outpatient treatment is not safe and psychiatric  
736 hospitalization is recommended for diagnostic evaluation and treatment <sup>15</sup>. The preferred treatment  
737 setting is a mother–baby joint admission unit, but these units are not available worldwide <sup>200, 201</sup>.  
738 Alternatives are the admission of the mother only to a hospital with expertise in perinatal  
739 psychiatric care or women’s mental health, or, if these facilities are unavailable, admission to a  
740 standard mental health inpatient setting, or based on a careful assessment of the safety of both the  
741 mother and the infant, intensive home treatment where available and with appropriate  
742 supervision<sup>16</sup>. The effect of these approaches on long term outcomes of the mother and baby are  
743 being investigated<sup>15</sup>.

744 The management of postpartum psychosis is dependent on psychiatric history. For women with  
745 known severe psychiatric illness with non-perinatal episodes, reviewing the nature and  
746 effectiveness of past treatments and restarting previous effective treatment is important.  
747 Management of women without a history of bipolar disorder or other severe psychiatric disorder are  
748 summarized in Figure 2. The main treatment goals for women without a prior history of bipolar

749 disorder, psychosis or other severe psychiatric disorder include the limitation of the current episode  
750 and the prevention of a bipolar disease course with multiple episodes. Accordingly, management in  
751 the first year postpartum should focus on full recovery (that is, complete symptom remission and  
752 social and vocational functioning). In the absence of guidelines and controlled trials, treatment  
753 recommendations are based on results from naturalistic cohort studies and expert consensus  
754 groups<sup>24, 202</sup>. The largest study (consisting of 68 patients) demonstrated the efficacy of a stepwise  
755 sequence of short-term benzodiazepines, antipsychotics and lithium, and showed high remission  
756 rates (remission in 98.4% of women) in the acute phase<sup>203</sup>. Moreover, this study demonstrated  
757 that lithium monotherapy is protective against relapse of psychosis, depression and mania within  
758 one year. The second largest study described successful ECT treatment in 34 patients with  
759 postpartum psychosis of whom many had symptoms of catatonia<sup>204</sup>. The effectiveness of lithium  
760 and ECT is supported by case reports<sup>205</sup>. Successful treatment with antipsychotics has been  
761 described in case reports<sup>206, 207</sup>, but antipsychotic monotherapy did not show efficacy in a cohort  
762 study<sup>203</sup>. Together, lithium monotherapy might be the preferred initial intervention for postpartum  
763 psychosis but adjunctive treatment with benzodiazepines or antipsychotics is useful for the acute  
764 treatment of agitation, mania and psychotic symptoms, given the well documented effectiveness in  
765 non-perinatal populations. Several antipsychotics are used for the treatment of severe PPDs  
766 including risperidone, quetiapine and olanzapine<sup>208</sup>. ECT is the primary treatment for patients with  
767 severe catatonic or depressed psychotic features, or if patient's prefer this therapy<sup>15</sup>.

768 Anticonvulsants (that is, antiepileptic medications) are also used less frequently as mood  
769 stabilizers in the treatment of bipolar disorder because of concerns of teratogenicity. However,  
770 valproate use during pregnancy and lactation is associated with neural tube defects and  
771 neurocognitive developmental delays in the offspring<sup>24, 209</sup>. Thus, valproate use is not advised  
772 during the perinatal period, unless the risk/benefit assessment determines a prior efficacy in  
773 particular women. By contrast, lamotrigine is used for the treatment of bipolar–depression (not  
774 bipolar–mania) and is not associated with an increased risk of congenital malformations in

775 offspring.<sup>210</sup> A review of recent studies demonstrated that lamotrigine had no adverse outcomes  
776 on infant IQ or neurodevelopment.<sup>211</sup>

777 The management of a breastfeeding woman with a severe psychiatric episode is challenging  
778 due to concerns about the exposure of breastmilk to pharmacological therapies and the need  
779 for sleep preservation in the mother<sup>15</sup>. The use of lactation inhibitors should be avoided. In some  
780 countries, the mother is recommended to breastfeed only if extensive psychiatric support and  
781 access to a pediatric professional that can monitor the infant are available. Moreover, the mother  
782 and her partner should be educated about the risks of breastfeeding with pharmacotherapy. In  
783 other countries, a more individualized approach (for example, NICE guidelines) based on previous  
784 responses to medication, preferences regarding breastfeeding and psychopathology<sup>24</sup>, other than  
785 avoidance of breastfeeding if lithium (rather than an antipsychotic) is used. Small case series have  
786 provided information regarding the safety of lithium in lactation<sup>212, 213</sup>. When possible, level of  
787 lithium in the infants serum should be closely monitored; on average, the serum level of lithium is  
788 25% of the maternal levels, but the range can vary and dehydration can lead to toxic levels<sup>15,213</sup>.  
789 No adverse effects were reported in ten infants of breastfeeding mothers who received ECT<sup>204</sup>.

790

## 791 **[H1] Quality of life**

792 The symptoms and morbidity of postpartum depression are often reported in the academic  
793 literature, but this offers only a rather constricted view of the quality of life (QOL) of women with  
794 PPDs.<sup>125,214 215</sup> Nevertheless, the classic core symptoms of a depressive disorder would be  
795 expected to decrease the subjective quality of both an individual's inner life experience (anhedonia,  
796 sadness, hopelessness, thoughts of death), and their functioning (psychomotor retardation or  
797 agitation, disturbed sleep). Anxiety is frequently co-morbid and this further influences quality of life  
798 with persistent worry symptoms.



799 Definitions vary, but QoL is a broader multidimensional construct which commonly incorporates  
800 two central aspects: emotional well-being (including, frequency and intensity of joy, sadness,  
801 affection) and life evaluation (how satisfied one is with one's life, for example, housing,  
802 employment). Health, and the ability to function, as an essential component of QoL is referred to as  
803 health-related quality of life (HRQoL). Maternal QoL during the postpartum period also affects her  
804 infant's current and future quality of life. Many mothers with postpartum depression have difficulty  
805 interacting with their infants in a positive way<sup>216</sup>, such as making less eye contact, showing less  
806 synchronous responsiveness, being uninvolved and showing restricted affect during mother-infant  
807 interactions.<sup>217-219</sup> Infant attachment security is a key predictor of child outcomes, including  
808 neurological, psychological and social outcomes over the course of development<sup>220</sup>. In addition,  
809 children of mothers with perinatal depression might have poorer psychological outcomes when  
810 they reach 18 years of age<sup>221</sup>. Some women have an intrusive engagement style that might lead to  
811 long-term difficulties in child social, cognitive and behavioural domains<sup>222</sup>. Women with postpartum  
812 psychosis face even more many parenting challenges often including disrupted attachment, which  
813 impact on the quality of life for mother and child.<sup>223</sup>

814 One of the most widely used generic measures of QOL is the 36 item short-form (SF-36)<sup>224, 225</sup>,  
815 which has eight health domains that measure limitations in physical or usual role activities due to  
816 health issues, limitations in usual role or social activities due to emotional issues, pain, mental  
817 health, vitality, and general health perceptions. The SF-36 has been used in over 1,000  
818 publications and for over 130 disorders including those that occur during the postpartum period,  
819 and both the full and short versions of this scale have been validated by numerous studies<sup>225, 226</sup>.  
820 However, few measures of QOL have been developed specifically for use in the postpartum  
821 period<sup>227</sup>. Only <sup>228</sup> three instruments for use during the postpartum period were reported in one  
822 systematic review: The Mother-Generated Index (MGI)<sup>229</sup>, the Maternal Postpartum Quality of Life  
823 Questionnaire (MAPP-QOL)<sup>230</sup> and the Rural Postpartum QOL scale (RPQoL)<sup>231</sup>. The MGI requires  
824 women to specify domains of their life that have been affected by the birth of their baby, either

825 positive or negative, and to then score these out of 10.<sup>232</sup> The most common changes reported  
826 were tiredness, less personal time, less time with partner or other family members, a worse  
827 relationship with partner or other family members, physical complaints, low self-esteem, financial  
828 worries, negative feelings towards the baby, more housework, poor sex life, decreased pleasure  
829 from baby, less sense of happiness or fulfilment<sup>229</sup>.

830 Not surprisingly, people with depressive illness in general report lower scores on generic QoL and  
831 HRQoL measures, as do women experiencing postpartum depressed mood.<sup>233 234</sup> Small studies in  
832 postpartum women also suggest QoL is amenable to intervention. Improvements in QOL are not  
833 fully explained by improvements in the severity of depressive symptom suggesting that  
834 interventions should go beyond the mere reduction of symptom severity and consider other factors  
835 that contribute to QoL as targets for intervention. . Maternal-specific measures of QOL could be  
836 integrated into postpartum depression screening programs or routine postnatal care<sup>235, 236</sup>. Indeed,  
837 QoL measures that allow a women to identify which areas of her life are most important to her,  
838 could be used to allow women to indicate where she would like to see improvements.<sup>229</sup> Emerging  
839 studies have highlighted the beneficial effect of social support on QOL<sup>237</sup>, in addition to risk factors  
840 for reduced QOL such as younger age and lower socio-economic status in women with postpartum  
841 depression<sup>238</sup>.

842 PPDs have economic considerations and can affect quality-adjusted life years (QALY)<sup>142</sup>. QALY  
843 takes into account how treatment affects quality and quantity of life, and accordingly, QoL  
844 measurement is necessary for studies of cost-effectiveness of treatments. Compelling data from  
845 the London School of Economics<sup>239</sup> demonstrated the high economic costs of PPDs and the need  
846 to address the loss of QALYs of women and their children, by treatment and prevention of these  
847 disorders. This finding is particularly pertinent given the short and long-term effects of postpartum  
848 depression. Indeed, as most women recover from postpartum mood disorders, these disorders can  
849 become chronic in a subgroup of women<sup>240</sup>. One study demonstrated that, for each one-year

850 cohort of births, perinatal depression, anxiety and psychosis cost the UK around £8.1 billion in the  
851 long-term.<sup>239</sup>

852

## 853 **[H1] Outlook**

854 The postpartum period is a vulnerable time for onset of psychiatric illness. Indeed, postpartum mood  
855 and anxiety are heterogeneous and might be triggered by biopsychosocial factors including a  
856 vulnerability to the robust endocrine and immune-related changes that occur at childbirth. The  
857 heterogeneity of these disorders requires a thoughtful approach to assessment and treatment  
858 planning that includes the clinical presentation, family and personal psychiatric history, other  
859 psychosocial risk factors (including history of trauma), and awareness of potential biological or  
860 genetic contributions that might influence risk and vulnerability.

861 The precise vulnerability that leads to some women developing PPDs is currently unknown and novel  
862 research approaches are needed to identify the underlying pathophysiology of both prepartum and  
863 postpartum anxiety, depression and psychosis. This will require a multi-faceted approach in  
864 preclinical, clinical and translational research, to determine the mechanisms behind the neurobiology  
865 and physiological correlates of PPDs, and the observed peripartum mood and mothering behaviors.  
866 Additionally, these strategies must address the differences in the timing of symptom onset and the  
867 diverse types of symptoms.

868 Given the morbidity and mortality of postpartum psychosis, episodes of psychosis might be best  
869 considered to represent women with a bipolar disorder diathesis with a puerperal trigger.

870 Understanding this trigger will be beneficial and should allow the development of new treatments  
871 and, ultimately, enable the prevention of psychosis or prevent unfavourable outcomes in women at  
872 high risk. Effective evidence based treatment approaches are available for psychosis and  
873 depression, including psychopharmacology, psychotherapy and ECT and circadian manipulation.

874 However, postpartum depression and postpartum psychosis require different and targeted

875 treatment approaches and therefore, bipolarity must be considered in the evaluation and  
876 management of all women with postpartum mood and anxiety disorders. In addition, primary  
877 treatment goals should include the limitation of the current episode and the prevention of future  
878 episodes (including unipolar or bipolar disease with multiple episodes, and chronic anxiety).  
879 Whether there a continuum of severity between postpartum depression and postpartum psychosis,  
880 or whether these disorders represent different conditions with different aetiological factors requires  
881 further study.

882 A potential barrier to the engagement and retention of women in the treatment of postpartum mood  
883 disorders is stigma. Understand this stigma and the fear that women have regarding postpartum  
884 mood disorders is essential. The voices of women with postpartum mood disorders must be  
885 incorporated into the development of services to ensure the needs of women, their infants and  
886 families are met<sup>241, 242</sup>.

887 To date, the amount of research provides an important road map for PPDs in general, and  
888 guidelines for screening or identification and treatment for perinatal depression in many countries  
889 gives a strong mandate to improve mental health care for all women in the perinatal period. Thus,  
890 developing effective strategies in low, middle and high income countries that allow the delivery of  
891 targeted therapies to women with different clinical phenotypes and severity of PPDs is imperative.  
892 In addition, whether the current ICD-10 and DSM-5 classification systems are adequate for  
893 detecting specific phenotypes or diagnostic groups of patients should be evaluated. We should  
894 also consider that PPDs might be phenotypically different than psychiatric disorders that begin  
895 during pregnancy. Indeed, disorders that occur postpartum might have unique characteristics in  
896 epidemiology, pathophysiology, psychosocial contributions, prevention and management than  
897 disorders that occur during pregnancy.

898 In summary, PPDs are morbid and costly disorders. Advocating for early identification and  
899 screening that begins in pregnancy to identify women at risk, in addition to timely and effective

900 treatments of PPDs is essential. Given the recent advances in knowledge, this an incredibly  
901 exciting time for research in perinatal mood disorders. New approaches might allow the  
902 identification of the underlying causes of postpartum mood disorders, which could lead efforts to  
903 identify women at risk and personalize treatment. Although genetic, biological and hormonal  
904 signals likely have an important role in risk of these disorders, psychosocial contributions including  
905 the current impact of lifetime stressors must be part of comprehensive work-up and treatment plan.  
906 The social determinants of postpartum mood disorders, such as poverty, domestic violence, poor  
907 housing, and insecure migrant status, should also be assessed as part of routine practice of  
908 maternal health care for all women. Finally, we must recognize that maternal mental health is  
909 necessary for the physical and mental health of mothers, infants and families<sup>17</sup> and advocating and  
910 protecting this population is our obligation.

911

912

913 **Display items**

914

915 **Box 1. Paternal postpartum depression.**

916 Fathers can also experience depression after the birth of a child. Indeed, in men, the prevalence of  
917 depression after the birth of a child is greater than at other times during life<sup>243</sup>. Although the  
918 literature of paternal depression is much smaller than that for maternal depression, the available  
919 literature demonstrates that paternal depression increases the risk for long-term adverse outcomes  
920 in the child due to potential impairments in parenting<sup>244, 245</sup>. In addition, a strong link between  
921 maternal depression and paternal depression has been reported. Pregnant women who had  
922 partners with depression during their pregnancy had worse depression symptom severity during  
923 the first six months postpartum<sup>246</sup>. Thus, including fathers in health assessments during the  
924 postpartum period and screening fathers for postpartum psychiatric disorders at similar time  
925 intervals as maternal screening is important. Efforts aiming to improve the overall health and  
926 functioning of the family unit will lead to best outcomes for the child<sup>243, 245</sup>.

927

928

929

930 **Box 2. Postpartum mental illness in low-income and middle-income countries**

931 In resource-poor settings, women of reproductive age typically have socio-economic and health  
932 challenges that interplay in mutually reinforcing ways.<sup>247, 248</sup> For example, low levels of education,  
933 low gender status, food insecurity, domestic abuse and lack of access to social and health services  
934 leave women and girls vulnerable to maternal mortality and chronic morbidities, including common  
935 mental health disorders.<sup>249, 250</sup> Indeed, the prevalence of common perinatal mental disorders in low-  
936 and lower-middle-income countries is higher than in high-income countries. One systematic review  
937 and meta-analysis<sup>21</sup> demonstrated a weighted mean prevalence of 15.6% in pregnant women and  
938 19.8% in women after childbirth. The most strongly associated factors for perinatal mental  
939 disorders are socio-economic disadvantage, unintended pregnancy, younger age, unmarried  
940 status, lacking intimate partner empathy and support, hostile inlaws, partner violence, insufficient  
941 emotional and practical support, a history of mental health problems and in certain settings, a  
942 female infant.<sup>21</sup>

943

944 **[H1] Considerations for management**

945 Mental health prevention and treatment investments in high-income countries is more than US\$50  
946 per year per person, compared with less than US\$2 in most LMIC,<sup>251</sup> resulting in a profound  
947 paucity of mental health providers in these settings. The intervention coverage for common mental  
948 disorders (including those that occur during the perinatal period) ranges from 7% to 28% in  
949 LMICs.<sup>252</sup> The low monetary allocation represents, in part, poor appreciation by decision makers of  
950 the effect of mental illness on population disability and socio-economic development, low levels of  
951 political will and capacity, and competing health and development priorities.<sup>253</sup> Interventions that  
952 are most likely to succeed in LMICs would, therefore, need to adopt a systems strengthening,  
953 integrated and low-cost approach. Examples showing promise have integrated mental health into  
954 primary care, maternal and child health services or into the routine community based delivery of

955 health services (carried out by trained lay workers, or primary care healthworkers, using a task-  
956 sharing approach)<sup>254</sup>. Emerging evidence supports the benefit of including poverty alleviation  
957 strategies in to mental health interventions.<sup>255</sup>

958

## 959 **[H1] Types of interventions**

960 A systematic review and meta-analysis of evidence of common perinatal mental disorder trials from  
961 LMIC, reported similar relative risk outcomes in studies carried out in high-income countries<sup>256</sup>. For  
962 the 13 trials selected, the pooled effect size for maternal depression was  $-0.38$ . In this review,  
963 trials that demonstrated positive results used several culturally adapted treatment paradigms,  
964 either singly or in combination. The Thinking Health Program in Pakistan was a cognitive behaviour  
965 therapy (CBT) intervention delivered in homes in a semi-rural setting by Lady Health Workers.<sup>257</sup>  
966 Uptake of the intervention leveraged the belief, within multi-generational households, that the  
967 intervention with the mother would improve the infant's well-being . In an urban, deprived setting in  
968 Chile, midwives and nurses were trained to deliver eight weekly structured psychoeducational  
969 group sessions. These sessions included information about symptoms and treatments, some  
970 problem solving strategies, behavioural activation strategies (such as scheduling pleasurable  
971 activities) and some cognitive techniques using postnatal examples<sup>258</sup> .

972 Subsequently, a trial in Zimbabwe<sup>259</sup> used peer counsellors to deliver six weeks of group problem  
973 solving therapy adapted for the local setting to postnatal women with depression. In this study,  
974 family members were co-opted to support the mothers through strategies identified in the problem  
975 solving and a specific treatment element that explored community resources and support systems  
976 was included. Six weeks after the intervention, the drop in mean EPDS score was greater in the  
977 PST group than the control group who received antidepressant therapy. No difference in outcomes  
978 between women with or without HIV was reported.

979



980 **Box 3. Targeting risk factors for postpartum depression.**

981 Assessing the common psychosocial risk factors for postpartum depression might have the  
982 following functions:

- 983 • Assisting in the initiation of targeted interventions or determining rational management  
984 decisions to mitigate the risks across several global settings through
  - 985 ○ risk reduction interventions (for example, referring women for social grants, to  
986 domestic violence support groups or to a women’s shelter and provision of  
987 integrated interventions for the mood disorder, which also addresses domestic  
988 violence<sup>260</sup>
  - 989 ○ activation of protective factors, (such as interpersonal therapy for relationship  
990 difficulties or activating social support networks)<sup>175, 260</sup>.
- 991 • Assisting in screening of women who have an increased risk of postpartum depression but  
992 do not currently have the disorder.
- 993 • Assisting in timely referral for support in women with suspected postpartum depression and  
994 complicated psychosocial risk factors who are reluctant to endorse symptoms during  
995 screening due to stigma and poor contextual validity of the screening tool in some global  
996 settings, among other reasons<sup>261</sup>. This approach acknowledges that there may be several  
997 contextual factors contributing to false negative mental health screening results.

998

999

1000 **Box 4. General management guidelines for non-psychotic psychiatric disorders**

- 1001 1. Identify somatic comorbidities and optimize their management.
- 1002 2. Check the mode of delivery, if complications were present and if delivery was experienced  
1003 as traumatic. In the case of post-traumatic stress symptoms, consider specific treatments.
- 1004 3. Assess for suicidal thoughts and intrusive thoughts of harm toward the baby. Consider the  
1005 safety of the baby and whether the mother can provide care for the baby if she is alone or if  
1006 other adult supervision is required.
- 1007 4. Ask the mother of her attitude towards her baby and observe maternal-child interactions.  
1008 Consider specific treatments with signs of problematic interactions or bonding.
- 1009 5. Review the feeding pattern of the baby. Address problems with breast or bottle-feeding.
- 1010 6. Provide strategies to preserve sleep, such as finding another person to feed the infant at  
1011 night.
- 1012 7. Assess psychiatric history before delivery. Review the nature and effectiveness of past  
1013 treatments, and restart previous effective treatment when appropriate.

1014

1015

1016

1017 **Figure 1, Mechanisms of postpartum psychiatric disorders.**

1018 Several factors have been implicated in the aetiology of postpartum psychiatric disorders, including  
1019 both postpartum depression and postpartum psychosis. These factors include psycho-social  
1020 factors and biological factors that are specific to pregnancy and the postpartum period, such as  
1021 drastic alterations in gonadal sex steroids and impaired mother-infant interactions. Whether the  
1022 aetiology of psychiatric disorders occurring in prenatally, during pregnancy or during the  
1023 postpartum period is different requires future study.

1024

1025 **Figure 2. Management of first onset postpartum psychiatric disorders.**

1026 Management of postpartum psychiatric disorders should take into account the diagnosis (such as  
1027 psychosis, anxiety or depression), symptom severity and, with regards to mood and anxiety  
1028 disorders, whether the mother is breastfeeding.

1029

1030 **Bibliography: [Au: There are 5 pairs of duplicated references, can you please fix these using**  
1031 **your reference manager:-- YES, this has been fixed. Thanks for letting me know!**

- 1032 • **27 and 147-- corrected**
- 1033 • **16 and 19-- corrected**
- 1034 • **30 and 162-- corrected**
- 1035 • **11 and 22—corrected.**
- 1036 • **20 and 56] --corrected**

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